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How This Manual is Organized

1. **Installing GeneLinker™**. Topics relating to installing, upgrading or uninstalling GeneLinker™.
3. **Using GeneLinker™**. Detailed descriptive and procedural topics covering all of GeneLinker™’s functionality.

Additional Sources of Information

**Readme.txt**
This file contains last minute additions to the documentation.

**Tips**
Most GeneLinker™ dialogs have a **Tips** button. Clicking a **Tips** button displays a brief hint about the functionality invoked by the dialog.

**Online Help**
GeneLinker™ has comprehensive online help built into the product. The content of the online help is the same as this printed manual.

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Installing GeneLinker™

System Specification

Overview

GeneLinker™ Gold requires a system that meets or exceeds the following specification:

- Microsoft Windows® NT 4.0 Service Pack 6a, Windows® 2000, XP, 95, 98 and ME. Windows® 2000, NT and XP are the preferred platforms as they are more stable and manage memory more effectively
- 256 MB RAM (512 MB RAM recommended)
- PII 400 MHz processor or better
- 500 MB hard disk space

GeneLinker™ Platinum is typically pre-installed on an IBM system that meets or exceeds the following specification:

- Microsoft Windows® 2000 Professional
- 2.5 GB of RAM
- Single Intel Xeon-2200 2.2 GHz processor
- NVIDIA 64MB Video card
- 18.2 GB Hard Drive
- 48X IDE CD-ROM
- 10/100 Ethernet card
- 3.5 inch 1.44MB Floppy drive

Network Requirements

- For floating licenses (Floating Server, Floating Client), GeneLinker™ requires a TCP/IP network, and that the TCP/IP protocol be installed on both the license server and the user workstations. In addition, one of the three protocols SNMP, NetBEUI, or IPX/SPX must be installed on both the server and the workstations (GeneLinker™ uses the protocol service to determine the hostid of the system). Any mix of the three protocols on the server and on different workstations is acceptable. By default, many of these protocols are available.
- For other licenses (Licensed Client (node-locked), Demo), there are no network requirements.

We recommend that license servers (for floating licenses) be installed on machines that are running the Windows® NT or Windows® 2000 operating system.

Related Topics:

GeneLinker™ Database
**GeneLinker™ Database**

**Overview**
GeneLinker™ stores all of its dataset, experiment, gene, gene list, and annotation data in a database on the local file system under the GeneLinker™ directory (MMC) in a folder named Repository. GeneLinker™ currently supports a MySQL, DB2, or Oracle database. The MySQL source code is provided on the GeneLinker™ CDROM in the MySQLSrc directory.

**MySQL**
The default database used by GeneLinker™ is MySQL. If you are using this database, you are not required to install, configure, or maintain the database in any way. When GeneLinker™ is started, it will start the database, and when GeneLinker™ is shut down, it will shut down the database.

**DB2 and Oracle**
If you choose to use a DB2 or Oracle database, then you will have to install DB2 or Oracle on the GeneLinker™ computer and create a valid account for GeneLinker™ to use. You will have to start and stop the database manually. See Setting Up a DB2 GeneLinker™ Database for details of the DB2 setup process. See Setting Up an Oracle GeneLinker™ Database for details of the Oracle setup process.

**Notes**
- The GeneLinker™ database should not be tweaked or configured outside of GeneLinker™.
- It is recommended that you **do not** use the GeneLinker™ database with any other application or data. Doing so could result in an unusable, corrupted database.
- The GeneLinker™ uninstall procedure has an option to keep or remove the database.
- As an example, a typical file size would be approximately 0.5 Megabytes for a dataset consisting of 1000 genes by 100 samples.

**Related Topics:**
- Setting Up a DB2 GeneLinker™ Database
- Setting Up an Oracle GeneLinker™ Database
- Saving

**Setting Up a DB2 GeneLinker™ Database**
Overview
Using a DB2 GeneLinker™ database requires some preliminary setup.

Actions
1. If you do not already have access to a running DB2, install one. Visit the following site for full details: http://www.ibm.com/software/data/db2/
2. As the database administrator, create a database in DB2 called, for example, BIO_DB.
3. Create an account (user name and password) for accessing the BIO_DB database.
4. Configure your DB2 installation so that the BIO_DB database is accessible using the above account on the computer where GeneLinker™ is installed.
5. Run the DB2ConfigurationUtility.bat application found in the Maintenance folder of the GeneLinker™ installation folder. You will be prompted for the name of the database (BIO_DB in this example), the user name, and password.
   Warning: this password appears in plain text in the GeneLinker™ configuration file (GeneLinker.conf). Please take whatever precautions are required to secure this file or use a unique password for this application (to limit the risk if this password becomes known to others).

If there are any problems during step 5 (for example, you mistype the name of the database), then GeneLinker™'s configuration will not be changed.

Note that a DB2 GeneLinker™ database cannot be shared by multiple users. Attempting to do so will corrupt the database and cause valuable information to be lost.

Related Topic:
GeneLinker™ Database

Setting Up an Oracle GeneLinker™ Database

Overview
Using an Oracle GeneLinker™ database requires some preliminary setup.

Actions
1. If you do not already have access to a running Oracle database, install one. Visit the following site for full details: http://www.oracle.com/ip/deploy/database/oracle9i/
2. As the database administrator, create a database in Oracle called, for example, BIO_DB.
3. Create an account (user name and password) for accessing the BIO_DB database.
4. Configure your Oracle installation so that the **BIO_DB** database is accessible using the above account on the computer where GeneLinker™ is installed.

5. Run the **OracleConfigurationUtility.bat** application found in the **Maintenance** folder of the GeneLinker™ installation folder. You will be prompted for the name of the database (**BIO_DB** in this example), the user name, and password.

   **Warning:** this password appears in plain text in the GeneLinker™ configuration file (GeneLinker.conf). Please take whatever precautions are required to secure this file or use a unique password for this application (to limit the risk if this password becomes known to others).


   If there are any problems during step 5 (for example, you mistype the name of the database), then GeneLinker™’s configuration will not be changed.

   **Note** that an Oracle GeneLinker™ database cannot be shared by multiple users. Attempting to do so will corrupt the database and cause valuable information to be lost.

**Related Topic:**

GeneLinker™ Database

**Installation Procedure**

**Overview**

If you are upgrading GeneLinker™ Gold to Version 3.1, please follow the instructions in Upgrading GeneLinker™ Gold.

If you are upgrading GeneLinker™ Platinum to Version 2.1, please follow the instruction in Upgrading GeneLinker™ Platinum.

Please follow the installation process appropriate to your license type.

**Licenses**

GeneLinker™ license types.

- **A Demonstration Client** is a time-limited single license for a single copy of GeneLinker™ to run on a single computer.

- **A Licensed Client (node-locked)** is a single license for a single copy of GeneLinker™ to run on a single computer.

- **Floating License Server / Floating Client** license types provide a network solution for multiple users. When GeneLinker™ is started on a client workstation, it requests a license from the GeneLinker™ license server. If a license is available, GeneLinker™ will run on the client workstation.

See License Overview for further information on licenses.
All License Types Start Here
GeneLinker™ uses an installer program to make the installation process simple.

1. Insert the GeneLinker™ CD into your drive. The installation process should start automatically. **Skip to step 7 if you see the installation welcome dialog on your screen.**

2. With the GeneLinker™ CD in your drive, click the Windows **Start** button.

3. Select **Run**.

4. Navigate to the appropriate directory on the GeneLinker™ CD-ROM.

5. Double-click on the file **setup.exe**. The installation process initializes.
6. The **Welcome** dialog is displayed.

7. Click **Next** to continue.

8. It is recommended that you close any other applications you may be running. Click **Next** to continue.

9. Read the license agreement displayed in the dialog and click **Yes** to continue.
10. Read the ReadMe.Txt file displayed in the dialog and click **Next** to continue. If you are installing GeneLinker™ Platinum, skip to step 12.

11. Select the type of license you have.
   - If you have a demo or a single, node-locked license, click **Licensed Client**.
   - If you have a floating license and your machine is not to be the license server, click **Floating Client**.
   - If you have a floating license and your machine is to be the license server, click **License Server**.
12. If the information shown in the dialog is incorrect, type over the provided name and company information. Click **Next** to continue.

13. If the default destination folder is not where you want GeneLinker™ installed, click **Browse** and select the correct folder. Click **Next** to continue.

14. If the default program folder is not where you want the program icon placed, select another folder. Click **Next** to continue.

15. The installation system information is displayed for you to read. Click **Next** to continue.
16. The GeneLinker™ files are transferred onto your computer.

17. The GeneLinker™ license manager is configured.

18. Click **Finish**. The **Setup** dialog closes.

19. At this point, the installation process is complete. You may need to change the license information within GeneLinker™ depending on the type of license you have.

- If you have a **Demonstration Client** or a **Floating Client** license, GeneLinker™ is ready for use.

- If you have a single, node-locked license (**Licensed Client**) or a floating **License Server** license, the license information that was installed needs to be changed. Please follow the instructions in the topic linked to in the table below.

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**Related Topics:**
Starting the Program
If you have an expired Demonstration Client license:
If your demo license expires, please contact Molecular Mining Corporation (MMC) sales to purchase GeneLinker™.
Updating Demo License to Licensed Client
Updating Demo License to License Server
Demo License Time Extension

If your license changes:
Changing from Licensed Client to License Server

If your system or server changes:
Licensed Client: Configuration Change
Licensed Client: Moving from One Computer to Another
License Server: Moving from One Computer to Another
License Server: Configuration Change
Updating Floating Client after Server Move

**Upgrading GeneLinker(TM)**

**Gold**

**Upgrading GeneLinker™ Gold**

**Overview**
Please follow these instructions for upgrading GeneLinker™ Gold to Version 3.1.

- **If your current version of GeneLinker™ Gold is less than Version 2.5**, you will need to Uninstall the old version of GeneLinker™ before installing the new one. If you try to do the upgrade without uninstalling the old version first, you will see the message, 'The GeneLinker™ data repository on this computer predates GeneLinker™ Gold 2.5 and cannot be upgraded by this installer. Before installing this new version of GeneLinker™, you must first remove the old version using Add/Remove Programs from the Control Panel.'

- **If you have a floating client license**, this upgrade should be performed only after the license server has been upgraded.

GeneLinker™ Gold uses an installer program to make the upgrade process simple. If you are running GeneLinker™ Gold, please exit the application before starting the upgrade process.

**Actions**
1. Insert the GeneLinker™ CD into your drive. The upgrade process should start automatically. If you have GeneLinker™ running, you will be prompted to exit it. **Skip to step 7 if you see the welcome dialog on your screen.**
2. With the GeneLinker™ CD in your drive, click the Windows **Start** button.
3. Select **Run**.
4. Navigate to the appropriate directory on the GeneLinker™ CD-ROM.

5. Double-click on the file **setup.exe**. The upgrade process initializes.

6. The **Welcome** dialog is displayed.
7. Click **Next** to continue. A message is displayed. If there is sufficient space on your disk, a backup of your data will be made. If there is insufficient disk space for the backup, the following message is displayed, "Before running GeneLinker™ Gold 3.1, we recommend strongly that you make a backup copy of the folder which holds your GeneLinker™ data: <path of repository folder>. This folder takes up about <size of repository> of disk space. Your data repository will be upgraded automatically to a new format the first time you run GeneLinker™ Gold 3.1. The new, upgraded repository is not compatible with earlier versions of GeneLinker™."

8. Click **OK**.
9. The GeneLinker™ Gold 3.1 files are copied to your computer. If you have a demo license, a message is displayed indicating a new demonstration license has been installed.

10. Click **OK**.

11. Click **Finish**. The **Setup** dialog closes.

12. At this point, the installation part of the upgrade process is complete. You may need to change the license information within GeneLinker™ depending on the type of license you have.

   - If you have a **Demonstration Client** or a **Floating Client** license, GeneLinker™ Gold 3.1 is ready for use once the computer has been rebooted.
   - If you have a single, node-locked license (**Licensed Client**) or a floating **License Server** license, the license information that was installed needs to be changed. Please follow the instructions in the topic linked to in the table below.

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Related Topic:
Starting the Program

Platinum

Upgrading GeneLinker™ Platinum

Overview
Please follow these instructions for upgrading GeneLinker™ Platinum to Version 2.1.

- **If your current version of GeneLinker™ Platinum is less than Version 1.2**, you will need to Uninstall the old version of GeneLinker™ before installing the new one. If you try to do the upgrade without uninstalling the old version first, you will see the message, ’The GeneLinker™ data repository on this computer predates GeneLinker™ Platinum 1.2 and cannot be upgraded by this installer. Before installing this new version of GeneLinker™, you must first remove the old version using Add/Remove Programs from the Control Panel.’

GeneLinker™ Platinum uses an installer program to make the upgrade process simple. If you are running GeneLinker™ Platinum, please exit the application before starting the upgrade process.

Actions
1. Insert the GeneLinker™ CD into your drive. The upgrade process should start automatically. If you have GeneLinker™ running, you will be prompted to exit it. **Skip to step 7 if you see the welcome dialog on your screen.**
2. With the GeneLinker™ CD in your drive, click the Windows **Start** button.
3. Select **Run**.
4. Navigate to the appropriate directory on the GeneLinker™ CD-ROM.
5. Double-click on the file **setup.exe**. The upgrade process initializes.

6. The **Welcome** dialog is displayed.
7. Click **Next** to continue. A message is displayed. If there is sufficient space on your disk, a backup of your data will be made. If there is insufficient disk space for the backup, the following message is displayed, "Before running GeneLinker™ Gold 3.0, we recommend strongly that you make a backup copy of the folder which holds your GeneLinker™ data: <path of repository folder>. This folder takes up about <size of repository> of disk space. Your data repository will be upgraded automatically to a new format the first time you run GeneLinker™ Gold 3.0. The new, upgraded repository is not compatible with earlier versions of GeneLinker™."

8. Click **OK**.
9. The GeneLinker™ Platinum 2.1 files are copied to your computer. If you have a demo license, a message is displayed indicating a new demonstration license has been installed.

10. Click **OK**.

11. Click **Finish**. The Setup dialog closes.

12. At this point, the installation part of the upgrade process is complete. You may need to change the license information within GeneLinker™ depending on the type of license you have.

   • If you have a **Demonstration Client** license, GeneLinker™ Platinum 2.1 is ready for use once the computer has been rebooted.

   • If you have a single, node-locked license (**Licensed Client**), the license information that was installed needs to be changed. Please follow the instructions in the topic linked to in the table below.

<table>
<thead>
<tr>
<th>License Type</th>
<th>Procedure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Licensed Client</td>
<td>Updating Demo License to Licensed Client</td>
</tr>
</tbody>
</table>
Uninstalling GeneLinker(TM)

Uninstallation Procedure

Overview
Use this procedure to remove the GeneLinker™ application from your computer. If GeneLinker™ is running, close it before you begin to uninstall.

Actions
1. Click the Windows Start button. Under Settings, click Control Panel.
2. On the Control Panel, double-click Add/Remove Programs.
3. Click on GeneLinker. The program is highlighted.
4. Click the Change/Remove button next to GeneLinker™. The Reinstall or Remove dialog is displayed.
5. Click the Remove option to select it. Click Next. The Confirm File Deletion dialog is displayed.
6. Click **OK** to remove the application from your system. A dialog is displayed giving you the option to remove or delete your data.

Removing (Deleting) the Repository

- Deleting the repository completely removes all genes, datasets that have been imported, experiments, and gene lists. **If you want to preserve your working data, do not delete the repository.**

7. If you want to delete the repository, check the **Remove GeneLinker's data repository** box.
8. Click **Continue**.

**Related Topic:**

Installation
Welcome to GeneLinker™

Thank you for choosing GeneLinker™ as your gene expression analysis system. The GeneLinker™ family of products are designed to help you discover underlying patterns in the data generated by modern high-throughput gene expression measurement techniques; the first step in discovering new relationships among genes.

Introduction

This tour describes the GeneLinker™ main window and outlines the program's major functionality groups (e.g. data import, preprocessing, clustering, visualization, and for platinum - classification). The fastest way to learn to use GeneLinker™ is to finish this tour and then run the tutorials.

Terminology

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
</table>
| Dataset   | A dataset is either a raw or preprocessed set of expression values for a number of genes over a number of samples. A dataset can have reliability measurements or variables associated with it. For a complete description see Datasets Overview and Reliability Measures.  
  - A standard dataset contains a single value for each gene for every sample (some may be replicate measurements within or between chips; in an incomplete dataset, one or more values are null or missing).
  - A two-color dataset contains two values for each gene for every sample. One value is the treatment expression level and the other is the control expression level. See Two-Color Data. |
| Experiment| An experiment is a dataset that has had its gene or sample order organized by the application of an experiment process such as clustering. |
| Variable  | In GeneLinker™, a variable is a column of data other than gene expression values used to differentiate samples. See Variables Overview.
  A variable can store:
  - Phenotypic observations about the samples.
    e.g. malignant vs. benign.
  - Predictions of phenotypes by a trained classifier.
    e.g. predicted malignant vs. predicted benign.
  - Information about experimental conditions. |
e.g. high dose vs. low dose; time the sample was taken; animal A vs. animal B vs. animal C, etc.

**GeneLinker™ Tour - Main Window Layout**

**Overview**

GeneLinker™ runs in one main window. At the top of the window is the menu bar and the toolbar. The work area is divided into three panes (outlined in red): the navigator, the description pane, and the plots pane. At the bottom is the status bar.

![GeneLinker™ Main Window Layout](image)

**The Navigator (upper left)**

The navigator organizes your data and gives you access to it. All items listed in the navigator have already been saved into the GeneLinker™ database. There are three tabs in this pane, each listing a specific type of data.

- **The Experiments** tab displays a hierarchical tree of your datasets and experiments. Each item in the tree is tagged with an icon to indicate its type (e.g. dataset, hierarchical clustering experiment, principal components experiment, etc.).

- **The Genes** tab displays an alphabetical listing of all your genes.

- **The Gene Lists** tab displays an alphabetical listing of all of your gene lists. Clicking a tab brings it to the front. Clicking an item in the navigator highlights it and makes it the selected item. Information about the selected item is displayed in the description pane. Program functions are applied to the selected item.
The Description Pane (lower left)
The description pane displays information about the item selected in the navigator, or a gene selected in a table or plot. This information can include the name of the item, the number of genes and samples it contains, its creation date, parameters used in its creation (if it is an experiment), and so forth.

The Plots Pane (right)
The plots pane is the place for visualizing your data and experiments. When you use the table viewer or a create a plot, it is displayed in the plots pane. The plots in the plot pane can be arranged by dragging them or by using the Cascade Windows item on the Window menu.

Shortcuts and Tips
GeneLinker™ was designed for ease of use. Right-clicking an item (such as a dataset, or gene in the navigator or on a plot) displays a shortcut menu giving you quick access to its functions.

Most dialogs (such as normalization or filtering) have a Tips button. Clicking Tips displays a brief description of the function and how to use it. For example:

If you want to know what function an icon invokes, hover the mouse over the icon for a moment. A tooltip is displayed naming the function.

GeneLinker™ Tour - Clustering and PCA

Clustering / PCA and Visualization

Introduction to Clustering
Clustering is used to group biological samples or genes into separate clusters based on their statistical behavior. The main objective of clustering is to find similarities between experiments or genes (given their expression ratios across all genes or samples,
respectively), and then group similar samples or genes together to assist in understanding relationships that might exist among them.

**Clustering**

- Apply K-Means, Jarvis-Patrick, or agglomerative hierarchical clustering to your dataset, or perhaps try a Self-Organizing Map (SOM). The results of each clustering experiment is listed in the **Experiments** navigator under the dataset it was based on. Each experiment result item is tagged with an icon to indicate the experiment type.

- Visualize the Clustering Experiment Results - GeneLinker™ has an extensive set of plots that can be used to visualize the results of clustering hopefully revealing interesting or significant patterns.

**Introduction to Principal Component Analysis**

Component Analysis is an unsupervised or class-free approach to finding the most informative or explanatory features in data. In particular, **Principal Component Analysis (PCA)** substantially reduces the complexity of data in which a large number of variables (e.g. thousands) are interrelated, such as in large-scale gene expression data obtained across a variety of different samples or conditions. PCA accomplishes this by computing a new, much smaller set of **uncorrelated variables** which best represent the original data. PCA is a powerful, well-established technique for data reduction and visualization. 2D and 3D PCA plots often place objects with similar patterns near each other.

**Principal Component Analysis (PCA)**

- Apply PCA by genes or by samples. Again, the experiment results are listed in the **Experiments** navigator tagged with the PCA icon.

- Visualize the PCA Results - GeneLinker™ offers a variety of 2D plots and a 3D Score plot to give a clear picture of the hidden structure in the data.

**Platinum**

**GeneLinker™ Tour - Platinum SLAM™ Classification**

**Please note**: these functions are introduced within a conceptual 'workflow' for the purpose of introduction only. Within GeneLinker™, you are free to apply any appropriate function to your data at any time.

1. **Import Gene Expression Data**
A training dataset (expression values with known classes) is required to train an artificial neural network (ANN) classifier. A test dataset can be imported to test a trained classifier. The two datasets must be studies of the same phenomenon (i.e. the variable type for both is the same, e.g. SRBC Tumors).

2. Import Variable Data
Import the classes (e.g. EWS, NB, BL, RMS) for the training dataset.

3. Discretize the Expression Data
Expression data is continuous. To apply the SLAM™ data mining algorithm, the data must first be discretized.

4. Apply SLAM™ Association Mining and Visualize the Results
SLAM™ (Sub-Linear Association Mining) is a technology that finds hidden linear and non-linear correlations in discretized gene expression data. The SLAM™ association viewer displays the results of running SLAM™ and allows you to work with the results.

5. Create Gene List
As an aid to supervised learning, a gene list is created from the genes (features) identified as significant by SLAM™. If necessary, this gene list can be used to filter the test dataset to ensure it contains the same genes as the training dataset.

6. Create an ANN Classifier and View Training Results
Creating an ANN classifier is the process of exposing a committee of neural networks to data with known classes of a particular type. The training results can be displayed in a classification plot or an MSE plot.

7. Classify Data and Visualize the Classification Results
Classification is the process of using a trained classifier to predict the classes of the test dataset.

Platinum
GeneLinker™ Tour - Platinum IBIS Classification

Overview
IBIS (Integrated Bayesian Inference System) is a system that is able to predict class membership for a gene expression dataset containing measurements for the same phenomenon as the dataset used to train the IBIS classifier. One of the major strengths of the IBIS method is its ability to reveal nonlinear and non-monotonic associations
between pairs of genes and their concerted response to a particular stimulus such as a drug.

**Platinum Classification and Prediction Using IBIS**

**Please note:** these functions are introduced within a conceptual 'workflow' for the purpose of introduction only. Within GeneLinker™, you are free to apply any appropriate function to your data at any time, in any order.

1. **Import Data**  
   A *training dataset* (expression values with known classes) is required for creating an IBIS classifier. A *test dataset* can be used to test the classifier. The two datasets must be studies of the same phenomenon (the variable type for both is the same).

2. **Import Variable Data**  
   Import the class observations for the training dataset.

3. **Preprocess Your Data**  
   GeneLinker™ offers a variety of preprocessing options which can be applied one or more times to a dataset. You can then view the preprocessed data as you would raw data (table viewer or color matrix plot).

4. **Optionally, Perform an IBIS Search**  
   The IBIS search process creates a list of proto-classifiers, one for each gene or gene pair. Each proto-classifier consists of the gene/gene pair identifier, an accuracy value, and the MSE value. The proto-classifier list can be viewed in the IBIS search results viewer.

5. **Create a Classifier and View Results**  
   You can create a Linear Discriminant Analysis (LDA), Quadratic Discriminant Analysis (QDA), or a Uniform/Gaussian Discriminant Analysis (UGDA) classifier from a proto-classifier (IBIS search results), or from any gene or gene pair. The results can be viewed in an IBIS Gradient plot.

6. **Classify Data and Visualize Results**  
   Classification is the process of using a trained classifier to predict the classes of data (of the same type). An IBIS classifier can be applied to a dataset that contains the gene or gene pair used to create the classifier. The results can be viewed in a Classification plot or an IBIS Gradient plot.

**GeneLinker™ Tour - Common Functions**

**Creating Gene Lists**
A gene list is a list of one or more genes. Gene lists can be used to filter datasets to create smaller datasets for detailed study, or to share gene information with colleagues.

**Lookup Gene in a Public Database**
Select a gene in the **Genes** navigator or on a plot and lookup information about it in a public database. The gene information is displayed in your web browser.

**Recording Your Work - Annotations and Reports**
You can annotate your genes, datasets or experiments. These annotations are included within appropriate GeneLinker™ reports.

GeneLinker™ can generate a report on a specific item such as a gene, dataset, or experiment. Another type of report that can be generated is a workflow report. It includes all of the steps from the raw data to the selected experiment item.

**Exporting Data and Images**
A dataset can be exported to a text file.
Images can be exported to .png files.

**GeneLinker™ Tour - Conclusion**

**Overview**
You have now completed the introductory product tour. You have been introduced to the GeneLinker™ main window, concepts and workflows. The next step in mastering GeneLinker™ is to run the tutorials. Each tutorial leads you through an analysis of a real dataset exercising the majority of GeneLinker™’s powerful functionality.

**Related Topics:**
List of Tutorials

**Product Information**

**GeneLinker™ Product Suite**

**Overview**
**GeneLinker™ Gold** is the first member of the GeneLinker™ family of products developed by Molecular Mining Corporation (MMC). This application gives you powerful tools to explore the data gathered from your gene expression experiments. With GeneLinker™ Gold, you can preprocess your data, perform clustering experiments, or principal components analysis and view the results of those experiments in many
different plots and charts.

**GeneLinker™ Platinum** is the breakthrough product developed by MMC. GeneLinker™ Platinum contains all the functionality of GeneLinker™ Gold plus many additional features including the proprietary SLAM™ technology. SLAM™ (Sub-Linear Association Mining) is an extremely fast, scalable association-mining algorithm that uses a novel sampling and binning scheme employing various hypothesis testing methods. This new technology breaks the combinatorial barriers that previously prevented the discovery and measurement of statistically interesting higher-order correlations in gene expression datasets. SLAM™ can be applied to gene-gene and gene-phenotype interactions. It can also be used in the construction of predictive models relating any of: expression, proteomics, SNPs/haplotypes, toxicity response, therapeutic response, environmental, clinical outcomes, etc.

**GeneLinker™ Diamond** is an enterprise-wide software solution for the analysis of gene expression datasets. This innovative product offers all of your users the complete functionality of GeneLinker™ Platinum with the added benefit of a unified data source. The GeneLinker™ Diamond relational database repository of all of your genes, gene lists, datasets and experiments makes all of your data and discoveries immediately available to all of your scientists.

**Related Topics:**
- GeneLinker™ Tour
- Feature List

**GeneLinker™ Feature List**

**Overview**

**Designed for ease of use, GeneLinker™ features:**
- Straightforward interface to import spotted microarray, Affymetrix® chip, or similar data including two-color GenePix data;
- Tabbed pane navigator that provides hierarchical views of all datasets and experiments (tagged with parameter settings), genes, and gene lists;
- Description pane that displays information about the selected dataset, experiment, gene, or gene list;
- Relational database (MySQL, DB2 or Oracle) for storage of GeneLinker™ data.
- Automatic saving of experiment results.
- HTML-based reporting (single experiment or entire workflow);
- Advanced image capture.

**Designed to help in data exploration, GeneLinker™ features:**
- Table view or color matrix plot of datasets (raw or preprocessed);
• Estimation/elimination of missing data values;
• Value removal;
• Advanced filtering and gene prioritization based on N-Fold induction and repression and difference measures;
• Preprocessing and data normalization capabilities (e.g. scaling, transformation, Lowess);
• F-Test with results viewer;
• Summary statistics chart;
• Hierarchical clustering of genes or samples using single, average, or complete linkage with distance metric options including Euclidean, Manhattan, Pearson Correlation, etc;
• Non-hierarchical clustering of genes or samples using K-Means or Jarvis-Patrick methods;
• Self Organizing Map clustering with plots;
• Principal Component Analysis with 2D plots and 3D Score plot;
• A wide variety of plots including Scatter, Coordinate, Centroid, Cluster, Matrix Tree, etc. with user-selectable data range, color schemes, and shared selection;
• Profile Matching to one or more reference genes;
• Annotations editor/viewer;
• Direct links to external data sources such as GenBank, UniGene, Affymetrix, etc.
• Gene list creation and filtering.

Platinum
GeneLinker™ Platinum builds on the functionality introduced in GeneLinker™ Gold
• Patented SLAM™ association mining technology to aid in feature identification for use in supervised learning;
• Supervised Learning (training of neural networks to predict gene expression data classes) with informative plots.
• IBIS Classification (Integrated Bayesian Inference System) including IBIS Search (with viewer), classifier creation from search results or a selected gene or gene pair.
• Visualize IBIS classifier in an IBIS Gradient plot.
• Classification using an ANN or an IBIS classifier.

Related Topics:
GeneLinker™ Tour
Tutorials

Tutorials/Use Case Scenarios

Tutorial 1: Gene Expression During Rat Spinal Cord Development
• This tutorial covers data import and transposition, normalization, renaming experiments, K-Means clustering, matrix tree, centroid, and cluster plots, generating experiment and workflow reports, and exporting images.

Tutorial 2: Analysis of NCI60 Data
• This tutorial covers importing and preprocessing data, renaming datasets, estimating missing values, agglomerative hierarchical clustering, matrix tree plots, color matrix plots, resizing and customizing plots, and generating reports.

Tutorial 3: Jarvis-Patrick Clustering
• This tutorial covers estimating missing values, normalization, performing Jarvis-Patrick clustering analysis on the datasets from the first two tutorials, and displaying data in a matrix tree plot.

Tutorial 4: Self-Organizing Maps (SOMs)
• This tutorial covers importing data, using the table viewer, the summary statistics chart, value removal, filtering, normalization, using Self-Organizing Maps to cluster Leukemia data, visualizing SOM results in a SOM plot and in a cluster plot.

Tutorial 5: Principal Component Analysis (PCA)
• This tutorial demonstrates how to use Principal Component Analysis as a method of extracting more information from data. The tutorial covers data import and displaying PCA results in various plots including: scree, loadings line, color matrix, score (raw and normalized) and 3D score (raw and normalized) plots.

Sample Workflow Using Spotted Array N-Fold Culling With Log Transformation
• This workflow is used for ratio (Cy3/Cy5) data to filter out genes that do not show a large induction or repression in any sample in the dataset, and then to log normalize the data so that inductions and repressions have equal but opposite sign.

Platinum Tutorial 6: Learning to Distinguish Cancer Classes
• This tutorial demonstrates how to train GeneLinker™ Platinum's artificial neural networks ANNs) to distinguish between sample classes. As an example, data on four similar tumor types is studied. Program features covered include importing variables, the SLAM™ association-mining technology (algorithm and viewer), creating gene lists for filtering, filtering, classification, and classification plots.

Platinum Tutorial 7: IBIS Classification
• This tutorial demonstrates how to search for a gene to use as an IBIS classifier. One IBIS classifier is produced using Linear Discriminant Analysis (LDA) and a second is produced using Quadratic Discriminant Analysis (QDA). An IBIS Gradient plot is used to analyze the results of the classifier creation.

Tutorial 8: Affymetrix Data
• This tutorial demonstrates how to use Affymetrix data in GeneLinker™.

Tutorial 1: Gene Expression During Rat Spinal Cord Development
Tutorial 1: Introduction

Welcome to the first tutorial. This tutorial introduces you to clustering by walking you through a simple analysis of a real dataset. You will be shown how to normalize the data, cluster it, and then visualize the clustering results in different types of plots.

Skills You Will Learn:
How to import gene expression data from a file into the GeneLinker™ database.
How to use the table viewer.
How to normalize a dataset.
How to perform clustering experiments.
How to display plots.
How to generate a report and export an image.

Dataset Information
This tutorial uses a dataset described in a 1998 paper (see URL http://www.pnas.org/cgi/content/abstract/95/1/334) by Xiling Wen, Stefanie Fuhrman, George S. Michaels, Daniel B. Carr, Susan Smith, Jeffrey L. Barker and Roland Somogyi, 'Large-scale temporal gene expression mapping of central nervous system development.' Proc. Nat. Acad. Sci. USA, Vol. 95, pp.334-339, January 1998. You may find it useful to have a copy of the paper on hand -- either on your screen, or printed out -- while working through this tutorial. In this tutorial this paper is referred to as 'Wen et al.', or simply 'Wen'.

The raw data represent RT-PCR product ratios (sample/control densities from gel images), averaged over three measurements. This expression study was designed to discover relationships between members of important gene families during different phases of rat cervical spinal cord development, assayed over nine time points before (E=embryonic) and after birth (P=postnatal). The selection covers a range of developmental markers and intercellular signaling genes, involving neurotransmitters and growth factors.

Wen et al. first clustered the genes 'from the combined 17 dimensional vectors of nine expression values (ranging between 0 to 1) and eight slopes (ranging between -1 and +1; slopes were calculated based on a reduced time interval of 1, not taking into account the variable time intervals). [They] included slopes to take into account offset but parallel patterns.' Computing this difference information (which they call 'slope') cannot be done entirely within GeneLinker™. For the purpose of this tutorial, slopes are ignored, and the software is used only to investigate the expression levels.

Tutorial Length
This tutorial should take about an hour, depending on how long you spend investigating the data, and how fast your machine is. Note that if you must stop part way through the tutorial, simply exit the program by selecting Exit from the File menu. The data and experiments you have performed to that point are saved automatically by GeneLinker™.
The next time you start GeneLinker™, you can continue on with the next step in the tutorial.

**Tutorial 1: Step 1 Start GeneLinker™ and Import the Data**

**Start GeneLinker™**
1. Double-click the GeneLinker™ program icon on your desktop to start the application.
   • See GeneLinker Tour - Main Window Layout for a brief introduction to the GeneLinker™ program window.
   • In the upper left pane (navigator), you will see three tabs: Genes, Gene Lists and Experiments. They give you three views of the data in the GeneLinker™ database. Clicking a tab brings that view to the front.

**Import the Gene Expression Data**
1. Click the Experiments tab to display the Experiments navigator. All datasets and experiments present in the database are listed here in a hierarchical tree.
2. If the dataset Spinal_cord is present, skip the rest of this step and continue with step 2 - View and Normalize the Data.
3. Click the Import Gene Expression Data toolbar icon (far left on toolbar - to discover what function an icon invokes, hover the mouse pointer over it for a couple of seconds. A tooltip is displayed naming the function), or select Import from the File menu and Gene Expression Data from the sub menu. The Data Import dialog is displayed.

4. GeneLinker™ uses a template to interpret or parse the data values as they are read in from the data file. The installed default for the template is Tabular. If the Template listed on the Data Import dialog is NOT Tabular, click the Template Change button. This displays the Import Templates dialog. Click Tabular and click Select. The Data Import dialog is updated showing Tabular as the template.
5. You now have to tell GeneLinker™ where the data file is located. Click the Source File Change button. The Open dialog is displayed.
6. Navigate to the GeneLinker™ Tutorial folder (if necessary) and click the file Spinal_cord.txt. The file is highlighted.
7. Click Open. The Data Import dialog is updated with the source file.

8. Ensure that the Gene Database is set to GenBank (use the drop-down list to choose GenBank if necessary). If you import a file that has gene identifiers other than GenBank, set the Gene Database to match your data. For the Spinal_cord dataset, GenBank is correct.
9. Click Import. The Import Data dialog is displayed.
GeneLinker™ examines the file and offers to transpose it. Within GeneLinker™, datasets have the genes in columns and the samples in rows.

When importing data using a Tabular template, GeneLinker™ assumes that the more numerous dimension of your data represents genes (most microarray experiments involve more genes than samples). If this is so (as in this tutorial), then clicking OK is all that is required.

Note: the options Use Sample Names and Use Gene Names are checked and disabled in the Import Data dialog box. GeneLinker™ has recognized that in this dataset, the first row and column contain alphameric labels. Gene expression data is always numeric, hence the disabled checkboxes.

10. Click OK. The data is imported and an item named Spinal_cord is added to the Experiments navigator. This represents your raw data, which is now available to perform experiments on using the various GeneLinker™ functions.

   Note: when a dataset is imported, it is assigned a unique name. If the incoming dataset has the same name as an existing one, it is renamed automatically by the program (a numeric identifier is appended to the original name). For example, if you import Spinal_cord.txt again, it will be assigned the name Spinal_cord 1.

**Tutorial 1: Step 2 View and Normalize the Data**

The table viewer displays a spreadsheet-like view of the data in a dataset.
View the Data with the Table Viewer

1. If the Spinal_cord dataset in the Experiments navigator is not already highlighted, click it.

2. Click the Table View toolbar icon, or select Table View from the Explore menu, or right-click the item and select Table View from the shortcut menu. The data is displayed in table form in the right-hand pane (plots pane).

3. Click the right scrollbar arrow at the bottom of the table viewer to scroll right about 6 or 8 genes so you see the genes L1, NFL, and NFM.

Note: NFL expression ranges up to 14.92 and NFM up to 27.69 over the control, while L1 never gets above 0.96 of the control concentration. While the difference between strongly expressed and weakly expressed genes is interesting, it's not what we're currently after. Instead, normalize each gene by dividing by its maximum expression ratio.

To learn more about how to use the table viewer, please see Table Viewer Functions.

Normalize the Data

GeneLinker™ offers multiple normalization, filtering, and other data preprocessing techniques which can be applied one or more times (in various combinations) to a dataset. In this tutorial, the data is normalized by dividing by the maximum. Please see Normalization Overview for details on all of the normalization operations.

1. If the Spinal_cord dataset in the Experiments navigator is not already highlighted, click it.

2. Click the Normalize toolbar icon, or select Normalize from the Data menu, or right-click the item and select Normalize from the shortcut menu. The first Normalization parameters dialog is displayed.
3. Double-click the Other Transformations radio button, or ensure Other Transformations is selected and click Next. The second Normalization dialog is displayed.

4. Double-click the Divide by Maximum radio button, or ensure Divide by Maximum is selected and click Finish. The Experiment Progress dialog is displayed.

The dialog is dynamically updated as the normalization operation is performed. Upon successful completion, a new Normalization item is added to the Experiments navigator, attached to and below the Spinal_cord raw dataset. It is named something like Normalization (2002-08-01 16:04:50) - using the current date and time.
To learn about cancelling an operation or experiment, please see Cancelling an Operation or Experiment.

Tutorial 1: Step 3 View Parameters and Rename Experiment

View Experiment Parameters
Thinking ahead, what would happen if you tried out six different normalizations on the same dataset today, and then came back in tomorrow and wanted to re-examine those results? How would you determine which node on the experiment tree corresponds to a particular analysis sequence?

- You can always determine which parameters generated a certain node on the experiment tree by right-clicking it and selecting Show Parameters from the shortcut menu (or by clicking the experiment and selecting Show Parameters from the Tools menu). Try this now.

- When you click on an item in the Experiments navigator, look at the information displayed about it in the Description Pane (lower left). It is similar in content to the Parameters dialog.

Rename an Experiment
Default names are provided for all datasets and experiments based on either the name of the file being imported, or on the type of experiment being performed. Any item listed in the navigator can be renamed at any time. This gives you the opportunity to apply your own naming convention to the data.

1. Right-click the Normalization item that was just generated in the Experiments navigator, and select Rename Experiment from the shortcut menu. A box is drawn around the item with a blinking cursor at the end of it.

2. Press (and hold) the <Backspace> key to delete the program-generated name, and type in something significant to you (e.g. ‘Divided by max’ or ‘maxdiv’). Press <Enter> to accept this new name.

Note: GeneLinker™ saves all files automatically. Once an item is visible in the Experiments navigator, it has already been saved to the database.
Tutorial 1: Step 4 Perform Hierarchical Clustering

In this step of the tutorial, you will perform a hierarchical clustering experiment on the normalized data to reveal its intrinsic structure. For complete details on the clustering operations available in GeneLinker™, please see Clustering Overview.

**Perform Hierarchical Clustering:**
1. If the renamed normalization dataset in the **Experiments** navigator is not already highlighted, click it.
2. Click the **Hierarchical Clustering** toolbar icon, or select **Hierarchical Clustering** from the **Clustering** menu, or right-click the item and select **Hierarchical Clustering** from the shortcut menu. The **Hierarchical Clustering** parameters dialog is displayed.

![Hierarchical Clustering Parameters Dialog]

3. Set dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clustering Orientation</td>
<td>Cluster Genes</td>
</tr>
<tr>
<td>Distance Measurement: Between Data Points</td>
<td>Euclidean</td>
</tr>
<tr>
<td>Distance Measurement: Between Clusters</td>
<td>Average Linkage</td>
</tr>
</tbody>
</table>

4. Click **OK**. The clustering operation is performed and upon successful completion, a new Gene Hierarchical Clustering experiment is added to the **Experiments** navigator under the normalized dataset. You can rename it if you wish.

If you have automatic visualizations enabled in your user preferences, a matrix tree plot of the clustering results is displayed.

Tutorial 1: Step 5 Create a Matrix Tree Plot

GeneLinker™ has an excellent set of plots for examining your data. These are described in detail in the Plots section of the online manual.

If the matrix tree plot is already displayed, there is no need to recreate it. Read the Interpretation section below for information about the plot.
Create a Matrix Tree Plot

1. Double-click the hierarchical clustering experiment just created in the Experiments navigator. The item is highlighted and a matrix tree plot of the selected item is displayed.

OR

1. If the hierarchical clustering experiment just created in the Experiments navigator is not already highlighted, click it.

2. Click the Matrix Tree Plot toolbar icon, or select Matrix Tree Plot from the Clustering menu, or right-click the item and select Matrix Tree Plot from the shortcut menu. A matrix tree plot of the selected item is displayed.

A tree plot can take up a lot of space on your screen. You may want to maximize the GeneLinker™ window if it's not already maximized, and/or stretch the pane displaying the tree plot as wide as possible. **Note** that you can increase the width of the plots pane and reduce the width of the navigator pane by clicking-and-dragging the frame between them with the mouse.

Interpretation

In the hierarchy just created, note that at the extreme left of the plot is a cluster of several genes that are highly expressed early in the embryonic stage, at days E11, E13 and E15. This cluster includes the established early developmental markers nAChRd, G67I86, G67I80/86, nestin and nAChRe, as well as SC6, PDGFB, Ins1, keratin, SC7 and trk (see Wen et al. for explanation of gene name abbreviations).

Another cluster of genes with slightly broader expression profiles, but still mostly embryonic, appears at the extreme right of the plot (use the scrollbars to view the right of the plot). This cluster includes nAChRa6, PDGFR, MK2, NT3, GDNF, TH, cellubrevin, cyclin B, Brm, Ka1, and is enriched in members of the insulin-like growth factor signaling pathway.
pathway, IGFR1, IGF II, IGFR2, the latter being a receptor/ligand gene pair. Between them, these clusters map well to Wen's 'Wave 1'. Note that the combined clusters contain another receptor/ligand pair, PDGFb and PDGFR.

Just to the left of the right-most group is a cluster of nearly constantly expressed genes, easily picked out by eye as a nearly-solid mass of red. This cluster includes 'housekeeping' genes such as actin, TCP, SOD, CCO1 and CCO2, and maps well to Wen's 'Constant' class.

- Examine the tree plot for other groups with similarly simple characterizations, such as high expression in the adult mouse (Wen's Wave 4) or in the perinatal timepoints (Wen's Wave 3).

There are two reasons why the early-expressed genes don’t all appear side-by-side:

1. In the normalization and metric used above, the genes in the cluster including PDGFR, GDNF, and cellubrevin are mathematically closer to the constant genes than to the very early genes such as PDGFb, Ins1, and keratin. The mathematics don't always reflect qualitative ideas about similarity. However, if you try different normalizations and metrics you will obtain different clusterings. For example, if you try Scaling between 0 and 1 (instead of Divide by Maximum as you did above) you will find that the 'constant' cluster disappears, because this will magnify each gene's range of expression so that none will appear to be constant.

2. There is some arbitrariness in the construction of a tree diagram. At each branch point, GeneLinker™ must decide which branch to draw to the left and which branch to draw to the right. Consequently, the subcluster on the extreme right of our tree is no further mathematically from the subcluster on the extreme left than any other subcluster in the right half of the plot.

**Tutorial 1: Step 6 Perform Partitional Clustering**

From visual examination of a hierarchical clustering, Wen et al. identified five groups or 'waves' plus a small number of outliers or 'other' genes. This step will demonstrate that GeneLinker™ can be used to get a similar clustering, using the K-Means clustering function.

The key feature of K-Means clustering is that you choose a priori the number of clusters you think the data should be divided into. This number is the 'K' in K-Means.

The K-Means algorithm uses the same Euclidean Average-Linkage distance metric used for hierarchical clustering earlier.

Perform Partitional Clustering

1. If the renamed normalization item in the Experiments navigator is not already highlighted, click it.

2. Click the Partitional Clustering toolbar icon, or select Partitional Clustering from the Clustering menu, or right-click the item and select Partitional Clustering from the shortcut menu. The Partitional Clustering parameters dialog is displayed.
3. Set dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clustering Orientation</td>
<td>Cluster Genes</td>
</tr>
<tr>
<td>Distance Measurements: Between Data Points</td>
<td>Euclidean</td>
</tr>
<tr>
<td>Distance Measurements: Between Clusters</td>
<td>Average Linkage</td>
</tr>
<tr>
<td>Algorithm Properties: Type</td>
<td>K-Means</td>
</tr>
<tr>
<td>Algorithm Properties: Number of Means</td>
<td>5</td>
</tr>
<tr>
<td>Algorithm Properties: Random Seed</td>
<td>999</td>
</tr>
</tbody>
</table>

3. Click **OK**. The clustering operation is performed and upon successful completion, a new Gene Partitional Clustering experiment is added to the **Experiments** navigator under the original dataset. Rename it if you like.

If you have automatic visualizations enabled in your user preferences, a matrix tree plot of the clustering results is displayed. You can close this plot when you are finished looking at it.

**Use of the Random Seed Parameter**

In normal use, setting the random seed is neither necessary nor recommended. In a tutorial you set the random seed to a consistent value so that you will obtain precisely the same results that we depict and discuss, which makes the tutorial easier to understand. When you are not following a tutorial, you should generally not adjust the random seed at all.

The random seed setting may affect irrelevant details, such as the labelling and ordering of clusters. In other cases the random seed may affect relevant details, such as which genes occur together in clusters. Because of this latter possibility, it is sometimes worth repeating an experiment with different random seeds to see what the effects are. (In step 7 see 'The Centroid Plot: Variability in K-Means Clustering' below.) GeneLinker™ helps with this by setting a new random seed every time an operation is carried out, so you don’t need to.

On occasion you may need to determine whether a certain variation in results is due to the random element, or some other cause. For this reason you are able to set the random seed to a fixed value, thus controlling that source of variation.
Tutorial 1: Step 7 Create a Centroid Plot

Create a Centroid Plot
1. If the partitional clustering item in the Experiments navigator is not already highlighted, click it.
2. Select Centroid Plot from the Clustering menu, or right-click the item and select Centroid Plot from the shortcut menu. A centroid plot of the dataset is displayed.

The Centroid Plot is so named because each line represents the centroid or average element of a cluster. It is conceptually identical to the average waves plotted in Figure 3a of Wen et al. You should be able to see a clear visual resemblance between the clusters shown here, the clusters you just computed, and Wen’s clusters. Comparing just the figure above with Wen, note the following:

- A ‘constant’ cluster (4);
- a cluster (2) with an early maximum, similar to Wen’s Wave 1;
- a cluster (1) with a maximum at the ‘A’ (adult) timepoint, similar to Wen’s Wave 4, and
- two other clusters (3 and 5) with maxima at intermediate timepoints.

The Centroid Plot: Variability in K-Means Clustering
The colors and the cluster numbering in your Centroid Plot will probably be different from the above image, since clusters are arbitrarily labeled and colored.

More importantly, though, the line shapes will probably be slightly different. An important point about K-Means clustering is there is a random element in it. K-Means first randomly allocates items to clusters, and then systematically moves one item at a time from cluster to cluster in such a way as to minimize distances within clusters and maximize distances between clusters. However, there is no guarantee that all random starting allocations will lead to the same final clustering, only that the final clustering will
have reasonably low intracluster distances compared to the inter-cluster distances. This can be viewed as the cost of obtaining clusters quickly, but you can also look at it as a tool to show how meaningful your clusters are. If you rerun K-Means clustering a few times and get wildly different results, your data probably does not have any significant natural divisions, and you should probably not read anything into the clusters it produces. Conversely, if you rerun K-Means clustering twice and get similar results, the corresponding clusters are probably well-separated and meaningful.

For more information on clustering, refer to Clustering Overview.

**Tutorial 1: Step 8 Create a Cluster Plot**

There are several ways to examine cluster membership in detail. One is to create a Matrix Tree Plot as you did for the hierarchical clustering. In the case of partitional clustering, the ‘tree’ is flat, not hierarchical. Another way is to create a Cluster Plot from the clustering item in the **Experiments** navigator.

**Create a Cluster Plot:**

1. If the partitional clustering item in the **Experiments** navigator is not already highlighted, click it.

2. Select **Cluster Plot** from the **Clustering** menu, or right-click the item and select **Cluster Plot** from the shortcut menu. A cluster plot of the dataset is displayed.

![Cluster Plot](image)

A Cluster Plot of the entire dataset shows a line for each gene (because genes, not samples, were clustered). Each line is colored according to the cluster it belongs to. As you can see, the plot is fairly busy and not terribly informative even for a moderate amount of data like this. It is more informative to plot the individual clusters.
To Plot an Individual Cluster:
1. Click on the Centroid plot to make it the active window.
2. Click on a cluster name in the legend to highlight it and its line in the Centroid Plot. You can also click on the line itself, but with other lines nearby this may be difficult. For the purposes of this tutorial, select only one cluster (cluster 1 for the image below).
   - To select multiple clusters, press and hold the <Ctrl> key and click on cluster names in the legend.
   - To select a series of clusters, press and hold the <Shift> key and click on the first and last cluster names in the series.
3. Select Cluster Plot from the Clustering menu, or right-click on the plot and select Cluster Plot from the shortcut menu. A cluster plot of the selected cluster is displayed.

The new Cluster Plot shows the individual gene profiles for the genes in the selected cluster only, and also shows their names in the legend on the right. This illustration shows Cluster 1 from the Centroid Plot above. If you compare the genes present in the picture above with those in Wen’s Wave 4, you will see considerable but not perfect overlap.
   - See if there is a similar cluster in your clustering of the data. What genes does it have in common with the example shown here, and with Wen’s Wave 4?

Tutorial 1: Step 9 Generate Report and Export Image

Sometimes you may wish to have something printed on paper or saved in a file to show your colleagues or retain for your records.
Create an Experiment Report
1. If the partitional clustering experiment (the one produced in step 6 of this tutorial) in the Experiments navigator is not already highlighted, click it.
2. Select Generate Report from the File menu. The Save As dialog is displayed.
3. Provide information about where to store the file and under what name (or accept the provided defaults), and click Save. An experiment report is produced that describes the clustering parameters used and lists all the clustered items (genes) by their cluster membership, along with some summary statistics on the clusters. Reports are generated in HTML format. Once the report has been generated (and saved), GeneLinker™ starts up your default web browser (specified in your User Preferences) and displays the report.

Create a Workflow Report
1. Click the hierarchical clustering experiment (from step 4) in the Experiments navigator.

2. Select **Generate Workflow Report** from the **File** menu. The **Save As** dialog is displayed.

3. Again, provide information about where to store the file and under what name (or accept the provided defaults) and click **Save**. A workflow report is generated. It contains the same information as the experiment report, and also describes the entire 'descent' of the data, from the raw dataset down to the node being reported on. For example, a workflow report on this clustering experiment also summarizes the originating dataset and the normalization parameters used.

Workflow reports are generated in HTML format. Once the report has been generated (and saved), GeneLinker™ starts up your default web browser (specified in your User Preferences) and displays the report.
Export an Image

1. Click on a plot to make it the active window.
2. Select **Export Image** from the **File** menu or right-click on the plot and select **Export Image** from the shortcut menu. The **Save** dialog is displayed.

3. Enter a **File name**.
4. Select a file format (PNG, SVG, or PDF) from the **Files of type** drop-down list.
5. Click **Save**. The image is saved to a file of the specified type in the specified location. A message is displayed in the status bar when the image file save operation is complete.

**Tutorial 1: Conclusion**

When you are finished, you can close all the open plots either by clicking on the 'x' box in the upper-right hand corner of each, or by selecting **Close All** from the **Window** menu.

**Where To Go From Here**

- Go through the other tutorials.
- Read the Online Help to learn more about the various functions of GeneLinker™.
- Further explore GeneLinker™ by using additional features.
- Load up your favorite dataset and try out all the buttons and menu items.
- Don't forget to right-click on things like plots - many details of graphics can be customized.

**Tutorial 2: Clustering of NCI60 Dataset**

**Tutorial 2: Introduction**
This tutorial leads you through the process of preparing a dataset that has missing values, clustering it, and then visualizing the clustering results.

Skills You Will Learn:
How to import gene expression data from a tabular file into the GeneLinker™ database.
How to import a gene list.
How to import a variable (class labels).
How to estimate missing values.
How to rename a dataset in the Experiments navigator.
How to perform a hierarchical clustering experiment.
How to view experiment results in a matrix tree plot.
How to generate a report and export an image.

Dataset Information
The National Cancer Institute (NCI) maintains a set of 60 human cancer cell lines (NCI60). They are used in cDNA microarray studies to assess the gene expression profiles, as well as in screening anti-cancer drugs Reference 1.

The purpose of this tutorial is to demonstrate GeneLinker™ analysis and how it creates new perspectives on important biomedical relationships. A number of GeneLinker™ functions are used to go through the analysis in a step-by-step fashion. The approach is similar to that in Reference 1.

The data consists of expression measurements for 1416 differentially expressed genes (normalized log(Cy3/Cy5)) for 60 cancer cell lines. This is referred to in Reference 1 and in this tutorial as the t-matrix. Other NCI60 datasets, including the gene expression data for all 9,703 genes (all_genes), drug activities against the 60 cell lines (A-matrix and A118-matrix), and the gene-drug correlation data (AT-matrix), are not discussed here. Please see Reference 1 and Reference 2 for more detailed discussions of the original experiments and data.

Tutorial Length
This tutorial should take about 20 minutes, depending on how long you spend investigating the data, and how fast your machine is.

If you must stop part way through the tutorial, exit the program by selecting Exit from the File menu. The data and experiments you have performed to that point will be saved automatically by the application. The next time you start GeneLinker™, you can continue on with the next step in the tutorial.

Tutorial 2: Step 1 Start GeneLinker™ and Import the Data

Start GeneLinker™
1. Double-click the GeneLinker™ program icon 🖌️ on your desktop to start the
Import the Data
1. Click the Import Gene Expression Data toolbar icon, or select Import from the File menu and Gene Expression Data from the sub menu. The Data Import dialog is displayed.

2. GeneLinker™ uses a template to interpret or parse the data file being imported. Ensure that the template listed on the dialog is Tabular.

3. The next step is to identify the name and location of the data source file. Click the button to the right of the Source File box. The Open dialog is displayed.

4. The tutorial data files are located in the Tutorial folder. This is the folder listed in Look in, so you do not need to navigate to it. Click the file t_matrix.csv, and click Open. The Data Import dialog is updated with the file name.

5. Ensure the Gene Database is set to GenBank. The IMAGE Consortium clone IDs in the original data file have been mapped to GenBank accession numbers in the tutorial data file by taking the 5’ accession number if there is one, and taking the 3’ accession number otherwise. For an example of how to use IMAGE clone ids as gene identifiers, see Tutorial 6.

6. Click Import. The Import Data dialog is displayed.
GeneLinker™ examines the file and offers to transpose it. Within GeneLinker™, datasets have the genes in columns and the samples in rows.

When importing data using a Tabular template, GeneLinker™ assumes that the more numerous dimension of the data represents genes (most microarray experiments involve more genes than samples). If this is so (as in this tutorial), then clicking OK is all that is required.

**Note:** the options Use Sample Names and Use Gene Names are checked and disabled. GeneLinker™ has recognized that in this dataset, the first row and column contain alphabetic labels. Gene expression data is always numeric, hence the disabled checkboxes.

7. Click OK. The data is imported into the database and a dataset item named t_matrix is added to the Experiments navigator. This represents your raw data, which is now available to perform experiments on using the various GeneLinker™ functions.

### Tutorial 2: Step 2 Estimate Missing Data Values

The NCI60 studies rejected some data due to low signal or for quality control reasons. GeneLinker™ has functionality for eliminating genes that meet a specified threshold number of missing values, and for estimating missing values.

**Estimate Missing Data Values**

1. If the t_matrix dataset in the Experiments navigator is not already highlighted, click it.
2. Click the **Estimate Missing Values** toolbar icon 📐, or select **Estimate Missing Values** from the **Data** menu, or right-click the item and select **Estimate Missing Values** from the shortcut menu. The **Estimate Missing Values** dialog is displayed.

![Estimate Missing Values dialog]

3. Set dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remove Genes That Have Missing Values</td>
<td>30</td>
</tr>
<tr>
<td>Replacement Technique</td>
<td>Nearest Neighbors</td>
</tr>
<tr>
<td>Distance Metric</td>
<td>Euclidean</td>
</tr>
<tr>
<td>Number of Nearest Neighbors</td>
<td>3</td>
</tr>
</tbody>
</table>

4. Click **OK**. The **Experiment Progress** dialog is displayed.

![Experiment Progress dialog]

The dialog is dynamically updated as the Estimate Missing Values operation is performed. Upon successful completion, a new Estimated: #mv < 30 | median complete dataset is added to the **Experiments** navigator under the original dataset. This new dataset has the **complete dataset** icon 📐 before its name. (An incomplete dataset has the **incomplete dataset** icon 📐.)

**Note:** in addition to estimating missing values, GeneLinker™ provides facilities for normalizing and filtering data. Use of these functions is described in detail in the preprocessing section of the help. The dataset we are using was suitably normalized by the original authors.
Tutorial 2: Step 3 Rename the Dataset

Default names are provided for all datasets and experiments based on either the name of the file being imported, or on the type of experiment being performed. Any item listed in the navigator can be renamed at any time. This gives you the opportunity to apply your own naming convention to the data.

Rename a Dataset
1. Right-click the Estimated: #mv < 30 | median dataset in the Experiments navigator, and select Rename Experiment from the shortcut menu. A box is drawn around the dataset name, with a blinking cursor at the end of the name.
2. Press the <Backspace> key to delete the program-generated name.
3. Type in something significant to you (e.g. 3-nearest-neighbors estimation).
4. Press <Enter> to accept this new name. The experiment is renamed with the new name.

Please note that GeneLinker™ saves all files automatically. Once an item is visible in the Experiments navigator, it has already been saved to the GeneLinker™ database. The renaming facility is for convenience. For instance, the name recommended in this example allows you to see at a glance, the type of missing value estimation which produced that dataset. This would be particularly valuable were you, for instance, comparing different methods for missing value estimation. The parameters used to generate every dataset are captured automatically by GeneLinker™ and can always be viewed by selecting the item and examining the Description Pane in the lower left of the application window.

5. Do that now: The dataset is still highlighted. Look at the information provided in the Description Pane. Among other things, notice that there are 1374 genes in this dataset.
6. Click the parent dataset, ‘t_matrix’, and examine the information about it in the **Description Pane**. Notice that there are **1375** genes in the parent dataset.

![t_matrix]

The Estimate Missing Values operation filtered out one gene because it had more missing values than we wanted. In the next step we will demonstrate one way of identifying that filtered gene.

**Tutorial 2: Step 4 Display Color Matrix Plots**

In this step, we use the Shared Selection feature to see which gene was filtered out during missing value estimation.

1. Double-click the 3-nearest-neighbors estimation dataset in the **Experiments** navigator. The item is highlighted and a color matrix plot of the dataset is displayed.

![Color Matrix Plot: 3-nearest-neighbors estimation]

2. Click the first gene name on the plot. The gene name is highlighted.
3. Use the scrollbar on the bottom of the plot to scroll to the far right.
4. Press and hold down the <Shift> key and click the last gene name on the plot. All of the gene names are highlighted.
5. Double-click the t_matrix dataset in the **Experiments** navigator. The item is highlighted and a color matrix plot of it is displayed.

Notice that the genes that you selected on the first plot are highlighted in the new plot. This facility called shared selection helps you locate selected genes on any table or plot in which they appear.

6. Scroll slowly to the right. You will see one gene that is not highlighted. This is the gene that was filtered out when you estimated missing values.
7. Click the non-highlighted gene (gene H12289). The gene is highlighted and the rest are un-highlighted. Look at the information about the gene in the **Description Pane**.

In the next step, we will import a gene list that contains additional information about the genes in the dataset.

**Tutorial 2: Step 5 Import a Gene List**

File `t_matrix_genelist.csv` contains descriptions for each gene in the dataset. The way to bring these descriptions into GeneLinker™ is to perform a gene list import. Since the genes already exist in the GeneLinker™ database (genes are imported when you import a dataset), they are not imported again when you import a gene list. Instead, the existing genes are updated with the additional information in the gene list file. The gene list itself is imported into the **Gene Lists** navigator. For complete details on this process, please see Importing a Gene List.

You may wish to examine the file ‘`t_matrix_genelist.csv`’ in a spreadsheet, or by using an editing tool. The file contains, in the first column, gene identifiers matching those appearing in the expression data file. (Order is not important.) The second column may contain a gene symbol or short gene name (if one is known) and the third column contains a longer description of the gene.
Import a Gene List

1. Review the information about the filtered gene in the Description Pane.

2. Select Import from the File menu and Gene List from the sub menu. The Open dialog is displayed.

3. Since the gene list file does not have the extension .txt, you will need to change the Files of type selection. Use the drop-down list to select All files (*.*) This displays all of the files in the Tutorial folder (including the gene list file t_matrix_genelist.csv).

4. Click the file t_matrix_genelist.csv. The file name is highlighted.

5. Click Open. The Import Gene List dialog is displayed.

6. Ensure GenBank is set in the Gene Database drop-down list.

7. Click OK. The gene list and gene descriptions are imported into the GeneLinker™ database. A new gene list item is added to the Gene Lists navigator.
There is no requirement that the gene list match any particular expression dataset. A gene list is simply that: a list of genes which can include descriptions. Gene lists provide a means to import symbols and descriptions into GeneLinker™ to be associated with gene identifiers.

Whenever a single gene is selected in a GeneLinker™ view, the Description Pane in the lower left corner of the GeneLinker™ window displays what information has been imported about that gene: The database identifier, the database type (e.g. GenBank, Unigene, Affymetrix, Custom), and the symbol and the gene description if any have been imported.

8. Click the filtered gene (H12289) on the t-matrix color matrix plot. The gene is highlighted.
9. Look at the Description Pane just below the navigator. Note the additional information about the gene that was added by importing the gene list.

![GeneLinker screenshot]

**Tutorial 2: Step 6 Perform Hierarchical Clustering**

**Perform Hierarchical Clustering**

1. Click the 3-nearest neighbors dataset in the Experiments navigator (Click the Experiments tab to display the Experiments navigator). The item is highlighted.
2. Click the Hierarchical Clustering toolbar icon, or select Hierarchical Clustering from the Clustering menu, or right-click the item and select Hierarchical Clustering from the shortcut menu. The Hierarchical Clustering dialog is displayed.

3. Set parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>GeneLinker Gold 3.1 / GeneLinker Platinum 2.1</td>
<td>65</td>
</tr>
</tbody>
</table>
Clustering Orientation | Cluster Samples
---|---
Data Measurements: Between Data Points | Pearson Correlation
Data Measurements: Between Clusters | Average Linkage

- Note that Agglomerative (the default option) is set as the Type parameter in the Algorithm Properties group.

4. Click OK. The clustering operation is performed, and upon successful completion, a new Sample Hierarchical Clustering experiment is added to the Experiments navigator under the original dataset.

GeneLinker™ provides many different clustering algorithms, and there are other clustering methods listed under Partitional Clustering. Genes can be clustered in addition to samples by using the same command sequence as above but changing the choice of clustering orientation from Samples to Genes.

If you have automatic visualizations enabled in your user preferences, a matrix tree plot of the clustering results is displayed.

**Tutorial 2: Step 7 Create a Matrix Tree Plot**

GeneLinker™ has an excellent set of plots for examining your data. These are described in detail in the Plots section of the online manual.

If the matrix tree plot is already displayed, there is no need to recreate it. Read the sections below the image for information about the plot.

**Create a Matrix Tree Plot**

1. Double-click the Sample Hierarchical Clustering experiment in the Experiments navigator. It is tagged with the Hierarchical Clustering icon. The item is highlighted and a matrix tree plot of the selected item is displayed. The gene names appear as the column headings and the sixty cancer cell lines are labels for the rows.

Or

1. If the Sample Hierarchical Clustering experiment in the Experiments navigator is not already highlighted, click it.

2. Click the Matrix Tree Plot toolbar icon, or select Matrix Tree Plot from the Clustering menu, or right-click the item and select Matrix Tree Plot from the shortcut menu. A matrix tree plot of the selected item is displayed.
Matrix tree plots can be manipulated, resized or customized. Please give the following a try:

**To Scroll a Matrix Tree Plot**
Use the scrollbars to move the plot. Clicking an arrow moves the plot one color tile width at a time. To move more rapidly, click and drag the scroll thumb.

**To Identify a Gene or Sample and See the Expression Value**
Hover the mouse cursor over the colored tile for which you want to know the value. A tooltip appears displaying the gene name, sample name, and gene expression value. The tooltip disappears as you move the pointer off that tile.

**To Change the Color or Scale of the Gradient**
1. Double-click the plot legend. The **Customize** dialog is displayed.

   ![Customize dialog](image)

2. Set the parameters to customize the plot.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Data Range:</strong></td>
<td>Type a new value into the <strong>Minimum</strong> and/or the <strong>Maximum</strong> field(s) and press &lt;Enter&gt; or use the scroll arrows to set the value(s). The plot is re-drawn using the new values.</td>
</tr>
</tbody>
</table>
**Use actual range**

Click the **Use actual range** button to set the minimum and maximum for the display from the actual minimum and maximum values in the dataset. The plot is re-drawn using the Actual Range values.

**Palette**

Click a new color scheme in the **Palette** drop-down list. The plot is dynamically re-drawn using the new colors.

3. Click **OK** to keep the new settings, or click **Cancel** to revert to the previous ones.

**To Resize the Plot**

1. Click **Resize** at the top of the plot. The **Resize** dialog is displayed.

2. Use the sliders to set the width and/or height of the color tiles. The column and/or row labels are not displayed if you set the width or height too small.

3. Click the icon in the upper corner of the **Resize** dialog to close it.

**To See Only the Dendrogram (with Sample Labels)**

- Right-click on the plot, and select **Hide Color Matrix** from the shortcut menu. The color matrix is removed from view leaving the dendrogram side-by-side with the cell line labels.
- Right-click on the plot again and select **Show Color Matrix** to bring the color matrix back.

When you are finished examining the plots, you can close them.

**Tutorial 2: Step 8 Import Cancer Class Variable**

For complete details on variables, please see Variables Overview.

1. Click the t_matrix dataset in the **Experiments** navigator. The item is highlighted.
2. Select **Import** from the **File** menu and **Variable** from the sub menu. The **Import Variable** dialog is displayed.
   - The **Dataset** name is displayed at the top of the dialog and the number of samples in the dataset is listed under the name.
3. To set the source file for the variable data, click the ... button to the right of the Source File box. The Open dialog is displayed.

4. The tutorial data files are located in the Tutorial folder. This is the folder listed in the Look in box, so you do not need to navigate to it. Click the file t_matrix_classes.csv.

5. Click Open.

- The Source File name is displayed with the number of observations and classes in the file listed underneath.
- The default Variable Name and Description are displayed.
- The Create Variable Type dialog is displayed because there are no existing variable types.
6. Enter **NCI60 Cancer Classes** into the **Name** box on the **Create Variable Type** dialog.

7. Click **OK**. The variable type is created and is listed in the **Choose a Variable Type** box on the **Import Variable** dialog.

8. The **Preview** allows you to view which sample belongs to which class and the total number of entries for each class. Click **Preview**. When you are finished examining the contents of the Preview, click **Close** to close it.

9. Enter **Cancer Classes** for the **Variable Name**.
10. Click Import. The variable data is imported into the database, and in the Experiments navigator, the t_matrix dataset icon is marked with the variable tag.

**Tutorial 2: Step 9 Color Samples by Class**

**To Color the Samples by Class**

We will need to refresh the Matrix Tree Plot in order to view the new class variable on it.

1. Close all the open plots by selecting Close All from the Window menu.
2. Double-click the Sample Hierarchical Clustering experiment in the Experiments navigator. The item is highlighted and a new matrix tree plot is displayed.
3. Click the **Color by Variable** button at the top of the plot. A block of color appears to the left of each row indicating which cancer class that sample belongs to. This makes it easy to compare a sample clustering to known classes.

4. To see the key of colors, click the **Color Manager** button on the plot, or select **Color Manager** from the **Tools** menu. The **Color Manager** dialog is displayed.
5. On the **Color Manager** dialog, click the **Variables** tab. The **Variables** pane is displayed.

6. Ensure **NCI60 Cancer Classes** is selected in the **Variable Type Classes** drop-down list.

7. You can change the color mapped to any class using the **Color Manager**. Click the color box to the left of the ME class. The **Pick a Color** dialog is displayed.

8. Click a dark blue color swatch. You can choose colors from swatches, or by their HSV (hue, saturation, and value) or RGB (red, green, blue) descriptions. The color is displayed in the **Recent** list.

9. Click **OK**. The dialog closes and the new color is applied to the ME class on the matrix tree plot.

10. Click the icon in the upper right corner of the **Color Manager** to close it.

**Tutorial 2: Step 10 Generate Report and Export Image**

**Generate an Experiment Report**

1. Click the Sample Hierarchical Clustering experiment in the **Experiments** navigator. The item is highlighted.

2. Select **Generate Report** from the **File** menu. The **Save As** dialog is displayed.
3. Type in a file name or use the provided default.
4. Click **Save**. A report of the clustering results is created in HTML format. It is saved to your disk and your browser is started displaying the report.

**Note:** the length of the report is proportional to the size of the dataset.

**Export an Image**
1. Right-click in the matrix tree plot and select **Hide Color Matrix** in the shortcut menu.
2. With the color matrix turned off, right-click on the plot and select **Export Image** from the shortcut menu. The **Save** dialog is displayed.
3. Navigate to the folder where you want the image file saved.
4. Type in a **File name** for the image file.
5. Select an image file format from the **Save as type** drop-down list. The options are .png, .svg and .pdf. See Exporting an Image for full details.
6. Click **Save**. GeneLinker™ exports an image file of the specified type to the specified location.

Other methods for visualizing clustered data are available, such as a **Centroid Plot** or **Cluster Plot**. Creating these is described in detail in Tutorial 1.

**Tutorial 2: Conclusion**

**Discussion of the Results**

The matrix tree plot from clustering the cancer cell lines is included here as the following:

Figure 1. Clustering of the cancer cell lines according to gene expression profiles

Colon, renal, and CNS cancers, leukemias and melanomas all form fairly homogeneous clusters with these genes in this metric. Ovarian cancers show somewhat more disparity. The two prostate cancer samples show no strong association with any other group nor with each other, and the lung cancers seem to have almost no cohesion at all in this space. The breast cancers are scattered as well, two of them clustering with the melanomas, two with the CNS cancers, two beside the colon cancers, and one more in a heterogeneous cluster which also includes a prostate, two ovarian, two lung, one renal and one CNS cancer, and one melanoma cell line.

Note that 'BR:MDA-N' and 'BR:MDA-MB-435' form a sub-cluster inside the melanoma cluster. This is also indicated in Reference 1. GeneLinker™ confirms that several cancer cell lines (such as 'ME:LOX IMVI', 'RE:SN12C' and 'OV:OVCAR-8') do not cluster according to their origins, as was also found by Reference 1.

Note the similarity between the clustering of the t_matrix and the results presented in Figure 1 and Fig. 2a in Reference 1. Slight variations in the clustering parameters account for the differences.

When you are finished, you can close all the open plots either by clicking on the 'x' box.
in the upper-right hand corner of each, or by selecting **Close All** from the **Window** menu.

**Summary**

This tutorial demonstrated how to obtain and preprocess the dataset from the NCI60 studies, how to import the data, how to estimate missing values and how to do clustering calculations. A Matrix Tree Plot of the clustering of gene expression was created.

There are other commands in GeneLinker™ for handling data, analyzing data and visualizing analysis results. These are illustrated in other tutorials included in the release.

**References**

**Reference 1**


A copy of the paper can be obtained at: http://discover.nci.nih.gov/nature2000/

**Reference 2**


**Where To Go From Here**

- Go through the other tutorials provided.
- Read the online Help to learn more about the various functions of GeneLinker™.
- Further explore GeneLinker™ by using additional features.
- Load up your favorite dataset and try out all the buttons and menu items.
- Don't forget to right-click on things like plots - many details of graphics can be customized.
- Visit the Molecular Mining website at http://www.molecularmining.com/ for the latest information on GeneLinker™ enhancements and additional products.

**Tutorial 2: Figure 1 - Clustering of the cancer cell lines according to gene expression profiles**
Tutorial 3: Jarvis-Patrick Clustering

Tutorial 3: Introduction

This tutorial introduces you to data normalization and Jarvis-Patrick partitional clustering. The results of the clustering experiments are viewed in a matrix tree plot.

Skills You Will Learn:
How to import gene expression data from a file into the GeneLinker™ database.
How to normalize data.
How to estimate missing values.
How to perform a partitional clustering experiment.
How to view experiment results in a matrix tree plot.

Jarvis-Patrick Partitional Clustering

Also known as mutual nearest neighbors clustering, Jarvis-Patrick clustering is a very fast non-stochastic clustering method. It has seen considerable use in the cheminformatics community, but has not been widely used in gene expression analysis until now.
Jarvis-Patrick clustering depends on two user-configurable parameters: the number of nearest Neighbors to Examine, and the number of those neighbors that must be shared in order for the two items (genes, for instance) to be clustered together. The two items must also be among each other’s nearest neighbors. The appropriate values to use for these parameters depend on the data being clustered and the objective of the analysis. Starting with one or two common neighbors out of five or six nearest neighbors tends to produce a manageable number of clusters on datasets of 100-200 items. The larger the list of Neighbors to Examine, the more likely it is that common neighbors will be found to join any two items, and so increasing this number tends to lead to fewer and larger clusters. Conversely, the more common neighbors are required, the fewer joins are found, and this tends to lead to more and smaller clusters.

A typical Jarvis-Patrick clustering contains a wide variety of cluster sizes. There are usually a significant number of singleton genes in any Jarvis-Patrick clustering, along with a small number of very large clusters, and a smattering of fairly tight clusters containing between 1 and 10 genes. As well, the clusters are not constrained to be as globular as in, for example, average-linkage K-Means clustering. When combined with the number of singletons, this means that a centroid plot will often not illustrate the clusters’ characteristics very clearly. Instead, using a Matrix Tree Plot is recommended for a comparative overview of the clusters.

Assumptions

This tutorial assumes you have already completed Tutorial 1 and Tutorial 2 thus having the Spinal_cord and t_matrix datasets in the Experiments navigator. If the Spinal_cord and/or t_matrix datasets are missing, follow the Data Import procedure in Tutorials 1 and 2 to import them.

Tutorial Length

This tutorial is split into two parts: part A deals with the Spinal_cord dataset and part B deals with the t_matrix dataset. The entire tutorial should take about 20 minutes, depending on how long you spend investigating the data, and how fast your machine is.

If you must stop part way through the tutorial, simply exit the program by selecting Exit from the File menu. The data and experiments you have performed to that point are saved automatically by GeneLinker™. The next time you start GeneLinker™, you can continue on with the next step in the tutorial.

Tutorial 3A: Step 1 Normalize the Data

Normalize the Data

1. If the Spinal_cord dataset in the Experiments navigator is not already highlighted, click it.
2. Click the Normalize toolbar icon , or select Normalize from the Data menu, or right-click the item and select Normalize from the shortcut menu. The first Normalization parameters dialog is displayed.
3. Double-click the **Other Transformations** radio button, or click it and click **Next**. The second **Normalization** dialog is displayed.

4. Double-click the **Scaling between 0 and 1** radio button, or click it and click **Finish**. The **Experiment Progress** dialog is displayed.

The dialog is dynamically updated as the normalization operation is performed. Upon successful completion, a new Norm: Scaled min to max dataset is added to the **Experiments** navigator under the original dataset.
Tutorial 3A: Step 2 Perform Partitional Clustering

Perform Partitional Clustering
1. If the new Norm: Scaled min to max dataset in the Experiments navigator is not already highlighted, click it.

2. Click the Partitional Clustering toolbar icon, or select Partitional Clustering from the Clustering menu, or right-click and select Partitional Clustering from the shortcut menu. The Partitional Clustering parameters dialog is displayed.

3. Set dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clustering Orientation</td>
<td>Cluster Genes</td>
</tr>
<tr>
<td>Distance Measurements: Between Data Points</td>
<td>Euclidean</td>
</tr>
<tr>
<td>Algorithm Properties: Type</td>
<td>Jarvis-Patrick</td>
</tr>
<tr>
<td>Algorithm Properties: Neighbors to Examine</td>
<td>6</td>
</tr>
<tr>
<td>Algorithm Properties: Neighbors in Common</td>
<td>2</td>
</tr>
</tbody>
</table>

4. Click OK. The clustering operation is performed and upon successful completion, a new J-P (6,2): genes | Euclid | average experiment is added to the Experiments navigator under the original dataset.

If you have automatic visualizations enabled in your user preferences, a matrix tree plot of the clustering results is displayed.

Tutorial 3A: Step 3 Create a Matrix Tree Plot

If the matrix tree plot is already displayed, there is no need to recreate it. Read the sections below the image for information about the plot.
Create a Matrix Tree Plot
1. Double-click the J-P (6,2): genes | Euclid | average experiment in the Experiments navigator. The item is highlighted and a matrix tree plot is displayed.

OR
1. If the J-P (6,2): genes | Euclid | average experiment in the Experiments navigator is not already highlighted, click it.
2. Click the Matrix Tree Plot toolbar icon, or select Matrix Tree Plot from the Clustering menu, or right-click and select Matrix Tree Plot from the shortcut menu. A matrix tree plot of the experiment is displayed.

Scroll the Matrix Tree Plot
1. Use the bottom scrollbar to scroll to the far right of the plot. The 'comb' under the grid of color tiles illustrate cluster membership.
   • At the far right of the Matrix Tree Plot are seven singleton genes including SC2, EGFR and trkB, which were also nominated as outliers by Wen et al. using FITCH clustering and a divide-by-max normalization.
   • Just to the left of that you can see four very tight clusters: three characterized by late expression maxima, and one (SC6 and nAChRd) by an early expression maximum. These are shown in the figure just above.
   • Three groups to the left of the singletons is a cluster of six genes including three mGlu receptors, all highly expressed in the late embryo and perinatal timepoints.
   • Two groups to the left of that is a large cluster (41 genes) including a large number of neurotransmitter receptors: three of four serotonin (5HT) receptors; three acetylcholine receptors (plus acetylcholinesterase); NMDA1/2B/2C; mGluR3/4/5/7; GABA receptors GRa1/2/3/4/5 and GRg1/2/3. This cluster’s expression profiles are characterized by minimal expression in the E11 and E13 timepoints, followed by fairly uniform expression thereafter.
2. Use the bottom scrollbar to scroll back to the left of the plot.
At the far left is a second large cluster (47 genes) covering a wide variety of genes.

**Tutorial 3B: Step 1 Estimate Missing Values**

By clustering the NCI60 t_matrix dataset, you can get an idea of the speed of Jarvis-Patrick clustering. First, missing values in the dataset must be estimated. If you have completed Tutorial 2 (you have a 3-nearest-neighbors dataset under the t_matrix dataset in the Experiments navigator already) skip to Step 2 Perform Partitional Clustering.

**Estimate Missing Values**

1. If the t_matrix dataset in the Experiments navigator is not already highlighted, click it.
2. Click the Estimate Missing Values toolbar icon, or select Estimate Missing Values from the Data menu, or right-click the item and select Estimate Missing Values from the shortcut menu. The Estimate Missing Values dialog is displayed.
3. Set dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remove Genes That Have Missing Values</td>
<td>30</td>
</tr>
<tr>
<td>Replacement Technique</td>
<td>Nearest Neighbors Estimation</td>
</tr>
<tr>
<td>Distance Metric</td>
<td>Euclidean</td>
</tr>
<tr>
<td>Choice of Median or Mean</td>
<td>3</td>
</tr>
</tbody>
</table>

4. Click OK. The Estimate Missing Value operation is performed and upon successful completion, a new complete Estimated: #mv <30 | medians dataset is added to the Experiments navigator under the original dataset.

**Tutorial 3B: Step 2 Perform Partitional Clustering**

**Perform Partitional Clustering**

1. If the 3-nearest-neighbors (or Estimated: #mv <30 | median) dataset in the Experiments navigator is not already highlighted, click it.

2. Click the Partitional Clustering toolbar icon 🗻, or select Partitional Clustering from the Clustering menu, or right-click the item and select Partitional Clustering from the shortcut menu. The Partitional Clustering parameters dialog is displayed.
3. Set dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clustering Orientation</td>
<td>Cluster Genes</td>
</tr>
<tr>
<td>Distance Measurements: Between Data Points</td>
<td>Euclidean</td>
</tr>
<tr>
<td>Algorithm Properties: Type</td>
<td>Jarvis-Patrick</td>
</tr>
<tr>
<td>Algorithm Properties: Neighbors to Examine</td>
<td>6</td>
</tr>
<tr>
<td>Algorithm Properties: Neighbors in Common</td>
<td>2</td>
</tr>
</tbody>
</table>

4. Click OK. The partitional clustering operation is performed and upon successful completion, a new J-P (6,2) genes | Euclid | average experiment is added to the Experiments navigator under the original dataset.

If you have automatic visualizations enabled in your user preferences, a matrix tree plot of the clustering results is displayed.

**Tutorial 3B: Step 3 Create a Matrix Tree Plot**

If the matrix tree plot is already displayed, there is no need to re-create it. Read the sections below the image for information about the plot.

**Create a Matrix Tree Plot**

1. Double-click the J-P (6,2) genes | Euclid | average experiment in the Experiments navigator. The item is highlighted and a matrix tree plot is displayed.

OR

1. If the J-P (6,2) genes | Euclid | average experiment in the Experiments navigator is not already highlighted, click on it.

2. Click the Matrix Tree Plot toolbar icon or select Matrix Tree Plot from the Clustering menu, or right-click the item and select Matrix Tree Plot from the shortcut menu. A matrix tree plot of the dataset is displayed.
Resize the plot

You can use the **Resize** function to reduce the size of the plot. You will still be able to identify the genes and samples associated with particular color tile by hovering the mouse pointer over the color tile and reading the tooltip which appears.

1. Click the **Resize** button at the top of the plot. The **Resize** dialog is displayed.

2. Reduce the **height** until you can see all of the samples and the clustering comb at the same time. Click the 'x' in the upper right corner of the **Resize** dialog to dismiss it. The reduced height causes the sample labels to not be displayed on the plot.

3. Click the **Find** toolbar icon. The **Find** dialog is displayed.
4. Type the gene name **W47225** into the **Find what** box.
5. Click **Find**. The plot scrolls to the right so that the gene **W47225** (an EST highly similar to interleukin-1 beta) is visible and highlighted.

Notice the strong resemblance between W47225 (IL1B) and its immediate neighbor W46667 (another EST), both highly overexpressed in melanoma LOXIMVI (sample 1). Also in that cluster are a number of ESTs and SIDs.

When you are finished, you can close all the open plots either by clicking on the 'x' box in the upper-right hand corner of each, or by selecting **Close All** from the **Window** menu.

---

### Tutorial 3: Conclusion

### References


### Where To Go From Here

- Go through the other tutorials provided.
- Read the Online Help to learn more about the various functions of GeneLinker™.
- Further explore GeneLinker™ by using additional features.
- Load up your favorite dataset and try out all the buttons and menu items.
- Don't forget to right-click on things like plots - many details of graphics can be customized.
- Visit the Molecular Mining website at http://www.molecularmining.com/ for the latest information on GeneLinker™ enhancements and additional products.
Tutorial 4: Self Organizing Maps (SOMs)

Tutorial 4: Introduction

This tutorial introduces you to Self-Organizing Maps (SOMs). The results of the SOM clustering is viewed in a SOM plot. This tutorial uses Leukemia data to demonstrate how SOMs can be used. The Self-Organizing Map (SOM) is a clustering method with its roots in Artificial Neural Networks [Kohonen2001]. SOMs have been used in the literature to explore several different gene expression datasets [for example, Golub1999; Tamayo1999; Toronen1999; and Hill2000].

Skills You Will Learn:
How to import gene expression data from a file into the GeneLinker database.
How to display summary statistics about a dataset.
How to remove values and genes with missing values.
How to normalize data.
How to perform a SOM clustering experiment.
How to view SOM experiment results in a SOM plot.

How SOMs Work
SOMs work somewhat like K-Means clustering but are a little richer. With K-Means, you choose the number of clusters to fit the data into. For a SOM you choose the shape and size of a network of clusters to fit the data into. In a SOM, we call these clusters 'nodes'. In GeneLinker™, the nodes are arranged in a rectangular grid for which you need to choose the height and the width. Much like for K-Means clustering, you should choose an initial size based on what you suspect about the number of classes in your data.

Like K-Means, a SOM initially populates its nodes or clusters by randomly sampling the data (or randomly generating points in the data space, depending on the initialization option you choose), and then refines the nodes in a systematic fashion. Unlike K-Means clustering, however, a SOM will not force there to be exactly as many clusters as there are nodes, because it is possible for a node to end up without any associated cluster items when the map is complete. A further difference with K-Means clustering is that the SOM automatically provides some information on the similarity between nodes - i.e., how strongly the certain nodes resemble each other.

Overview of the Tutorial Data
Golub et al. (1999) reported on a dataset of gene expression patterns from leukemia patients. The problem was to distinguish acute myeloid leukemia (AML) from acute lymphoblastic leukemia (ALL). They additionally considered the question of whether the cell type (B-cell or T-cell) could be distinguished.

Gene expression levels for 72 patients were measured using Affymetrix™ equipment. This data is available from the website of the Whitehead Institute at MIT. A formatted
version of the data is provided with GeneLinker™.

Tutorial Length
This tutorial should take about 30 minutes, depending on how long you spend investigating the data, and how fast your machine is.

If you must stop part way through the tutorial, simply exit the program by selecting Exit from the File menu. The data and experiments you have performed to that point are saved automatically by GeneLinker™. The next time you start GeneLinker™, you can continue on with the next step in the tutorial.

Tutorial 4: Step 1 Import the Data

Import the Data
1. Click the Import Gene Expression Data toolbar icon or select Import from the File menu and Gene Expression Data from the sub menu. The Data Import dialog is displayed.

2. Set the Gene Database to Affymetrix using the drop-down list.
3. The next step is to identify the name and location of the data source file. Click the button to the right of the Source File box. The Open dialog is displayed.
4. The tutorial data files are located in the Tutorial folder. This is the folder listed in Look in, so you do not need to navigate to it. Click the file aml_all.csv.

5. Click Open. The Data Import dialog is updated with the file name.
6. Click **Import**. The **Import Data** dialog is displayed.

GeneLinker™ examines the file and offers to transpose it. Within GeneLinker™, datasets have the genes in columns and the samples in rows. When importing data using a Tabular template, GeneLinker™ assumes that the more numerous dimension of your data represents genes (most microarray experiments involve more genes than samples). If this is so (as in this tutorial), then clicking **OK** is all that is required.

**Note:** the options **Use Sample Names** and **Use Gene Names** are checked and disabled in the **Import Data** dialog box. GeneLinker™ has recognized that in this dataset, the first row and column contain alphabetic labels. Gene expression data is always numeric, hence the disabled checkboxes.

7. Click **OK**. The dataset is imported into GeneLinker™ and a new item, aml_all, is added to the **Experiments** navigator.

**Tutorial 4: Step 2 View the Data**
View the Data with the Table Viewer

1. If the aml_all dataset in the Experiments navigator is not already highlighted, click it.
2. Click the Table View toolbar icon, or right-click the item and select Table View from the shortcut menu. This dataset is large (7129 genes), so displaying the data in the table viewer may take a few seconds.

Note: each sample is numbered according to the supplementary material provided by the Whitehead Institute, and is further labeled by its cancer class (AML or ALL). AML samples are further labeled by cell type (B-cell or T-cell).

Tutorial 4: Step 3 Display Summary Statistics

Display Summary Statistics

1. If the aml_all dataset in the Experiments navigator is not already highlighted, click it.
2. Click the Summary Statistics toolbar icon, or select Summary Statistics from the Statistics menu. The Summary Statistics chart is displayed.

- Notice the large number of negative values in what is considered to be count data.

Tutorial 4: Step 4 Remove Negative Values
Remove Negative Values

1. If the aml_all dataset in the Experiments navigator is not already highlighted, click it.
2. Select Remove Values from the Data menu, or right-click on the item and select Remove Values from the shortcut menu. The Remove Values dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Removal Technique</td>
<td>by Expression Value</td>
</tr>
<tr>
<td>Expression Value</td>
<td>Set the comparison type to &lt;=</td>
</tr>
<tr>
<td></td>
<td>Set the threshold value to 0</td>
</tr>
</tbody>
</table>

4. Click OK. The Experiment Progress dialog is displayed.

The dialog is dynamically updated as the Remove Values operation is performed. Upon successful completion, a new incomplete dataset (containing strictly positive values) is added to the Experiments navigator under the original dataset.

Tutorial 4: Step 5 Remove Genes that have Missing Values

Remove Genes that have Missing Values

1. If the Removed: v <= 0.0 dataset in the Experiments navigator is not already highlighted, click it.
2. Click the Estimate Missing Values toolbar icon , or select Estimate Missing Values from the Data menu, or right-click the item and select Estimate Missing Values from the shortcut menu. The Estimate Missing Values dialog is displayed.
3. Move the **Remove Genes That Have Missing Values** slider until the value is set to 1. This will cause all genes with at least one missing value to be removed. The rest of the dialog is grayed out since there will be no missing values left to estimate.

4. Click **OK**. The gene elimination operation is performed, and upon successful completion, a new Estimated: #mv < 1 | median dataset is added to the **Experiments** navigator under the original dataset.

**Tutorial 4: Step 6 Normalize the Data**

**Normalize the Data**

1. If the Estimated: #mv < 1 | median dataset in the **Experiments** navigator is not already highlighted, click it.

2. Click the **Normalize** icon  or select **Normalize** from the **Data** menu, or right-click the item and select **Normalize** from the shortcut menu. The first **Normalization** dialog is displayed.
3. Double-click **Logarithm** or ensure **Logarithm** is selected and click **Next**. The second **Normalization** dialog is displayed.

4. Double-click the **base 2** radio button or ensure the **base 2** button is selected and click **Finish**. The normalization operation is performed, and upon successful completion, a new Norm: log2 dataset is added under the Estimated: #mv < 1 | median dataset the **Experiments** navigator.

**Tutorial 4: Step 7 Display Summary Statistics**

**Display Summary Statistics**

1. If the Norm: log2 dataset in the **Experiments** navigator is not already highlighted,
click it.

2. Click the **Summary Statistics** icon, or select **Summary Statistics** from the **Statistics** menu. The Summary Statistics chart is displayed.

![Normalization Histogram](image)

3. The Summary Statistics chart shows an approximately normal distribution reflecting the roughly log-normal shape of the normalized data.

**Tutorial 4: Step 8 Create a SOM Experiment**

**Create a SOM Experiment**

1. If the Norm: log2 dataset in the **Experiments** navigator is not already highlighted, click it.

2. Click the **Self-Organizing Map** toolbar icon, or select **Self-Organizing Map** from the **Clustering** menu, or right-click the item and select **Self-Organizing Map** from the shortcut menu. The **Self-Organizing Map** parameters dialog is displayed.
3. Set dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orientation</td>
<td>Samples</td>
</tr>
<tr>
<td>Distance Metric</td>
<td>Pearson Correlation</td>
</tr>
<tr>
<td>Map Dimensions: Height</td>
<td>2</td>
</tr>
<tr>
<td>Map Dimensions: Width</td>
<td>2</td>
</tr>
<tr>
<td>Algorithm Properties: Random seed</td>
<td>99</td>
</tr>
</tbody>
</table>

4. Accept all the other defaults and click **OK**. The SOM operation is performed and a new SOM: samples | 2x2 | Pearson item is added to the **Experiments** navigator under the original dataset.

If you have automatic visualizations enabled in your user preferences, a SOM plot is displayed.

We are using a low number of nodes in this SOM because we are only looking for a small number of classes among the samples, namely: AML or ALL, and possibly the cell type, B or T.

**Note on use of the Random seed parameter:** In normal use, setting the random seed is neither necessary nor recommended. In a tutorial you set the random seed to a consistent value so that you will obtain precisely the same results as we depict and discuss, which makes the tutorial easier to understand. When you are not following a tutorial, you should generally not adjust the random seed at all.

The random seed setting may affect irrelevant details, such as the orientation and labelling of the SOM map. In other cases the random seed may affect relevant details, such as which genes occur together in clusters. Because of this latter possibility, it is sometimes worth repeating an experiment with different random seeds to see what the effects are. **GeneLinker™** helps with this by setting a new random seed every time an operation is carried out, so you don't need to.

On occasion you may need to determine whether a certain variation in results is due to
the random element, or some other cause. For this reason you are able to set the random seed to a fixed value, thus controlling that source of variation.

**Tutorial 4: Step 9 Create a SOM Plot**

If the SOM Plot is already displayed, there is no need to recreate it. Read below the image for information about the plot.

**Create a SOM Plot**
1. Double-click the SOM: samples | 2x2 | Pearson experiment in the **Experiments** navigator. The item is highlighted and a SOM plot of the selected item is displayed.

OR
1. If the SOM: samples | 2x2 | Pearson experiment in the **Experiments** navigator is not already highlighted, click it.
2. Select **SOM Plot** from the **Clustering** menu, or right-click the item and select **SOM Plot** from the shortcut menu. A SOM plot of the selected item is displayed.

![SOM Plot](image)

**A Tour of the Plot**

The 'architecture' of the SOM, which you input as Height and Width values in the example above, forms the heart of the plot. Each node of the SOM is depicted as a small solid circle. These are arranged in an array, in this case, of 4 nodes (= 2x2).

Each node is also surrounded by an open circle of varying size. The radius of this open circle indicates the number of cluster items associated with each node (e.g. the number of samples, if you clustered samples).

- Hover the mouse pointer over the node for about 2 seconds. A tooltip appears showing the number of items in that cluster, and the cluster name (e.g. 'Cluster #1').
- Click on one of the gray circles to select that cluster.

In the right-hand pane is the list of items in the selected cluster, and in the lower pane is
a characteristic profile of that cluster.

**Similarity Between Nodes**

Each node in a SOM is defined by its reference vector, and the similarity or distance between these reference vectors is part of the plot. This similarity is represented two ways:

1. By the coloration of the background behind the array of nodes.
2. By the lines linking adjacent nodes.

By default, the background color scheme uses dark blue to represent high similarity and white to indicate low similarity. Thus groups of similar nodes can be recognized as dark blue areas separated by light blue areas. Conversely, the lines linking adjacent nodes are colored light to represent high similarity, and dark to represent low similarity, so they should stand out against the background.

- If you forget this convention, you can look up the significance of the color scheme by right-clicking anywhere in the main SOM display and choosing **Customize** from the shortcut menu.

You can see that in our example the most similar pair of neighboring nodes is the pair at the bottom, Clusters #1 and #2.

- Click on Cluster #4 (the upper right node) to see what samples cluster there. From the sample names shown in the right-hand pane of the SOM display, you can see that this cluster is composed entirely of ALL samples drawn from T cells. Cluster #3 to its left is purely composed of AML samples, while Clusters #1 and #2 are principally made up of ALL samples from B cells - as might be expected from their high similarity mentioned above.

**Node Membership**

Display a line graph showing all the items in the cluster by clicking a node and selecting **Cluster Plot** from the **Clustering** menu, or by right-clicking a node and selecting **Cluster Plot** from the shortcut menu.
Tutorial 4: Conclusion

Discussion of the Results:
If you create new SOMs of the same data but with different random seeds, you should find slightly different distributions of samples each time. However, you should also find that there are certain features that do not change. For instance, there are consistently a small cluster of ALL-T samples, two clusters dominated by ALL-B samples, and a cluster of AML samples. The position of each of these clusters in the SOM will change, and certain samples will move from one cluster to another. Note, however, that certain samples do seem to cluster together consistently. For instance, sample AML-66 has a tendency to cluster with ALL-B samples. This indicates that sample AML-66 has a gene expression profile more like those of other ALL-B samples than of other AML samples, under this clustering protocol. This sample might therefore be considered a candidate for further investigation. A good first step would be to repeat the analysis varying other parameters such as the gene filtering method, the normalization, and the type of metric, to determine whether the interesting observation holds.

When you are finished, you can close all the open plots either by clicking on the 'x' box in the upper-right hand corner of each, or by selecting Close All from the Window menu.

References:
1. The basic reference on SOMs from the machine-learning perspective is Teuvo Kohonen Self-Organizing Maps, 2nd edn. (Berlin: Springer, 1997). Contains no discussion of application to gene expression data.
'Molecular Classification of Cancer: Class Discovery and Class Prediction by Gene Expression Monitoring' [Science 286: 531, 1999] applied 2x1 and 4x1 SOMs to the first 38 samples of the AML/ALL dataset.


Where To Go From Here

- Go through the other tutorials provided.
- Read the Online Help to learn more about the various functions of GeneLinker™.
- Further explore GeneLinker™ by using additional features.
- Load up your favorite dataset and try out all the buttons and menu items.
- Don't forget to right-click on things like plots - many details of graphics can be customized.
- Visit the Molecular Mining website at http://www.molecularmining.com/ for the latest information on GeneLinker™ enhancements and additional products.

Tutorial 5: Principal Component Analysis

Tutorial 5: Introduction

This tutorial introduces you to Principal Component Analysis (PCA). You will be shown how to perform the PCA experiment and then visualize the results in different types of plots.

Skills You Will Learn:
How to import gene expression data from a file into the GeneLinker™ database.
How to perform a PCA experiment.
How to visualize the results of a PCA experiment in various plots.
How to use the 3D plot functions.

Principal Component Analysis

A number of recently published analyses of gene expression data have centered their attention on Principal Component Analysis (PCA) as a method of extracting more information from data. We will study this application using the yeast elutrition dataset.
studied by Alter, Brown & Botstein [Alter2000].

The traditional application of PCA is to reduce the dimensionality of data. In gene expression experiments, where there are typically thousands of variables, it can be extremely useful to collapse the genes into a smaller set of principal components. This makes most types of plots easier to interpret, which can help to identify structure in the data.

In Alter et al, they discuss a dataset that explores the gene expression over time in yeast during an elutriation study. They include 14 measurements at half-hour intervals. One of the goals of the study was to verify whether there were cyclic patterns in gene expression that were commensurate with the yeast cell cycle. A related question was whether the genes known to be involved in various stages of the cell cycle would show time-shifted expression waves.

**Tutorial Length**

This tutorial should take about 30 minutes, depending on how long you spend investigating the data, and how fast your machine is.

If you must stop part way through the tutorial, simply exit the program by selecting Exit from the File menu. The data and experiments you have performed to that point are saved automatically by GeneLinker™. The next time you start GeneLinker™, you can continue on with the next step in the tutorial.

**Tutorial 5: Step 1 Import the Data**

**Import the Data**

1. Click the Import Gene Expression Data toolbar icon, or select Import from the File menu and Gene Expression Data from the sub menu. The Data Import dialog is displayed.

![Data Import Dialog](image)

2. Set the Gene Database field to Custom using the drop-down list. The gene ids in the Elutriation dataset are SGD ids.
3. GeneLinker™ uses a template to interpret data files being imported. Ensure that the template is Tabular.
4. The next step is to identify the name and location of the data source file. Click the button to the right of the Source File box. The Open dialog is displayed.
5. The tutorial data files are located in the Tutorial folder. This is the folder listed in Look in, so you do not need to navigate to it. Click the file Elutriation.csv and click Open. The Data Import dialog is updated with the file name.

6. Click Import. The Import Data dialog is displayed.

GeneLinker™ examines the file and offers to transpose it. Within GeneLinker™, datasets have the genes in columns and the samples in rows.
When importing data using a Tabular template, GeneLinker™ assumes that the more numerous dimension of your data represents genes (most microarray experiments involve more genes than samples). If this is so (as in this tutorial), then clicking **OK** is all that is required.

**Note:** the options **Use Sample Names** and **Use Gene Names** are checked and disabled in the Import Data dialog box. GeneLinker™ has recognized that in this dataset, the first row and column contain alphameric labels. Gene expression data is always numeric, hence the disabled checkboxes.

7. Click **OK**. The data is imported and an item named Elutriation is added to the **Experiments** navigator. This represents the raw, publicly available data which has already been normalized.

**Tutorial 5: Step 2 Principal Component Analysis**

Principal Component Analysis

1. If the Elutriation dataset in the **Experiments** navigator is not already highlighted, click it.

2. Click the **Principal Component Analysis** toolbar icon or select **Principal Component Analysis** from the PCA menu, or right-click the item and select **Principal Component Analysis** from the shortcut menu. The **PCA** parameters dialog is displayed.

3. You may choose to perform PCA calculation on either **Genes** or **Samples**. The typical use of PCA is to reduce the genes to a smaller number of 'variables' as in this tutorial. Ensure that **Genes** is selected. (In other applications, where the samples are being thought of as 'variables' or measurements for particular genes, you would select **Samples**).

4. Click **OK**. The **Experiment Progress** dialog is displayed.

The dialog is dynamically updated as the PCA calculation is performed. Upon successful completion, a PCA: genes item is added to the **Experiments** navigator under the original dataset.

If you have automatic visualizations enabled in your user preferences, a 3D Score Plot is displayed.
Tutorial 5: Step 3 Display a Scree Plot

Principal components can be used to determine how many real dimensions there are in the data. There is a particular mathematical meaning to number of dimensions, but an intuitive understanding can be achieved by considering the amount of variation in the data that is explained by various principal components. If a small number of components accounts for most of the variation in the data, then the other components can be thought of as noise variables.

Determining which principal components account for which parts of the variance can be done by looking at a Scree Plot.

Display a Scree Plot
1. If the PCA: genes experiment in the Experiments navigator is not already highlighted, click it.
2. Select Scree Plot from the PCA menu, or right-click the item and select Scree Plot from the shortcut menu. A scree plot of the PCA results is displayed.

Interpretation:
The Scree Plot has two lines: the lower line shows the proportion of variance for each principal component, while the upper line shows the cumulative variance explained by the first N components. The principal components are sorted in decreasing order of variance, so the most ‘important’ principal component is always listed first. In this dataset the first two principal components explain much more of the variance in the data (roughly 25% and 20% respectively) than do any of the subsequent principal components (all less than 10%). In this data, most of the important biological behavior is somehow being captured in these two components, leading us to take a closer look at them and their meaning in the context of the yeast cell cycle.
Tutorial 5: Step 4 Display a Loadings Line Plot

Visualize the Principal Components
The principal components are new variables made up of combinations of the original data variables, in this case, genes. Each component is some linear combination of the original gene variables, and often looking at which genes or gene families have a large contribution to a principal component can be an indication of shared function of behavior, similar to the inferences that can be made using clustering.

Three plots are available to view the coefficients or loadings: Loadings Scatter Plots, Loadings Line Plots and Loadings Color Matrix Plots. Loadings Scatter Plots with many thousands of variables tend to be non-informative: they are better suited to PCA on smaller gene sets or on samples. As a results, we will focus our attention on the other two plot styles.

Display a Loadings Line Plot
1. If the PCA: genes experiment in the Experiments navigator is not already highlighted, click it.
2. Select Loadings Line Plot from the PCA menu, or right-click the item and select Loadings Line Plot from the shortcut menu. A loadings line plot of the PCA results is displayed.

- If you want the plot to be wider, right-click on the plot and select Resize from the shortcut menu to set the desired dimensions of the plot.

Interpretation
Even in this traditional Loadings Line Plot it is difficult to see much structure. In particular, the first two principal components, which are of most interest because of their
ability to explain most of the variance in the data, are quite difficult to see in this plot. A Loadings Line Plot can be more helpful when PCA is done by samples or if a relatively small number of genes is being studied.

**Tutorial 5: Step 5 Display a Loadings Color Matrix Plot**

To get a finer resolution of the coefficients, it can be more effective to look at a Loadings Color Matrix Plot. These represent exactly the same numbers that were in the Loadings Line Plot, but they are displayed in a way that is easier to interpret when large numbers of variables are present.

**Display a Loadings Color Matrix Plot**

1. If the PCA:genes item in the Experiments navigator is not already highlighted, click it.

2. Click the **Loadings Color Matrix Plot** toolbar icon, or select **Loadings Color Matrix Plot** from the PCA menu, or right-click the item and select **Loadings Color Matrix Plot** from the shortcut menu. A loadings color matrix plot of the PCA results is displayed.

**Note**: This plot initially displays genes' or samples' rows in descending numerical order as established by the loadings on the first principal component (PC1). You can change the display order of rows by clicking the respective up/down arrow at the top of each PC column in the color matrix. For each PC, you may choose to sort the genes in descending order of absolute value, simple descending order, or ascending order. This allows you to identify easily genes which are most strongly correlated or anti-correlated with the first principal component, for example.

3. To see more gene components at once, click **Resize** and move the **Height** slider to the far left (minimum).
Tutorial 5: Step 6 Display a Score Plot

Visualize the Projection of the Samples
In Alter et al. it was clear that there were cyclic patterns in the data, visible across different genes. The next question was whether this cyclic behavior could be seen in the time progression of the samples. One way to study this is to look at the score plot of the Principal Component Analysis. In particular, since the first two principal components of the genes seem to show this cyclic property, and they account for the majority of the variance in the data, we would like to examine the projection of the samples over time onto these two most important components.

Display a Score Plot
1. If the PCA:genes experiment in the Experiments navigator is not already highlighted, click it.
2. Select Score Plot from the PCA menu, or right-click the item and select Score Plot from the shortcut menu. A score plot of the PCA results is displayed.

• The scatter plot displays a point for every sample in the dataset and it can be difficult to interpret, especially with respect to the units. However, if you look carefully at the points and their distribution you will see that there is a pattern to the data.
3. On the right hand side of the Score Plot in the legend, click the first data point, e_0m. The name is highlighted as is its point in the bottom of the plot.
4. Press the <down arrow> to select successive samples (e_30m, e_60m, etc) and watch as the highlighted point walks clockwise around the plot.

This general clockwise layout of the points as they lie in time is another indicator that a cyclic behavior is being captured by the first two principal components. To better see
this pattern, normalize the Score Plot:

5. Click the **Raw Data/Normalize Score Plot** button in the upper right of the score plot window. The score plot is updated to show a normalized version of the data.

![Score Plot](image)

**Interpretation**

In this plot, the original samples are again projected onto the new variables or principal components. The difference is that the projections have been normalized so the values in the plot reflect how similar each sample is to a given principal component. Alter referred to this as the correlation between a sample and a principal component. Using this type of plot we can make more direct comparisons of the amount each principal component represents of each sample. Again, we can see the points that fall successively in time also follow each other in a clockwise direction around the unit circle.

In both the raw and normalized versions of the score plot, the 300 minute sample (e_300m) seems to break the circular pattern. In such cases, where one or two point seem to be anomalous, or break a general pattern in the data, it can be helpful to study these exceptional points using other sources of information. For example, with PCA, we do not need to limit ourselves to the first two principal components.

**Tutorial 5: Step 7 Display a 3D Score Plot**

**Display a 3D Score Plot**

1. Double-click the PCA:genes experiment in the **Experiments** navigator. The item is highlighted and a 3D score plot of the selected item is displayed showing the first three PCs.

   OR

1. If the PCA:genes experiment in the **Experiments** navigator is not already highlighted,
click it.

2. Click the **3D Score Plot** toolbar icon, or select **3D Score Plot** from the PCA menu or right-click the item and select **3D Score Plot** from the shortcut menu. A 3D score plot of the selected item is displayed showing the first three PCs.

![3D Score Plot](image)

- Notice that this view is similar to the 2-dimensional plots from before, but with the depth of the points reflecting their scoring relative to the third principal component.

3. In the right-hand list of points (legend), click the point 'e_300m'. The item and its point are highlighted.

**Rotate the plot**

1. Click on the plot and slowly drag the mouse to the left to spin the plot until it is similar to the one below.
Interpretation

This plot brings out a dramatic difference between the measurements at 300 minutes relative to the other measurements. Not only do the gene expression levels at this time seem not share the same cell cycle patterns as the other time points, this time point has very different properties, reflected in the abnormally high score in the third principal component. This indicates that something fundamentally different occurred during this measurement, with either experimental error or some type of significant biological change being the natural candidates.

2. Click the Home button in the upper right of the plot. This returns the plot to its original orientation.

3. Click the Raw Data/Normalize 3D Score Plot button in the upper right of the score plot window. The 3D score plot is updated to show a normalized version of the data. Rotate the plot as above.
Interpretation

In a score plot, the later principal components, which represent less of the overall variance, can seem visually less significant than the first few principal components. This appearance can be deceptive and lead you to neglect the real impact or separation due to later principal components. To compensate, score plots may be normalized so that each principal component has the same range (-1 to +1). When normalization is applied to the Elutriation data, the separation of time point e_300m along PC3 is even more visible than in the original plot.

Tutorial 5: Conclusion

Summary

In this tutorial we have taken a yeast cell cycle dataset with a strong cyclic behavior and examined it through Principal Component Analysis. During this survey we have considered three important elements of PCA: the variances in the data (Scree Plot), the relationship between the genes and the components (Loadings Line Plot and Loadings Color Matrix Plot), and the projection of the samples in the new components (Score Plot - Raw Data and Normalized). The Scree Plot indicated that the first two principal components captured most of the behavior of the data. The Loadings and Score Plots brought into relief the periodicity of the yeast cell cycle, both in genes and in time.

When you are finished, you can close all the open plots either by clicking on the 'x' box in the upper-right hand corner of each, or by selecting Close All from the Window menu.
References

Where To Go From Here
• Go through the other tutorials provided.
• Read the Online Help to learn more about the various functions of GeneLinker™.
• Further explore GeneLinker™ by using additional features.
• Load up your favorite data set and try out all the buttons and menu items.
• Don't forget to right-click on things like plots - many details of graphics can be customized.
• Visit the Molecular Mining website at http://www.molecularmining.com/ for the latest information on GeneLinker™ Gold enhancements and additional products.

Tutorial 6: Learning to Distinguish Cancer Classes

Tutorial 6: Introduction

This tutorial introduces you to data mining and prediction. You will use the integrated SLAM™ technology to mine a dataset for sets of gene associations. A gene list will be created from the most interesting features (genes). You will create and evaluate an ANN classifier.

Skills You Will Learn:
How to import gene expression data from a file into the GeneLinker™ database.
How to import variable class data.
How to discretize expression data.
How to run SLAM.
How to use the SLAM association viewer.
How to create a gene list.
How to create, evaluate, and predict classes using an ANN classifier.

Scientific Background
This tutorial is a reanalysis of the data reported by Khan, Wei, Ringnér et al. in Nature Medicine (2001) [Ref.1]. We refer to this paper simply as 'Khan' in this tutorial.

The object of the paper and of this tutorial is to learn to distinguish, at the molecular level, between types of small round blue-cell tumors (SRBCTs) such as Ewing sarcoma (EWS), Burkitt lymphoma (BL), neuroblastoma (NB) and rhabdomyosarcoma (RMS).
These tumors are difficult to distinguish by visual methods, and respond to different treatments.

The data is available on the World Wide Web as supplementary material, at http://www.thep.lu.se/pub/Preprints/01/lu_tp_01_06_supp.html. The authors pre-filtered the data for a minimal level of expression, leaving measurements for 2308 genes.

**Tutorial Workflow**

The purpose of the workflow covered by this tutorial is to select a small number of genes (called features) that as a set are able to predict the cancer type of a given tissue sample. Once this small set of genes has been selected by SLAM™, a committee of artificial neural networks (ANNs) is trained using the expression levels of only those genes.

Feature selection and ANN training take place on the same set of data, called the **training dataset**. The samples in this dataset have known classes, so the ANN training is done under the supervision of this available knowledge. Once the ANN committee has been trained, it can be used on new data of the same phenomenon (SRBCTs), to predict the classes of its samples. This new data is called the **test dataset**.

This tutorial demonstrates how a combination of SLAM™ and a committee of trained ANNs can be used to effectively classify difficult-to-distinguish cancers using as few as eight genes.

**What You Will Learn:**

1. How to run SLAM™ and use the results to create gene lists.
2. How to train artificial neural networks (ANNs)
3. How to use trained ANNs to distinguish and predict sample classes.

**Tutorial Length**

This tutorial should take about an hour, depending on how long you spend investigating the data, and how fast your machine is.

If you must stop part way through the tutorial, simply exit the program by selecting Exit from the File menu. The data and experiments you have performed to that point are saved automatically by GeneLinker™. The next time you start GeneLinker™, you can continue on with the next step in the tutorial.

**Platinum**

**Tutorial 6: Step 1 Import the Data**

**Import the Data**

Two datasets need to be imported to perform this tutorial. The first is 'Khan_training_data' and the second is 'Khan_test_data'. Follow the procedure for importing the first dataset and then repeat it for the second (using the correct dataset file name).

1. Click the **Import Gene Expression Data** toolbar icon , or select **Import** from the
File menu and Gene Expression Data from the sub menu. The Data Import dialog is displayed.

2. Set the Gene Database to Custom. (On the second import, you should find the dialog retains the setting you gave it on the first import, so no need to reset it.)
   - The identifiers in this dataset are clone ids from the IMAGE Consortium (http://image.llnl.gov). Since they are neither GenBank, UniGene nor Affymetrix identifiers, use the Custom database slot for these. Later in the tutorial we will look up the genes in the GenBank database via their IMAGE identifiers.

3. Click the Source File button. The Open dialog is displayed.

4. Click the file 'Khan_training_data.csv'. (For the second import, 'Khan_test_data.csv').

5. Click Open. The Data Import dialog is updated with the file name.

6. Click Import. The Import Data dialog is displayed.
7. Click **OK**. The dataset is imported and a new item is added to the **Experiments** navigator. *Repeat the import process for the second dataset.*

For detailed information on importing data, see Data Import Step 1: Selecting a Template.

**Platinum**

**Tutorial 6: Step 2 Import Variable Data**

For complete information on variables, see Variables Overview.

Variable (class) data for both Khan datasets needs to be imported. The first class data file is Khan_training_classes.csv and the second is Khan_test_classes.csv. Follow the procedure to import the first and then repeat it to import the second using the additional information in parentheses.

**Import Variable Data**

1. Click the Khan_training_data dataset (Khan_test_data for the second import) in the **Experiments** navigator. The item is highlighted.

2. Select **Import** from the **File** menu, and **Variable** from the sub menu. The **Import Variable** dialog is displayed.
   - The **Dataset** name is displayed at the top of the dialog and the number of samples in the dataset is listed under the name.
3. Click the **Source File ...** button. The **Open** dialog is displayed.

4. Click the file Khan_training_classes.csv (Khan_test_classes.csv for the second import). The item is highlighted.

5. Click **Open**.
   - The **Source File** name is displayed with the number of observations and classes in the file listed underneath.
   - The default **Variable Name** and **Description** are displayed.
6. The **Preview** allows you to view which sample belongs to which class and the total number of entries for each class. Click **Preview**. When you are finished examining the contents of the Preview, click **Close** to close it.

7. Type **training classes** into the **Variable Name** field overwriting what was there (**test classes** for the second import).

*For the second import, skip to #12 below - no need to create the variable type again.*

8. For the first import, click **New Variable Type**. The **Create Variable Type** dialog is displayed.

- This variable type is used to group together all the observations and predictions of SRBC tumor types. For further discussion of variables and variables types, see Variables Overview. Once we have created the variable type **tumor type**, we will import variables of that type describing (first) the tumor type of the training data, and (second) the tumor type of the test data.

9. Type **SRBC Tumors** into the **Name** field, overwriting the default name.

10. Click **OK**. The **Import Variables** dialog is updated with the new variable type.
Note: the number of samples (listed under the Dataset name at the top of the dialog) equals the number of observations listed below the Source File. It is essential that these numbers match - that is, there is a class value for each and every sample.

11. Click Import. The variable (class) data is imported and the Khan_training_data (Khan_test_data) dataset in the Experiments navigator is tagged with the variable information indicator icon.

For detailed information on variable import, see Importing Variables.

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**Tutorial 6: Step 3 Discretize the Data**

The first step in our analysis of this dataset is to use SLAM™ to look for associations between multiple genes and the tumor type.

SLAM™ finds associations between genes based on identical patterns of gene expression. For example, if Gene A is HIGH whenever Gene B is LOW, SLAM™ identifies an association between Gene A and Gene B. Because the number of possible patterns is enormous, particularly when looking for patterns between five or ten genes rather than just two, we need a fast, simple means of comparing expression levels. By discretizing the data, it becomes possible to compare expression levels in terms of a small number of discrete categories (e.g. HIGH/MEDIUM/LOW) rather than continuous values. This speeds up the comparison process by many orders of magnitude.

**Discretize the Data**
1. Click the Khan_training_data dataset in the Experiments navigator. The item is highlighted.

2. Click the Discretize toolbar icon or select Discretize Data from the Predict menu, or right-click the item and select Discretize Data from the shortcut menu. The Discretization parameters dialog is displayed.

---

**Operation Type**

- **Quantile Discretization** means dividing the data into equally-populated groups. Thus 3-way quantile discretization per gene will yield a roughly equal number of high (2), medium (1) and low (0) values for each gene.

- **Range Discretization** makes the groups cover equal ranges. For example, if the gene had values ranging from 0.0 to 24.0, a 3-way range discretization would consist of values between 0 and 8, 8 and 16, and 16 and 24, and the three groups might be quite differently populated.

**Number of Bins**

Choosing the number of bins is a balancing act. The more bins you use, the less information is discarded by the discretization. But the more bins there are, the fewer associations SLAM™ will find.

Accept the default parameters (Quantile discretization, Per Gene, and 3 bins).

3. Click **OK**. The Experiment Progress dialog is displayed. It is dynamically updated as the discretization operation is performed.

Upon successful completion, a new Discretized: 3 bins/gene | quantile dataset is added under Khan_training_data in the Experiments navigator.

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**Platinum**

**Tutorial 6: Step 4 Run SLAM**

**Associations in Data**

Sub-Linear Association Mining (SLAM™) is a method for finding associations in discrete
data. An association is a set of variables (genes) and values which occur together in a dataset at a rate higher than that expected by chance. For instance, it might happen that in kidney tissue repression of gene A results in the up-regulation of genes B and C, and down-regulation of gene Q. In this case, we would expect to find an association in the dataset like this:


**Note:** this says nothing about how B, C, and Q are regulated when A is not repressed, or when a different tissue is being considered.

Such sets of variables have several potential uses. In GeneLinker™, they are used to identify key sets of genes which might be predictive of a given sample classification. This use, called *feature selection*, is vital to making predictions because of the enormous number of genes in a microarray experiment which are typically not connected to the class of interest.

### The SLAM™ Parameters

Imagine you are searching for a book in a library, and you know it’s Dewey Decimal number. One way you could find it would be to start at 100.00 and walk along the shelves until you get to the number of your book. This is not very efficient. Instead, you might walk around at random and glance at numbers now and then, making a random sampling of what books are near you at any given time. This is a surprisingly efficient strategy, and SLAM™ uses something like it to find associations in gene expression data.

Two of the parameters in the dialog above relate to SLAM™’s random sampling behavior. One is the **Number of Iterations**. This is the number of random subsets of your data SLAM™ uses to find associations. The higher the number of iterations, the more and better associations will be found, but the longer the algorithm will take to run.

The second parameter is the **Random Seed**. This controls the sequence of random numbers that are used by the algorithm to select subsets. If the seed is set to the same value, and SLAM™ is run again, it will produce identical results. Running SLAM™ on the same data with different random seeds will produce similar, but not identical, results, because slightly different subsets will have been selected from the data.

The **Representative Variable** is the variable you want to classify on. Datasets may have several variables associated with them (cancer type, tissue type, gender, etc.) and you can use SLAM™ to search for features that discriminate between values of any variable.

**Support** is the number of subsets an association must appear in before it is considered significant. Associations with less than the minimum support will not be reported.

**Matthews Number** is a measure of how good an association is at discriminating between classes. Perfect discrimination is represented by a Matthews number of 1. Useful values are typically between 0.5 and 0.7.

### Run SLAM™ on the Discretized Data

1. If the newly created Discretized: 3 bins/gene | quantile dataset in the Experiments navigator is not highlighted, click it.
2. Click the SLAM™ toolbar icon 🎨, or select SLAM from the Predict menu, or right-click the item and select SLAM from the shortcut menu. The SLAM™ parameters
dialog is displayed.

![SLAM dialog image]

3. Set the dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Representative Variable</td>
<td>training classes</td>
</tr>
<tr>
<td>Number of Iterations</td>
<td>30000</td>
</tr>
<tr>
<td>Support</td>
<td>4</td>
</tr>
<tr>
<td>Matthews Number</td>
<td>0.7</td>
</tr>
<tr>
<td>Results</td>
<td>Return all results found.</td>
</tr>
<tr>
<td>Random Seed</td>
<td>999 (see Note below)</td>
</tr>
</tbody>
</table>

4. Click OK. The SLAM™ operation is performed. *This may take fifteen minutes or so*, on an IBM box as described in the System Specification. Upon successful completion, a new item (SLAM) is added under the Discretization item in the **Experiments** navigator.

If you have automatic visualizations enabled in your user preferences, the SLAM Association Viewer is displayed.

**Note on Use of the Random Seed Parameter**

In normal use, setting the random seed is neither necessary nor recommended. In a tutorial you set the random seed to a consistent value so that you will obtain precisely the same results as we depict and discuss, which makes the tutorial easier to understand. When you are not following a tutorial, you should generally not adjust the random seed at all.

In SLAM™, the random seed can be thought of as prescribing the starting point for the search for associations. If SLAM™ is allowed to run long enough, it will find all of an enormous set of associations which inhabit any given dataset, but the smaller you set the number of iterations, the greater will be the effect of the random seed. Conversely, the random seed matters less and less as the number of iterations grows greater. It is usually better to set the iteration number high and let SLAM™ run overnight than to do repeated runs with different random seeds.
If the SLAM association viewer is already displayed, there is no need to recreate it. Read the sections below the image for information about the SLAM Association Viewer.

**View SLAM™ Results**

1. Double-click the newly created SLAM: training classes | 30000 | 4| 0.7 item in the Experiments navigator. The item is highlighted and the SLAM association viewer is displayed.

OR

1. If the newly created SLAM: training classes | 30000 | 4| 0.7 item in the Experiments navigator is not already highlighted, click it.
2. Click the Association Viewer toolbar icon, or select Association Viewer from the Predict menu, or right-click the item and select Association Viewer from the shortcut menu. The SLAM™ association viewer is displayed.

The SLAM™ Association Viewer has three functional areas:

**Associations:**

The **Associations** list displays a list of all the associations found during the SLAM™ run. To sort the list by a particular characteristic, click on the column header for that characteristic. Clicking again on the same header reverses the order of the sort (ascending or descending).

The Associations list can be sorted by:

- Matthews statistic (a measure of the predictive power of the association),
- Support (the number of samples in the dataset which match the pattern),
- Class, or
- The number of genes in the association.

**Genes:**
The **Genes** list box in the upper right lists the genes in the checked associations. A gene list can be created from the checked genes in the **Genes** box. The gene list can be used to identify interesting genes (features) for use in supervised learning experiments. **Note:** only one copy of a gene name is listed in the **Genes** list box. The **Count** column indicates the number of associations the gene occurs within.

**Association Filter**

Since SLAM™ can potentially find hundreds or even thousands of associations, some methods are provided in the Association Filter group for reducing the number of associations displayed. You can display only associations with a Matthews statistic above an adjustable cutoff, or you can display only associations containing certain genes, or not containing certain genes.

**Platinum**

**Tutorial 6: Step 6 Create a Gene List**

The next objective is to find genes that are key indicators or **features** which can be used to discriminate between cancer classes. The first step is to create a gene list from the discovered associations using the **Create Gene List** function built into the SLAM™ Association Viewer.

**Create a Gene List**

1. If you changed the sorting of the association list, click the **Matthews** column header until the associations are sorted in decreasing order of Matthews statistic (this is the default order for associations).
2. Click the top checkbox in the **Associations** list. Then press and hold down the **<Shift>** key and click the checkbox beside the highest association involving class BL. This selects the top eleven associations and adds their 123 genes to the **Genes** list box displayed to the right of the **Associations** list. This captures at least some associations for three of the four classes we are trying to distinguish.

Because classes with few samples (such as class BL in this dataset) tend to generate associations with many genes, these 11 associations have given us 123 genes in the **Genes** list box. This is too many features to use for training a classifier when we only have 63 samples. Using closer to 1/10th as many features as samples is a much better idea, so we will now reduce the number of genes.
3. In the **Genes** list box, click the **Count** column header so that the genes are sorted in descending order of number of associations (the column header contains a small down arrowhead). Notice that only the top 8 genes occur more than once in these 11 associations.

4. Click the **Uncheck All** button below the **Genes** list box.

5. Click the checkbox to the left of the top gene in the **Genes** list box. Then press and hold down the <Shift> key and click the checkbox beside the eighth gene. This selects the 8 genes with a count greater than 1. The text below the **Genes** list box says '8 of 123 genes selected'.

6. Click the **Create Gene List** button. The **Create a Gene List** dialog is displayed.

7. In the **Name** field, type **Tutorial 6 list** and in the **Description** field, type **8 genes from top 11 associations**.

8. Click **Save**. The new gene list is added to the **Gene Lists** navigator.
   - Click the **Gene Lists** tab in the navigator to see the list of gene lists.
   - Click the **Experiments** tab to return to the **Experiments** navigator.

9. Click the **Close** icon in the upper right corner of the SLAM™ Association Viewer.
In this step, new datasets, containing only the expression values for the genes in the gene list, are created from the training and test datasets by the process of gene list filtering. This step ensures that the dataset used to train the ANN classifier contains the same genes as the test dataset.

**Note:** gene list filtering does not change the order of genes in a dataset, and for classifying with an ANN classifier, the test dataset must contain not only the same genes as the training dataset, but they must also be in the same order and without any extra genes.

**Filter Original Datasets Using the Gene List**

Follow the procedure for the 'Khan_training_data' dataset and then repeat it for 'Khan_test_data'.

1. Click the 'Khan_training_data' ('Khan_test_data' for the second filter) item in the **Experiments** navigator. The item is highlighted.
2. Click the [Filter](#) toolbar icon, or select **Filter Genes** from the **Data** menu, or right-click the item and select **Filter Genes** from the shortcut menu. The **Filter Genes** parameters dialog is displayed.

3. Set dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filtering Operation</td>
<td>Gene List Filtering</td>
</tr>
<tr>
<td>Filtering Operation Type</td>
<td>Keep only genes that are in this list</td>
</tr>
<tr>
<td>Gene List</td>
<td>Tutorial 6 List</td>
</tr>
</tbody>
</table>

4. Click **OK**. The gene list filtering operation is performed, and a new item (Filter Genes) is added under the 'Khan_training_data' ('Khan_test_data') item in the **Experiments** navigator.

- Since the classifier that is to be created must have the same inputs (genes) to work on when it makes predictions as it does when it is trained, the training and test datasets are filtered the same way. If this is not done, the classifier may produce nonsensical predictions. It is not strictly necessary to filter both the training and test data at the same time. You could filter the test data after you have created a classifier, but before running the classifier on the test data.
ANN Classifier Structure
GeneLinker™’s Artificial Neural Networks consist of three layers of nodes or neurons.

The input layer is connected to the output layer via a hidden, or internal, layer. The input layer has a single node per gene, so if you have eight genes that you want to train the ANNs on, GeneLinker™ automatically builds networks with eight input nodes. The output layer has a single node per class, so if the data have four classes, GeneLinker™ automatically builds a network with four output nodes. The number of nodes in the hidden layer should be greater than or equal to the number of nodes in the input layer, and fewer than twice the number of nodes in the input layer. Too many nodes in the hidden layer results in poor training performance, and too few results in poor classification performance.

Because individual ANNs can sometimes perform poorly on certain inputs, having a committee architecture improves the reliability of classification. Typically 10 is a reasonable number of committee members, with the requirement that 80% of committee members agree for a classification to be made. For a complete description of all of the parameters for creating an ANN committee classifier, please see Creating an ANN Classifier.

Create an ANN Classifier
1. Click the Filtered:keep {Tutorial 6 list} item under the Khan_training_data item in the Experiments navigator. The item is highlighted.
2. Click the Create Classifier toolbar icon , or select Create Classifier from the Predict menu, or right-click the item and select Create Classifier from the shortcut menu. The Create Classifier parameters dialog is displayed.
3. Set dialog parameters.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Representative Variable</td>
<td>training classes</td>
</tr>
<tr>
<td>Training Parameters: Hidden Units</td>
<td>5</td>
</tr>
<tr>
<td>Miscellaneous: Random Seed</td>
<td>999 (See Note below)</td>
</tr>
</tbody>
</table>

4. Accept the default values for the all other parameters and click OK. The Create Classifier operation is performed, and a new item (ANN: training classes | 8-5-4 | N=10 | 0.0010 | 10) is added under the Khan_training_data Filtered: keep {Tutorial 6 list} item in the Experiments navigator.

If you have automatic visualizations enabled in your user preferences, the Classification plot showing training results is displayed.

Training Parameters
The number of classifiers (10) is arbitrary. The number of hidden units (5) is more significant. Using more hidden units than there are input classes (i.e. 4 in this example) is a little risky but not wrong. In this case the number of hidden units is the number of classes we're really dealing with: 4 SRBCTs plus 1 class for the non-SRBCT samples in the test dataset.

Note: For reasons discussed in 'Tutorial 6: Step 5 Run SLAM', setting the random seed is neither necessary nor recommended in normal use. In the Create Classifier function, the random seed determines how the samples are divided up into subsets for training the component learners (committee members). It also determines how the individual learners (neural nets) are initialized. The random seed generally only affects predictions for borderline or ambiguous samples, which the committee also helps diagnose.

For a discussion of the other parameters in this dialog, see Create Classifier.

It is possible to view the results of the classifier training at this point (see Classifier Plot Training Results), but it is even more informative to go on and test the classifier using data it has not already seen.

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Tutorial 6: Step 9 Classify Test Data

We now further test our classifier by predicting the classes of some samples which it has not already seen. These are in the Khan_test_data dataset. We have already filtered it, so we have a subset containing exactly those same genes we have just used to train the classifier.

Classify New Samples
1. Click the Filtered: keep {Tutorial 6 list} item under Khan_test_data in the Experiments navigator.
2. Click the Classify toolbar icon  , or select Classify from the Predict menu, or right-click the item and select Classify from the shortcut menu. The Classify parameters...
dialog is displayed.

3. Set dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>Type in a name for the variable which will contain the predicted classes of the test data. <strong>Predictions</strong> is a suitable name in this instance. (If you were planning on doing multiple different predictions, you might want to give it a more distinctive name.)</td>
</tr>
<tr>
<td>Description</td>
<td>If you wish, click in the field and type in a long, informative description to the prediction being carried out.</td>
</tr>
<tr>
<td>Classifier</td>
<td>This displays a subset of the <strong>Experiments</strong> navigator containing those classifiers that can be applied to the dataset. Click on the <strong>ANN: training classes</strong> item beneath the Khan_training_data heading (the classifier just trained).</td>
</tr>
</tbody>
</table>

4. Click **OK**. The Classify function is performed, a new variable is added to the dataset family, and a new Classify item (named Predictions) is added the **Experiments** navigator under the Filter Genes item.

   If you have automatic visualizations enabled in your user preferences, the Classification plot is displayed showing the classification results.

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**Tutorial 6: Step 10 Display a Confusion Matrix**

**View the Classify Results**

1. If the newly created Predictions (or whatever name you gave the new classification in the previous step) item in the **Experiments** navigator is not already highlighted, click it.

2. Select **Variable Manager** from the **Tools** menu. The **Variables** dialog is displayed.
You see a list of the variables GeneLinker™ currently has associated with the Khan_test_data dataset family. Each variable has a name, a type, and whether it was imported (Observed) or generated by a classifier (Predicted).

3. Click on **test classes**. It is highlighted.
4. Hold down the `<Ctrl>` key and click on the Predictions item. Both variables are highlighted.
5. Click **Show Confusion Matrix** at the bottom of the dialog. The Confusion Matrix plot is displayed.

**Description of the Confusion Matrix**

The confusion matrix is an array which summarizes the comparison between two variables relating to a dataset. Typically the variables are an observation and a prediction. Each row in the confusion matrix represents an observed class, each column represents a predicted class, and each cell counts the number of samples in the intersection of those two classes. Entries on the diagonal of the matrix (in dark green) count the correct calls or predictions. Entries off the diagonal (in red, if there are any) count the misclassifications.

At the top of the confusion matrix display are two bars representing the overall accuracy of the prediction and the error rate.

Observations labelled 'Unknown' are not included in calculating the accuracy of the learner, since they are taken to represent cases where the scientist really does not know the class of the sample. Therefore any prediction made by GeneLinker™ in these cases can neither be counted as correct or incorrect.

In contrast, a prediction of 'Unknown' from GeneLinker™ means that the program could not confidently assign a class to the sample. Such a prediction is counted as an error if
there is an observed class available for the sample (that is, a class other than 'Unknown').

This behaviour of the confusion matrix summary can be modified by checking or un-checking the box at the left of each row and the head of each column. You can also use the checkboxes, for example, to restrict the accuracy summary to consider only two classes of a multi-class problem.

**Discussion of the Example Data**

Five samples in this test data do not belong to any of the four training classes: TEST-3, TEST-5 and TEST-11 are other cancers, and TEST-9 and TEST-13 are normal muscle tissue. They are labelled 'Unknown' in this tutorial and are represented by the last row in the confusion matrix above. Four of these five non-SRBCT samples are predicted to belong to one or the other of the training classes, which illustrates an important point: *the classifier cannot be relied upon to detect classes which lie outside the domain of the training data. It tries, but it does not always succeed.*

This is an important point about machine learning, and worth reinforcing with an imaginary example from human learning. Suppose a young child had seen lots of dogs, but never seen a wolf – not even a picture. When first presented with a picture of a wolf, the child will very likely proclaim 'Dog!' The child would probably do the same with a picture of a fox. Machine learners are no smarter, and in fact tend to be less able to distinguish outlying cases. When training a machine learner, it is important that the samples chosen for training represent all the classes that the learner will eventually be expected to distinguish.

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**Tutorial 6: Step 11 Display a Classification Plot**

**Display a Classification Plot**

1. If the Predictions item (or whatever you named it) in the Experiments navigator is not already highlighted, click it.

2. Select Classification Plot from the Predict menu, or right-click the item and select Classification Plot from the shortcut menu. The Classification Plot is displayed showing the predicted classes, the raw votes of the component classifiers and other information.

3. From the Comparison Variable drop-down list box in the upper right corner, select test classes. Some of the rectangles in the view turn red, signifying misclassifications.
Interpretation
This is a very rich display, and it may take some experience before you are able to interpret it easily.

Each row represents a sample. On the left of each row is a Sample name and Prediction or predicted class. The rest of the display consists of boxes representing the outputs of the artificial neural networks for each of the possible classes for that sample.

Each column represents a class. The colors of the boxes are significant:

- A box highlighted in dark green is the predicted class for that sample.
- A box highlighted in red is the true class of that sample if one is known. (See the discussion in Step 10 about observations of ‘Unknown’.) The class of a sample that has a dark green box and a red box has been predicted incorrectly. If the classifier predicts the sample class correctly, or if the correct value is not known, only a dark green box appears.
- A box that is colored gray represents neither the predicted class nor the true class.
- If GeneLinker™ refuses to make a prediction for a sample, it will have 'Unknown' listed under prediction and no dark green box.
- If the sample's true class is 'Unknown', it will not have a red box. (This will not happen when viewing training data since true classes must be known for all training samples.)

Hence the number of red boxes in the display indicates the number of misclassifications. Reducing the rate of misclassifications is discussed below.

Component Classifier Votes
Inside each box is a representation of the votes of each of the neural networks in the committee. Each of 10 neural networks was trained on a different 90% of the training data. Each of the horizontal rectangles in the view above represents the output of all 10
neural networks for a given class on a given sample. If all 10 neural networks are in agreement (i.e. have the same output value) then there will be a solid bar - at the right end if they all have high output (i.e. that is the sample's class), at the left end if they all have low output (i.e. that is not the sample's class).

Class Prediction Process
The class prediction (or call) is done by a simple vote. For a given sample, each neural network votes for the class with the highest output. If $2/3$ (default setting) of the networks agree on a single class, we call that a prediction. In any other case, no prediction is made and the sample is labelled 'Unknown'.

Example:
- Look at TEST-10 in the image above. Because $2/3$ of the neural networks could not agree on which class it was, 'Unknown' was entered as the prediction. However, there is more information about TEST-10 in the display than just its misclassification.
- Look at the outputs for class BL: the box in the second column. There is a solid gray bar at the left end of the histogram - this indicates that the ANN outputs for that class were uniformly zero. None of the neural networks gave any weight to classifying the sample as BL. Under class EWS, the results were almost the same: one or two ANNs gave a result only marginally greater than zero. In other words, the ANNs were unanimous that the sample did not fall into the BL or EWS classes.
- The ANN outputs for the other two classes are mixed - some ANNs voted for NB and some for RMS. In the context of the input genes, we conclude that the sample more nearly resembles RMS and NB than it does EWS or BL. In other words, the sample lies somewhere near the decision boundary between classes RMS and NB.
- As the red box indicates, the true class for this sample is RMS. Perhaps if we have set the voting threshold lower - around 50% - then the classifier would have made a prediction of RMS for this sample.
- The other sample which was not given a prediction (or predicted to be 'Unknown', if you wish) was TEST-11. Interestingly, TEST-11 was one of the five test samples which did not fall into the original four training classes. TEST-11 was a non-SRBCT cancer sample.

Reasons For Misclassifications:
There are often no misclassifications in the training data – artificial neural networks are fairly powerful and adaptable learners. If there are misclassifications, however, it may be for one of several possible reasons:
- We may be using a set of genes which do not discriminate between the sample classes.
- The training set may be unbalanced. That is, it may have too many examples of one class and not enough of another.
- We may have set the number of hidden units in the neural networks too small.
- The data may contain errors such as mislabelled samples or incorrect measurements.
• The voting threshold may be set too low.
• The stopping criteria may have been set too loose (maximum iterations too small).
The above reasons may affect either training or test results. If the training results are excellent but the test results are poor, it may be for one of the following additional reasons:
• The test data may be drawn from a significantly different population than the training data (such as the non-SRBCTs in the example above).
• The test data may not have been normalized in a similar fashion to the training data.
• The test dataset may have been filtered with different genes than the training dataset. (GeneLinker™ checks only that the number of genes used in training and prediction is the same, not their identities).
• We may have set the number of hidden units in the neural networks too large.
• We may have too many features (genes) for the number of samples in the training set.
• The stopping criteria may have been set too tight (maximum iterations too large).
These last three conditions correspond to a condition called ‘overtraining’. You can think of this as analogous to a child learning a certain set of examples by rote, but failing to be able to generalize from the examples to new cases. When a neural network is either given too much memory for detail (too many hidden nodes or input nodes) or is forced to learn the input examples too well (stopping criteria too tight), then it may simply ‘memorize’ the training data to the detriment of generalizing well on test data.

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Tutorial 6: Step 12 Set URL for Lookup Gene Operation

Set URL for Lookup Gene Operation
You can create different sets of genes and evaluate the discriminant power of each by training and testing a new classifier using each gene list. You might create these alternate gene lists by running SLAM™ longer, by choosing different genes from the SLAM™ output, or from your existing knowledge of which genes participate in a given process or disease state. One way to determine what is known about a gene is to use the Lookup Gene function of GeneLinker™. If you imported your expression data using GenBank or UniGene identifiers, you can look them up simply by choosing the Lookup Gene icon. It is enabled whenever you have a gene or a gene list selected.
If you don't have GenBank or UniGene identifiers associated with your expression data, you may still be able to look up genes directly from GeneLinker™. The dataset for this tutorial, for example, uses IMAGE Consortium clone ids. Steps 12 and 13 demonstrate how GeneLinker™ can look up genes via their clone ids.

1. Select Preferences from the Tools menu. The User Preferences dialog is displayed.
2. Click the **Gene Database** tab. The **User Preferences** dialog is updated.

3. Under **Lookup Gene Database URLs**, click in the text box next to **Custom**. The text in the box is highlighted.

4. Either:
   a) Use the right arrow key to move the cursor right until you see MMC_ID.
   b) Use the mouse to highlight MMC_ID.
   c) Type "**IMAGE:MMC_ID**" including the quotation marks.

   Or:
   a) Press <Delete>. The text box is cleared.
   b) Copy and paste the following URL into the text box (all on a single line):
   
   ```text
   ```
   
   • All that changes is that the string **MMC_ID** becomes "**IMAGE:MMC_ID**".
   
   **Note** the addition of the quotation marks.
• **Note** that the URL must remain on a single line. Any line break you see in the tutorial text is due to word wrap in the GeneLinker™ Help viewer. Be sure to type the URL in on a single line.

• The actual gene identifier (e.g. **207274**) is substituted for the sub-string **MMC_ID** when you perform a Lookup Gene operation on that gene.

5. Click **OK**.

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**Tutorial 6: Step 13 Lookup Genes**

**Lookup Gene 207274**

1. Click the Filtered: keep {Tutorial 6 list} dataset under the khan_training_data item in the **Experiments** navigator (created in Step 7: Filter Datasets Using a Gene List). The item is highlighted.

2. Click the **Color Matrix Plot** toolbar icon, or select **Color Matrix Plot** from the **Explore** menu, or right-click the item and select **Color Matrix Plot** from the shortcut menu. A color matrix plot of the dataset is displayed.

3. Click the gene 207274 (2nd from left). The gene is highlighted.

4. Click the **Lookup Gene** toolbar icon, or select **Lookup Gene** from the **Tools** menu. Your HTML browser is launched displaying the GenBank entry for the selected gene. IMAGE close 207274 is insulin-like growth factor II (human).
Tutorial 6: Conclusion

Conclusion
In this tutorial, you learned about the SLAM™ algorithm and how to use it to select a small set of genes (features) that can be used to train a committee of artificial neural networks (ANNs) to predict the classes of new samples. For further information, please see ANN Classification and Prediction Overview.

When you are finished, you can close all the open plots either by clicking on the 'x' box in the upper-right hand corner of each, or by selecting Close All from the Window menu.

References
Reference 1:

Where To Go From Here
• Go through the other tutorials.
• Read the Online Help to learn more about the various functions of GeneLinker™.
• Further explore GeneLinker™ by using additional features.
• Load up your favorite dataset and try out all the buttons and menu items.
• Don't forget to right-click on things like plots - many details of graphics can be customized.
• Visit the Molecular Mining website at http://www.molecularmining.com/ for the latest information on GeneLinker™ enhancements and additional products.

Tutorial 7: IBIS

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Tutorial 7: Introduction

Overview
IBIS (Integrated Bayesian Inference System) offers powerful search capabilities into your data. It can identify non-linear and combinatorial patterns of gene expression that characterize different toxicity responses, disease states, or treatment outcomes. Furthermore, it can be used to build classifiers that can identify these patterns in new samples.

IBIS is used most commonly as a search tool, to identify single genes and small gene sets that show interesting expression patterns relative to the sample classification. We will work through an example related to personalized medicine. We will attempt to identify patterns of basal gene expression that are predictive of drug response, using the NCI 60 data from the Developmental Therapeutics Program and the Genomics and Bioinformatics Group, both from the National Cancer Institute, National Institutes of Health. In this experiment 60 cancer cell lines from various tissues had their basal gene expression level measured. Each cell line was also exposed to a number of anti-cancer treatments, and the GI50 was measured. A valuable question to ask is whether the pre-treatment basal expression can be used to predict the effectiveness of a compound. This would provide a molecular basis for selecting appropriate therapies. IBIS can help to answer these types of questions by identifying gene expression patterns that are characteristic of effective or ineffective compounds.

IBIS has a number of different parameters that allow you to search for different types of biologically plausible relationships in the data. We will start with identifying simple but perhaps less predictive patterns, and introduce more effective models.

The simplest type of predictive gene expression patterns involve only a single gene, and are linear in nature. These patterns are often expressed as rules, such as when PSA levels are high, prostate cancer is likely. IBIS can be used to identify these types of patterns.

Tutorial Length
This tutorial should take about 45 minutes, depending on how long you spend investigating the data, and how fast your machine is.

If you must stop part way through the tutorial, simply exit the program by selecting Exit from the File menu. The data and experiments you have performed to that point are saved automatically by GeneLinker™. The next time you start GeneLinker™, you can
continue on with the next step in the tutorial.

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Tutorial 7: Step 1 Import the Data

Import the dataset NCI60_basal_expression.csv. This file contains the basal expression levels for 1041 genes in 60 cancer cell lines. The data are normalized log ratios.

Import the Data
1. Click the Import Gene Expression Data toolbar icon , or select Import from the File menu and Gene Expression Data from the sub menu. The Data Import dialog is displayed.

2. If the Template listed on the dialog is not Tabular, click the Template Change button, select Tabular and click Select. The Data Import dialog is updated with the Tabular template.
3. Ensure the Gene Database is set to GenBank. Use the drop-down list to set it if needed.
4. Click the Source File Change button. The Open dialog is displayed. If necessary, navigate to the tutorial folder.

5. Click the file NCI60_basal_expression.csv. The file name is highlighted. Click Open. The Data Import dialog is updated with the file name information.
6. Click **Import**. The **Import Data** dialog is displayed.

7. Since the data is already in the correct orientation, and GeneLinker™ has already identified the existence column header names, just click **OK**. The data is imported and a new item entitled **NCI_basal_expression** is added to the **Experiments** navigator.

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**Tutorial 7: Step 2 Import Variable Data**

**Overview**

Import the variable **NCI60_thiopurine_response.csv**. This file contains, for each cell line in the expression dataset, whether that cell line was inhibited by the application of thiopurine. We consider a cell line to be inhibited ('High Response') if its GI50 measurement is at least 10 times below the average, indicating a reasonable level of cell-line specific inhibition. Otherwise, the cell line is classed as 'Low Response'.
Actions

1. If the **NCI60_basal_expression** dataset item in the **Experiments** navigator is not already highlighted, click it.

2. Select **Import** from the **File** menu and **Variable** from the sub menu. The **Import Variables** dialog is displayed.

3. Click the **Source File ...** button. The **Open** dialog is displayed.

4. Click the file **NCI60_cmpd_response.csv**. The file name is highlighted. Click **Open**. The **Import Variables** dialog is updated with the **Source File** name and the number of observations and classes below it.
5. Type **Thiopurine** into the **Variable Name** field.

6. Click the **New Variable Type** button. The **Create Variable Type** dialog is displayed.

7. Type **High/Low** into the **Name** field. Click **OK**. The **Import Variables** dialog is updated.
8. The **Preview** allows you to view which sample belongs to which class and the total number of entries for each class. Click **Preview**. When you are finished examining the contents of the Preview, click **Close** to close it.

9. Click **Import**. The variable information is imported and the NCI60 basal expression dataset item in the **Experiments** navigator is tagged with the variables icon.

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**Tutorial 7: Step 3 Perform IBIS 1D LDA Search**

**Overview**

Perform an IBIS Linear Discriminant Analysis (LDA) search. This search should be relatively quick. The IBIS search process evaluates the accuracy of each gene (in the 1D case) when used as a linear discriminator.

A discriminator is a feature that distinguishes between classes. A linear discriminator can be thought of as a straight line drawn between classes. For example, when two football teams line up for the kickoff at the start of the game, they can be separated by a straight line at center field. After play begins, however, there is not likely to be any straight line which can be drawn that is likely to have all the players from one team on one side and the other team on the other.

Occasionally there may be a simple curved line which can be drawn between the players – or the classes. A quadratic discriminator and a Gaussian discriminator are two simple types of discriminators which can yield curved lines.
Actions

1. If the NCI60_basal_expression dataset item in the Experiments navigator is not already highlighted, click it.

2. Select IBIS Classifier Search from the Predict menu, or right-click the item and select IBIS Classifier Search from the shortcut menu. The IBIS Classifier Search dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Representative Variable</td>
<td>Thiopurine</td>
<td>Training variable.</td>
</tr>
<tr>
<td>Classifier Type</td>
<td>Linear</td>
<td>Linear, Quadratic, or Uniform/Gaussian.</td>
</tr>
<tr>
<td>Dimension</td>
<td>1 (singleton gene)</td>
<td>1D or 2D</td>
</tr>
<tr>
<td>Minimum Standard Deviation</td>
<td>0.1</td>
<td>Use the minimum standard deviation to capture your estimate of the error in the measurements. With too small a value, you will find degenerate looking patterns that are not believable. With too large a value, you risk missing important patterns due to over-smoothing the classifier.</td>
</tr>
<tr>
<td>Committee Size</td>
<td>60</td>
<td>Number of component classifiers in the IBIS classifier.</td>
</tr>
<tr>
<td>Committee Votes Required</td>
<td>40 of 60 (66%)</td>
<td>Threshold for making a class prediction.</td>
</tr>
<tr>
<td>Random Seed</td>
<td>999</td>
<td>Initial value for the random number generator.</td>
</tr>
</tbody>
</table>

4. Click OK. The IBIS LDA search is performed and a new item Thiopurine IBIS search LDA 1D is added to the Experiments navigator under the original dataset.

   If you have automatic visualizations enabled in your user preferences, the IBIS Search Results Viewer is displayed.
Overview
The IBIS Search Results Viewer consists of a 3-column listing. The first column contains gene identifiers, the second contains MSE values, and the third contains accuracy figures. The results are initially sorted by the accuracy values. Both the MSE and accuracy values are indications of the ability of the classifier (gene) to separate the inhibited cell lines from the uninhibited cell lines, given the treatment of Compound A. The MSE values reflect the level to which the data matched the linear model, with lower values being better, while the accuracy values reflect the predictive accuracy of a linear model in separating the inhibited from uninhibited cell lines. When comparing two genes that have the same accuracy value, the one with the lower MSE is generally to be preferred.

Actions
If the IBIS Search Results Viewer is already displayed, skip to #2 below the image.
1. Double-click the Thiopurine IBIS search LDA 1D item in the Experiments navigator (or click the item and select IBIS Search Results from the Predict menu). The item is highlighted and the IBIS Search Results Viewer is displayed.

2. Click the MSE column header. The genes or proto-classifiers are re-sorted according to their mean-squared error.

3. Click the Accuracy header. The genes are once again sorted by their accuracy as
classifiers.

Discussion
The IBIS Search Results Viewer has three columns of information. The first column contains gene identifiers, the second contains cross-validation accuracy scores, and the third contains mean squared error (MSE) values. The results are initially sorted by accuracy. Both the MSE and accuracy values are indications of the ability of the classifier (gene) to separate the high-response samples (cell lines) from the low response samples. The MSE values reflect how well the data match the linear model, with lower values being better. Accuracy values reflect the predictive accuracy of a linear model in separating the high responses from low responses.

When comparing two genes that have the same accuracy value, the one with the lower MSE is generally to be preferred. You will find, though, that accuracy and MSE tend to be highly correlated, a high accuracy generally indicating a low MSE and vice versa.

Let us examine the top gene AA046755, which has an accuracy of 82% and an MSE of 0.18. We will display the actual gene expression measurements for this gene superimposed on the output of the IBIS linear classifier to get a sense of which samples are correctly and incorrectly classified.

Platinum
Tutorial 7: Step 5 Display IBIS Gradient Plot

Actions
1. Click the top gene (AA046755) in the IBIS Search Results Viewer. The gene is highlighted.
2. Click Gradient Plot. The Classifier Gradient Plot is displayed.

Discussion
In this plot, we see three areas. The most important area for now is the scatter plot in
The left-right position of each point on the plot represents the expression level of gene AA046755 in one of the 60 cell lines. Because this is 1D IBIS, only one dimension of the plot is meaningful: The horizontal axis. The height of each point is assigned randomly to minimize visual overlap, so be careful not to impute any meaning to the vertical position of the points.

Each point is colored according to the cell line’s observed response to thiopurine, as shown in the legend at the bottom left. The background of the scatter plot is a color gradient that corresponds to the IBIS classifier’s prediction, in the same basic color scheme as the point coloring. We can see which samples are incorrectly classified by comparing the color of the points to the color of the background. We can see that down-regulation of AA046755 (negative values) occur more frequently with high response to thiopurine. The line where high response crosses over to low response – where blue crosses over to red – is at about a log ratio of -1.

When we imagine the complexity of a cell’s response to a treatment, it is unsurprising that we cannot achieve perfect separation using a single gene and a linear classifier. IBIS allows you to explore relationships between pairs of genes, and to use non-linear classifiers to identify patterns.

Platinum

Tutorial 7: Step 6 Perform IBIS 2D LDA Search

Overview
Perform an IBIS 2-dimensional search over gene pairs. A 2-dimensional search takes longer than the 1-dimensional search performed previously. IBIS can examine every possible pair of genes in the dataset (1041 * 1040 / 2 = 541320 pairs) and evaluate the MSE and accuracy of each classifier (gene pair) on that data.

For the purposes of this tutorial, we will use the 1D IBIS results to filter down the number of genes that will be searched by 2D IBIS. However, if we were to simply choose the best 1D classification genes, we would expect that two-dimensional combinations of them would also produce fairly good classification just because the individual genes were already fairly good. So instead we shall use 2D IBIS to examine the genes that are not good 1D predictors, to see if there are cases where combinatorial effects are prominent.

Actions

1. Click the IBIS Search Results: 1D LDA window to bring it to the front. If you closed the window, double-click on the Thiopurine IBIS search results item in the Experiments navigator. The IBIS Results Viewer is displayed.
2. Ensure that the results are sorted by **Accuracy** (the default).

3. Click the checkbox to the left of the top gene (AA046755) so that it is checked.

4. Scroll down until accuracy values of 67% and 65% are visible. Press and hold the <Shift> key and click the checkbox to the left of the last gene with an accuracy of 67% (H26883). This checks every gene from the top gene down to this one.

5. Click the **Create Gene List** button. The **Create a Gene List** dialog is displayed.

6. For the **Name**, type in **>67% accuracy**.

7. Click **Save**. A gene list is created and added to the **Gene Lists** navigator.

7. Click the NCI60 basal expression dataset item in the **Experiments** navigator. The item is highlighted.

8. Click the **Filter Genes** toolbar icon, or select **Filter Genes** from the **Data** menu, or right-click the item and select **Filter Genes** from the shortcut menu. The **Filter Genes** dialog is displayed.
9. Select **Gene List Filtering** from the **Filtering Operation** drop-down list.

10. Select **Remove all genes that are in this list**.

11. Select the gene list > 67% accuracy from the **Gene List** drop-down list.

12. Click **OK**. A new Filtered: removed {> 67% accuracy} dataset is added to the **Experiments** navigator. It contains the 110 genes which had less than 67% accuracy as 1D linear predictors of thiopurine response.

13. If the new Filtered: removed {> 67% accuracy} dataset in the **Experiments** navigator is not already highlighted, click it.

14. Click the **IBIS Classifier Search** toolbar icon, or select **IBIS Classifier Search** from the **Predict** menu, or right-click the item and select **IBIS Classifier Search** from the shortcut menu. The menu. The **IBIS Classifier Search** dialog is displayed.

---

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Setting</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Representative Variable</strong></td>
<td>Thiopurine</td>
</tr>
<tr>
<td><strong>Classifier Type</strong></td>
<td>Linear</td>
</tr>
<tr>
<td><strong>Dimension</strong></td>
<td>2 (gene pairs)</td>
</tr>
<tr>
<td><strong>Minimum Standard Deviation</strong></td>
<td>0.1</td>
</tr>
<tr>
<td><strong>Committee Size</strong></td>
<td>60</td>
</tr>
<tr>
<td><strong>Committee Votes Required</strong></td>
<td>40 of 60 (66%)</td>
</tr>
<tr>
<td><strong>Random Seed</strong></td>
<td>999</td>
</tr>
</tbody>
</table>
4. Click **OK**. The IBIS 2D LDA search is performed and a new item IBIS Search Results LDA 2D is added to the **Experiments** navigator under the original dataset. This typically takes 5 to 10 minutes depending on the speed and load of your machine.

If you have automatic visualization enabled in your user preferences, the IBIS Search Results Viewer is displayed.

**Platinum**

**Tutorial 7: Step 7 View IBIS 2D LDA Search Results**

**Overview**

This plot is similar to the one for the 1D results seen earlier. The changes are in the **Genes** column, where instead of having single genes, each entry is a pair of genes. Also, there is a new **Genes** list box at the right, allowing you to view and sort the unique genes found in multiple proto-classifiers.

**Actions**

If the IBIS Search Results Viewer is already displayed, skip to #2 below the image.

1. Double-click the **IBIS Search Results LDA 2D** item in the **Experiments** navigator. The item is highlighted and the **IBIS Results Viewer** is displayed.

**Discussion**

In the IBIS 2D LDA results we see that our accuracy values range as high as 83%. So
using genes which were filtered so as to omit the best individual genes, we can still obtain classification accuracies comparable to those obtained with single genes, which were in this case as high as 83%. This highlights the potential of combinatorial classifiers and predictors.

Platinum

Tutorial 7: Step 8 Display IBIS Gradient Plot

Overview
This plot is similar to the one shown for the 1D LDA results, except that now two genes are used in the scatter plot. The vertical dimension signifies the expression level of one of the genes, rather than random jitter. Furthermore the gradient behind the scatter plot now reflects the two dimensional nature of the classification pattern. We shall examine a gene pair with an easily interpreted pattern.

Actions
1. Click the MSE column header in the IBIS Results Viewer. The search results are sorted by mean square error.
2. Click the top item, the gene pair H59368 and W51913 with accuracy 78% and MSE 0.1657. The item is highlighted.
3. Click Gradient Plot. The IBIS Gradient Plot is displayed.

Discussion
This gene pair depicts an AND relationship: If basal expression of W51913 is low AND basal expression of H59368 is high, then response to thiopurine tends to be high (blue). This rule has 78% accuracy, determined by leave-one-out cross-validation which was the result of setting the number of committees equal to the number of samples. Furthermore, since the genes involved were not individual predictors with >67% accuracy, the predictive power of this relationship is at least partly a combinatorial
effect: One cannot get the same result by considering the genes independent of one another.

**Platinum**

**Tutorial 7: Conclusion**

In general, it is best to start identifying simpler patterns in the data first. This usually means using IBIS with single genes and Linear Discriminant Analysis (LDA) to begin with. Only if the accuracy or MSE values are unsatisfactory should you try Quadratic Discriminant Analysis (QDA) and Uniform/Gaussian Discriminant Analysis (UGDA) as well as gene pairs.

Remember that single gene IBIS searches are always relatively quick, even for tens of thousands of genes. However, when looking for patterns over gene pairs, the run time will be multiplied by the number of genes in the dataset again. For instance, if running 1D IBIS took 1 minute on 500 genes, then 2D IBIS will take about 500 minutes (8 hours) on the same data. Effective filtering of genes is an important step to make gene pair searches practical.

Use the minimum standard deviation to capture your estimate of the error in the measurements. With too small a value, you will find degenerate looking patterns that are not believable. With too large a value, you risk missing important patterns due to over-smoothing the classifier.

When you are finished, you can close all the open plots either by clicking on the 'x' box in the upper-right hand corner of each, or by selecting Close All from the Window menu.

**Where To Go From Here**

- Go through the other tutorials.
- Read the Online Help to learn more about the various functions of GeneLinker™.
- Further explore GeneLinker™ by using additional features.
- Load up your favorite dataset and try out all the buttons and menu items.
- Don't forget to right-click on things like plots - many details of graphics can be customized.

**Tutorial 7: Appendix: Minimum Standard Deviation in IBIS**

This appendix describes the choice and effect of the Minimum Standard Deviation parameter in IBIS.

**Minimum Standard Deviation Too Small**

In some datasets IBIS will find patterns like the one shown below:
The points for the class colored red nearly fall all on one straight line. If too small a value is chosen for the Minimum Standard Deviation, QDA or UGDA IBIS will create very narrow region covering those points and compute a very high accuracy.

However, the likelihood that such a classifier reflects biological reality is exceedingly small if the width of the class region is smaller than the random variation in gene expression inherent in the system.

Similarly an LDA classifier could compute an unrealistically high accuracy by forming a class boundary between samples which are separated by less than the natural random variation in expression in the genes.

**Minimum Standard Deviation Too Large**

On the other hand significant effects can be obscured by setting the Minimum Standard Deviation too large. Consider the same dataset as depicted above, only this time with a larger Minimum Standard Deviation.

It is reasonable to suppose that the pattern here might be significant (up to the limitations of the number of samples). But as the Minimum Standard Deviation is increased, the region predicted as ‘red’ gets increasingly broad – and eventually circular – until the legitimate linear correlation between the two genes for the red class samples is lost. At the same time, the accuracy score for these genes as predictors goes down rapidly, as the broadening of the prediction region takes in more and more blue samples. Therefore setting the Minimum Standard Deviation much larger than the natural variation in the expression values can result in real patterns going undetected.

**Default Value**

GeneLinker computes a suggested Minimum Standard Deviation each time the IBIS Classifier Search dialog box is opened. The suggested or default value is computed from a random sample of the data, and so the number may be different each time.
Because the Minimum Standard Deviation only has an effect in rare cases, and because the random variation in the default value is small, it is not usually necessary to change the default value. If you believe you have a case like one of those described above you may wish to use a fixed estimate of the standard deviation for all IBIS runs. You may also wish to try several different values to see what effect they have on the classification accuracy and Mean Squared Error.

Tutorial 8: Affymetrix Data

Tutorial 8: Introduction

This tutorial leads you through the process of importing and performing experiments on Affymetrix MAS 5.0 data.

Skills You Will Learn:
How to import Affymetrix MAS 5.0 gene expression data into the GeneLinker™ database.
How to import a gene list.
How to set the gene display name.
How to import a variable (class labels).
How to remove genes by reliability measure.
How to estimate missing values.
How to perform an F-Test and view the results.
How to create a gene list.
How to perform gene list filtering.
How to perform a hierarchical clustering or a principal component analysis experiment.
How to display and manipulate a matrix tree plot and a 3D score plot.

Dataset Information

Tutorial Length
This tutorial should take about 20 minutes, depending on how long you spend investigating the data, and how fast your machine is. Note that if you must stop part way through the tutorial, exit the program by selecting Exit from the File menu. The data and experiments you have performed to that point will be saved automatically by the application. The next time you start GeneLinker™, you can continue on with the next step in the tutorial.

Tutorial 8: Step 1 Import Affymetrix Data
1. Click the **Import Gene Expression Data** toolbar icon, or select **Import** from the **File** menu and **Gene Expression Data** from the sub menu. The **Data Import** dialog is displayed.

2. Click the button next to the **Template**. The **Import Templates** dialog is displayed.

3. Click **Affymetrix 5.0**.

4. Click **Select**. The **Data Import** dialog is updated with the new template and the dialog changes conformation to support importing from multiple data files in a single folder.

5. Set the **Gene Database** to **Affymetrix** by selecting it from the drop-down list.

6. The **Source Folder** by default is the Tutorial folder. Click the ... button to the right. This displays the **Open** dialog.
7. Click the **Affymetrix** folder. The folder name is highlighted.

8. Click **Select Folder**. The **Data Import** dialog is updated with the new **Source Folder**.

9. In the **Source Files** list, click the file **Chip1.txt**. The file is highlighted.

10. Click the **right arrow** button at the top between the **Source Files** and the **Import Files** lists. The Chip1.txt file is transferred into the **Import Files** list.

11. Click the **right arrow** button five more times to transfer the next five files into the **Import Files** list.
Each data file contains gene expression values for one sample. The files are imported from top to bottom, with the top file becoming the first sample in the dataset, the second file becoming the second sample, and so on to the last. This means that it is essential that the files listed in the Import Files list be in sample order before you click Import. The buttons to the right of the Import Files list can be used to reorder the files. In this tutorial it is not necessary to do this since the files are already in the correct order for import.

12. Click Import. After several seconds, the Import Data dialog is displayed.
Within GeneLinker™, datasets have the genes in columns and the samples in rows.

**Note:** the options **Use Sample Names** and **Use Gene Names** are checked and disabled in the **Import Data** dialog. GeneLinker™ has recognized that in this dataset, the first values are alphameric gene labels. Gene expression data is always numeric, hence the disabled checkboxes. GeneLinker™ has derived the sample names from the sample data files names.

13. Click **OK**. The data is imported and a new dataset (Chip 1) is added to the **Experiments** navigator.

![Chip 1 dataset](image)

The dataset name is derived from the first sample file name. If you like, you can rename the dataset by right-clicking on the dataset in the navigator, selecting **Rename Experiment** from the shortcut menu and typing in a new name.

**Tutorial 8: Step 2 Import Gene List**

A gene list is imported to bring in additional meta-data about the genes in the dataset.

1. Click the Chip1 dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Table View** toolbar icon, or right-click the dataset and select **Table View** from the shortcut menu. A table view of the dataset is displayed.

![Table View](image)

3. On the table viewer, move the mouse pointer until it is on the border between the first and second gene names. The pointer becomes a two-headed arrow. Click and drag to the right to widen the columns in the table until the gene names are completely displayed. Click on the fourth gene (AFFX-MurIL2_at).
4. Look at the **Description pane** in the lower left corner of the window. It displays the information about the gene that is currently in the database.

5. Close the table view by clicking the icon in its upper right corner.

6. Select **Import** from the **File** menu and **Gene List** from the sub menu. The **Open** dialog is displayed.

7. Double-click the **Affymetrix** folder.

8. Set the **Files of type** to **All Files (*.*)**.

9. Click the file **Hum-U95a.csv**.

10. Click **Open**. The **Import Gene List** dialog is displayed.

11. The **Gene Database** is correctly set to **Affymetrix**, so all you need to do is click **OK**. The gene list is imported and is added to the **Gene Lists** navigator.

12. Click the **Experiments** tab in the navigator. The **Experiments** navigator is displayed.

13. Click the Chip1 dataset in the **Experiments** navigator. The item is highlighted.

14. Click the **Table View** toolbar icon, or right-click the dataset and select **Table View**.
from the shortcut menu. A table view of the dataset is displayed.

<table>
<thead>
<tr>
<th></th>
<th>AFFX-MurFAS_at</th>
<th>AFFX-MurFAS_at</th>
<th>AFFX-MurFAS_at</th>
<th>AFFX-MurFAS_at</th>
<th>AFFX-MurFAS_at</th>
<th>AFFX-MurFAS_at</th>
<th>AFFX-MurFAS_at</th>
<th>AFFX-MurFAS_at</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chip1</td>
<td>2.9</td>
<td>2.5</td>
<td>2.4</td>
<td>19.0</td>
<td>106.8</td>
<td>666.6</td>
<td>723.5</td>
<td>246.8</td>
</tr>
<tr>
<td>Chip2</td>
<td>7.9</td>
<td>7.5</td>
<td>5.6</td>
<td>40.8</td>
<td>161.6</td>
<td>464.4</td>
<td>102.2</td>
<td></td>
</tr>
<tr>
<td>Chip3</td>
<td>7.8</td>
<td>2.4</td>
<td>0.8</td>
<td>7.5</td>
<td>89.3</td>
<td>140.7</td>
<td>88.1</td>
<td>458.3</td>
</tr>
<tr>
<td>Chip4</td>
<td>7.3</td>
<td>5.8</td>
<td>0.7</td>
<td>23.7</td>
<td>106.8</td>
<td>228.8</td>
<td>116.8</td>
<td>389.8</td>
</tr>
<tr>
<td>Chip5</td>
<td>7.7</td>
<td>1.6</td>
<td>0.8</td>
<td>11.4</td>
<td>40.3</td>
<td>158.4</td>
<td>72.6</td>
<td>122.1</td>
</tr>
<tr>
<td>Chip6</td>
<td>6.5</td>
<td>7</td>
<td>9.8</td>
<td>12.9</td>
<td>191.2</td>
<td>194.9</td>
<td>75.9</td>
<td>201.9</td>
</tr>
</tbody>
</table>

15. Click on the fourth gene name (AFFX-MurFAS_at) on the table view. The gene is highlighted.

16. Look at the **Description Pane**. The information about the gene that was in the gene list has been added to the database.

![Description Pane](image)

**Tnfrsf6**

*Description*: Tumor necrosis factor receptor superfamily, member 6

*Annotations*: 0

*Created*: 2002-11-26 19:37:28

---

**Tutorial 8: Step 3 Set Gene Display Name**

1. Select **Preferences** from the **Tools** menu. The **User Preferences** dialog is displayed.

![User Preferences](image)

2. Click the **Gene Database** tab. The **Gene Database** pane is displayed.
3. Set the **Gene Display Name** to **Gene Name**.
4. Click **OK**. Your preferences are updated.

**Tutorial 8: Step 4 Import a Variable**

1. If the Chip1 dataset in the **Experiments** navigator is not already highlighted, click it.
2. Select **Import** from the **File** menu and **Variable** from the sub menu. The **Import Variables** dialog is displayed.

- The **Dataset** is set to **Chip1**. The number of samples it contains is listed below it.
3. Click the **Source File** button. The **Open** dialog is displayed.

4. Double-click the **Affymetrix** folder. The files in the Affymetrix folder are displayed.

5. Click the file **affy_var.txt**. The file is highlighted.

6. Click **Open**.
   - The **Source File** name is displayed with its number of observations and classes listed below.
   - The default **Variable Name** and **Description** are displayed.

7. The **Preview** allows you to view which sample belongs to which class and the total number of entries for each class. Click **Preview**. When you are finished examining the contents of the Preview, click **Close** to close it.

8. Enter **Affy Variable** in the **Variable Name** text box.

9. Optionally, enter a new description for the variable in the **Description** text box.

10. Click the **New Variable Type** button. The **Create Variable Type** dialog is displayed.
11. Enter **Affy Example** for the **Name** of the new variable type, and optionally a **Description**.

12. Click **OK**. The new variable type is displayed in the **Choose a Variable Type** list on the **Import Variables** dialog.

13. Click **Import**. The variable data is imported into the database, and in the **Experiments** navigator, the Chip1 dataset icon is marked with the variable tag.

**Tutorial 8: Step 5 Remove Genes With Poor Reliability**

1. If the Chip1 dataset in the **Experiments** navigator is not already highlighted, click it.

2. Select **Remove Values** from the **Data** menu, or right-click the dataset in the navigator and select **Remove Values** from the shortcut menu. The **Remove Values** dialog is displayed.
3. Select **By Reliability Measure** for the **Removal Technique**. The dialog is updated.

4. Set the **Reliability Measure** threshold to **0.101** by moving the slider or using the arrow keys on your keyboard.

5. Click **OK**. The operation is performed, and upon successful completion, a new Removed: $p > 0.101$ incomplete dataset is added to the **Experiments** navigator.

**Tutorial 8: Step 6 Estimate Missing Values**

1. If the new incomplete Removed: $p > 0.101$ dataset in the **Experiments** navigator is not already highlighted, click it.

2. Click the **Estimate Missing Values** toolbar icon, or select **Estimate Missing Values** from the **Data** menu. The **Estimate Missing Values** dialog is displayed.
3. Set the **Remove Genes That Have Missing Values** threshold to 2.
4. Click the radio button next to **Nearest Neighbors** in the **Replacement Technique** group.

The default distance metric **Euclidean** is correct.

5. Set the **Number of Nearest Neighbors** to 5.
6. Click **OK**. The operation is performed, and upon completion, a new complete Estimated: #mv <2 | nn=5 | Euclid dataset is added to the **Experiments** navigator.
Tutorial 8: Step 7 Perform F-Test and View Results

1. If the new complete Estimated: #mv <2 | nn=5 | Euclid dataset in the Experiments navigator is not already highlighted, click it.
2. Select ANOVA from the Statistics menu. The ANOVA dialog is displayed.

3. Click OK. The F-Test is performed and a new F-Test: affy_var dataset is added to the Experiments navigator.
4. If you have automatic visualizations enabled in the user preferences, the ANOVA Viewer is displayed. If not, double-click the new F-Test: affy_var dataset in the Experiments navigator to display the ANOVA Viewer.

In step 3 of this tutorial, you set the gene display name to gene name in your user preferences. The gene names are what you currently see in the ANOVA Viewer. In this step you will change the gene display name setting to see Affymetrix gene identifiers displayed in the ANOVA Viewer.

5. Click the icon in the upper right corner of the ANOVA Viewer to close it.
6. Select **Preferences** from the **Tools** menu. The **User Preferences** dialog is displayed.

7. Click the **Gene Database** tab. The **Gene Database** pane is displayed.

8. Set the **Gene Display Name** to **Affymetrix**.

9. Click **OK**. Your preferences are updated.

10. Double-click the new F-Test: affy_var dataset in the **Experiments** navigator. The **ANOVA Viewer** is displayed.
11. Click the first gene checkbox. The gene is highlighted and a checkmark appears in the checkbox.

12. Press and hold down the <Shift> key and scroll down until you see the p-value 0.0497 (gene 34378_at).

13. Click the checkbox for gene 34378_at. All the genes from the first to that gene are highlighted and checked. Release the <Shift> key.

14. Click Create Gene List. The Create Gene List dialog is displayed.
15. Type **Affy Gene List** into the **Name** text box. Optionally, you may type in a description.
16. Click **Save**. The gene list is created and is added to the **Gene Lists** navigator.
17. Click the icon in the upper right corner of the ANOVA Viewer to close it.

**Tutorial 8: Step 8 Gene List Filtering**

1. Click the Estimated: #mv <2 | nn=5 | Euclid dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Filter Genes** toolbar icon , or select **Filter Genes** from the **Data** menu. The **Filter Genes** dialog is displayed.
3. Select **Gene List Filtering** from the **Filtering Operation** drop-down list. The option **Keep only genes that are in this list** is selected by default. This is correct for this tutorial.
4. Select the gene list Affy Gene List from the **Gene List** drop down list.
5. Click **OK**. The filtering operation is performed, and upon successful completion, a new Filtered: keep {Affy Gene List} dataset is added to the **Experiments** navigator.

**Tutorial 8: Step 9 Hierarchical Clustering**

1. If the new Filtered: keep {Affy Gene List} dataset in the **Experiments** navigator is not already highlighted, click it.
2. Click the **Hierarchical Clustering** toolbar icon , or select **Hierarchical Clustering**
from the **Clustering** menu. The **Hierarchical Clustering** dialog is displayed.

![Hierarchical Clustering dialog](image)

3. The default values are correct, so just click **OK**. The hierarchical clustering operation is performed, and upon successful completion, a new Hier: genes | Euclid | average experiment is added to the **Experiments** navigator.

If automatic visualizations are enabled in your user preferences, a matrix tree plot is displayed.

**Tutorial 8: Step 10 Display Matrix Tree Plot**

If the matrix tree plot is already displayed, skip to #2.

1. Double-click the Hier: genes | Euclid | average experiment in the **Experiments** navigator. The item is highlighted and a matrix tree plot is displayed.

![Matrix tree plot](image)

2. Click the **Color by Variable** button 🌈. Blocks of color are displayed to the right of the
sample names colored according to the class of each sample.

3. Click the first gene on the plot. The gene is highlighted. Look at the Description Pane. Information about the gene is displayed.

4. Click the icon in the upper right corner of the plot to close it.

Tutorial 8: Step 11 Principal Component Analysis

1. Click the Filtered: keep {Affy Gene List} dataset in the Experiments navigator. The item is highlighted.

2. Click the Principal Component Analysis toolbar icon , or select Principal Component Analysis from the PCA menu. The PCA dialog is displayed.

3. The orientation is set to Genes by default, so just click OK. The PCA operation is...
performed, and upon successful completion, a new PCA: genes experiment is added to the **Experiments** navigator.

If you have automatic visualizations enabled in your user preferences, a 3D score plot is displayed.

**Tutorial 8: Step 12 Display 3D Score Plot**

If the 3D Score Plot is already displayed, skip to #2.

1. Double-click the PCA: genes experiment in the **Experiments** navigator. The item is highlighted and a 3D score plot is displayed.

2. Click the **Color by Variable** button. The points on the plot are colored by their respective classes.
Tutorial 8: Conclusion

In this tutorial you learned how to import Affymetrix MAS 5.0 gene expression data, a gene list and variable data into the GeneLinker™ database. Next, genes were removed by reliability measure and missing values were estimated. You performed an F-test, viewed the results, created a gene list and performed gene list filtering. Finally, you performed a hierarchical clustering and a principal component analysis experiment and viewed the results in appropriate 2D and 3D plots.

Where To Go From Here

- Go through the other tutorials.
- Read the Online Help to learn more about the various functions of GeneLinker™.
- Further explore GeneLinker™ by using additional features.
- Load up your favorite dataset and try out all the buttons and menu items.
- Don't forget to right-click on things like plots - many details of graphics can be customized.
- Visit the Molecular Mining website at http://www.molecularmining.com/ for the latest information on GeneLinker™ enhancements and additional products.
Sample Workflow Using Spotted Array N-Fold Culling With Log Transformation

Overview
This workflow is used for ratio (Cy3/Cy5) data to filter out genes that do not show a large induction or repression in any sample in the dataset, and then to log normalize the data so that inductions and repressions have equal but opposite sign. You must specify the value for the N-fold filtering operation. For example, if you specify 2, then genes that show a value of 2 or greater (induction) or a value of 1/2 or less (repression) remain in the dataset after filtering. This operation discards genes that do not show significant expression changes. Following filtering, a log normalization operation is used to give inductions and repressions equal but opposite sign. In our example above, log2 2 = 1 and log2 1/2 = -1.

Note: selecting a value of less than or equal to 0.0 is not allowed.

Actions
1. Click the Perou dataset in the Experiments navigator (if the Perou item is not there, import the Perou dataset). The item is highlighted. Note that the Description Pane (under the Navigator) reports the number of genes/samples (approximately 5600 genes in this example).
2. Click the Filter toolbar icon, or select Filter Genes from the Data menu, or right-click the item and select Filter Genes from the shortcut menu. The Filter Genes parameters dialog is displayed.
3. Select Spotted Array N-Fold Culling from the Filtering Operation drop-down list.
4. Set the Induction/repression threshold to 3.0.
5. Click OK. The Experiment Progress dialog is displayed.

The dialog is dynamically updated as the filtering operation is performed. Upon successful completion, a new filtered dataset is added to the Experiments navigator pane under the original dataset.
• Setting the threshold value to 3.0 in this example reduces the number of genes down to approximately 460.

6. Click the filtered dataset in the Experiments navigator. The dataset is highlighted.

7. Click the Normalize toolbar icon ☰, or select Normalize from the Data menu, or right-click the item and select Normalize from the shortcut menu. The first Normalization parameters dialog is displayed.

8. Double-click the Logarithm radio button, or ensure Logarithm is selected and click Next. The second Normalization dialog is displayed.

9. Double-click the base 2 radio button, or ensure base 2 is selected and click Finish. The normalization operation is performed and upon successful completion, a new normalization item is added to the Experiments navigator pane under the filtered dataset.

10. At this point you can try applying Hierarchical or K-Means partitional clustering on
the data. (Right-click the item in the **Experiments** navigator and make selections from the shortcut menu.)

**Related Topics:**
- Performing Agglomerative Hierarchical Clustering
- Performing K-Means Clustering
Using GeneLinker(TM)

Installing GeneLinker™

Getting Started With GeneLinker™

Using GeneLinker™

| How to install, upgrade, or uninstall GeneLinker™. | Product Tour and comprehensive Tutorials. | Detailed descriptive and procedural topics. |

How to Find Information

- Display the Main Program Functions List and follow the links.
- Expand the chapters in the table of contents to display specific topics.
- Type in or search for a keyword in the index.
- Troubleshooting and Technical Support.

This manual applies to both the Gold and Platinum versions of GeneLinker™. See General Formatting Conventions for version identification information.

Demonstration Versions

The demonstration version of GeneLinker™ Gold and GeneLinker™ Platinum gives you access to all of the powerful functionality of the purchased version.

- The only limitation of a demonstration version compared to a purchased version is that demonstration licenses expire after a short time.
- Run through all of the tutorials (tutorial 6 Learning to Distinguish Cancer Classes and tutorial 7 IBIS Classification are only available in GeneLinker™ Platinum).
- Please contact our sales staff for a demonstration or pricing information. We would love to hear from you!

Molecular Mining Corporation
(617) 547-6373
or send an email to:
sales@molecularmining.com
### Main Program Functions List

<table>
<thead>
<tr>
<th>Data</th>
<th>Preprocessing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Importing Gene Expression Data</td>
<td>Eliminating &amp; Estimating Missing</td>
</tr>
<tr>
<td>Variables</td>
<td>Values</td>
</tr>
<tr>
<td>Genes and Gene Lists</td>
<td>Filtering</td>
</tr>
<tr>
<td>Exporting Data</td>
<td>Normalization</td>
</tr>
<tr>
<td></td>
<td>Value Removal</td>
</tr>
<tr>
<td>Clustering</td>
<td>Other Functions</td>
</tr>
<tr>
<td>K-Means Clustering</td>
<td>Principal Components Analysis</td>
</tr>
<tr>
<td>Jarvis-Patrick Clustering</td>
<td>3D Score Plot</td>
</tr>
<tr>
<td>Agglomerative Hierarchical Clustering</td>
<td>Annotations</td>
</tr>
<tr>
<td>Self-Organizing Maps (SOMs)</td>
<td>Generating Reports</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>Plots</td>
<td>Plot Functions</td>
</tr>
<tr>
<td>Matrix Tree Plot</td>
<td>Shared Selection</td>
</tr>
<tr>
<td>Centroid Plot</td>
<td>Profile Matching</td>
</tr>
<tr>
<td>Summary Statistics Chart</td>
<td>Color By Gene Lists or Variables</td>
</tr>
<tr>
<td>SOM Plot</td>
<td>Exporting Images</td>
</tr>
</tbody>
</table>

| Platinum SLAM                             | Platinum IBIS                      |
| ANN Classification & Prediction Overview   | IBIS Overview                      |
| SLAM Association Viewer                   | IBIS Search                        |
| Classification Plot                       | IBIS Gradient Plot                 |

### About GeneLinker and This Manual

**Acknowledgements**

This product includes software developed by the Apache Software Foundation [http://www.apache.org/](http://www.apache.org/).

- The complete license is available in MMC/GeneLinker Gold/ApacheLicense.txt for GeneLinker™ Gold and in MMC/GeneLinker Platinum/ApacheLicense.txt for GeneLinker™ Platinum.

This product also includes Sitraka's JClass product. Sitraka can be found on the web at: [http://www.sitraka.com/software/jclass/](http://www.sitraka.com/software/jclass/).

- The complete license is available in MMC/GeneLinker Gold/JClassLicense.txt for GeneLinker™ Gold and in MMC/GeneLinker Platinum/JClassLicense.txt for GeneLinker™ Platinum.
As part of our compliance with the MySQL license agreement, the source for MySQL has been included on the GeneLinker™ CD-ROM.

Disclaimer

Overview

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Links to External Sites

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Data Backup

GeneLinker™ makes every effort to ensure that your GeneLinker™ database will not be corrupted, but we still recommend the use of third-party backup solutions that would allow you to recover older versions of your GeneLinker™ database. The GeneLinker™ database resides in the Repository folder in the directory where you installed GeneLinker™:

- the MMC/GeneLinker Gold(Platinum)/Repository folder
- and in the DB2 or Oracle database, if either is used as the GeneLinker™ database instead of the default MySQL database.
Related Topics:
GeneLinker™ Tour
GeneLinker™ Product Suite

Audience Assumptions

Overview
It is assumed that you are familiar with the basics of running a Windows® application including navigation and file management.
While some background information is provided, it is assumed that you have a working knowledge of the terminology and techniques used in molecular biology, as well as basic familiarity with data mining goals and statistical techniques.

Related Topic:
Disclaimer
GeneLinker™ Functions List

General Formatting Conventions

Overview
General Formatting Conventions Used in the GeneLinker™ Online Manual
- Each topic has one or more of the following sections: Overview, Actions, Related Topics.
- All menu and menu item names appear in bold.
- Buttons, icons, and tab headings appear in bold.
- Window, dialog, and field names are displayed in bold.
- Keyboard keys to be pressed are denoted in angle brackets (e.g. <Enter> key).

Version Identification (Gold, Platinum)
- Platinum specific topics are marked with a green-and-platinum stripe in the left margin and the word Platinum (in platinum) in the top line.
- Gold specific topics are marked with a green-and-gold stripe in the left margin and the word Gold (in gold) in the top line.
- Mixed-version topics have a blank first line and an empty left margin.

Within a mixed version topic, a section that is for Platinum only begins with a platinum banner containing the word ‘Platinum’ in white. Where appropriate, the banner contains a title for the section.
Help Window Functions

Overview

The Help window is divided vertically into two separate areas, or *panes*. The left pane displays the table of contents or index, and the right pane displays information about the topic selected in the left pane.

Actions

Table of Contents

- To display the table of contents, click the **Contents** tab.
- To open or close a book under the **Contents** tab, double-click on the **book** icon.
- To open a book under a book, click the **plus** icon beside it.
- To close a book under a book, click the **minus** icon beside it.
- Click on a topic to display its contents in the right pane.

Index

- To display an alphabetical keyword index, click the **Index** tab.
- Scroll through the keywords in the list and click one of interest. The topic associated with that keyword is displayed in the right pane.
- To find a word (or part of a word) in the index, type the word (or part of a word) into the **Find** box at the top of the index and press <Enter>. **Note:** if you search on more than one word at a time, please use whole words only. If you use partial words in a multi-word search, the search may fail to find the topic.

Related Topics:

- GeneLinker™ Tour
- GeneLinker™ Function List

Starting GeneLinker and Setting Preferences

Starting the Program

Actions

During the installation process the GeneLinker™ program icon ‑ is placed on your computer's desktop. Double-click this icon to start the application. **Note:** if you have a large amount of data, it may take a few minutes for GeneLinker™ to load it into the database.
**GeneLinker™ Gold 3.0 Upgrade from GeneLinker™ Gold 2.x**

If you have upgraded from GeneLinker™ Gold 2.x to GeneLinker™ Gold 3.0, the data repository is upgraded to the new format the first time you run the new version of GeneLinker™. A message is displayed.

**Related Topic:**
Exiting the Program

**Changing Your User Preferences**

**Overview**
This facility allows GeneLinker™ to remember your preferences from one session to the next.

**Actions**
1. Select **Preferences** from the **Tools** menu. The **User Preferences** dialog is displayed.

2. Click the **General** tab to display the general preferences pane.
3. Set the parameters.

<table>
<thead>
<tr>
<th>Element</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>User Name</td>
<td>Your user identifier that appears in annotations and reports containing annotations.</td>
</tr>
<tr>
<td>Web Browser</td>
<td>The path to your preferred HTML browser.</td>
</tr>
<tr>
<td>Enable automatic</td>
<td>If this checkbox is checked then whenever any analysis experiment (any experiment other than data import, normalization, filtering, or missing value estimation)</td>
</tr>
<tr>
<td>visualizations</td>
<td></td>
</tr>
<tr>
<td>PCA Components to Display</td>
<td></td>
</tr>
<tr>
<td>Histogram Bins for Summary Statistics</td>
<td></td>
</tr>
</tbody>
</table>
completes, the default visualization that is associated with the experiment will be opened automatically as soon as the experiment completes. By default this preference is checked.

Enable Shared Selection
If this checkbox is checked, items such as genes and samples that are selected in one visualization will also be highlighted in other visualizations. By default this preference is checked.

PCA Components to Display
The default number of principal components to display in a loadings line plot or loadings color matrix plot (*display only - does not affect the calculation*).

Histogram Bins for Summary Statistics
The default number of bins for the Summary Statistics chart.

4. Click the **Gene Database** tab to display the gene database pane.

5. Set the parameters.

<table>
<thead>
<tr>
<th>Element</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gene Display Name</td>
<td>The default type of gene identifier used for display.</td>
</tr>
<tr>
<td>Lookup Gene Database URL: Affymetrix</td>
<td>Database URL for looking up a gene with an Affymetrix gene identifier. See <strong>Affymetrix URL Format</strong> below.</td>
</tr>
<tr>
<td>Lookup Gene Database URL: GenBank</td>
<td>Database URL for looking up a gene with a GenBank gene identifier. See <strong>GenBank URL Format</strong> below.</td>
</tr>
<tr>
<td>Lookup Gene Database URL: UniGene</td>
<td>Database URL for looking up a gene with a UniGene gene identifier. See <strong>UniGene URL Format</strong> below.</td>
</tr>
<tr>
<td>Lookup Gene Database URL: Custom</td>
<td>The URL used to access another gene database. Use the correct URL format for the database you are accessing.</td>
</tr>
</tbody>
</table>

6. Click **OK** to save changes to the settings or click **Cancel** to keep the previous values.
For more information about forming query strings, refer to Linking to PubMed and other Entrez databases:

Affymetrix URL Format:
- https://www.netaffx.com/index2.jsp

GenBank URL Format:

**Note** the use of the term **MMC_ID**. This term must appear in the URL. The application will replace this term with the identifier of a gene.

**For example**, if the gene being queried has the identifier **AF098020**, then the application will use the following URL to obtain information about that gene:

UniGene URL Format:

**Note**: the use of the terms **MMC_ORGANISM** and **MMC_ID**. These terms must appear in the URL. The application will replace these terms with the appropriate components of a UniGene gene identifier.

**For example**, if the gene being queried has the UniGene identifier **Ht.9573**, then the application will use the following URL to obtain information about that gene:

**Related Topics:**
- Lookup Gene
- Principal Component Analysis
- Summary Statistics

**Saving**

**Overview**

**Experiments Navigator Items**

Datasets and experiments do not have to be explicitly saved. When an experiment is run, the results are immediately and automatically saved to the GeneLinker™ database. The completion of this is indicated by the appearance of an item in the **Experiments** navigator as a 'child' under the original dataset.

If you want to back up or access your data for use in another application, simply export
the data to a file.

**Annotations**
Annotations do not have to be explicitly saved either. Once any addition or change is made to an annotation, the database is updated automatically. An annotation icon appears next to any item in a navigator tree that has an annotation.

**Profile Matching Results**
Profile Matching results are saved when you answer yes to the save profile match prompt. You also have the opportunity to save an unsaved profile match when you exit GeneLinker™.

**Gene Lists**
Gene lists are saved when they are created. Click the **Save List** button and provide a file name.

**Exiting the Program**

**Actions**
1. Ensure any required data/experiment information has been exported or reported on, as appropriate.
2. It is not necessary to save any datasets or experiments. GeneLinker™ automatically saves all datasets and experiments to its database in the Repository folder in the GeneLinker™ directory on your disk as you work.
3. Select **Exit** from the **File** menu. The GeneLinker™ application closes.

**Related Topics:**
- GeneLinker™ Database
- Annotating Data
- Creating a Gene List

**Application Interface**

**The Navigator**
Overview

The upper left pane of the GeneLinker™ window is called the Experiments, Genes, or Gene Lists navigator depending on which of the tabs is selected. GeneLinker™ displays the Experiments tab by default. All items listed in the navigator have already been saved to the GeneLinker™ database.

- The **Experiments** tab displays all your datasets and experiments in a hierarchical tree.
- The **Genes** tab displays an alphabetical list of all of the genes in your repository.
- The **Gene Lists** tab shows an alphabetical list of all of your gene lists.

**Navigator (Experiments, Genes, Gene Lists)**

<table>
<thead>
<tr>
<th>Icon</th>
<th>Type of Item</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image" alt="Complete dataset" /></td>
<td>A complete dataset (raw, preprocessed).</td>
</tr>
<tr>
<td><img src="image" alt="Incomplete dataset" /></td>
<td>An incomplete dataset.</td>
</tr>
<tr>
<td><img src="image" alt="Dataset with variable information" /></td>
<td>A complete dataset with variable information.</td>
</tr>
<tr>
<td><img src="image" alt="Incompleteness with variable information" /></td>
<td>An incomplete dataset with variable information.</td>
</tr>
<tr>
<td><img src="image" alt="F-Test results" /></td>
<td>F-Test results item.</td>
</tr>
<tr>
<td><img src="image" alt="Hierarchical clustering" /></td>
<td>A hierarchical clustering experiment.</td>
</tr>
<tr>
<td><img src="image" alt="Partitional clustering" /></td>
<td>A partitional clustering experiment.</td>
</tr>
<tr>
<td><img src="image" alt="SOM experiment" /></td>
<td>A SOM experiment.</td>
</tr>
<tr>
<td><img src="image" alt="PCA experiment" /></td>
<td>A PCA experiment.</td>
</tr>
<tr>
<td><img src="image" alt="Discretized data" /></td>
<td>Discretized data.</td>
</tr>
<tr>
<td><img src="image" alt="SLAM results" /></td>
<td>SLAM results.</td>
</tr>
<tr>
<td><img src="image" alt="Trained ANN classifier" /></td>
<td>Trained ANN classifier.</td>
</tr>
<tr>
<td><img src="image" alt="IBIS search results" /></td>
<td>IBIS search results.</td>
</tr>
<tr>
<td><img src="image" alt="Trained IBIS classifier" /></td>
<td>Trained IBIS classifier.</td>
</tr>
<tr>
<td><img src="image" alt="Classification results" /></td>
<td>Classification results.</td>
</tr>
<tr>
<td><img src="image" alt="Annotated item" /></td>
<td>An annotated item (icon to the right of the name).</td>
</tr>
</tbody>
</table>

*An experiment item* (clustering, SOM, PCA, etc.) is tagged with an icon appropriate to the process that created it.

**Default Dataset or Experiment Names**

Imported files, see Import for an explanation of where the dataset name comes from.
For all generated datasets or experiments, GeneLinker™ provides a default name. The default name is based on the type of process (and its parameter settings) used to create it.

### Example Dataset/Experiment Names

| Removed: v <= 7.6 | Removed: p > 0.65 |
| Estimated: #mv < 5 | mean | Estimated: #mv < 8 | nn=2 | euclid |
| Estimated: #mv < 5 | v=17.078 |
| Filtered: keep {myGeneList} |
| Filtered: remove {your Favourite Gene List} |
| Filtered: n-fold with n >= 2.5 |
| Filtered: range #=256 |
| Norm: log2 |
| Norm: Sample scaling: divide, mean=6.7 |
| Norm: Sample scaling: subtract mean |
| Norm: LinReg: [16-ALL B] | {likelyC56} |
| Norm: Lowess: window=0.25 |
| Norm: Neg ctrls: {u14-P inhibitors} | median | all samples |
| Norm: Pos ctrls: {some other gene list} | mean | each sample |
| Norm: Divided by max |
| F-test: "my Variable name here" |
| K-W test: "my Variable name here" |
| Hier: genes | Euclid | single |
| K-means, k=4: samples | Chebych | complete |
| J-P (4, 2): samples | Manhatn | avg |
| SOM: samples | 5x4 | Spear |
| PCA: genes |
| Discretized: 6 bins/sample | quantile |
| SLAM: "my Rep Variable #2" | 10,000 | 2 | 0.6 |
| ANN: "leukemia-Dr D" | 16-5-3 | N=10 | 0.001 | 15 |
| IBIS search: "Awl or AML test" | LDA | 1D |
| IBIS: "leukemia-Dr C" | QDA | 2D | N=10 |
| IBIS: "leukemia-Dr A" | UGDA, ALL | 1D | N=10 |
| myNewVariableName |
| Profile: {avg custom} | Spear |

### Related Topics:
- Using the Experiments Navigator
- Using the Genes Navigator
- Using the Gene Lists Navigator

### Navigator Pane Functions
Using the Experiments Navigator

Overview
The Experiments navigator displays a hierarchical tree listing of all of the datasets and experiments you have in your GeneLinker™ database. Clicking the Experiments tab brings the Experiments navigator to the front.

Actions
Expanding the Tree
• Click the plus icon beside the item. The item’s sub-experiments are displayed.

Collapsing the Tree
• Click the minus icon beside the item. The item's sub-experiments are hidden.

Toggling Between the Expanded and Collapsed State
• Double-click the item name. In the expanded state, the branch collapses; in the collapsed state, the branch expands.

Selecting an Item
• Click the item name. The item is highlighted and information about it is displayed in the Description pane just below the navigator pane.

Displaying the Shortcut Menu
• Right-click an item. A shortcut menu is displayed. Select an item on the shortcut menu to invoke the function.

Scrolling
• Clicking on the scrollbar at the side or bottom of the pane (when they are visible) moves the display.

<table>
<thead>
<tr>
<th>Double-Click an Item</th>
<th>Function Invoked</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dataset (complete or incomplete, raw data)</td>
<td>Color Matrix Plot</td>
</tr>
</tbody>
</table>
preprocessed, discretized, with or without variables, etc.)

<table>
<thead>
<tr>
<th>Experiment Type</th>
<th>Plot Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clustering experiment (hierarchical or partitional)</td>
<td>Matrix Tree Plot</td>
</tr>
<tr>
<td>SOM experiment</td>
<td>SOM Plot</td>
</tr>
<tr>
<td>PCA experiment</td>
<td>3D Score Plot</td>
</tr>
<tr>
<td>F-Test results</td>
<td>F-Test Viewer</td>
</tr>
<tr>
<td>Platinum: SLAM results</td>
<td>SLAM Association Viewer</td>
</tr>
<tr>
<td>Platinum: Classification results</td>
<td>Classification Plot</td>
</tr>
<tr>
<td>Platinum: IBIS search results</td>
<td>IBIS Search Results Viewer</td>
</tr>
<tr>
<td>Platinum: IBIS classifier</td>
<td>Classifier Gradient Plot</td>
</tr>
</tbody>
</table>

**Related Topics:**
- The Navigator Pane
- Renaming Datasets or Experiments
- Viewing Experiment Parameters

## Viewing Experiment Parameters

### Overview
When reviewing an experiment, you can examine the parameters with which it was run.

### Actions
1. Click a dataset or an experiment in the **Experiments** navigator. The item is highlighted.
2. Look at the information displayed in the **Description Pane** just below the navigator.
3. Select **Show Parameters** from the **Tools** menu, or right-click the item and select **Show Parameters** from the shortcut menu. The **Parameters for** dialog is displayed.

**Related Topics:**
- The Navigator
- The Description Pane
Renaming a Dataset or Experiment

Overview
It is possible to rename a dataset or experiment listed in the Experiments navigator.

Actions
1. Click a dataset or experiment in the Experiments navigator. The item is highlighted.
2. Select Rename Experiment from the Edit menu, or right-click the item and select Rename Experiment from the shortcut menu. The item name is bounded in an edit box.
3. Overtype the existing name with a new unique name.
4. Press <Enter> when finished to accept the new name.

Related Topics:
The Navigator
The Description Pane

Deleting a Dataset or Experiment

Overview
Deleting a dataset or experiment from the Experiments navigator deletes it from the database. This action does not delete any genes or gene lists from the database. Deleting a dataset or experiment closes all tables or plots of it.

Actions
1. Right-click a dataset or experiment in the Experiments navigator. The item is highlighted and the shortcut menu is displayed.
2. Select Delete Experiment from the shortcut menu. A confirmation dialog is displayed.
3. Click **Delete**. The dataset or experiment is deleted from the **Experiments** navigator and from the database. Any tables or plots showing the deleted item are closed.

**Related Topics:**
- The Navigator Pane
- Using the Experiments Navigator

### Using the Genes Navigator

#### Overview
The **Genes** navigator pane displays an alphabetical list of all of the genes you have in your GeneLinker™ database. Clicking the **Genes** tab brings the **Genes** navigator to the front.

![GeneList](gene.png)

#### Actions

**Selecting a Gene**
- Click the gene name in the **Genes** navigator. The gene is highlighted and information about it is displayed in the **Description** pane just below the navigator.

**Displaying the Shortcut Menu**
- Right-click a gene name to display the shortcut menu. Select an item on the menu to invoke the function.

**Locating a Particular Gene**
- Click in the **Locate** text field above the gene list and type in the name of the gene. As you type, the closest match is highlighted in the list of genes.

<table>
<thead>
<tr>
<th>Double-Click an Item</th>
<th>Function Invoked</th>
</tr>
</thead>
</table>

GeneLinker Gold 3.1 / GeneLinker Platinum 2.1 189
Related Topics:
- The Navigator
- The Description Pane
- Lookup Gene

Using the Gene Lists Navigator

Overview
The Gene Lists navigator pane displays an alphabetical list of all of the gene lists you have in your GeneLinker™ database. Clicking the Gene Lists tab brings the Gene Lists navigator to the front.

Actions
Displaying the Genes in a Gene List
- Click on the plus icon beside it to expand the list of genes under the gene list name.

Editing the Properties of a Gene List
- Double-click on a gene list name, or click on a gene list name and then click the edit gene list properties button just above the list of gene lists.

Saving a Gene List
- Click on a gene list name and then click the save gene list button just above the list of gene lists.

Deleting a Gene List
- Click on a gene list name and then click the delete gene list button just above the list of gene lists.

<table>
<thead>
<tr>
<th>Double-Click an Item</th>
<th>Function Invoked</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gene list name</td>
<td>Edit Gene List</td>
</tr>
<tr>
<td>Gene</td>
<td>Lookup Gene</td>
</tr>
</tbody>
</table>

Double-Click an Item | Function Invoked
--- | ---
Gene list name | Edit Gene List
Gene | Lookup Gene
The Description Pane

Overview

The Description pane is located in the lower left of the main window. It shows information about the item highlighted in the navigator pane, or a gene highlighted in a table or on a plot. This information can include:

- Name of dataset, experiment, gene (name possibly truncated), or gene list,
- Gene ID type,
- Gene description,
- Creation date/time,
- Annotations count,
- Gene list description,
- Number of genes,
- Number of samples.

In GeneLinker™, we refer to a dataset which has both treatment and control values stored as Two-Color Data. In the description pane for such a dataset it will say Two Channels Available: Yes. If the description pane does not say this, then GeneLinker™ does not have the required two values for each spot and cannot treat the data as Two-Color Data. If you believe you imported two-color data but the description pane says Two Channels Available: No, re-examine your data and your choice of a data import template. Two-Color Data can be imported using GenePix, Quantarray and Scanarray templates, but not all templates of those types import two-color data.
**Actions**

**Changing the Height of the Description Pane**
- Click and hold the border between the navigator and the description pane. Drag up to increase the description pane height, drag down to shrink it.

**Hiding the Description Pane**
- Click down arrow on top border of **Description** pane. The navigator is extended to the bottom of the window. The **Description** pane (below the navigator) is reduced to a thick border with an up and a down arrow on it.

**Restoring the Description Pane**
- Click on the up arrow on the thick border that is the **Description** pane below the navigator. The **Description** pane is restored to the size it was before it was hidden.

**Related Topics:**
- Viewing Experiment Parameters

**The Plots Pane**

**Overview**
The right pane of the GeneLinker™ main window is called the **Plots** pane. This is where all tables, charts and plots are drawn. Each table, chart or plot is a separate window.
Actions

Bringing a Plot to the Front

• Click on the plot.

Arranging the Plot Windows

• Select Cascade Windows from the Window menu.

Closing a Plot Window

• Click on the plot and then select Close from the Window menu, or click the \( \times \) icon in the upper right corner of the plot.

Closing All the Plot Windows

• Select Close All from the Window menu.

Related Topics:

Creating a Table View of Gene Expression data
Creating a Color Matrix Plot
Creating a Summary Statistics Chart
The Toolbar

Overview
The toolbar is located at the top of the GeneLinker™ window under the menu bar. The toolbar icons give you quick access to most of the program functionality.

This image is of the GeneLinker™ Platinum toolbar. The GeneLinker™ Gold toolbar has all the same icons except the Platinum specific ones (see list below).

The top of the following icon list corresponds to the left of the toolbar; the bottom of the list corresponds to the right of the toolbar. Click an item to view detailed information about that function.

- Data Import Step 1: Selecting a Template
- Create Gene List from Selection
- Find
- Annotate
- Estimate Missing Values
- Filter Genes
- Normalize
- Table View
- Color Matrix Plot / Loadings Color Matrix Plot (for a PCA experiment)
- Variable Viewer
- Summary Statistics
- Hierarchical Clustering
- Partitional Clustering
- Self Organizing Map
- Matrix Tree Plot
- Two Way Matrix Tree Plot
- Principal Components Analysis
- 3D Score Plot
- Discretize Data
- SLAM
- Create ANN Classifier
- IBIS Search
- Classify
- Lookup Gene
- Profile Matching
- Help

Toolbar Features
The GeneLinker™ toolbar icons are context sensitive. That is, only the icons representing functions appropriate for the selected item are enabled.

- An enabled icon is drawn in color.
• A disabled icon is grayed-out appearing to be embossed into the toolbar. When the mouse pointer passes over an enabled toolbar icon, the icon is drawn with a border. Also, its description appears in the main window status bar. When the mouse pointer hovers over a toolbar icon for a short time, a tooltip naming the icon function is displayed. At the far right of the toolbar is the molecule spinner 🌀. The molecule spinner spins when GeneLinker™ is performing a task.

The toolbar icons cannot be moved, rearranged or otherwise customized.

Actions
1. Click on an item in the Experiments, Genes, or Gene Lists navigator, or select one or more items on a plot. The icons representing functions appropriate to that item are enabled (drawn in color).
2. Click on an enabled toolbar icon to apply that function to the selected item.

Related Topic:
Keyboard Shortcuts

The Menus

File Menu

Overview
The File menu items provide access to the data, image saving, and reporting facilities of GeneLinker™. Exit closes the application.

<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Import: Gene Expression Data</td>
<td>Import data from formatted text files into the repository.</td>
</tr>
<tr>
<td>Import: Gene List</td>
<td>Import a gene list file.</td>
</tr>
<tr>
<td>Import: Variable</td>
<td>Import variable information for a dataset.</td>
</tr>
<tr>
<td>Export Data</td>
<td>Save the selected data as a comma-separated value (.csv) file, for use in other programs.</td>
</tr>
<tr>
<td>Menu Item</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Export Image</td>
<td>Save the selected plot to an image file.</td>
</tr>
<tr>
<td>Generate Report</td>
<td>Generate a report for the selected experiment.</td>
</tr>
<tr>
<td>Generate Workflow Report</td>
<td>Generate a workflow report that includes the entire branch of the <strong>Experiments</strong> tree, from the root dataset to the selected experiment.</td>
</tr>
<tr>
<td>Exit</td>
<td>Exit GeneLinker™. Note that all datasets and experiments listed in the <strong>Experiments</strong> tree are saved automatically by the program.</td>
</tr>
</tbody>
</table>

**Related Topics:**
- Importing Gene Expression Data
- Exporting a PNG Image
- Generating a Report
- Exporting to DecisionSite

**Edit Menu**

**Overview**
These menu items provide access to editing tools.

<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Create Gene List from Selection</td>
<td>Create a gene list from the highlighted selection in a table view or plot.</td>
</tr>
<tr>
<td>Find</td>
<td>Find a specific gene in a table or plot. See Find for more information.</td>
</tr>
<tr>
<td>Find Next</td>
<td>Find the next occurrence of a gene in a table or plot. See Find Next for more information.</td>
</tr>
<tr>
<td>Find Previous</td>
<td>Find the previous occurrence of a gene in a table or plot. See Find Previous for more information.</td>
</tr>
<tr>
<td>Annotate</td>
<td>Opens the annotations editor allowing you to add, change, delete, or view annotations. See Annotations Overview.</td>
</tr>
<tr>
<td>Rename Experiment</td>
<td>Rename the selected experiment.</td>
</tr>
<tr>
<td>Delete Experiment...</td>
<td>Delete the selected experiment or dataset and all of its sub-experiments.</td>
</tr>
</tbody>
</table>
Related Topics:
Creating a Gene List from Within GeneLinker™
Annotations Overview

View Menu

Overview
These menu items provide tools for customizing the active plot.

<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Customize</td>
<td>Customize the appearance of a plot.</td>
</tr>
<tr>
<td>Resize</td>
<td>Resize a plot.</td>
</tr>
<tr>
<td>Zoom</td>
<td>Zoom a SOM plot.</td>
</tr>
</tbody>
</table>

Related Topic:
Configuring Plot Components

Data Menu

Overview
These menu items provide access to editing tools.

<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remove Values</td>
<td>Remove values from the selected dataset, above, at or below the specified threshold.</td>
</tr>
<tr>
<td>Estimate Missing Values</td>
<td>Fill in the missing values in the selected incomplete dataset.</td>
</tr>
<tr>
<td>Filter Genes</td>
<td>Filter the genes from the selected experiment.</td>
</tr>
<tr>
<td>Normalize</td>
<td>Normalize the data from the selected experiment.</td>
</tr>
</tbody>
</table>

Related Topics:
Overview of Estimating Missing Values
Filtering Overview
Normalization Overview

Statistics Menu

Overview
These menu items provide access to statistics tools.

<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reliability</td>
<td>View the reliability measures associated with the selected dataset in a spreadsheet-like format.</td>
</tr>
<tr>
<td>Measures</td>
<td></td>
</tr>
<tr>
<td>F-Test</td>
<td>Generate p-values for genes in the selected dataset based on a grouping variable.</td>
</tr>
<tr>
<td>F-Test Viewer</td>
<td>View a table of the results of the F-Test.</td>
</tr>
<tr>
<td>Summary Statistics</td>
<td>View the Summary Statistics for a dataset. The Summary chart is a histogram that shows the distribution of the data values among a number of bins (20 is the default). The Summary Statistics text display lists the minimum and maximum values, mean, median, standard deviation, co-efficient of variance and the number of data and missing values.</td>
</tr>
</tbody>
</table>

Related Topics:
Creating a Table View of Reliability Data
F-Test
Summary Statistics

Explore Menu

Overview
These menu items provide access to editing tools.
<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Table View</td>
<td>View the data in the selected dataset in a spreadsheet-like format.</td>
</tr>
<tr>
<td>Color Matrix Plot</td>
<td>View the results of the selected experiment as a Color Matrix Plot.</td>
</tr>
<tr>
<td>Scatter Plot</td>
<td>View a pair of selected genes or samples in a Scatter Plot.</td>
</tr>
<tr>
<td>Intensity-Bias Plot</td>
<td>View a sample to determine if Lowess normalization is needed.</td>
</tr>
<tr>
<td>Coordinate Plot</td>
<td>View the results of the selected experiment as a Coordinate Plot.</td>
</tr>
<tr>
<td>Variable Viewer</td>
<td>View the variable data associated with a dataset.</td>
</tr>
</tbody>
</table>

**Related Topics:**
- Creating a Table View of Expression Data
- Creating a Color Matrix Plot
- Variable Viewer

**Clustering Menu**

**Overview**
These menu items provide tools for manipulating the experiment selected in the Experiments navigator pane.

<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hierarchical Clustering</td>
<td>Cluster the data using a hierarchical clustering method (e.g. agglomerative clustering). Hierarchical clusters may include other clusters, forming a tree-like structure.</td>
</tr>
<tr>
<td>Partitional Clustering</td>
<td>Cluster the data using a partitional clustering method (e.g. K-Means, Jarvis-Patrick clustering). Partitional clusters are flat or non-hierarchical. They do not contain other clusters.</td>
</tr>
</tbody>
</table>
Left Table:

| Self-Organizing Map | A SOM can be used to explore the groupings and relations within data by projecting the data onto a 2D image that clearly indicates regions of similarity. A SOM can also be used to view clusters. |
| Export Partitional Cluster | Exports the selected cluster from a partitional clustering plot to a file at the specified location. The file contains gene or sample names with their cluster identifiers. |
| Matrix Tree Plot | View the results of the selected experiment as a Dendrogram Plot or a Partitional Plot that shows the clustering relationships of the genes or samples. |
| Two Way Matrix Tree Plot | View the results of two clustering experiments simultaneously - one on genes, and the other on samples. Both must be derived from the same original dataset. |
| Centroid Plot | View the results of the selected clustering experiment as a Centroid Plot (each line corresponds to the profile of a cluster centroid). |
| Cluster Plot | View the results of the selected experiment as a Cluster Plot with items colored according to cluster membership. |
| SOM Plot | View SOM results via the composition of a proximity-gradient map, a list of the items (genes/samples) contained in a specific cluster and (optionally) a profile plot. |

Related Topics:

- Clustering Overview
- Self-Organizing Maps

Right Table:

<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Principal Component Analysis</td>
<td>PCA can be used to reduce the complexity of multivariate data in which a large number of variables (e.g., thousands) are interrelated, such as in large-scale gene expression data obtained across a variety of different samples or conditions.</td>
</tr>
<tr>
<td>Scree Plot</td>
<td>View PCA results in a Scree Plot. It is a simple line segment plot that shows the fraction of total variance in the data as explained or</td>
</tr>
</tbody>
</table>
represented by each PC.

<table>
<thead>
<tr>
<th>Score Plot</th>
<th>View the PCA results in a Score Plot. It is a scatter plot with the x axis representing a user-selected PC. The y axis represents another user-selected PC. The plot contains points that represent the original 'samples' (e.g., projected Samples if PCA by Genes (the 'variables'), projected Genes if PCA by Samples (the 'variables')) projected onto the user-selected PCs. By default, the Score Plot shows data on the first two PCs.</th>
</tr>
</thead>
<tbody>
<tr>
<td>3D Score Plot</td>
<td>View the PCA results in a 3D Score Plot. It is a scatter plot with the x, y and z axes representing user-selected PCs. The plot contains points that represent the original 'samples' (e.g., projected Samples if PCA by Genes (the 'variables'), projected Genes if PCA by Samples (the 'variables')) projected onto the user-selected PCs. By default, the 3D Score Plot shows data on the first three PCs.</td>
</tr>
<tr>
<td>Loadings Line Plot</td>
<td>View PCA results in a Loadings Line Plot. It displays the individual elements of the PCs in Principal Components Analysis, allowing you to see the relative influence of genes or samples on the PCs.</td>
</tr>
<tr>
<td>Loadings Scatter Plot</td>
<td>View PCA results in a Loadings Scatter Plot. The loadings of a given PC represent the relative extent to which the original 'variables' (genes or samples, depending on the Orientation selected for the PCA) influence the PC. The Loadings Scatter Plot displays these loadings compared to one another in a scatter plot of one selected PC vs. another selected PC.</td>
</tr>
<tr>
<td>Loadings Color Matrix Plot</td>
<td>View PCA results in a Loadings Color Matrix Plot. Re-order genes in plot by selecting a PC and an ordering (ascending, descending, absolute descending).</td>
</tr>
</tbody>
</table>

**Related Topics:**
- Overview of Principal Component Analysis (PCA)
- Creating a 3D Score Plot

## Platinum

### Predict Menu

**Overview**

These menu items provide tools for manipulating the experiment selected in the Experiments navigator.
<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Discretize Data</td>
<td>Create a dataset of discrete values reflecting the expression levels of the original data.</td>
</tr>
<tr>
<td>SLAM</td>
<td>Use the SLAM technology to find associations in a dataset.</td>
</tr>
<tr>
<td>Association Viewer</td>
<td>View a listing of the associations found by SLAM. The association viewer can also be used to create gene lists.</td>
</tr>
<tr>
<td>Create ANN Classifier</td>
<td>Use a dataset with known variables to train an ANN classifier.</td>
</tr>
<tr>
<td>Classification Plot</td>
<td>View the results of training an ANN classifier or classification of a dataset using either an ANN or an IBIS classifier.</td>
</tr>
<tr>
<td>Mean Squared Error Plot</td>
<td>Display a plot of the mean squared error of training an ANN classifier.</td>
</tr>
<tr>
<td>IBIS Classifier Search</td>
<td>Search a dataset for potential gene or gene pair IBIS classifiers.</td>
</tr>
<tr>
<td>IBIS Results Viewer</td>
<td>Display a table of IBIS proto-classifiers with statistics.</td>
</tr>
<tr>
<td>Create IBIS Classifier</td>
<td>Create an IBIS classifier from IBIS search results or a gene or gene pair.</td>
</tr>
<tr>
<td>Classifier Gradient Plot</td>
<td>Display an IBIS gradient plot of training or classification results.</td>
</tr>
<tr>
<td>Classify</td>
<td>Use a trained classifier (ANN or IBIS) to classify a dataset (predict a variable).</td>
</tr>
</tbody>
</table>

**Related Topics:**
- ANN Classification and Prediction Overview
- IBIS Overview

**Tools Menu**

**Overview**
These menu items provide access to the GeneLinker™ tool set.
Menu Item | Description
---|---
Lookup Gene | Lookup the selected gene in a specific gene database. Selecting this item spawns an external web browser displaying information about the selected gene. (The gene database web address (URL) is configurable via the Preferences item on the Edit menu).
Variable Manager | Displays a list of variables associated with a dataset.
Color Manager | Manages the colors for plots.
Profile Matching | Sort genes in a plot using a user specified expression profile as a reference.
Show Parameters | View the parameters used in the selected experiment.
License Information | Update the GeneLinker™ product license information.
Preferences | Edit your user preferences. Refer to Changing Your User Preferences for more information.

**Related Topics:**
- Color Manager
- Profile Matching
- License Information Overview

**Window Menu**

**Overview**

This menu provides tools for manipulating the windows that appear within the application’s main window. It also displays a list of open windows, any of which you may click to bring it to the front to view.

<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Close</td>
<td>Close the active window.</td>
</tr>
</tbody>
</table>
Help Menu

Overview
This menu provides access to help and company/product information.

<table>
<thead>
<tr>
<th>Menu Item</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GeneLinker Help</td>
<td>Show the online help table of contents.</td>
</tr>
<tr>
<td>View Printable Version of Help</td>
<td>Spawns Acrobat reader to show the help PDF.</td>
</tr>
<tr>
<td>Visit Molecular Mining</td>
<td>Spawn web browser displaying the MMC Web Site.</td>
</tr>
<tr>
<td>GeneLinker™ Technical Support</td>
<td>Spawn web browser displaying the MMC technical support page.</td>
</tr>
<tr>
<td>About</td>
<td>Show details about GeneLinker™ and your system.</td>
</tr>
</tbody>
</table>

Related Topic:
Help Window Functions

Data: Expression Measurements and Variables

Datasets Overview

Overview
GeneLinker™ imports three different kinds of data: expression data, variables, and gene lists. Of these three, only expression data is absolutely essential, which is why it is imported separately from the other two. However, variables and gene lists are very useful if they are available. Please see Variables Overview and Gene Lists Overview for more information.

The basic requirement for all GeneLinker™’s analysis capabilities is a set of expression values for a number of genes over a number of samples. In GeneLinker™ we refer to
this imported data as a root dataset because it lies at the root of a data family, a hierarchy or tree of datasets appearing in the Experiments navigator. (Like many trees in computer programs, these family trees of related datasets grow from the top left to the right and down.)

A root dataset can have any - or none - of the following characteristics associated with it:

Two-Color Data: Data from experiments involving paired dyes (red-green or Cy3-Cy5) can be treated specially by GeneLinker™. Please see Two-Color Data for more information.

Reliability Measures: Each spot or measurement may have associated with it a measure of its reliability or quality. Please see Reliability Measures for more information.

Variables: Each sample in a dataset may have associated with it a variety of phenotypes, experimental factors, treatments or conditions. Please see Variables Overview for more information.

Missing Values: Data may be missing for some genes in some samples, perhaps due to quality control filtering or due to minor version changes between different microarrays. For more information about the handling of missing values, please see Overview of Estimating Missing Values.

There are several mathematical distinctions among expression data which you should be aware of. Here are the most common mathematical classes of data and their significant characteristics.

Abundance Data

Synonyms: Count data, positive abundance data.

Example: Affymetrix data, CodeLink data.

Characteristics: All values are positive (or zero) since this type of data answers the question how many of <something> are there? The <something> might be molecules, but more likely it is some instrumental proxy, like phosphor intensity, which must also be non-negative. The histogram of count data for mRNA abundance is usually strongly peaked near the theoretical minimum of zero and tails off to the right.

Problems:

Zero values are theoretically possible (there may be none of a given thing there), but can cause numerical difficulties when doing various things like converting to ratios (division by zero is undefined) or taking logarithms (log zero is also undefined). Since instrumental measurements of very small values are usually unreliable in practice, it is often a good idea to eliminate zeroes in count data and replace them with some small positive value which lies near or below the instrumental detection limit.

Negative values may occur, but are generally symptomatic of a problem which ought to be fixed. For instance, much abundance data is computed by subtracting a background count from a foreground count. If the background exceeds the foreground, a negative value occurs which should be corrected. A common interpretation of this circumstance is unknown value, probably small.
Ratio Data

Example: Data from two-color experiments. GenePix, Genomic Solutions, Quantarray, ScanArray data.

Characteristics: All values are (theoretically) positive. Ratios are always defined with respect to some baseline or control sample. The histogram for mRNA ratios typically looks a lot like an abundance histogram, strongly tailed to the right. If the data were not too noisy and you could zoom in very tightly you might see that the histogram is peaked at 1.0 instead of near 0.

Data described as Two-Color Data by GeneLinker™ displays and is processed as ratio data. All Two-Color Data is ratio data, but not all ratio data is Two-Color Data.

Problems:

Ratio data can have negative values just like abundance data, most frequently because they are derived from abundances which have the background-subtraction problems described above. Zeros can also occur, and infinities as well if a zero happens to occur in the denominator (control sample) of a given treatment/control pair.

Related to the problem of zeros and infinities is the problem of large unreliable values. If the control value for a given sample is not actually zero, but nonetheless very small and unreliable, then the ratio may be deceptively large – and even more unreliable. It is extremely difficult to diagnose this problem when one only has the ratios to work with, so the user is advised to be careful of this in their data generation and upstream data processing.

See Also: Two-Color Data.

Log Ratio

Example: Usually generated by performing logarithm on imported ratio data. Common in published datasets (e.g. NCI60).

Characteristics: Values are positive and negative. The histogram for mRNA log ratios is typically a symmetric bell curve with a peak near zero.

Problems:

Logarithms cannot be computed for negative or zero values, so many of the problems are absent from log ratio data because they have been of necessity addressed upstream.

The problem of unreliable large ratios can nonetheless propagate into log ratio data undetected if care is not taken.

Frequently, zeroes or negatives in the ratio data are converted to missing values in the log ratio data derived therefrom.

Log Abundance

It is not uncommon to take the logarithm of abundance data without first nominating a baseline and taking ratios.

Example: Performing a log normalization on Affymetrix data yields log abundance data.

Characteristics: Values are positive and negative. The histogram for mRNA log
abundance is typically a bell curve.

Problems:

Logarithms cannot be computed for negative or zero values, so many of the problems described for the other data types are absent from log abundance data because they have been of necessity addressed upstream.

Frequently, zeroes or negatives due to background subtraction in the abundance data are converted to missing values in log abundance data derived therefrom.

Related Topics:

- Renaming Datasets or Experiments
- Viewing Experiment Parameters
- How to Import Expression Data

Importing Expression Data

How to Import Expression Data

Overview

Importing expression data into GeneLinker™ is a four-step process.

1. Choose a template that matches the format of the data in your file(s).
   
   The template to choose usually has the same name as the software which generated your data files, although there may be several to choose between in some cases. See Selecting a Template for Data Import for more information.

2. Select the source files in which GeneLinker™ should look for the data. This process is slightly different depending on whether you have all your data in one file, or whether it is spread across several files. Selecting a Template for Data Import gives you directions appropriate to your situation.

3. Ensure that the gene database matches the gene identifiers in the data. This may be done either before or after you select the source files. See Selecting the Gene Database Type for more information.

4. After GeneLinker™ has read the source files, a preview of the data is presented on the Import Data dialog so you can verify that the imported dataset is correct before it is saved to GeneLinker™’s database.

Note: In GeneLinker™, we refer to a dataset which has both treatment and control values stored as Two-Color Data. In the description pane for such a dataset it will say Two Channels Available: Yes. If the description pane does not say this, then GeneLinker™ does not have the required two values for each spot and cannot treat the data as Two-Color Data. If you believe you imported two-color data but the description pane says Two Channels Available: No, re-examine your data and your
choice of a data import template. Two-Color Data can be imported using GenePix, Quantarray and Scanarray templates, but not all templates of those types import two-color data.

Related Topics:
  - Two-Color Data
  - Selecting a Template for Data Import
  - Selecting the Gene Database Type
  - Importing One File Containing All Samples
  - Importing Multiple Files With One Sample Each

File Formats and Templates

Importing Data from Tabular Files

Overview
A Tabular file is a single file of expression values for multiple samples or chips. This is a generic format, not specific to any particular microarray software. If your data is not in one of the other formats described in Selecting a Template for Data Import, then you should use tabular format.

You can transform your data into tabular format in a number of ways, but the simplest is to use a spreadsheet program (like Microsoft Excel®, for example). Cut-and-paste your expression measurements into a simple table, and then export the table to an intermediate file. In order for it to import properly into GeneLinker™, the intermediate file should have the following characteristics:

- The data must all be in one text file (DOS®/Windows®, UNIX, or Macintosh).
- The data must be in a table. That is, it must be organized into rows of equal lengths and columns of equal lengths.
- By default GeneLinker™ expects the rows of the file to represent samples and the columns genes, but this is not required. If the data file represents genes as rows and samples as columns, then you can orient it properly by ensuring the Transpose box is checked during the verification step of the data import process.
- The first row should contain column names. The first column should contain row names. Absent column or row names may cause parts of your data to be misinterpreted.
- A single character must delimit fields. Example delimiter characters are the comma or the tab character. Comma-delimited is recommended over tab-delimited. For best results ensure your data is in a .csv file before importing. In a Comma Separated Values (.csv) file, each record (row) is stored as text with a comma delimiter separating each field and a carriage return/line feed character pair marking the end of each record (row).
- At least one row and one column of data must be present. These are in addition to
the row and column names.

- Missing values are signified by leaving blank space or no space between a consecutive pair of column delimiters. Alternatively, missing values may be signified by the string 'NA'.
- Anything preceding the first column separator in the first row will be ignored. That is, the upper left cell may contain anything, or nothing.

Example of a CSV data file with 4 genes and 3 samples:

<table>
<thead>
<tr>
<th></th>
<th>G1</th>
<th>G2</th>
<th>G3</th>
<th>G4</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>1.1</td>
<td>1.2</td>
<td>1.3</td>
<td>1.4</td>
</tr>
<tr>
<td>S2</td>
<td>2.1</td>
<td>2.2</td>
<td>2.3</td>
<td>2.4</td>
</tr>
<tr>
<td>S3</td>
<td>3.1</td>
<td>3.2</td>
<td>3.3</td>
<td>3.4</td>
</tr>
</tbody>
</table>

Example of a CSV data file with missing values:

<table>
<thead>
<tr>
<th></th>
<th>G1</th>
<th>G2</th>
<th>G3</th>
<th>G4</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>1.1</td>
<td>1.2</td>
<td>1.3</td>
<td>1.4</td>
</tr>
<tr>
<td>S2</td>
<td>2.1</td>
<td></td>
<td>2.3</td>
<td></td>
</tr>
<tr>
<td>S3</td>
<td></td>
<td>NA</td>
<td>3.3</td>
<td>3.4</td>
</tr>
</tbody>
</table>

Merging replicate genes:
If you have replicate spots (genes) on each chip, you may choose to have GeneLinker™ merge these into a single average measurement. The spread between the replicates will be converted into a reliability measure. For more background on this process, read Merging Within-Chip Replicate Measurements.

In order to do this, you have to select the template that properly describes the organization of your data. If you have a table in which each column represents a gene and each row a sample, then use the Tabular Merge Replicate Columns template. If you have a table in which each row represents a gene and each column a sample, then use the Tabular Merge Replicate Rows template.

Reliability Measures:
If you have some other source for reliability measures, you can import them into GeneLinker™ along with your expression data. Use the Tabular with Reliability Measures template.

The reliability measures must be in a tabular file of identical shape to your gene expression data file. If your gene expression data file is named FileName.ext then your reliability measures must be in a file named FileName_rm.ext in the same folder. GeneLinker™ expects that reliability measures will be between 0 and 1 inclusive, and that values close to 0 will indicate highly reliable data.

See Reliability Measures for more information.

Related Topics:
Selecting a Template for Data Import
Importing One File Containing All Samples

Importing Data from Affymetrix MAS 4.0 Files

Overview
The data files must be in Affymetrix MAS 4.0 tabular file format.

<table>
<thead>
<tr>
<th>Probe Set</th>
<th>Positive</th>
<th>Negative</th>
<th>Pairs Used</th>
<th>Log Avg Diff</th>
<th>Abs Call</th>
</tr>
</thead>
<tbody>
<tr>
<td>Condition A</td>
<td>AFFX-Blc-B_5st</td>
<td>5</td>
<td>3</td>
<td>20</td>
<td>1.8</td>
</tr>
<tr>
<td>Condition A</td>
<td>AFFX-Blc-B_4st</td>
<td>10</td>
<td>3</td>
<td>20</td>
<td>2.0</td>
</tr>
<tr>
<td>Condition A</td>
<td>AFFX-Blc-B_3st</td>
<td>9</td>
<td>4</td>
<td>20</td>
<td>0.0</td>
</tr>
<tr>
<td>Condition A</td>
<td>AFFX-Blc-B_2st</td>
<td>14</td>
<td>1</td>
<td>20</td>
<td>4.0</td>
</tr>
</tbody>
</table>

Import Process
Multiple files are processed into a single dataset. The sample order of the imported dataset is determined by the order of the source sample data files listed in the Import Data dialog.

- The file headers are discarded.
- Gene identifier information is retrieved from the Probe Set column of the first file and is stored as an Affymetrix Identifier.
- Gene expression data is retrieved from the Avg Diff column and the reliability measure is translated from the Present/Absent/Marginal P/A/M flags (P=0.0; M=0.5; A=1.0) of each file in the order they are placed in the Import Data dialog.

Related Topics:
Selecting a Template for Data Import
Importing Multiple Files With One Sample Each

Importing Data from Affymetrix MAS 5.0 Files

Overview
The data files must be in Affymetrix MAS 5.0 tabular file format.

<table>
<thead>
<tr>
<th>Stat</th>
<th>Stat</th>
<th>Signal</th>
<th>Detection</th>
<th>Detection p-value</th>
<th>Descriptions</th>
</tr>
</thead>
<tbody>
<tr>
<td>AFFX-MurIL2</td>
<td>20</td>
<td>20</td>
<td>61.3</td>
<td>A</td>
<td>0.000006 M 16782 Mouse interleukin</td>
</tr>
<tr>
<td>AFFX-MurIL1</td>
<td>20</td>
<td>20</td>
<td>736.3</td>
<td>A</td>
<td>0.016324 M 17887 Mouse interleukin</td>
</tr>
<tr>
<td>AFFX-MurIL4</td>
<td>20</td>
<td>20</td>
<td>57.1</td>
<td>A</td>
<td>0.000024 M 25322 Mus musculus ii</td>
</tr>
<tr>
<td>AFFX-MurIL3</td>
<td>20</td>
<td>20</td>
<td>59.4</td>
<td>A</td>
<td>0.000006 M 16782 Mouse interleukin</td>
</tr>
<tr>
<td>AFFX-BaaBA</td>
<td>20</td>
<td>20</td>
<td>5543.9</td>
<td>P</td>
<td>0.000317 J04429 C. elegans babA gene</td>
</tr>
<tr>
<td>AFFX-BaaAL</td>
<td>20</td>
<td>20</td>
<td>15841.5</td>
<td>P</td>
<td>0.000006 J04429 C. elegans babAL gene</td>
</tr>
<tr>
<td>AFFX-BaaB</td>
<td>20</td>
<td>20</td>
<td>4095.3</td>
<td>P</td>
<td>0.000141 J04429 C. elegans babB gene</td>
</tr>
<tr>
<td>AFFX-BaaC</td>
<td>20</td>
<td>20</td>
<td>2898.2</td>
<td>P</td>
<td>0.001141 J04429 C. elegans babC gene</td>
</tr>
<tr>
<td>AFFX-BaaD</td>
<td>20</td>
<td>20</td>
<td>14150.4</td>
<td>P</td>
<td>0.000141 J04429 C. elegans babD gene</td>
</tr>
<tr>
<td>AFFX-BaaE</td>
<td>20</td>
<td>20</td>
<td>12962.8</td>
<td>P</td>
<td>0.000006 J04429 C. elegans babE gene</td>
</tr>
<tr>
<td>AFFX-CraE</td>
<td>20</td>
<td>20</td>
<td>88016.1</td>
<td>P</td>
<td>0.000381 J04429 C. elegans babE gene</td>
</tr>
<tr>
<td>AFFX-CraC</td>
<td>20</td>
<td>20</td>
<td>14800.4</td>
<td>P</td>
<td>0.000004 J03453 Bacillus subtilis P</td>
</tr>
<tr>
<td>AFFX-CraD</td>
<td>20</td>
<td>20</td>
<td>18340.6</td>
<td>P</td>
<td>0.000044 J03453 Bacillus subtilis P</td>
</tr>
<tr>
<td>AFFX-CraB</td>
<td>20</td>
<td>20</td>
<td>915.8</td>
<td>A</td>
<td>0.507996 J04429 C. elegans babB gene</td>
</tr>
</tbody>
</table>
In MAS 5, the data should be exported from the *Pivot Tab* in tab delimited .txt format. Ensure that the exported files all contain the *Signal* and *Detection p-value* columns.

**Import Process**

Multiple files are processed into a single dataset. The sample order of the imported dataset is determined by the order of the source sample data files listed in the **Import Data** dialog.

- The file headers are discarded.
- **Gene identifier** information is retrieved from the *first* column of the first file and is stored as an **Affymetrix Identifier**.
- Gene *expression data* is retrieved from the *Signal* column and the *reliability measure* is retrieved from the *Detection p-value* column of each file in the order they are placed in the **Import Data** dialog.

**Related Topics:**

- Selecting a Template for Data Import
- Importing Multiple Files With One Sample Each

**Importing Data from CodeLink XML Files**

**Overview**

The data files must be in the CodeLink PROFILE XML file format.

CodeLink may associate up to three XML files with each slide or sample: A PATTERN file, a PROFILE file and an ID file. The PROFILE file contains the expression data which GeneLinker™ imports.

Example PROFILE.XML viewed with Microsoft Internet Explorer:
When selecting files for import, you need only select the PROFILE.XML files as in the picture below. The PATTERN and ID files should not be selected.

Import Process

Multiple files are processed into a single dataset. The sample order of the imported dataset is determined by the order of the source sample data files listed in the Data Import dialog as shown above.

You should use the GenBank gene database type when importing CodeLink data.

Characteristics of the CodeLink Import Template

GeneLinker Gold 3.1 / GeneLinker Platinum 2.1
The CodeLink import template has the following characteristics.

1. GenBank accession numbers are used as gene identifiers. These are obtained by stripping the reporter name of its '_PROBE\n' extension. Although the systematic names are also GenBank accession numbers, they are sometimes non-unique: That is, two different probes may be mapped to a single systematic name. In order to preserve the distinct identities of the probes GeneLinker™ uses the reporter names. If the systematic names are desired, they can be imported as descriptions via gene list import.

2. GeneLinker™ reads the normalized iod value as the expression value. These values are already background-subtracted and normalized by division by the median value of the DISCOVERY probes on the slide.

Related Topics:
- Selecting a Template for Data Import
- Importing Multiple Files With One Sample Each

Importing Data from dChip xls Files

Overview
The data files must be in the dChip tabular file format.

<table>
<thead>
<tr>
<th>Probe ID</th>
<th>Gene Accession</th>
<th>Description</th>
<th>Affy Identifier</th>
<th>NAP call</th>
<th>GBM 597</th>
<th>GBM 597 NAT</th>
<th>NAP call PILO 633 PILO 633 GBM 660</th>
<th>GE</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>400 P</td>
<td>423.237 P</td>
<td>4964 P</td>
<td>345 P</td>
<td>333 6262 P</td>
<td>274 3438 P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>441 P</td>
<td>4964 P</td>
<td>417.45 P</td>
<td>3652 62 P</td>
<td>352.62 P</td>
<td>259 3108 P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>332 4 M</td>
<td>380.668 P</td>
<td>428 74 P</td>
<td>3526 71 P</td>
<td>333.0168 P</td>
<td>292 4272 P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>329 9 P</td>
<td>379.038 P</td>
<td>478.84 P</td>
<td>3592 12 P</td>
<td>259 3108 P</td>
<td>292 4272 P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>306 2 P</td>
<td>349.635 P</td>
<td>433.3 P</td>
<td>3482 69 P</td>
<td>292 4272 P</td>
<td>292 4272 P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>306 6 P</td>
<td>302.967 P</td>
<td>416.3 P</td>
<td>3491 39 P</td>
<td>329 0234 P</td>
<td>292 4272 P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>366 3 P</td>
<td>366 209 P</td>
<td>476.22 P</td>
<td>3557 18 P</td>
<td>340 1238 P</td>
<td>292 4272 P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>387 2 P</td>
<td>342.927 P</td>
<td>491.38 P</td>
<td>3473 63 P</td>
<td>347 9078 P</td>
<td>292 4272 P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>386 8 P</td>
<td>311.838 P</td>
<td>487.88 P</td>
<td>3517 19 P</td>
<td>359 3737 P</td>
<td>292 4272 P</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AFFX-503</td>
<td>J0423 E c</td>
<td>381 9 A</td>
<td>480.47 A</td>
<td>3408 98 A</td>
<td>394 4893 A</td>
<td>292 4272 P</td>
<td>292 4272 P</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Import Process
One or two files are processed into a single dataset. For Affy chips that are broken across two files, such as HU133A/B use the DCHIP paired xls files template and select both files in the pair. For unpaired files use the DCHIP single xls file template.

- Gene identifier information is retrieved from the first column of the first file and is stored as an Affy Identifier.
- For paired chips, samples are ordered according to their order in the first file. Samples that are present in one file but not the other will have missing values.
for the file they are missing from.

Related Topics:
Selecting a Template for Data Import
Importing Multiple Files With One Sample Each

Importing Data from GenePix Files

Overview
The data files must be in the Axon .gpr file format.

<table>
<thead>
<tr>
<th>Sample Order</th>
</tr>
</thead>
<tbody>
<tr>
<td>The sample order of imported datasets is determined by the order of the source sample data files listed in the Import Data dialog.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Template</th>
<th>Result of Import</th>
</tr>
</thead>
<tbody>
<tr>
<td>GenePix</td>
<td>Multiple files are processed into a single dataset.</td>
</tr>
<tr>
<td>GenePix Merge Replicates</td>
<td>Multiple files are processed into a single dataset.</td>
</tr>
<tr>
<td>GenePix Green/Red</td>
<td>Multiple files are processed into a single ratio dataset (treatment/control).</td>
</tr>
<tr>
<td>GenePix Red/Green</td>
<td>Multiple files are processed into a single ratio dataset (treatment/control).</td>
</tr>
</tbody>
</table>
If you are importing using one of the two-color data templates (the dye colors are listed as treatment/control in the template name), all data values <0 are replaced with missing values (null values). Between-chip replicate measurements are imported as samples with the same names.

When the import process is complete, a dataset that is the ratio of treatment/control is added to the Experiments navigator. A selected sample ratio can be displayed in an intensity-bias plot to determine whether Lowess normalization is appropriate for the dataset.

Import Process for GenePix and GenePix Merge Replicates

- The file headers are discarded.
- **Gene identifier** information is retrieved from the Name column of the first file and is stored as a GenBank Identifier.
- Gene expression data is retrieved from the Ratio of Medians column of each file in the order they are placed in the Import Data dialog.
- The resulting dataset is not be amenable to Lowess Normalization or Intensity-Bias plots. See Two-Color Data for more information.
- The GenePix Merge Replicates merges any duplicate gene identifiers and computes reliability measures from the spread. See Merging Within-Chip Replicate Measurements for more information.

Import Process for GenePix Green/Red

- The file headers are discarded.
- The RatioFormulation field is ignored.
- **Gene identifier** information is retrieved from the Name column of the first file and is stored as a GenBank Identifier.
- The control (green dye) expression data is calculated by subtracting the B532 Median column from the F532 Median column.
- The treatment (red dye) expression data is calculated by subtracting the B635 Median column from the F635 Median column.
- The resulting dataset is amenable to Lowess Normalization and Intensity-Bias plots.

Import Process for GenePix Red/Green

- The file headers are discarded.
- The RatioFormulation field is ignored.
- **Gene identifier** information is retrieved from the Name column of the first file and is stored as a GenBank Identifier.
- The control (red dye) expression data is calculated by subtracting the B635 Median column from the F635 Median column.
- The treatment (green dye) expression data is calculated by subtracting the B532 Median column from the F532 Median column.
- The resulting dataset is amenable to Lowess Normalization and Intensity-Bias plots.

**Related Topics:**
- Selecting a Template for Data Import
Importing Data from Genomic Solutions Files

Overview
The data files must be in the Genomic Solutions tabular file format.

<table>
<thead>
<tr>
<th>Gene Name</th>
<th>Replicate ID</th>
<th>Replicate Ratio</th>
<th>Unique ID</th>
<th>Cx3 Volume</th>
<th>Cx5 Volume</th>
<th>Spot Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>HOLD 1</td>
<td>0</td>
<td>0.76</td>
<td>3001</td>
<td>1694714</td>
<td>1281422</td>
<td>0.76</td>
</tr>
<tr>
<td>HOLD 1</td>
<td>0</td>
<td>0.76</td>
<td>5115</td>
<td>171078</td>
<td>152089</td>
<td>0.77</td>
</tr>
<tr>
<td>HOLD 2</td>
<td>1</td>
<td>0.93</td>
<td>2901</td>
<td>140969</td>
<td>183759</td>
<td>1.03</td>
</tr>
<tr>
<td>HOLD 2</td>
<td>1</td>
<td>0.93</td>
<td>3003</td>
<td>1717572</td>
<td>1420295</td>
<td>0.83</td>
</tr>
</tbody>
</table>

Import Process
Multiple files are processed into a single dataset. The sample order of the imported dataset is determined by the order of the source sample data files listed in the Import Data dialog.

• The file headers are discarded.
• Gene identifier information is retrieved from the first column of the first file and is stored as a GenBank Identifier.
• Gene expression data is retrieved from the Spot Ratio column of each file in the order they are placed in the Import Data dialog.

Related Topics:
Selecting a Template for Data Import  
Importing Multiple Files With One Sample Each

Importing Data from Quantarray Files

Overview
The data files must be in the Quantarray file format.

File Header Section Example:
File Data Section Example:

Import Process

Multiple files are processed into a single two-color dataset. The sample order of the imported dataset is determined by the order of the source sample data files listed in the Import Data dialog.

Characteristics of the Quantarray Import Template

The Quantarray import template assumes the following about the format of the data files:

1. The data must be delimited by tab characters.
2. Gene identifiers are in the sixth column of the Data section.
3. The Measurements section is ignored.
4. Treatment and control channels are based on the information in the Image Info section of the Quantarray files. NOTE: All files must use the same channel (either ch1 or ch2) for the control channel. The channel used for control in all files is the channel labelled 'Control Image' in the last file in the import list. You can reorder the files in the import list using the black up- and down-arrow buttons on the Data Import dialog.
5. If the Image Info section is missing from the last file, then ch1 is used for the control channel and ch2 for the treatment channel.
6. It is assumed that the foreground and background counts are found in the Data section.
section in the columns headed ch1 Intensity, ch1 Background, ch2 Intensity and ch2 Background. The substrings 'ch1' and 'ch2' must match the lines in the Image Info section, if present.

7. GeneLinker™ stores the resulting ratios and associated intensities in a two-color dataset listed in the navigator. This makes it possible (for instance) to apply a Lowess correction to the dataset.

8. Spots for which the background count exceeds the foreground count are imported into GeneLinker™ as missing values. Negative ratios are not imported.

Related Topics:
- Selecting a Template for Data Import
- Importing Multiple Files With One Sample Each

Importing Data from ScanArray Files

Overview

The data files must be in the Perkin-Elmer ScanArray file format.

Sample Order

The sample order of imported datasets is determined by the order of the source sample data files listed in the Import Data dialog.

<table>
<thead>
<tr>
<th>Template</th>
<th>Result of Import</th>
</tr>
</thead>
<tbody>
<tr>
<td>ScanArray</td>
<td>Multiple files are processed into a single dataset.</td>
</tr>
<tr>
<td>ScanArray Merge Replicates</td>
<td>Multiple files are processed into a single dataset.</td>
</tr>
<tr>
<td>ScanArray Ch1/Ch2</td>
<td>Multiple files are processed into a single ratio dataset (treatment/control).</td>
</tr>
<tr>
<td>ScanArray Ch2/Ch1</td>
<td>Multiple files are processed into a single ratio dataset.</td>
</tr>
</tbody>
</table>
Import Process for ScanArray and ScanArray Merge Replicates

- The file headers are discarded.
- **Gene identifier** information is retrieved from the **Name** column of the first file and is stored as a **GenBank Identifier**.
- Gene **expression data** is retrieved from the **Ch2 Ratio of Medians** column of each file in the order they are placed in the **Import Data** dialog.

Import Process for ScanArray Ch1/Ch2

- The file headers are discarded.
- The **RatioFormulation** field is ignored.
- **Gene identifier** information is retrieved from the **Name** column of the first file and is stored as a **GenBank Identifier**.
- The **control (Ch2) expression data** is calculated by subtracting the **Ch2 B Median** column from the **Ch2 Median** column.
- The **treatment (Ch1) expression data** is calculated by subtracting the **Ch1 B Median** column from the **Ch1 Median** column.

Import Process for ScanArray Ch2/Ch1

- The file headers are discarded.
- The **RatioFormulation** field is ignored.
- **Gene identifier** information is retrieved from the **Name** column of the first file and is stored as a **GenBank Identifier**.
- The **control (Ch1) expression data** is calculated by subtracting the **Ch1 B Median** column from the **Ch1 Median** column.
- The **treatment (Ch2) expression data** is calculated by subtracting the **Ch2 B Median** column from the **Ch2 Median** column.

**Related Topics:**
- Selecting a Template for Data Import
- Importing Multiple Files With One Sample Each

**Selecting a Template for Data Import**

**Overview**

GeneLinker™ can read expression data files produced by a wide variety of other software. GeneLinker™ uses a template to interpret the contents of your data file or files.

**Data files containing one sample each:**

<table>
<thead>
<tr>
<th>Template Name</th>
<th>Template Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>GeneLinker Gold 3.1 / GeneLinker Platinum 2.1</td>
<td>219</td>
</tr>
<tr>
<td>Format</td>
<td>Description</td>
</tr>
<tr>
<td>---------------------------------</td>
<td>-----------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>Affymetrix 4.0</strong></td>
<td>Import Affymetrix MAS 4.0 data files.</td>
</tr>
<tr>
<td><strong>Affymetrix 5.0</strong></td>
<td>Import Affymetrix MAS 5.0 data files.</td>
</tr>
<tr>
<td><strong>CodeLink</strong></td>
<td>Import CodeLink XML files.</td>
</tr>
<tr>
<td><strong>dChip paired xls files</strong></td>
<td>Import dChip paired xls files.</td>
</tr>
<tr>
<td><strong>GenePix</strong></td>
<td>Import GenePix ATF data files.</td>
</tr>
<tr>
<td><strong>GenePix Green/Red</strong></td>
<td>Import GenePix ATF two-color data values (treatment=green/control=red).</td>
</tr>
<tr>
<td><strong>GenePix Merge Replicates</strong></td>
<td>Import GenePix ATF data files and generate reliability measures by merging replicates (see Merging Within-Chip Replicate Measurements).</td>
</tr>
<tr>
<td><strong>GenePix Red/Green</strong></td>
<td>Import GenePix ATF two-color data values (treatment=red/control=green).</td>
</tr>
<tr>
<td><strong>Genomic Solutions</strong></td>
<td>Import Genomic Solutions files.</td>
</tr>
<tr>
<td><strong>Genomic Solutions Merge Replicates</strong></td>
<td>Import Genomic Solutions data files and generate reliability measures by merging replicates (see Merging Within-Chip Replicate Measurements).</td>
</tr>
<tr>
<td><strong>Quantarray</strong></td>
<td>Import Quantarray data values into a two-color dataset.</td>
</tr>
<tr>
<td><strong>ScanArray</strong></td>
<td>Import ScanArray data files.</td>
</tr>
<tr>
<td><strong>ScanArray Merge Replicates</strong></td>
<td>Import ScanArray data files and generate reliability measures by merging replicates (see Merging Within-Chip Replicate Measurements).</td>
</tr>
<tr>
<td><strong>ScanArray TwoColor (Ch1/Ch2)</strong></td>
<td>Import ScanArray two-color data values (treatment=Ch1/control=Ch2).</td>
</tr>
<tr>
<td><strong>ScanArray TwoColor (Ch2/Ch1)</strong></td>
<td>Import ScanArray two-color data values (treatment=Ch2/control=Ch1).</td>
</tr>
</tbody>
</table>

**Data files containing all samples in one file:**

<table>
<thead>
<tr>
<th>Format</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Multi-Sample Data File</strong></td>
<td>Template Description</td>
</tr>
<tr>
<td><strong>DCHIP single xls file</strong></td>
<td>Import dChip single xls data file.</td>
</tr>
<tr>
<td><strong>Tabular</strong></td>
<td>Import tabular data from a single multi-sample data file.</td>
</tr>
<tr>
<td><strong>Tabular Merge Replicate Columns</strong></td>
<td>Import tabular data with genes represented by columns and generate reliability measures by merging replicate genes (see Merging Within-Chip Replicate Measurements). <strong>Be sure this is what you want. Tabular files more typically have genes in rows!</strong></td>
</tr>
<tr>
<td><strong>Tabular Merge Replicate Rows</strong></td>
<td>Import tabular data with genes represented by rows and generate reliability measure by merging replicate genes (see Merging Within-Chip Replicate Measurements).</td>
</tr>
<tr>
<td><strong>Tabular with Reliability Measures</strong></td>
<td>If you have generated reliability measures for tabular data independently of GeneLinker™, it is possible to import them along with your data. They must be in a tabular file of identical shape to your gene expression data file. If your gene expression data file is named <strong>FileName.ext</strong> then your reliability measures must be in a file named <strong>FileName_rm.ext</strong> in the same folder. GeneLinker™ expects that reliability measures will be between 0 and 1</td>
</tr>
</tbody>
</table>
inclusive, and that values close to 0 will indicate highly reliable data.

If you do not see a format in the lists above that matches the format of your data, your best course of action is to transform your data into Tabular format. See Importing Data from Tabular Files for more information.

**Actions**

1. Click the **Import Gene Expression Data** toolbar icon, or select **Import** from the **File** menu and **Gene Expression Data** from the sub menu. The **Data Import** dialog is displayed.

   ![Data Import Dialog](image)

2. Select a **Gene Database** from the drop-down list. This tells GeneLinker™ which type of gene identifier the genes being imported have (GenBank, Affymetrix, UniGene, or custom). If you need more information about this, see Selecting the Gene Database Type. You can also select the gene database after you have changed templates if you wish.

   The name of the *selected* template appears on the **Data Import** dialog. If this is the appropriate template for your data, go to either Importing One File Containing All Samples, or Importing Multiple Files With One Sample Each, as appropriate to the template you have selected.

   If the appropriate template name is not showing on the dialog, then continue:

3. Click the **Template Change** button. The **Import Templates** dialog is displayed.

   ![Import Templates Dialog](image)

4. Click the template that is appropriate for your data file(s). The template is highlighted.

5. To set the selected template as the default, click the checkbox next to **Make this the default template**. If you will be importing data of the same format repeatedly, you
should check this box so you will not need to re-select the same template each time you import data.

6. Click Select.

• If you selected one of the Tabular or DCHIP single xls file templates, the Data Import dialog is updated to permit you to specify a single data file to import, which should contain data about all the samples.

Go to Importing One File Containing All Samples.

• If you selected an Affymetrix, CodeLink, DCHIP paired xls files, GenePix, Genomic Solutions, Quantarray or ScanArray template, the Data Import dialog layout changes to permit you to select a set of single-sample data files from one folder.

Go to Importing Multiple Files With One Sample Each.

Note: gene identifiers have a length restriction of 25 characters. This means that on import of a dataset or a gene list, identifiers that are longer than 25 characters are truncated.

Related Topics:
- Importing One File Containing All Samples
- Importing Multiple Files With One Sample Each
- Merging Within-Chip Replicate Measurements

Selecting the Gene Database Type

Overview
Genes can be identified by a large number of different synonyms and looked up in a number of different databases. In order to provide database lookup of genes GeneLinker™ needs to know what database the imported gene identifiers refer to.

GeneLinker™ recognizes four different types of gene identifiers, corresponding to four different gene databases. These are:

1. Affymetrix identifiers: Referred to as probe set identifiers in Affymetrix literature. This is the Gene Database type to choose when you are importing data which originated on Affymetrix chips.

Examples: 100_g_at    41848_f_at    AFFX-BioB-3_at

See Affymetrix Identifiers for more information.

2. GenBank identifiers: GenBank accession numbers which refer to the GenBank sequence database maintained by NCBI.

Examples: AF111785    NM_002128    X12597

See GenBank Identifiers for more information.

3. UniGene identifiers: Cluster numbers which refer to the UniGene database
Examples: Hs.172028  Mm.3037  Rn.36437

See UniGene Identifiers for more information.

4. Custom identifiers: If your gene or spot identifiers do not fall into one of the categories above, we recommend you designate them as Custom identifiers. You may be able to instruct GeneLinker™ how to look up Custom identifiers by changing a setting in your User Preferences. See Changing Your User Preferences for more information.

Note: gene identifiers have a length restriction of 25 characters. This means that on import of a dataset or a gene list, identifiers that are longer than 25 characters are truncated.

Related Topics:
- How to Import Expression Data
- Importing One File Containing All Samples
- Importing Multiple Files With One Sample Each
- Lookup Gene

Importing Multiple Files With One Sample Each

Overview
It is assumed that you have already selected a multiple data files each containing a single sample type template (Affymetrix, CodeLink, DCHIP paired xls files, GenePix, Genomic Solutions, Quantarray, ScanArray) for data import (see Selecting a Template for Data Import or the appropriate Formats and Templates page). Follow the steps in this procedure to transfer your data from the files into the GeneLinker™ database.

If you selected a template that includes replicate merging, you may wish to read Merging Within-Chip Replicate Measurements for detailed information on this process.

For DCHIP paired xls files, there can be more than one sample per data file. In this case, samples are ordered according to their order in the first file. Samples that are present in one file but not the other will have missing values for the file they are missing from.

Actions
For these templates, the Data Import dialog looks like this:
• If you selected the incorrect template, Click the Template ... button to select the correct template.

• If the Gene Database is not correct, use the Gene Database drop-down list to set it to match the gene identifier type the genes in the data being imported have.

Select the Data Folder
All the data files for a given experiment must reside in a single folder.

• If the Source Folder listed on the Data Import dialog contains your data files and the data files are listed in the left list box, skip down to Choose Files for Import (below).

• If the Source Folder is incorrect, click the Source Folder ... button. The Open dialog is displayed

1. Navigate until the folder containing your data files is visible.
2. Click the folder name. The folder name is highlighted.
3. Click Select Folder. The Data Import dialog is updated with the selected folder name and the files in that folder are listed in the Source Files list box of the Data
Choose Files for Import

- **To select a single file**, click the file name.
- **To select multiple files**, press and hold the <Ctrl> key and click each file name.
- **To select a series of files**, press and hold the <Shift> key and click the first and last file names in the series.

1. Use the buttons between the list boxes to create an **Import Files** list in the right list box. The buttons between the left and right list boxes have the following functions:
   - The top button \(\rightarrow\) transfers the selected file(s) from the left to the right list box.
   - The second button \(\uparrow\) transfers the selected file(s) from the right to the left list box.
   - The third button \(\downarrow\) transfers **all** files (selected or not) from the left to the right list box.
   - The bottom button \(\leftarrow\) transfers **all** files (selected or not) from the right to the left list box.

2. **Order the import file list to be the sample order for the dataset that will be created.** Files are imported from the top of the list to the bottom. Use the buttons to order the files for import. The buttons to the right of the right list box have the following functions:
   - The top button \(\uparrow\) moves the selected file to the **top** of the list.
   - The second button \(\downarrow\) moves the selected file up one position in the list.
   - The third button \(\leftarrow\) moves the selected file down one position in the list.
   - The bottom button \(\rightarrow\) moves the selected file to the **bottom** of the list.
3. Click **Import**. The **Import Data** dialog is displayed.

- GeneLinker™ assumes that the number of genes is greater than the number of samples and orients the data so that the larger dimension (genes) is in columns. If this assumption is incorrect and the number of genes in your dataset is less than the number of samples, click the **Transpose** checkbox to pivot the data so that the larger dimension (samples) is in rows.
- If the first column and/or row contain text, GeneLinker™ uses the text as column
and/or row header names. If you have column and/or row names that are numeric, click the column and/or row name checkbox to indicate this to GeneLinker™.

4. When the data displayed in the Preview looks correct, click OK. Once the dataset has been successfully imported into the GeneLinker™ database, a new dataset item is added to the Experiments navigator.

**Notes**

If the name of the dataset being imported already exists in the Experiments navigator, the new dataset is given a new, unique name (a numerical identifier is appended to the original name) to make it distinct from the existing dataset.

If your data file is not in the correct format, the import process will fail. For complete file format details see:

- Importing Data from Affymetrix MAS 4.0 Files
- Importing Data from Affymetrix MAS 5.0 Files
- Importing Data from CodeLink XML Files
- Importing Data from dChip xls Files
- Importing Data from GenePix Files
- Importing Data from Genomic Solutions Files
- Importing Data from Quantarray Files
- Importing Data from Scanarray Files

**Related Topics:**

- Selecting a Template for Data Import
- Selecting the Gene Database Type
- Merging Within-Chip Replicate Measurements

**Importing One File Containing All Samples**

**Overview**

It is assumed that you have already selected a single, multi-sample data file type template (e.g. Tabular, DCHIP single xls file) for data import (see Selecting a Template for Data Import). Follow the steps in this procedure to transfer your data from the file into the GeneLinker™ database.

If you selected a template that includes replicate merging, you may wish to read Merging Within-Chip Replicate Measurements for more detailed information on that process.

**Actions**

Select the Data Folder and File, then Import.

1. Click the Source File ... button. The Open dialog is displayed.
2. Navigate to the correct folder and click the file to be imported. The file name is highlighted.

3. Click **Open**. The source file is listed on the **Data Import** dialog.

4. Select a **Gene Database** identifier from the drop-down list. This tells GeneLinker™ which type of gene identifier the genes being imported have. The options are GenBank, Affymetrix, UniGene and Custom.

5. Click **Import**. The **Import Data** dialog is displayed.
GeneLinker™ assumes that the number of genes is greater than the number of samples and orients the data so that the larger dimension (genes) is in columns. If this assumption is incorrect and the number of genes in your dataset is less than the number of samples, click the **Transpose** checkbox to pivot the data so that the larger dimension (samples) is in rows.

If the first column and/or row contain text, GeneLinker™ uses the text as column and/or row header names. If you have column and/or row names that are numeric, click the column and/or row name checkbox to indicate this to GeneLinker™.

<table>
<thead>
<tr>
<th></th>
<th>keratin</th>
<th>cellubrevin</th>
<th>nestin</th>
</tr>
</thead>
<tbody>
<tr>
<td>E11</td>
<td>1.7</td>
<td>5.75</td>
<td>2.53</td>
</tr>
<tr>
<td>E13</td>
<td>0.34</td>
<td>4.41</td>
<td>3.27</td>
</tr>
<tr>
<td>E15</td>
<td>0.52</td>
<td>1.19</td>
<td>5.2</td>
</tr>
<tr>
<td>E18</td>
<td>0.4</td>
<td>2.13</td>
<td>2.8</td>
</tr>
<tr>
<td>E21</td>
<td>0.68</td>
<td>2.3</td>
<td>1.5</td>
</tr>
<tr>
<td>P0</td>
<td>0.46</td>
<td>2.53</td>
<td>1.12</td>
</tr>
<tr>
<td>P7</td>
<td>0.32</td>
<td>3.89</td>
<td>0.53</td>
</tr>
<tr>
<td>P14</td>
<td>0.08</td>
<td>3.95</td>
<td>0.51</td>
</tr>
<tr>
<td>A</td>
<td>0.0</td>
<td>2.72</td>
<td>0.44</td>
</tr>
</tbody>
</table>
6. When the data displayed in the **Preview** looks correct, click **OK**. Once the dataset has been successfully imported into the GeneLinker™ database, a new dataset item is added to the **Experiments** navigator.

**Notes**

If the name of the dataset being imported already exists in the **Experiments** navigator, the new dataset is given a new, unique name (a numerical identifier is appended to the original name) to make it distinct from the existing dataset.

If your data file is not in the correct format, the import process will fail. For complete file format details see Importing Data from Tabular Files or Importing Data from dChip xls Files, as appropriate.

**Related Topics:**

- Selecting a Template for Data Import
- Selecting the Gene Database Type
- Merging Within-Chip Replicate Measurements

**Merging Within-Chip Replicate Measurements**

**Overview**

Certain import templates allow you to merge replicate genes occurring on the same chip into a single measurement. When this is done, GeneLinker™ uses the spread between the replicates to estimate a reliability measure for the resulting (average) measurement.

The statistical method used to merge replicate genes and generate a reliability measure is designed for use with small numbers of replicates (as few as two) and to give usable results even if there are missing data. To achieve this, the method assumes that the variability between the replicate measurements increases proportional to the abundance of the gene product, but otherwise has a roughly normal (Gaussian) distribution which is the same across all genes on the chip.

The figure below plots the difference between replicates against the average abundance (in arbitrary units) for a typical experiment with within-chip duplicate measurements. Notice that genes with greater abundance tend to have greater difference between the replicates.
By scaling the replicates according to the abundance, we obtain the plot in the figure below. Note how the scaled residuals tend to be large when the average abundance is near zero. This is to be expected since measurements near the detection threshold are relatively more error-prone.

The resulting distribution of residuals has the shape of a 'bell curve' but has very long tails representing measurements with abnormally high variation between the replicates. In statistical terms, this example has a very large kurtosis.
The integral of the tails of this distribution can be interpreted loosely as the probability of getting such an extreme residual by chance. We compute this probability and then take its complement in order to put this reliability measure on the same scale as the P-values many researchers are accustomed to. A value near zero means a reliable measurement; a value near one means an unreliable measurement.

Detailed Algorithm Used to Merge Within-Chip Replicate Measurements On Import

Here is a detailed description of the algorithm used to merge within-chip replicate measurements on import.

1. Read $x[\text{chip, gene, rep}]$ from datafile
2. Compute $\text{abundance}[\text{chip, gene}] = \text{mean}(x[\text{chip, gene,}])$
3. Save the abundance as the GeneLinker™ expression measurement
4. Compute $\text{resid}[\text{chip, gene, rep}] = (x[\text{chip, gene, rep}] - \text{abundance}[\text{chip, gene}]) / \text{abundance}[\text{chip, gene}]$. These are the residuals plotted in the Figures 2 and 3 above.
5. Compute $s = \text{stdev}(\text{resid}[; ; ;])$
6. Set $r[\text{chip, gene}] = \max(\text{abs}(\text{resid}[\text{chip, gene,}]))$ and compute the integral under the normal curve $N(0, s)$ between $-|r|$ and $+|r|$.
   - This step is quite conservative if you have more than three replicates, essentially taking the most extreme replicate as an indicator of the quality of the whole set.
7. Save this integral $p[\text{chip, gene}]$ as the GeneLinker™ reliability measure

If due to missing data there are no replicates for a given chip/gene pair, then that measurement is arbitrarily assigned a reliability measure of zero (perfectly reliable). Therefore measurements for which you have no reliability information will not be filtered out by the Value Removal by Reliability Measure operation.

Naturally the assumptions of this model may be tested if you have enough replicates for
each condition and gene. If you have more than three replicates and you feel this model is inappropriate, we recommend you use general-purpose statistical software to preprocess your data outside GeneLinker™, merging replicates before importing it in tabular format. You may eliminate unreliable measurements from the dataset before using the Tabular import template, or you may compute reliability measures and import them along with the expression data using the Tabular with Reliability Measures import template.

**Related Topics:**
- Creating a Table View of Reliability Data
- Removing Values by Reliability Measure

**Two-Color Data**

**Overview**

Many microarray experiments are carried out on paired samples, a *treatment* sample and a *control* sample, and the resulting expression levels measured on the same chip with two different fluorescent dyes. The most common fluorescent dyes used are Cy3 (green) and Cy5 (red), so these experiments are referred to as two-color experiments, Cy3/Cy5 experiments, or red/green experiments.

GeneLinker™ can carry out certain operations when it has both the treatment and control measurements, operations it cannot carry out if it has only the ratios. In GeneLinker™, we refer to a dataset which has both treatment and control values stored as **Two-Color Data**. In the description pane for such a dataset it will say **Two Channels Available: Yes. If the description pane does not say this, then GeneLinker™ does not have the required two values for each spot and cannot treat the data as Two-Color Data.** If you believe you imported two-color data but the description pane says **Two Channels Available: No**, re-examine your data and your choice of a data import template. Two-Color Data can be imported using GenePix, Quantarray and Scanarray templates, but not all templates of those types import two-color data. Please see the appropriate Formats and Templates pages for more information.

Certain operations are possible on Two-Color Data which are not applicable to regular data. These operations include Lowess Normalization and the Intensity-Bias Plot. When you make a table view, color matrix plot, or other visualization of a table with two channels available, the data displayed are the ratios.

**Related Topics:**
- Selecting a Template for Data Import
- Importing Data from GenePix Files
- Importing Data from Quantarray Files
- Importing Data from Scanarray Files
Reliability Measures

Overview
A reliability measure in GeneLinker™ is a numerical indication of the quality or reliability of a measurement of an individual gene’s expression in an individual sample. GeneLinker™ expects reliability measures to fall between 0 and 1, with 0 representing very reliable and 1 representing unreliable. This is patterned off the interpretation of p-values in traditional statistical tests, where small numbers indicate significance.

Reliability measures can come from several sources:
Some microarray analysis programs can generate an estimate of the measurement of each spot on each chip. For example, Affymetrix MAS 4.0 can export a Call with a value of Present (P), Marginal (M), or Absent (A) for each spot. Affymetrix MAS 5.0 can export a Detection p-value which lies between zero (definitely present) and one (definitely absent).
If you have microarray data which replicates genes on a single chip, some of GeneLinker™’s import templates can convert those replicated values into a merged (averaged) value and an associated reliability measure. See Merging Within-Chip Replicate Measurements for more information.
Finally, you can generate reliability measures yourself in tabular format and import them in concert with tabular data by choosing the Tabular With Reliability Measures import template.

Related Topics:
- Creating a Table View of Reliability Data
- Removing Values by Reliability Measure
- Importing One File Containing All Samples
- Importing Multiple Files With One Sample Each

Variables

Variables Overview

Overview
Definition of a Variable
In GeneLinker™, a variable is a column of data other than gene expression values used to differentiate samples. A variable can store:

- Phenotypic observations about the samples. e.g. malignant vs. benign.
- Predictions of phenotypes by a trained classifier.
e.g. predicted malignant vs. predicted benign.

- Information about experimental conditions.
  e.g. high dose vs. low dose; time the sample was taken; animal A vs. animal B vs. animal C, etc.

Variable File Formats

One-column: A one-column format file consists of the class name of each sample, one per line, in the same sample order as in the expression data file. **The first row must not contain a column header.**

Two-column: The two-column format has the sample names in the first column and the variable values (class names) in the second. The two-column format can be tab-separated or comma-separated. If you want class names which include commas, you must use two-column format with tab separators between the sample names and class labels. **The first row must contain column headers.**

<table>
<thead>
<tr>
<th>EWS</th>
<th>Sample Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>EWS</td>
<td>EWS-T1 EWS</td>
</tr>
<tr>
<td>EWS</td>
<td>EWS-T2 EWS</td>
</tr>
<tr>
<td>EWS</td>
<td>EWS-T3 EWS</td>
</tr>
<tr>
<td>EWS</td>
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</tr>
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<tr>
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<td>BL-C3 BL</td>
</tr>
<tr>
<td>BL</td>
<td>BL-C4 BL</td>
</tr>
</tbody>
</table>

Uses of a Variable

Variables can be used many ways in GeneLinker™.

- You can color the samples in certain plots by a variable.
- A variable can group replicates together for statistical differentiation using the F-Test. All members of the same group have the same variable value.
- SLAM™ can search for gene sets associated with the values of a variable.
- A variable can be used as training data for an ANN classifier or an IBIS classifier and a trained classifier can predict the values of a variable for new samples.
- Two variables of the same type can be compared using a confusion matrix.

Note on the Value 'Unknown'

Any GeneLinker™ variable may take on the special value of 'Unknown'. In the output of a trained classifier, this means that the classifier could not make a reliable prediction of the sample class. In other contexts, 'Unknown' is treated in the same manner as any other class. To reduce confusion we recommend that you use more informative class
labels and reserve 'Unknown' for the output of the classifier.

Variable Types

Variables which attempt to describe the same phenomenon are grouped together into a Variable Type. GeneLinker™ does not intuit which variables refer to the same phenomenon the way a person does, so you must define a variable type for each variable you import.

• For example, variables of type 'leukemia class' might have possible values of 'myeloblastic' and 'lymphoblastic'. Once you have created the variable type 'leukemia class', you could then import variables of that type like 'Diagnosis of pathologist A', 'Diagnosis of pathologist B', etc. You could then go on to train GeneLinker™ to classify the samples by leukemia type, and use GeneLinker™ to construct further variables like 'Prediction based on gene Q', ‘Prediction based on a set of 10 genes', and so on.

If you wished to study disease outcomes with the same expression dataset, you could define a new variable type 'outcome' which might have values such as 'survived' and 'died'. You could then import a variable of that type, train classifiers and attempt further predictions.

Observed vs. Predicted Variables

In GeneLinker™, imported variables are referred to as observed variables, and variables generated by a classifier are predicted. You can see the values of any or all of the variables associated with a given dataset using the Variable Viewer. You can edit, delete, compare or export variables using the Variable Manager.

Variable Indicator

In the Experiments navigator, a root dataset that has one or more variables associated with it has the variables tag on the icon next to its name. The same variables are associated with all the descendants of this dataset.

for a complete dataset.

for an incomplete dataset.

Variables and Classification

Variables are typically imported into GeneLinker™ for one of two purposes related to Classification: A variable may be a training target, providing known classes for training a classifier, or a variable may be a set of test results for comparison with the predictions of a trained classifier. Note that for a given prediction problem, both the training variable and the test variable must be imported as the same Variable Type.

Related Topics:

Importing Variables
Variable Viewer
Variable Manager
Variables in Supervised Learning
Importing Variables

Overview
See Variables Overview for a detailed discussion of variables.

Actions
1. Click a dataset in the Experiments navigator. The item is highlighted.
2. Select Import from the File menu and Variable from the sub menu. The Import Variables dialog is displayed.
   • The dataset that the variable information applies to is displayed as the Dataset. Variable information applies to all datasets in a branch of the Experiments navigator tree.
   • The number of samples in the dataset is shown under the dataset name.
   • All existing variable types are displayed in the Choose a Variable Type box.
   • All existing classes in the selected variable type are listed in the box on the right.
3. The Source File for the variable data is listed just below the Dataset. To set the source file, click the ... button. This displays the Open dialog.
4. Navigate to the correct folder and click on the variable data file name.
5. Click Open. The source file name is displayed on the Import Variables dialog and the number of observations in the file is listed. The number of observations in the file must match the number of samples in the dataset.
   • GeneLinker™ supplies a variable name and description. They are displayed at the bottom of the dialog.
   • If there are existing variable types, GeneLinker™ compares the classes in the new variable file to the classes of the existing types. If the classes are contained within an existing variable type, a message is displayed indicating this.
   • If no variable type exists, the Create Variable Type dialog is displayed. See the
section **Create Variable Type** below for instructions on how to do this.

**Preview**

To preview the contents of the new variable file, click the **Preview** button. The **Import Variable** preview dialog is displayed.

- The name of the variable file is displayed at the top.
- The class entries in the file are displayed under the **Preview** heading in the order they exist in the file. The scrollbar can be used to look through the entire list.
- The **Class Summary** on the right lists the names of all the classes and gives a count for each.
- Click **Close** to return to the **Import Variables** dialog.

**Create Variable Type**

To create a new variable type, click **New Variable Type** (or if there are no existing types, this dialog will be displayed automatically).

- Enter a name for the variable in the **Variable Name** text box.
- Optionally, enter a description for the variable in the **Variable Description** text box.
- Click **OK** to return to the **Import Variables** dialog.
- The 'unknown' class is automatically added to all new variable types. It will be listed on the **Import Variables** dialog.

6. Click **Import**. The variable data is imported into the database, and in the **Experiments** navigator, the dataset icon is marked with the variable tag (☑ for a complete dataset or ☐ for an incomplete dataset).
Related Topics:
- Variables Overview
- Variable Manager
- Variable Viewer

Variable Viewer

Overview
The variable viewer displays a list of all the variable types associated with the selected dataset. It also shows the relationships between the samples and the classes of the selected variable type(s).

Actions
1. Click on a dataset that has associated variable information (it is tagged with one of the variable icons - a complete dataset or an incomplete dataset) in the Experiments navigator. The item is highlighted.
2. Click the Variable Viewer toolbar icon, or select Variable Viewer from the Explore menu, or right-click the item and select Variable Viewer from the shortcut menu. The Variable Viewer is displayed.

Dataset Variables Table (left):
- The first column has checkboxes for selecting variable types to be displayed in the sample and class table. The second column lists all of the variable types associated with the dataset.

Sorting the Left Table by Variable Type
a) Click on the Variable column header. The table is sorted in ascending order and an upward pointing triangle is displayed in the column header.

b) Click on the Variable column header again to sort in descending order. A downward pointing triangle is displayed in the column header.
**Note:** sorting the left table does not affect the right table.

**Sample and Class Table (right):**
- The first column contains the index for the sample in the dataset. The sample names are listed in the second column. Each subsequent column, labelled with the variable type it describes, contains sample-specific class entries.

**Sorting the Right Table by Sample Index**
- a) Click on the **Sample index** column header. The table is sorted in ascending order and an upward pointing triangle is displayed in the column header.
- b) Click on the **Sample index** column header again to sort in descending order. A downward pointing triangle is displayed in the column header.

**Sorting the Right Table by Sample Name**
- a) Click on the **Sample name** column header. The table is sorted in ascending order and an upward pointing triangle is displayed in the column header.
- b) Click on the **Sample name** column header again to sort in descending order. A downward pointing triangle is displayed in the column header.

**Sorting the Right Table by Variable Type**
- a) Click on a variable type column header. The table is sorted in ascending order and an upward pointing triangle is displayed in the column header.
- b) Click on the same variable type column header again to sort in descending order. A downward pointing triangle is displayed in the column header.

**Related Topics:**
- Variables Overview
- Importing Variables

**Variable Manager**

**Overview**
The Variable Manager is used to view, edit, delete, or export variable data or to display a confusion matrix of variables associated with the selected dataset.

**Actions**
1. Click a dataset that has an associated variable (it is tagged with one of the variable icons - a complete dataset or an incomplete dataset) in the **Experiments** navigator. The dataset is highlighted.
2. Select **Variable Manager** from the **Tools** menu. The **Variable Manager** is displayed.

![Variable Manager Image]

**Editing a Variable**

a) Click on a variable name. The item is highlighted.

b) Click the **Edit** button. The **Edit Variable** dialog is displayed.

c) Type in a new name and/or description for the variable.

d) Click **OK** to keep the changes, or click **Cancel** to keep the original name and description.

**Deleting a Variable**

a) Click on a variable name. The item is highlighted.

b) Click the **Delete** button.

c) The variable is deleted

- If the variable being deleted is a prediction associated with a classification, the classification is also deleted. **Note:** the reverse is also true - that is, if you delete a classification the predicted variable is also deleted.

**Exporting a Variable**

a) Click on a variable name. The item is highlighted.

b) Click the **Export** button. The **Save As** dialog is displayed.
c) Type in a name for the data file or accept the file name GeneLinker™ generates.

d) Click **Save**. The variable data is exported to the file in two-column format. For example:

```
Sample  var1
EWS-T1  EWS
EWS-T2  EWS
EWS-T3  EWS
EWS-T4  EWS
BL-C05  BL
BL-C06  BL
BL-C07  BL
BL-C08  BL
BL-C09  BL
RMS-C01 RMS
RMS-C02 RMS
RMS-C03 RMS
RMS-C04 RMS
```

Related Topics:

- Displaying a Confusion Matrix
- Variables Overview

**Viewing, Renaming, Deleting**

**Creating a Table View of Gene Expression Data**

**Overview**

Datasets can be viewed by displaying them in a spreadsheet-like table. Genes are in columns and samples are in rows. If a gene does not have an identifier of the type specified for display in the user preferences, it is displayed in the column label using the gene identifier type that was imported for that gene.
• For a regular dataset, each cell in the table contains the expression level of that gene (gene name in column label) in that sample (sample name in row label).

• For a two-color dataset, each cell in the table contains a ratio expression level (Cy5/Cy3) of that gene in that sample.

• A missing value is blank.

• Selected column(s) or row(s) are displayed in dark blue with white text. See Interacting With the Table Viewer for full details on Table Viewer functions.

Actions
1. Click a dataset in the Experiments navigator. The item is highlighted.
2. Click the Table View toolbar icon , or select Table View from the Explore menu, or right-click the item and select Table View from the shortcut menu. A table view of the dataset is displayed.

Related Topics:
Interacting With the Table Viewer
Find
Creating a Gene List

Creating a Table View of Reliability Data

Overview
Reliability measures for a dataset can be viewed using the table viewer.

**Actions**
1. Click on a dataset that has reliability measures associated with it in the Experiments navigator. The item is highlighted.
2. Select Reliability Measures from the Statistics menu, or right-click the item and select Reliability Measures from the shortcut menu. A table view of the reliability data is displayed.

**Related Topics:**
- Reliability Measures
- Removing Values by Reliability Measure

**Table Viewer Functions**

**Overview**
The Table Viewer displays the gene expression values for the selected dataset. You can select a single, multiple, or a series of genes or samples for display in a Coordinate Plot or Summary Statistics chart. If you select a pair of genes or samples, you can display a Scatter plot. A selection of genes also can be used to create a gene list.

**Actions**
1. Click on a dataset in the Experiments navigator pane. The dataset is highlighted.
2. Click the Table View toolbar icon, or right-click the item and select Table View. The dataset is displayed in a table.

**Making Selections**
Genes are assumed to be in columns; samples are assumed to be in rows.
• **Selecting a single column or row:** click on the column or row header.

• **Selecting multiple columns or rows:** press and hold <Ctrl>, then click the column or row headers.

• **Selecting a series of columns or rows:** press and hold <Shift>, then click on the first and last column or row headers.
  
  • De-selecting an item within a series: release the <Shift> key and hold the <Ctrl> key and click on the item(s) to be de-selected. The rest of the series remains selected.

To use the highlighted items in a plot, right-click on the table viewer and select from the shortcut menu.

• If column(s) are selected, genes will plot (as series) across (all) samples.

• If row(s) are selected, samples will plot (as series) across (all) genes.

• You cannot selectively plot specific genes against specific samples (i.e. you cannot select columns and rows concurrently).

**Resizing the Columns**

The columns in the table viewer are equal in width, so when you perform a column width adjustment, it affects all columns equally. Note that on large datasets, resizing the columns can be slow.

1. Position the mouse cursor on the divider between two column names. The cursor is drawn as a two headed arrow.

2. Click and drag right to widen the columns, or drag left to shrink the columns.

**Related Topics:**

  - Data Import Step 1: Selecting a Template
  - Creating Gene Lists from Selections

**Creating a Color Matrix Plot**

**Overview**

A color matrix plot is used to visualize the values in a dataset. The plot consists of a legend at the top and a grid of colored cells, with the genes in the columns and the samples in the rows. The legend consists of a color gradient above an expression value scale. The default range for the scale is from the minimum to the maximum value contained within the dataset.

Missing values are colored using the color value at the mid-point of the scale and have a white ‘X’ drawn through the colored tile (this is only visible if the dimensions of the colored tiles are large enough to display it).

**Note:** you cannot create a color matrix plot for an experiment (clustered dataset). For those, create a Matrix Tree Plot.
**Actions**

1. Double-click a dataset (raw, preprocessed, discretized, etc.) in the **Experiments** navigator. The item is highlighted and a color matrix plot of the dataset is displayed.

   OR

1. Click a dataset in the **Experiments** navigator. The dataset is highlighted.

2. Click the **Color Matrix Plot** toolbar icon, or select **Color Matrix Plot** from the **Explore** menu, or right-click the item and select **Color Matrix Plot** from the shortcut menu. A color matrix plot of the dataset is displayed.

**Plot Indicators**

- As you move the mouse pointer over a gene or sample name, a gray bounding box is drawn around its column or row so you can easily see which tiles belong to it.
- The names of one or more selected genes or samples are highlighted in dark blue with white text. It is not possible to select genes and samples concurrently.

**Interacting With the Plot**

Selecting Items
Displaying a Gene Expression Value

**Plot Functions**

Profile Matching
Color by Gene Lists or Variables
Exporting an Image

**Customizing the Plot**

Changing the Gradient Color and Scale
Resizing Cells in a Color Grid
Toggling the Color Grid On or Off

**Related Topic:**

GeneLinker Gold 3.1 / GeneLinker Platinum 2.1
Preprocessing
Eliminating and Estimating Missing Values

Overview of Estimating Missing Values

Overview
Missing (null) values can lead to erroneous conclusions about data. Similarly, substitution of missing values may introduce inaccuracies and inconsistencies. Missing data values can negatively impact discovery results, and errors or data skews can proliferate across subsequent runs and cause a larger, cumulative error effect. As well, most analysis methods cannot be performed if there are missing values in the data.

Missing values may prevent proper classification, and poor substitution schemes for missing values may cause classification errors. If all the values substituted are determined by the most likely value, then the individual values are less likely to help define class (cluster) boundaries.

Actions
Two Step Process for Resolving Missing Values:
1. **Remove** (filter out) genes that have a minimum number of missing values.
   - Eliminate genes with a high number of missing values, since estimating high numbers of missing values may introduce bias to further analysis. The criteria to eliminate genes with missing values may be situation-dependent.
   - If you set the elimination threshold value to 1, all genes with missing values are removed.
2. **Replace** the remaining missing values. GeneLinker™ offers three techniques for estimating missing values:
   - Estimating values by a measure of central tendency;
   - Estimating missing values by nearest neighbors;
   - Replacing missing values with an arbitrary value.

Related Topics:
- Estimating Missing Values by a Measure of Central Tendency
- Missing Value Estimation by Nearest Neighbors
- Replacing Missing Values With an Arbitrary Value
Overview

The process of handling missing values consists of two steps: first, genes that have a minimum number of missing values are removed; and second, the remaining missing values are estimated using a measure of central tendency (mean or median).

On the Estimate Missing Values dialog, when the Remove Genes That Have Missing Values slider is set to 1, the rest of the dialog is grayed out. This is because all genes that have at least one missing value will be removed leaving no missing values to be estimated.

Actions

1. Click an incomplete dataset in the Experiments navigator. The item is highlighted.
2. Click the Missing Value Estimation toolbar icon, or select Estimate Missing Values from the Data menu, or right-click the item and select Estimate Missing Values from the shortcut menu. The Estimate Missing Values dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remove Genes That Have Missing Values</td>
<td>Set the threshold for culling genes prior to missing value estimation (1 = remove all genes that have missing values).</td>
</tr>
<tr>
<td>Replacement Technique</td>
<td>Select Measure of Central Tendency.</td>
</tr>
<tr>
<td>Options</td>
<td>Select which measurement to use: Median or Mean.</td>
</tr>
</tbody>
</table>

4. Click OK. The Experiment Progress dialog is displayed. It is dynamically updated as the Estimate Missing Values operation is performed. To cancel the Estimate Missing
Values operation, click the **Cancel** button.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the resulting dataset will be empty.
- Upon successful completion, a new dataset is added under the original dataset in the **Experiments** navigator.

**Related Topics:**
- Overview of Estimating Missing Values
- Nearest Neighbors Missing Value Estimation

### Nearest Neighbors Missing Value Estimation

**Overview**

The process of handling missing values consists of two steps: first, genes that have a minimum number of missing values are removed; and second, the remaining missing values are estimated using Nearest Neighbors estimation.

Nearest Neighbors estimation is a process by which missing values in a dataset are filled in with estimated values based on similarity between genes.

To estimate a missing value in a gene, the k genes with the closest profile (smallest distance) to the gene containing the missing value are determined. The missing value is then computed as a weighted average of the k values in that sample of the neighbors. **Note:** the k nearest neighbors can be computed only on complete datasets. Missing values have to be filled in with an initial approximation. The distance between two genes is computed using either Euclidean distance or Pearson Correlation.

The input to this function is an incomplete dataset; the output is a complete dataset. K is an integer representing the number of nearest neighbors to be taken into consideration.

On the **Estimate Missing Values** dialog, when the **Remove Genes That Have Missing Values** slider is set to 1, the rest of the dialog is grayed out. This is because all genes that have at least one missing value will be removed leaving no missing values to be estimated.

**Process Outline**

- All missing values in the selected dataset are initially approximated with their gene's mean.
- For each gene, the distances to all other genes are computed.
- For each gene, select the k genes with the smallest distances to it.
- Replace each value that was missing in the gene with the weighted average of the k
values belonging to the k nearest genes in the same sample.

**Actions**

1. Click an incomplete dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Estimate Missing Values** toolbar icon, or select **Estimate Missing Values** from the **Data** menu, or right-click the item and select **Estimate Missing Values** from the shortcut menu. The **Estimate Missing Values** dialog is displayed.

3. Set the parameters.

<table>
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<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Remove Genes That Have Missing Values</td>
<td>Set the threshold for culling genes prior to missing value estimation (1 = remove all genes with missing values).</td>
</tr>
<tr>
<td>Replacement Technique Options</td>
<td>Select <strong>Nearest Neighbors Estimation</strong>.</td>
</tr>
<tr>
<td></td>
<td>Set the <strong>Distance Metric</strong> to Euclidean or Pearson Correlation.</td>
</tr>
<tr>
<td></td>
<td>Set the <strong>Number of Nearest Neighbors</strong>.</td>
</tr>
</tbody>
</table>

4. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Estimate Missing Values operation is performed. To cancel the Estimate Missing Values operation, click the **Cancel** button.

Upon successful completion, a new complete dataset is added under the original dataset in the **Experiments** navigator.
Replacing Missing Values with an Arbitrary Value

Overview
The process of handling missing values consists of two steps: first, genes that have a minimum number of missing values are removed; and second, the remaining missing values are replaced with an arbitrary value.

On the Estimate Missing Values dialog, when the Remove Genes That Have Missing Values slider is set to 1, the rest of the dialog is grayed out. This is because all genes that have at least one missing value will be removed leaving no missing values to be estimated.

Actions
1. Click an incomplete dataset in the Experiments navigator. The item is highlighted.
2. Click the Estimate Missing Values toolbar icon or select Estimate Missing Values from the Data menu, or right-click the item and select Estimate Missing Values from the shortcut menu. The Estimate Missing Values dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
</table>

GeneLinker Gold 3.1 / GeneLinker Platinum 2.1
Remove Genes That Have Missing Values

Set the threshold for culling genes prior to missing value estimation (1 = remove all missing values).

Replacement Technique

Select Arbitrary Value for All Genes.

Options

Set the Replacement Value.

4. Click OK. The Experiment Progress dialog is displayed. It is dynamically updated as the Estimate Missing Values operation is performed. To cancel the Estimate Missing Values operation, click the Cancel button.

Upon successful completion, a new complete dataset is added under the original dataset in the Experiments navigator.

Related Topics:
Overview of Estimating Missing Values
Nearest Neighbors Missing Value Estimation

Filtering

Filtering Overview

Overview
Filtering provides a number of gene prioritization options. The processes generally take a large number of genes and apply selection criteria so that the output includes fewer genes.

Some methods remove all of the genes that do not meet specified criteria, while others allow you to specify the number of genes that will be left after the filtering.

Filtering and normalization processes can be applied one or more times to a dataset.

Note that for Affymetrix® data, it is recommended that genes with a high signal-to-noise ratio be used, since some experts believe that Affymetrix® values below 150 tend to be unreliable.

Complete and Incomplete Datasets

The only filtering operation that can be applied directly to an incomplete dataset is gene list filtering. If you do not have a gene list that contains one or more genes in the incomplete dataset, the gene list filtering option is disabled on the Filter Genes dialog. To resolve this, close the Filter Genes dialog, create an appropriate gene list, and then perform the gene list filtering operation.

To apply other filtering techniques to an incomplete dataset, the missing values first
need to be estimated or eliminated (resulting in a complete dataset). All filtering techniques can be applied to complete datasets.

**Note on N-Fold Culling**

N-Fold Culling cannot complete and displays a message if the minimum value for any gene is 0.0 ('The experiment could not be completed. Check that the operation and its parameters are appropriate to the data.') If the dataset contains negative values (but no zeroes) no error message is displayed, but N-Fold Culling may remove highly-changing genes.

**Both these problems can be avoided this way:**

Before applying N-Fold Culling, display a Summary Statistics chart of the dataset to see what its minimum value is. If it is zero or negative, then:

1. Use Remove Values to remove values less than some small threshold (e.g. the smallest positive value your equipment can meaningfully detect).
2. Use Missing Value Estimation to replace the removed values with some small positive constant (e.g. the same number used as a removal threshold).

**Filtering Techniques Available in GeneLinker™:**

- Maximum Culling
- Range Culling
- N-Fold Culling with N
- N-Fold Culling with a Specified Number of Genes
- Spotted Array N-Fold Culling
- Gene List Filtering
- F-Test

**Related Topic:**

- Overview of Estimating Missing Values

**Maximum Culling**

**Overview**

Maximum culling retains the specified number of genes that have the highest absolute values. The maximum value associated with each gene is calculated, and the specified number of genes with the highest expression values are retained. All others are culled.

**Actions**

1. Click a dataset in the **Experiments** navigator. The item is highlighted.
2. Click the Filter toolbar icon, or select **Filter Genes** from the Data menu, or right-click the item and select **Filter Genes** from the shortcut menu. The **Filter Genes** dialog is displayed.
3. Select Maximum Culling from the Filtering Operation drop-down list.
4. Set the number of genes to be retained in the Number of genes to keep field.
5. Click OK. The Experiment Progress dialog is displayed. It is dynamically updated as the Maximum Culling operation is performed. To cancel the Maximum Culling operation, click the Cancel button.

Upon successful completion, a new dataset is added under the original dataset in the Experiments navigator.

Related Topic:
Filtering Overview

Range Culling

Overview
Range culling retains the genes that have the largest ranges in values. The maximum and minimum expression values associated with each gene are calculated, and the range is calculated as the maximum - minimum.

The number of genes specified by the user that have the largest ranges are retained. All others are culled.

Actions
1. Click a complete dataset in the Experiments navigator. The item is highlighted.
2. Click the Filter toolbar icon, or select Filter Genes from the Data menu, or right-click the item and select Filter Genes from the shortcut menu. The Filter Genes dialog is displayed.
3. Select **Range Culling** from the **Filtering Operation** drop-down list.
4. Enter the number of genes that will be retained in the **Number of genes to keep** field.
5. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Range Culling operation is performed. To cancel the Range Culling operation, click the **Cancel** button.

Upon successful completion, a new dataset is added under the original dataset in the **Experiments** navigator.

**Related Topic:**
- Filtering Overview

**N-Fold Culling with N**

**Overview**
This operation allows you to specify a minimum n-fold change that must occur in a gene so that it is retained. For example, if you specified an n-fold of 2.5, any genes that do not show an n-fold increase over the samples of at least 2.5 would be culled.

The maximum and minimum expression values associated with each gene are calculated and the n-fold for that gene is calculated as the maximum/minimum.

N-Fold Culling is intended to be applied to positive abundance data, not to ratio data (for which you should use Spotted Array N-Fold Culling) or to log ratio data (for which you should use Range Culling).

**How to Handle Negative or Zero Values**
This operation cannot complete and displays a message if the minimum value for any gene is 0.0 ("The experiment could not be completed. Check that the operation and its parameters are appropriate to the data.") If the dataset contains negative values (but no zeroes) no error message is displayed, but N-Fold Culling may remove highly-changing genes.
Both these problems can be avoided this way:

Before applying N-Fold Culling, display a Summary Statistics chart of the dataset to see what its minimum value is. If it is zero or negative, then:

1. Use Remove Values to remove values less than some small threshold (e.g. the smallest positive value your equipment can meaningfully detect).
2. Use Missing Value Estimation to replace the removed values with some small positive constant (e.g. the same number used as a removal threshold).

Actions

1. Click a complete dataset in the Experiments navigator. The item is highlighted.
2. Click the Filter toolbar icon or select Filter Genes from the Data menu, or right-click the item and select Filter Genes from the shortcut menu. The Filter Genes dialog is displayed.
3. Select the N-Fold culling with N operation from the Filtering Operation drop-down list.
4. Enter the minimum n-fold change to be retained, in the N-Fold min/max ratio field.
5. Click OK. The Experiment Progress dialog is displayed. It is dynamically updated as the N-Fold Culling With N operation is performed. To cancel the N-Fold Culling With N operation, click the Cancel button.

Upon successful completion, a new dataset is added under the original dataset in the Experiments navigator.

Related Topic:
Filtering Overview

N-Fold Culling With a Specified Number of Genes

Overview

This operation allows you to retain a specified number of genes that have the highest n-
fold increases in their expression values.
The maximum and minimum expression values associated with each gene are calculated and the n-fold for that gene is calculated as the maximum/minimum. The number of genes specified that have the largest n-folds are retained. All others are culled.

N-Fold Culling is intended to be applied to positive abundance data, not to ratio data (for which you should use Spotted Array N-Fold Culling) or to log ratio data (for which you should use Range Culling).

How to Handle Negative or Zero Values
This operation cannot complete and displays a message if the minimum value for any gene is 0.0 ('The experiment could not be completed. Check that the operation and its parameters are appropriate to the data.') If the dataset contains negative values (but no zeroes) no error message is displayed, but N-Fold Culling may remove highly-changing genes.

Both these problems can be avoided this way:
Before applying N-Fold Culling, display a Summary Statistics chart of the dataset to see what its minimum value is. If it is zero or negative, then:
1. Use Remove Values to remove values less than some small threshold (e.g. the smallest positive value your equipment can meaningfully detect).
2. Use Missing Value Estimation to replace the removed values with some small positive constant (e.g. the same number used as a removal threshold).

Actions
1. Click a complete dataset in the Experiments navigator. The item is highlighted.
2. Click the Filter toolbar icon or select Filter Genes from the Data menu, or right-click the item and select Filter Genes from the shortcut menu. The Filter Genes dialog is displayed.

3. Select N-Fold Culling with Number of Genes in the Filtering Operation drop-down list.
4. In the Number of genes to keep field, type in the number of genes to be retained.
5. Click OK. The Experiment Progress dialog is displayed. It is dynamically updated as the N-Fold Culling With a Specified Number of Genes operation is performed. To cancel the N-Fold Culling With a Specified Number of Genes operation, click the Cancel button.
Upon successful completion, a new dataset is added under the original dataset in the **Experiments** navigator.

**Related Topic:**
- Filtering Overview

**Spotted Array N-Fold Culling**

**Overview**
This operation keeps all genes that have an n-fold induction or repression above a user specified value. Genes are kept if they have at least one value greater than or equal to x or one value less than or equal to 1/x.

**Note** that an x value of less than or equal to 0.0 is not allowed.

**Actions**
1. Click a complete dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Filter** toolbar icon, or select **Filter Genes** from the **Data** menu, or right-click the item and select **Filter Genes** from the shortcut menu. The **Filter Genes** dialog is displayed.
3. Select **Spotted Array N-Fold culling** from the **Filtering Operation** drop-down list.
4. Set the value of ‘x’ in the **Induction/repression threshold** field.
5. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Spotted Array N-Fold Culling operation is performed. To cancel the Spotted Array N-Fold Culling operation, click the **Cancel** button.
Upon successful completion, a new dataset is added under the original dataset in the Experiments navigator.

Related Topic:
Filtering Overview

Gene List Filtering (Subsetting)

Overview
Gene List filtering can be used to reduce the number of genes (features) for exploration and analysis. Gene list filtering can be applied to complete or incomplete datasets. To apply gene list filtering to a dataset, at least one gene list for that dataset must exist.

Actions
1. Click a dataset in the Experiments navigator. The item is highlighted.
2. Click the Filter toolbar icon, or select Filter Genes from the Data menu, or right-click the item and select Filter Genes from the shortcut menu.
   • For a complete dataset, this Filter Genes parameters dialog is displayed.

   ![Filter Genes dialog]

   • For an incomplete dataset, this Filter Genes dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Element</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Filtering Operation</td>
<td>Set this to Gene List Filtering (for incomplete datasets this is the only option).</td>
</tr>
<tr>
<td>Filtering Option</td>
<td>Set to keep or remove genes listed in the gene list.</td>
</tr>
<tr>
<td>Gene List</td>
<td>The name of the gene list to be used to filter the dataset.</td>
</tr>
</tbody>
</table>
Note: only the gene lists relevant to the dataset are visible in the drop-down list. If no gene lists are available for the selected dataset, this operation cannot be performed. Create a gene list for the dataset and then apply gene list filtering.

4. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Gene Filtering operation is performed. To cancel the Gene Filtering operation, click the **Cancel** button.

Upon successful completion, a new Gene Filtering item is added under the original item in the **Experiments** navigator.

**Related Topics:**
- Creating a Gene List
- Gene Lists Overview
- Supervised Learning

**Normalizing**

**Normalization Overview**

**Overview**

In GeneLinker™ the term normalization is used to describe scaling, translation, or any other numerical transformation of the data besides filtering. These transformations fall into three broad categories:

- You may need to correct for non-biological variations between different samples. For example, unintentional differences in hybridization procedures or between microarray chip manufacturing batches may cause systematic differences between samples. Normalizations which can help correct these sources of variation include mean scaling, median scaling, linear regression and control gene normalizations.

- Two-color data must be merged into ratios, and dye biases can also be corrected for at the same time.

- If you are going on to study the data by clustering, you may need to put different genes on a single scale of variation. Normalizations which may accomplish this include logarithm, standardization, division by maximum and scaling between 0 and 1.

Any number of these normalizations can be applied to dataset in succession. For instance, it may be appropriate to scale samples to correct for non-biological variations,
and then place genes on a common scale before clustering, association mining or supervised learning takes place.

Techniques for Correcting Non-Biological Variation Between Samples

- Linear Regression: This procedure scales the values relative to a baseline sample so that the best-fit slope of each sample is equivalent. All genes can be fitted, or only a user-selected set of ‘housekeeping’ genes.
- Division by Central Tendency (Mean): This procedure scales the expression values so that all samples have a common mean.
- Division by Central Tendency (Median): This procedure scales the expression values so that all samples have a common median.
- Positive and Negative Control Genes: In some experiments there may be one or more control genes whose values are expected to be constant. With multiple controls, the median or mean is calculated over all of the controls.
  - Normalization relative to negative controls subtracts the median or mean of the controls within the sample. Negative control genes are understood to be absent or below a detection threshold.
  - Normalization relative to positive controls divides each sample by the mean or median of the controls. Positive control genes are understood to be present in constant abundance in all samples.

Techniques for Adjusting Two-Color Data

- Lowess: The log-ratio expression values are adjusted by a locally-weighted linear regression on each sample to account for intensity-dependent dye bias.
- Logarithm: Gene expression values are replaced with the logarithm of their values. Taking the logarithm equalizes the influence of up- and down-regulated genes in ratio experiments.
- Subtraction of Central Tendency: This procedure transforms the expression values such that all samples have zero mean or median.

The Lowess normalization automatically merges the treatment and control channels into adjusted ratios. Any other operation on a two-color table automatically uses the unadjusted ratios.

Note: Lowess is the only normalization option for incomplete two-color datasets.

Techniques for Placing Different Genes on a Similar Scale

- Logarithm: Gene expression values are replaced with the logarithm of their values. In non-ratio experiments, taking the logarithm reduces the influence of high-abundance genes in comparison to low-abundance genes.
- Divide by Maximum: Gene expression values are scaled such that the largest value for each gene becomes one.
- Scaling Between 0 and 1: Gene expression values are scaled such that the smallest value for each gene becomes zero and the largest value becomes one. Also known as Min-Max Normalization.
- Standardize: Gene expression values are scaled such that each gene has an average of zero and a standard deviation of one.
Related Topics:
  Filtering Overview
  Clustering Overview

Linear Regression

Overview
This procedure scales the values across samples (gene chips) so that the slope of each sample is equivalent. This is done for all samples except the baseline.

This procedure fits a linear regression model using the intensities of the common genes in the baseline and each of the other samples. The inverse of the slope of the linear regression line becomes the (multiplicative) re-scaling factor for the current sample. The re-scaled intensity of the samples (other than baseline) becomes the original intensity multiplied by the re-scaling factor. This is done for all samples except the baseline. The baseline gets a re-scaling factor of 1.

Before clustering, it is recommended that standardization be performed after scaling using a baseline. Baseline scaling makes the intensities across chips equivalent, but genes may still differ in absolute intensity, and standardization can address this.

Actions
1. Click a complete dataset in the Experiments navigator. The item is highlighted.
2. Click the Normalize toolbar icon or select Normalize from the Data menu, or right-click the item and select Normalize from the shortcut menu. The first Normalization dialog is displayed.

3. Double-click the Sample Scaling radio button, or click it and click Next. The second
Normalization dialog is displayed.

4. Select **Linear Regression** as the **Scaling Type**.
5. Set the **Baseline Sample** from the drop-down list. If no baseline sample is selected, the sample displayed in the box is used for the normalization operation.
6. The **Control Genes** (housekeeping genes) can either be 'all genes in dataset' or the genes specified in a gene list.
   - If 'all genes in dataset' is selected, the operation that is performed is scaling using a baseline.
   - If a gene list is selected, the operation that is performed is scaling using housekeeping genes. For this option, the gene list must contain at least two genes from the dataset (min. required to calculate slope) and less than all the genes in the dataset (the control genes are always discarded prior to returning the normalized dataset).
   - If an appropriate gene list does not exist, click **Create Gene List**. The **Gene List Creator** dialog is displayed.

   a) Type in a **Name** for the list and optionally a **Description**.
   b) Click the checkboxes next to the genes to be included in the list.
   c) Click **Save**. The gene list is then displayed in the **Control Genes** list on the **Normalization** dialog.

7. Select the **Control Genes** from the drop-down list.
8. Click Finish. The Experiment Progress dialog is displayed. It is dynamically updated as the Sample Scaling Normalization operation is performed. To cancel the Sample Scaling Normalization operation, click the Cancel button.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the slope of the linear regression is zero or infinity (if a sample is constant).
- Upon successful completion, a new normalization dataset is added under the original dataset in the Experiments navigator.

Related Topics:
- Normalization Overview
- Clustering Overview
- Gene Lists Overview

Division by Central Tendency (Mean)

Overview
This procedure scales the values across samples (gene chips) so that the mean or total intensity of each sample is equivalent. This is done for all samples.

This scaling is useful if you have reason to believe that the total amount of mRNA measured in each sample should be approximately equivalent, but there may be non-biological sample-dependent factors influencing the raw measurements. For instance, if your data contains an entire genome but your experimental conditions are only expected to perturb a small number of genes then this type of scaling may be useful. Similarly, if you expect a large number of genes to be perturbed but both up- and down-regulation are equally likely, then the total amount of mRNA will probably be constant and this would be a reasonable operation.

The fewer non-responding genes there are in your dataset, the less sound is this scaling. For instance, if your data has been pre-filtered to retain only genes known to be affected by the experimental conditions, then this normalization may introduce undesirable distortions into your data. In the same vein, we recommend that you apply this normalization before applying any variation filtering.

This normalization is usually only meaningful if applied to count data. We do not recommend applying this normalization to ratio data or data which has already been subject to a logarithm transformation, both of which may yield zero or negative values. Applying mean scaling to samples with negative means may yield drastically distorted data. Applying mean scaling to samples with zero or near-zero means will cause GeneLinker™ to fail to complete the operation, and generate an error message.

Before clustering, it is recommended that standardization be performed after mean
scaling. Mean scaling makes the intensities across chips equivalent, but genes may still differ in absolute intensity, and standardization can address this.

**Actions**
1. Click a complete dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Normalize** toolbar icon, or select **Normalize** from the **Data** menu, or right-click the item and select **Normalize** from the shortcut menu. The first **Normalization** dialog is displayed.

3. Double-click the **Sample Scaling** radio button, or click it and click **Next**. The second **Normalization** dialog is displayed.

4. Select **Central Tendency** as the **Scaling Type**.
5. Set the **Central Tendency** to **Mean**.
6. Set the **Arbitrary New Mean** to the value to which the sample means should be scaled. The total intensity of each sample after scaling will be this number times the...
number of genes in the table.
7. Click **Finish**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Mean Scaling Normalization operation is performed. To cancel the Mean Scaling Normalization operation, click the **Cancel** button.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the mean of any sample is zero or near zero.
- Upon successful completion, a new normalization dataset is added under the original dataset in the **Experiments** navigator.

**Related Topics:**
- Normalization Overview
- Filtering Overview
- Clustering Overview

**Division by Central Tendency (Median)**

**Overview**
This procedure scales the values across samples (gene chips) so that the median of each sample is equivalent. This is done for all samples.

This scaling is useful if you have reason to believe that the most genes will be relatively unchanged, but there may be non-biological sample-dependent factors influencing the raw measurements. Similarly, if you expect a large number of genes to be perturbed but both up- and down-regulation are equally likely, then this would be a reasonable operation.

The greater the fraction of responding genes in your dataset, the less reliable is this scaling. For instance, if your data has been pre-filtered to retain only genes known to be affected by the experimental conditions, then this normalization may introduce undesirable distortions into your data. We therefore recommend that you apply this normalization before any variation filtering.

This normalization is usually only meaningful if applied to count data. We do not recommend applying this normalization to ratio data or data which has already been subject to a logarithm transformation, both of which may yield zero or negative values. Applying median scaling to samples with negative medians may yield drastically distorted data. Applying median scaling to samples with zero or near-zero medians will cause GeneLinker™ to fail to complete the operation, and generate an error message.

Median scaling is similar in principle to mean scaling, but the median is less susceptible to outliers and therefore preferred.

Before clustering, it is recommended that standardization be performed after median
scaling. Median scaling makes the scales of the chips approximately equivalent, but genes may still differ in scale and standardization can address this.

**Actions**

1. Click a complete dataset in the *Experiments* navigator. The item is highlighted.
2. Click the *Normalize* toolbar icon, or select *Normalize* from the *Data* menu, or right-click the item and select *Normalize* from the shortcut menu. The first *Normalization* dialog is displayed.

3. Double-click the *Sample Scaling* radio button, or click it and click **Next**. The second *Normalization* dialog is displayed.

4. Select *Central Tendency* as the *Scaling Type*.
5. Set the *Central Tendency* to *Median*.
6. Set the *Arbitrary New Median* to the value to which the sample medians should be scaled. The total intensity of each sample after scaling will be this number times the
number of genes in the dataset.

7. Click **Finish**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Median Scaling Normalization operation is performed. To cancel the Median Scaling Normalization operation, click the **Cancel** button.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the median of any sample is zero or near zero.
- Upon successful completion, a new normalization dataset is added under the original dataset in the **Experiments** navigator.

**Related Topics:**
- Normalization Overview
- Filtering Overview
- Clustering Overview

**Positive and Negative Control Genes**

**Overview**

In some microarray experiments, there may be one or more control genes that can be used to normalize between samples. With multiple controls, the median or mean is calculated over all of the controls.

The control genes are always discarded prior to returning the normalized dataset.

**Normalization Relative to Negative Controls**

For each sample, this is done by subtracting the median or mean of the negative controls within the sample. If you have only one control gene, the median or mean of the negative control is the value itself.

For example:

Gene i sample j - median of the negative control genes within sample j
Gene i sample k - median of the negative control genes within sample k

Below is an example that illustrates the application with three control genes for each sample:
Normalization Relative to Positive Controls
For each sample, this is done by dividing the median or mean of the positive controls within the sample. If you have only one control gene, the median or mean of the positive control is the value itself.
For example:
Gene i sample j / median of the positive control genes within sample j
Gene i sample k / median of the positive control genes within sample k
Refer to the above image.

Normalization Relative to Negative Controls Across Experiments
This is done by subtracting the median or mean of the negative control (one sample at a time) from all the values in the dataset.
For example:
Gene i sample j - median (all negative control genes across all samples)
Gene i sample k - median (all negative control genes across all samples)
Below is an image that illustrates the application with a single control for each sample:
Normalization Relative to Positive Controls Across Experiments
This is done by dividing the values in the dataset by the median value of the positive controls across all samples.
For example:
Gene i sample j / median (all positive control genes across all samples)
Gene i sample k / median (all positive control genes across all samples)
Refer to the above image.

**Actions**
1. Click a complete dataset in the Experiments navigator. The item is highlighted.
2. Click the Normalize toolbar icon, or select Normalize from the Data menu, or right-click the item and select Normalize from the shortcut menu. The first Normalization dialog is displayed.
3. Double-click the Positive and Negative Control Genes radio button, or click it and click Next. The second Normalization dialog is displayed.
4. For this operation, you must select or create a gene list of the control genes. The gene lists listed in the drop-down list are only those lists that are relevant to this dataset (that is, the list contains one or more genes that are in the dataset).

**To create a gene list:**

a) Click the Create Gene List button. The Gene List Creator dialog is displayed.

![Gene List Creator](image)

b) Type in a Name for the list and optionally a Description.

c) Click the checkboxes next to the genes to be included in the list.

d) Click Save. The gene list is then displayed in the Control Genes list on the Normalization dialog.

5. Select the Control type.

6. Select the Mean or Median for the Value.

7. Set the type of Range.

8. Click Finish. The Experiment Progress dialog is displayed. It is dynamically updated as the Control Genes Normalization operation is performed. To cancel the Control Genes Normalization operation, click the Cancel button.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the mean/median is zero or if the gene list contains all the genes in the dataset (the control genes are always discarded prior to returning the normalized dataset).

- Upon successful completion, a new normalization dataset is added under the original dataset in the Experiments navigator.

**Related Topics:**

Normalization Overview
Clustering Overview
Logarithm

Overview
This procedure transforms each gene using logarithms. Gene expression values are normalized by replacing them with the log (user selected base) of their values. For ratio data, log normalization makes inductions and repressions equal with opposite sign.

Actions
1. Click a complete dataset in the Experiments navigator. The item is highlighted.
2. Click the Normalize toolbar icon, or select Normalize from the Data menu, or right-click the item and select Normalize from the shortcut menu. The first Normalization dialog is displayed.
3. Ensure the Logarithm radio button is selected (this is the default) and click Next or double-click on the Logarithm radio button. The second Normalization dialog is displayed.
4. Double click the radio button next to the desired base, or click the radio button next to the desired base and click **Finish**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Log Normalization operation is performed. To cancel the Log Normalization operation, click the **Cancel** button.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the dataset contains values less than or equal to zero (they cannot be logged).
- If the operation succeeds, a new normalization dataset is added under the original dataset in the **Experiments** navigator.

**Related Topics:**
Normalization Overview  
Clustering Overview

**Divide by Maximum**

**Overview**
Gene expression values are normalized by dividing each value for a gene by the maximum value observed in any sample for that gene.

**Actions**
1. Click a complete dataset in the **Experiments** navigator. The item is highlighted.

2. Click the **Normalize** toolbar icon, or select **Normalize** from the **Data** menu, or right-click the item and select **Normalize** from the shortcut menu. The first **Normalization** dialog is displayed.

![Normalization Dialog](image)

3. Double-click the **Other Transformations** radio button, or click it and click **Next**. The second **Normalization** dialog is displayed.

![Normalization Dialog](image)

4. Double-click the **Divide by Maximum** radio button, or click it and click **Finish**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Divide by Maximum Normalization operation is performed. To cancel the Divide by Maximum Normalization operation, click the **Cancel** button.

![Normalization Dialog](image)
If the operation cannot complete an error message is displayed. The operation will fail, for example, if the maximum of a gene is zero.

Upon successful completion, a new normalization dataset is added under the original dataset in the Experiments navigator.

Related Topics:
Normalization Overview
Clustering Overview

Scaling Between 0 and 1

Overview
Gene expression values are normalized by subtracting the minimum value for each gene followed by dividing by the adjusted maximum value for that gene. This is also known as ‘Min. to Max. Scaling’.
This procedure scales all of the values for each gene so that they all fall in the range from 0 to 1. This can be done as part of the normalization process prior to running an experiment.

Actions
1. Click a complete dataset in the Experiments navigator. The item is highlighted.
2. Click the Normalize toolbar icon, or select Normalize from the Data menu, or right-click the item and select Normalize from the shortcut menu. The first Normalization dialog is displayed.
3. Double-click the **Other Transformations** radio button, or click it and click **Next**. The second **Normalization** dialog is displayed.

4. Double-click the **Scaling Between 0 and 1** radio button, or click it and click **Finish**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Scaling Between 0 and 1 Normalization operation is performed. To cancel the Scaling Between 0 and 1 Normalization operation, click the **Cancel** button.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the dataset contains a constant gene.
- Upon successful completion, a new normalization dataset is added under the
Gene expression values are normalized by subtracting the mean, followed by dividing the standard deviation for each gene.

This procedure standardizes each gene. The mean and standard deviation for each gene is calculated, and each value for the gene (x) is standardized using (x - mean)/standard deviation.

**Actions**

1. Click a complete dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Normalize** toolbar icon, or select **Normalize** from the **Data** menu, or right-click the item and select **Normalize** from the shortcut menu. The first **Normalization** dialog is displayed.
3. Double-click the **Other Transformations** radio button, or click it and click **Next**. The second **Normalization** dialog is displayed.
4. Double-click the **Standardize** radio button, or click it and click **Finish**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Standardize Normalization operation is performed. To cancel the Standardize Normalization operation, click the **Cancel** button.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the standard deviation of a gene is zero.
- Upon successful completion, a new normalization dataset is added under the original dataset in the **Experiments** navigator.

**Related Topics:**
- Normalization Overview
- Clustering Overview

**Overview of Lowess Normalization**

**Overview**

In experiments where two fluorescent dyes (red and green) have been used, intensity-dependent variation in dye bias may introduce spurious variations in the collected data. Lowess normalization merges two-color data, applying a smoothing adjustment that removes such variation.

**Lowess Normalization Characteristics**
- Lowess normalization may be applied to a **two-color** array expression dataset.
- All samples in the dataset are corrected independently.
• Lowess normalization can be applied to complete or incomplete datasets. If either the red or green intensity value is missing for a certain gene, there will be a missing value at the corresponding position in the log-ratio table which is generated.

**Lowess Normalization Method**

Lowess normalization assumes that the dye bias appears to be dependent on spot intensity. The adjusted ratio is computed by:

$$\text{log}(R/G) \to \text{log}(R/G) - c(A)$$

where $c(A)$ is the Lowess fit to the log($R/G$) vs log(sqrt($R \cdot G$)) plot.

If green has been chosen as the treatment dye and red as the control dye, then $R$ and $G$ are reversed in the above formula. Treatment and control dyes are designated when the data is imported into GeneLinker™.

Lowess regression, or locally weighted least squares regression, is a technique for fitting a smoothing curve to a dataset. The degree of smoothing is determined by the window width parameter. A larger window width results in a smoother curve, a smaller window results in more local variation.

Upon successful completion of the normalization, a new dataset with the Lowess-corrected ($R/G$) values (or ($G/R$) if appropriate) is stored in the repository and is added to the **Experiments** navigator. The result is a dataset of corrected ratios (not log ratios).

**Reference**


**Related Topics:**

- Lowess
- Subtraction of Central Tendency

**Lowess**
Overview

Lowess normalization is a method used to normalize a two-color array gene expression dataset to compensate for non-linear dye-bias. In this approach, the log-ratio for each sample is adjusted by the Lowess fitted value. The result is a dataset of corrected ratios (not log ratios). See Overview of Lowess Normalization for complete information.

Visualization

To determine whether or not Lowess normalization is appropriate for a dataset, display an intensity-bias plot of a sample ratio.

Actions

1. Click a two-color dataset in the Experiments navigator. The item is highlighted.
2. Click the Normalization toolbar icon , or select Normalize from the Data menu, or right-click the item and select Normalize from the shortcut menu. The first Normalization dialog is displayed.
3. Select Sample Scaling. The second Normalization dialog is displayed.
4. Select **Lowess**.

5. Set the **Window Width** parameter.

7. Click **Finish**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Lowess normalization operation is performed.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the mean of any sample is zero or near zero.
- Upon successful completion, a new normalization dataset is added under the original dataset in the **Experiments** navigator.

**Visualization**

An intensity-bias plot of the Lowess-corrected data can be made from the corrected data by creating a table view, selecting the desired row, and selecting **Intensity-Bias Plot** from the **Explore** menu as described above.

**Related Topics:**

- Creating an Intensity-Bias Plot of a Sample Ratio
- Subtraction of Central Tendency

**Subtraction of Central Tendency**

**Overview**

Subtraction of central tendency adjusts each sample in a dataset to have a median or
Subtraction of central tendency is typically used to adjust log-ratio values to result in a median (or mean) log-ratio of zero for each sample. This is appropriate, for instance, if the treatment and control dyes in a two-color experiment are incorporated with some bias independent of intensity.

Lowess normalization produces an adjustment almost identical to subtraction of a constant mean if the dye bias is, in fact, independent of intensity. But Lowess is not constrained to produce only a constant correction as subtraction of central tendency is, so it is more general. We therefore recommend Lowess normalization over subtraction of central tendency as a means of normalizing two-color datasets.

**Subtraction of Central Tendency Characteristics**

- All samples in the dataset are corrected independently.
- This normalization can be applied to complete or incomplete datasets. If either the red or green intensity value is missing for a certain gene, a missing value is placed at the corresponding position in the generated ratio dataset.

**Actions**

1. Click on a two-color dataset in the Experiments navigator. The item is highlighted.
2. Click on the Normalization toolbar icon, or select Normalize from the Data menu, or right-click the item and select Normalize from the shortcut menu. The first Normalization dialog is displayed.

3. Select Sample Scaling. The second Normalization dialog is displayed.
4. Select **Central Tendency** as the **Scaling Type**.
5. Set the **Central Tendency** operation to **Subtract**.
6. Set the **Subtract** central tendency type to **Mean** or **Median**.
7. Click **Finish**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the normalization operation is performed.

- If the operation cannot complete an error message is displayed. The operation will fail, for example, if the mean of any sample is zero or near zero.
- Upon successful completion, a new normalization dataset is added under the original dataset in the **Experiments** navigator.

**Visualization**

Once the normalization is complete, a scatter plot can be used to examine each corrected sample.

**Related Topics:**
- Creating an Intensity-Bias Plot of a Sample Ratio
- Lowess

**Creating an Intensity-Bias Plot of a Sample Ratio**

**Overview**
An intensity-bias plot can be used to view dye biases to determine whether normalization is required. An intensity-bias plot is a scatter plot of the log-ratio versus the log-intensity.

**Actions**
1. Click a two-color dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Table View** toolbar icon, or select **Table View** from the **Data** menu, or right-click the item and select **Table View** from the shortcut menu. A table view of the dataset is displayed.

3. Click on the name of a sample. The sample is highlighted.
4. Select **Intensity-Bias Plot** from the **Explore** menu. An intensity-bias plot of the highlighted sample is displayed.

**Related Topics:**
- Lowess
- Subtraction of Central Tendency

**Removing Values**

**Removing Values by Expression Value**
Overview

This function compares each value in the original dataset with the threshold using the specified comparison type (<=, =, >=). All values (v) that satisfy the condition (e.g.: v >= threshold) are replaced with missing values (null values) in the new dataset. If the original dataset is complete and some of its values are eliminated (they satisfy the condition), then the result is an incomplete dataset.

Value Representation

Values in datasets are real values and are represented as floating point numbers by the computer. Therefore, the threshold is actually a small range: ( threshold - 10exp(-7), threshold + 10exp(-7) ).

- A comparison of the form v = threshold performs the comparisons v >= threshold - 10exp(-7) and v <= threshold +10exp(-7). The value v passes the test if it meets both conditions.
- A comparison of the form v <= threshold, performs the comparison v <= threshold + 10exp(-7).
- A comparison of the form v >= threshold, performs the comparison v >= threshold - 10exp(-7).

If the parameters are set such that all of any gene’s values are removed, that gene will be completely removed (filtered) from the resulting dataset. No genes will be kept which are completely devoid of values. Therefore the resulting dataset may have fewer genes than the parent dataset in some cases.

Actions

1. Click a complete or incomplete dataset in the Experiments navigator. The item is highlighted.

2. Select Remove Values from the Data menu, or for an incomplete dataset, right-click the item and select Remove Values from the shortcut menu. The Remove Values parameters dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
</table>
Removal Technique | Select by Expression Value to perform value removal by the gene expression data values.
--- | ---
Expression Value | Set the comparison type and the threshold value.

4. Click OK. The Experiment Progress dialog is displayed. It is dynamically updated as the Value Removal operation is performed. To cancel the Remove Values operation, click the Cancel button.

Upon successful completion, a new dataset is added under the original dataset item in the Experiments navigator.

Related Topics:
- Cancelling an Operation
- Overview of Estimating Missing Values

Removing Values by Reliability Measure

Overview
This function is used to create missing values from unreliable gene expression values. Unreliability might be implied by a poor reliability measurement, or it might be implied by an extreme expression measurement.

This function can only be applied to top-level datasets that have associated reliability measurement data. The reliability measure may be a P-Value imported from an Affymetrix MAS 5.0 file, or computed on import from within-chip replicates.

The result of this operation can be either a complete or an incomplete dataset.

Actions
1. Click a complete or incomplete dataset with associated reliability measures in the Experiments navigator. The item is highlighted.
2. Select Remove Values from the Data menu, or right-click the item and select Remove Values from the shortcut menu. The Remove Values parameters dialog is displayed.
3. Click **by Reliability Value** as the **Removal Technique**. The dialog is updated.

4. Use the slider to set the reliability measure threshold. The reliability scale is from 1.0 (low reliability) to 0.0 (high reliability).

5. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the value removal operation is performed.

Upon successful completion, a new dataset is added under the original dataset item in the **Experiments** navigator.

- If the dataset you selected is not a top-level dataset, or if it does not have reliability data associated with it, the dialog is updated to indicate this. Click **OK** to exit this operation.
Related Topic:
Creating a Table View of Reliability Data

Statistics

Creating a Summary Statistics Chart

Overview
The Summary Statistics chart is a combination of a histogram plot of the values in a dataset (user selectable parameters) and a textual display of several key statistical values describing the dataset. This information could be used to see how many of the dataset’s values fall outside an expected range (possibly due to experimental error or other sources of ‘noise’). Another use could be to estimate whether the data values conform to an approximately normal or other sort of distribution. Since microarray data are almost never normal, this may be more useful after, for instance, log-transformation.

The numeric statistics given in the lower half of the display could be used to summarize and compare different datasets. For instance, the coefficient of variation is a one-number summary of how the data’s variation compares to its magnitude.

Histogram Chart
The histogram shows the distribution of the data values among a number of bins (15 is the default). A bin is a container for data values. Each bin has a minimum and a maximum bound. All data points that are greater than (and in the first bin equal to) the minimum bound and less than or equal to the maximum bound of a certain bin are placed into this bin.

The chart’s x-axis is labeled with the minimum bound for the first bin and the maximum bound for the last bin. If the minimum cutoff value is changed, the first bin is given a lower bound of -infinity. If the maximum cutoff value is changed, the last bin is given an upper bound of +infinity.

The chart’s y-axis is labeled with the frequency of data values. The sum of the
frequencies from all the bins equals the number of data values in the selected table, gene(s) or sample(s) (excluding missing values).

**Statistics Textual Display Items**
- minimum value
- maximum value
- mean
- median
- number of values (excluding missing ones)
- number of missing values
- standard deviation
- co-efficient of variance.

**Chart Parameters**
The chart parameters area is the place to specify the **number of bins**. Changing the number of bins causes the data range (minimum to maximum bound) for each bin to change. To have a smaller range per bin, increase the number of bins. Conversely, to have a larger range per bin, decrease the number of bins. **Note** that only integer values are accepted.

The chart parameters area is also the place to change the **cutoff values**. The minimum and maximum cutoff values are the upper bound of the first bin and the lower bound of the last bin respectively. When the **Manual** radio button is first clicked, the present cutoff value is displayed in the appropriate text box. To change the cutoff value, type over the displayed value.

The minimum and maximum cutoff values can be used to separate **outliers** from the main data by placing the outliers in bins outside the main data grouping. This is done by setting the minimum and maximum cutoff values at or near the outer bounds of the main grouping. For example, if the minimum cutoff value is set to .5 and the maximum cutoff value is set to 7.5, then all values less than or equal to .5 are grouped into one outlier bin that appears to the left of the ‘.5’ data co-ordinate label on the x-axis and all values greater than 7.5 are grouped into one outlier bin that appears to the right of the ‘7.5’ data co-ordinate label on the x-axis. All bins other than outlier bins maintain a contiguous linearity with respect to the x axis.

**Actions**
1. Click a complete or incomplete dataset in the **Experiments** navigator, or select gene(s) or sample(s) from a plot. The item is highlighted.
2. Click the **Summary Statistics** toolbar icon, or select **Summary Statistics** from the **Statistics** menu, or right-click the item and select **Summary Statistics** from the shortcut menu. The Summary Statistics chart is displayed.
Changing the Number of Bins
1. Parameters area. The minimum number of bins is 1 (without outlier bins), 2 (with 1 outlier) or 3 (with 2 outliers). The maximum number of bins is 1000. If you enter a value that is out of range, the Refresh button is disabled (grayed out).
2. Click the Refresh button to display the chart using the new parameters.
   • To change the default number of bins, see Changing Your User Preferences.

Changing the Cutoff Values
1. Click the Manual radio button and/or type the value into the First bin upper boundary and/or Last bin lower boundary text box. You do not have to change both.
2. Click the Refresh button to display the chart using the new parameters.
   Note: the Refresh button is disabled (grayed out) when the values (# of bins and cutoff values) match the current chart characteristics.

Exporting the Image
1. Click the histogram to make it the active window.
2. Select Export Image from the File menu, or right-click on the chart and select Export Image from the shortcut menu. The Save As dialog is displayed.
3. Navigate to the destination folder and fill in the name for the image file or accept the default name. The export image file includes the title, histogram, and summary statistics text. (For a complete dataset, the title could be the experiment name. For a single gene or sample, the gene or sample name could be used.)
   Note: When a report on a complete or an incomplete dataset is generated, the textual representation of the summary statistics is included within it.

Related Topics:
- Normalization Overview
- Filtering Overview
- Generating Reports
Overview of ANOVA

Overview

GeneLinker™ provides two different methods for performing a one-way Analysis of Variance, or ANOVA: The F-Test and the Kruskal-Wallis test. These methods are used to determine which genes vary most significantly between a set of conditions. If one has replicate chips measuring, for example, subjects treated with a drug and treated with a placebo, ANOVA can be used to rank the genes according to their change between the treatment and control conditions. ANOVA can be used to compare several conditions simultaneously, not just two at a time. ANOVA is most effective when all groups are the same size, each containing at least three samples (replicates).

When you carry out an ANOVA GeneLinker™ calculates a p-value for each gene. The p-value is the probability that the variation between conditions may have occurred by chance, so genes with smaller p-values are varying more significantly. The gene’s variation is less likely to have occurred by chance, and is conversely more likely to be connected to the difference in conditions. When you view an ANOVA result in GeneLinker™, the most significantly-varying genes – those with the smallest p-values – appear at the top of the list.

The conditions are specified by importing a variable, called the Grouping Variable. The different values of the Grouping Variable represent the different conditions between which significant variation may take place. For example if the Grouping Variable chosen looks like this:

A
A
A
B
B
B

then the first three samples will be considered replicates under one condition (A), and the second three samples will be considered replicates under another condition (B). The ANOVA will determine whether the variation between group A and group B is significantly greater than the (presumably random) variation within each group.

Note: If you do not have any replicates in your data, GeneLinker™ will display 'Undefined' for the p-value of every gene. 'Undefined' can also be computed for individual genes in certain circumstances, e.g. if there is no variation in the expression level of the gene.

A common use of the ANOVA is to remove invariant genes from a dataset. To do this:

1. Carry out an ANOVA.
2. Select the most significant genes in the ANOVA viewer. You may either choose a threshold p-value or choose some number of genes that is useful to you.
3. Create and save a gene list from this selection.

4. Use Gene List Filtering to generate a new data table containing only those genes which vary significantly.

See ANOVA Viewer for instructions on creating a gene list from ANOVA results.

Choosing between the F-Test and Kruskal-Wallis

The F-Test is a parametric test which is based on certain assumptions of normality about the data. The Kruskal-Wallis Test is a non-parametric test which makes no such assumptions. Because the Kruskal-Wallis Test uses only the rankings of the data points and not their absolute values, it is a less powerful test than the F-Test and may underestimate the significance of the changes in some genes (i.e. compute too large a p-value). If your data is approximately normal, or can be transformed so that it is, you should use the F-Test. If not, then use the Kruskal-Wallis Test.

Gene expression abundances are rarely normal, but are frequently log-normal. You can estimate the normality of your data visually using the Summary Statistics Chart in GeneLinker™. If the data is strongly skewed to the left, as in the first picture below, then you should first transform it using a Logarithm normalization. Viewing the Summary Statistics on the log-normalized data table should produce a normal histogram much like the one in the second picture. The second data table is suitable for application of the F-Test.

P-values and multiple testing

The p-value computed by GeneLinker™ is to be interpreted for each gene as the probability that the variation in that gene is random. When the test is being applied to thousands of genes – as is usually the case in microarray experiments – then even purely random data will contain some genes with small (significant) p-values. For example, if you choose to consider for further experimentation any gene with a p-value of less than 5% or 0.0500, then you can reasonably expect that about 5% of those genes are false positives, or genes which have obtained a small p-value by random chance. If you are using ANOVA as a gene filter and it is important to you to minimize the number of false positives, then you should choose a smaller p-value as a cutoff. For instance, if you are testing 1000 genes and want only a 50% chance of having one false positive in your gene list, then you should select only genes with p < 0.50/1000, or 0.0005. Be warned, however, that you will also be discarding genes which have real differential expression by so doing. i.e. you will increase the number of false negatives as you decrease the number of false positives. The systematically varying genes and the randomly varying genes will be intermixed in any real dataset. The only way to separate them better – the only way to decrease both the false positive rate and the false negative rate – is to do more experiments and obtain more replicates.
The simple adjustment of the p-value described above is technically known as a Bonferroni correction. The Bonferroni correction is rather conservative (i.e. severe) but has the virtue of simplicity. For more discussion of multiple testing corrections to microarray data, see for example S. Dudoit, Y. H. Yang, M. J. Callow and T. P. Speed, "Statistical methods for identifying differentially expressed genes in replicated cDNA microarray experiments" (2000), Stanford University Technical Report #578.

**F-Test Algorithm**

For a gene with \( M \) groups of samples, where each group \( I \) has \( N_i \) replicates \( (I = 1,2,...M) \) we want to determine if the gene has significantly changed between any pair of groups. The F-statistic is the ratio of two variances:

\[
F = \frac{\text{var}_1}{\text{var}_2}
\]

The null hypothesis is that the two variances are the same. The statistic follows a distribution parameterized by \( \nu_1 = n1 - 1 \) and \( \nu_2 = n2 - 1 \), where \( n1 \) and \( n2 \) are the number of samples in the populations used to calculate \( \text{var}_1 \) and \( \text{var}_2 \).

To use the F-test to filter genes, the F-statistic is first determined by calculating the total variations between and within samples. The result can be proven to follow the F-distribution.

\[
\text{variation}_\text{between}_\text{samples} = \left[ \sum_{i=1}^{M} \sum_{j=1}^{N_i} (Y_i - Y)^2 \right], \quad n1 = M -1
\]

\[
\text{variation}_\text{within}_\text{samples} = \left[ \sum_{i=1}^{M} \left( \sum_{j=1}^{N_i} (Y_{ij} - Y_i)^2 \right) \right], \quad n2 = (\sum_{i=1}^{M} N_i) - M
\]

The relevant F-statistic is then formed by taking:

\[
F = \left( \frac{\text{variation}_\text{between}_\text{samples}}{n1} \right) / \left( \frac{\text{variation}_\text{within}_\text{samples}}{n2} \right)
\]

The probability of this F-value arising from two identical distributions gives us a measure of the significance of the between-sample variation as compared to the within-sample variation. Small p-values indicate a low probability of the between-sample variation being due to sampling of the within-sample distribution, so small p-values indicate interesting genes.

**Kruskal-Wallis Algorithm**

The Kruskal-Wallis algorithm is analogous to the F-Test, except that instead of operating on the expression values directly it operates on the ranks of the expression values. That is, each gene first has its expression values sorted and a rank assigned to each value based on its position in the sorted list. The variances of the rank numbers within each group are computed, and the test proceeds as the F-Test described above.

**Related Topics:**
- Performing an ANOVA
- ANOVA Viewer
- Overview of Estimating Missing Values
Performing an ANOVA

Overview
This operation calculates p-values for the genes in a complete dataset. For details of the ANOVA algorithms, see Overview of ANOVA.

The input to this operation must be a complete dataset. If your dataset has missing values, see Overview of Estimating Missing Values for techniques available to eliminate or estimate missing values.

Actions
1. Click a complete dataset with variable information identifying the replicate samples in the Experiments navigator. The item is highlighted.
2. Select ANOVA from the Statistics menu, or right-click on the item and select ANOVA from the shortcut menu. The ANOVA dialog is displayed.

Note: if an appropriate grouping variable is not associated with the dataset, this is indicated on the dialog. In this situation, click Cancel and import an appropriate variable before trying again. See Overview of ANOVA for a discussion of appropriate variables.

3. Set the Operation (style of ANOVA) to F-Test or Kruskal-Wallis. See Overview of ANOVA for how to choose the right method.
4. Select the Grouping Variable from the drop-down list.
5. Click OK. The ANOVA operation is performed and upon successful completion, a new F-Test or Kruskal-Wallis Results item is added to the Experiments navigator under the original dataset.

The results can then be viewed using the ANOVA Viewer.

Related Topics:
- Overview of ANOVA
- ANOVA Viewer
- Overview of Estimating Missing Values

ANOVA Viewer
GeneLinker Gold 3.1 / GeneLinker Platinum 2.1
Overview

The ANOVA Viewer displays a list of the genes and their associated p-values from an F-Test Results or a Kruskal-Wallis Results item in the Experiments navigator. The list can be sorted, and genes can be selected for creating gene lists.

The first column of the viewer contains checkboxes indicating whether a gene is checked or not (unchecked). The second column contains index numbers. The index numbers are not associated with the genes, they merely indicate position within the current sort context. The third column contains gene names, and the fourth contains p-values.

Actions

1. Double-click an F-test Results or Kruskal-Wallis (K-W test) Results item in the Experiments navigator, or click the item and select ANOVA Viewer from the Statistics menu. The item is highlighted and the ANOVA Viewer is displayed.

Sorting the Genes

The default sort for the contents of the ANOVA Viewer is by ascending P-Value. This sort places the genes with the most significant P-values at the top of the list.

The list can be sorted by Gene (alphabetical, or reverse alphabetical) or by P-Value (ascending or descending).

Checking Genes

A checked box in the first column indicates that the gene is checked.

To check a gene, click on the empty checkbox next to the gene.

To uncheck a gene, click on the checked checkbox next to the gene.

To check a series of genes, press and hold the <shift> key and click on the first and
last gene checkboxes. All the genes between the first and last inclusive are selected.

Creating a Gene List
1. Check one or more genes.
2. Click Create Gene List. The Create a Gene List dialog is displayed.

3. Provide a Name and optionally a Description for the gene list.
4. Click OK. The gene list is created and a new item is added to the Gene Lists navigator.

Related Topics:
- Overview of ANOVA
- Performing an ANOVA
- Gene Lists Overview

Sample Merging

Sample Merging

Overview
This feature provides you with the capability to merge samples based on variables. Samples that have the same variable value (observation) are collapsed into a single representative sample using the mean or median. Variation within each group is captured in a deviation table that is associated with a sample merging experiment. The standard deviation is used if the samples are merged using the mean, and the absolute deviation around the median is used if the samples are merged using the median. This feature can be used to handle between chip replication (where different samples represent replicates of other samples). It can also be used to visually identify genes that either vary significantly or hardly at all for each class. This feature can also be used as a complement to classification. You can look at the profile of each class to help pick out features (genes) that might assist in creating a good classifier, or to see the average behavior of genes which have been picked out by other means such as SLAM or ANOVA.

Actions
1. Click a regular gene expression dataset in the Experiments navigator. The item is highlighted.
2. Select Sample Merging from the Statistics menu. The Sample Merging dialog is displayed.

![Sample Merging Dialog](image)

3. Set the Operation to Mean or Median.
4. Select the Sample Variable from the drop-down list.
5. Click OK.

The dataset is collapsed so that the new number of samples corresponds to the number of distinct variable values in the imported variable. The merged dataset has the variable that was used to identify the samples in each group attached to the resulting dataset.

The results can be viewed using the Sample Merging Viewer.

**Note:** you can import new variables against Sample Merging experiments. Variables are propagated upwards and downwards in the experiment tree. Descendent samples are marked as unknown if their observations for a given variable aren't unanimous.

**Related Topics:**
- Sample Merging Viewer
- Variable Import
- Summary Statistics

**Sample Merging Viewer**

**Overview**
The Sample Merging Viewer displays a profile plot of each sample with the deviations indicated using error bars. Each representative sample is plotted as a line with an expression value for each gene.

**Actions**
1. Double-click a Sample Merging item in the Experiments navigator, or click the item and select Sample Merging Viewer from the Statistics menu. The item is highlighted and the Sample Merging Viewer is displayed.
Cluster Analysis Characteristics:

- Numerical, statistical, and conceptual clustering.
- Agglomerative vs. divisive.
- Overlapping vs. disjoint clusters.
- Incremental vs. non-incremental.
• Flat vs. hierarchical representations.

In GeneLinker™, the following clustering methods are available:
• K-Means
• Jarvis-Patrick
• Agglomerative Hierarchical
• Self Organizing Maps
All of the above methods are applicable to both genes and samples.

Related Topic:
Distance Metrics Overview

Distance Metrics

Distance Metrics Overview

Overview
Distance Measurements Between Data Points
This parameter specifies how the distance between data points in the clustering input is measured. The options are:
• Euclidean: Use the standard Euclidean (as-the-crow-flies) distance.
• Euclidean Squared: Use the Euclidean squared distance in cases where you would use regular Euclidean distance in Jarvis-Patrick or K-Means clustering.
• Manhattan: Use the Manhattan (city-block) distance.
• Pearson Correlation: Use the Pearson Correlation coefficient to cluster together genes or samples with similar behavior; genes or samples with opposite behavior are assigned to different clusters.
• Pearson Squared: Use the squared Pearson Correlation coefficient to cluster together genes with similar or opposite behaviors (i.e. genes that are highly correlated and those that are highly anti-correlated are clustered together).
• Chebychev: Use Chebychev distance to cluster together genes that do not show dramatic expression differences in any samples; genes with a large expression difference in at least one sample are assigned to different clusters.
• Spearman: Use Spearman Correlation to cluster together genes whose expression profiles have similar shapes or show similar general trends (e.g. increasing expression with time), but whose expression levels may be very different.

Distance Measurements Between Clusters
This parameter specifies how the distance between clusters is measured. The options are:
• **Average Linkage**: The distance between two clusters is the average of the
distances between all the points in those clusters.

- **Single Linkage**: The distance between two clusters is the distance between the nearest neighbors in those clusters.
- **Complete Linkage**: The distance between two clusters is the distance between the furthest points in those clusters.

**Related Topics:**
- Overview of K-Means Clustering
- Overview of Jarvis-Patrick Clustering
- Overview of Agglomerative Hierarchical Clustering

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**Euclidean and Euclidean Squared**

**Overview**

**Euclidean Distance Metric**:
The Euclidean distance function measures the ‘as-the-crow-flies’ distance. The formula for this distance between a point \(X (X_1, X_2, \text{etc.})\) and a point \(Y (Y_1, Y_2, \text{etc.})\) is:

\[
d = \sqrt{\sum_{i=1}^{n} (x_i - y_i)^2}
\]

Deriving the Euclidean distance between two data points involves computing the square root of the sum of the squares of the differences between corresponding values.

The following figure illustrates the difference between Manhattan distance and Euclidean distance:

![Manhattan vs Euclidean](image)

**Euclidean Squared Distance Metric**
The Euclidean Squared distance metric uses the same equation as the Euclidean distance metric, but does not take the square root. As a result, clustering with the Euclidean Squared distance metric is faster than clustering with the regular Euclidean distance. The output of Jarvis-Patrick and K-Means clustering is not affected if Euclidean distance is replaced with Euclidean squared. However, the output of hierarchical clustering is likely to change.

**Related Topics:**
**Manhattan**

**Overview**

The Manhattan distance function computes the distance that would be traveled to get from one data point to the other if a grid-like path is followed. The Manhattan distance between two items is the sum of the differences of their corresponding components. The formula for this distance between a point \(X=(X_1, X_2, \text{ etc.})\) and a point \(Y=(Y_1, Y_2, \text{ etc.})\) is:

\[
d = \sum_{i=1}^{n} |x_i - y_i|
\]

Where \(n\) is the number of variables, and \(x_i\) and \(y_i\) are the values of the \(i\)th variable, at points \(X\) and \(Y\) respectively.

The following figure illustrates the difference between Manhattan distance and Euclidean distance:

![Manhattan vs Euclidean](image)

**Related Topics:**

- Euclidean and Euclidean Squared Distance Metric
- Distance Metrics Overview

**Pearson Correlation and Pearson Squared**

**Overview**

**Pearson Correlation**

Pearson Correlation measures the similarity in shape between two profiles. The formula for the Pearson Correlation distance is:

\[
d = 1 - r
\]

where

\[
r = \frac{Z(x) \cdot Z(y)}{n}
\]
is the dot product of the z-scores of the vectors x and y. The z-score of x is constructed by subtracting from x its mean and dividing by its standard deviation.

**Pearson Squared**

The Pearson Squared distance measures the similarity in shape between two profiles, but can also capture inverse relationships. For example, consider the following gene profiles:

In the figure on the left, the black profile and the red profile have almost perfect Pearson correlation despite the differences in basal expression level and scale. These genes would cluster together with either Pearson Correlation or Pearson Squared distance. In the figure on the right, the black and red profiles are almost perfectly *anti-correlated*. These genes would be placed in remote clusters using Pearson Correlation, but would be put in the same cluster using Pearson Squared.

The formula for the Pearson Squared distance is

\[ d = 1 - 2r \]

where \( r \) is the Pearson correlation defined above.

**Warning:** While most combinations of clustering algorithm and distance metrics provide meaningful results, there are a few combinations that are difficult to interpret. In particular, combining K-Means clustering with the **Pearson Squared** distance metric can lead to non-intuitive centroid plots since the centroid represents the mean of the cluster and Pearson Squared can group anti-correlated objects. In these cases, visually drilling into clusters to see the individual members through the use of Cluster Plots produce better results. Alternatively, the results of the clustering run can be visualized using the Matrix Tree Plot.

**Related Topics:**
- Clustering Overview
- Distance Metrics Overview

**Chebychev**

**Overview**

The Chebychev distance between two points is the maximum distance between the points in any single dimension. The distance between points \( X=(X_1, X_2, \text{ etc.}) \) and \( Y=(Y_1, \)
\(Y_2\), etc.) is computed using the formula:

\[
\text{Max}_i |X_i - Y_i|
\]

where \(X_i\) and \(Y_i\) are the values of the \(i\)th variable at points \(X\) and \(Y\), respectively. The Chebychev distance may be appropriate if the difference between points is reflected more by differences in individual dimensions rather than all the dimensions considered together.

**Note** that this distance measurement is very sensitive to outlying measurements.

**Related Topics:**
- Clustering Overview
- Distance Metrics Overview

**Spearman Rank Correlation**

**Overview**

Spearman Rank Correlation measures the correlation between two sequences of values. The two sequences are ranked separately and the differences in rank are calculated at each position, \(i\). The distance between sequences \(X = (X_1, X_2, \text{ etc.})\) and \(Y = (Y_1, Y_2, \text{ etc.})\) is computed using the following formula:

\[
1 - \frac{6 \sum_{i=1}^{n} (\text{rank}(X_i) - \text{rank}(Y_i))^2}{n(n^2-1)}
\]

Where \(X_i\) and \(Y_i\) are the \(i\)th values of sequences \(X\) and \(Y\) respectively. The range of Spearman Correlation is from -1 to 1. Spearman Correlation can detect certain linear and non-linear correlations. However, Pearson Correlation may be more appropriate for finding linear correlations.

**Related Topics:**
- Clustering Overview
- Distance Metrics Overview

**K-Means**

**K-Means Clustering Overview**

**Overview**

K-Means clustering generates a specific number of disjoint, flat (non-hierarchical) clusters. It is well suited to generating globular clusters. The K-Means method is numerical, unsupervised, non-deterministic and iterative.
K-Means Algorithm Properties

- There are always K clusters.
- There is always at least one item in each cluster.
- The clusters are non-hierarchical and they do not overlap.
- Every member of a cluster is closer to its cluster than any other cluster because closeness does not always involve the 'center' of clusters.

The K-Means Algorithm Process

- The dataset is partitioned into K clusters and the data points are randomly assigned to the clusters resulting in clusters that have roughly the same number of data points.
- For each data point:
  - Calculate the distance from the data point to each cluster.
  - If the data point is closest to its own cluster, leave it where it is. If the data point is not closest to its own cluster, move it into the closest cluster.
- Repeat the above step until a complete pass through all the data points results in no data point moving from one cluster to another. At this point the clusters are stable and the clustering process ends.
- The choice of initial partition can greatly affect the final clusters that result, in terms of inter-cluster and intracluster distances and cohesion.

K-Means Clustering in GeneLinker™

The version of the K-Means algorithm used in GeneLinker™ differs from the conventional K-Means algorithm in that GeneLinker™ does not compute the centroid of the clusters to measure the distance from a data point to a cluster. Instead, the algorithm uses a specified linkage distance metric. The use of the Average Linkage distance metric most closely corresponds to conventional K-Means, but can produce different results in many cases.

Advantages to Using this Technique

- With a large number of variables, K-Means may be computationally faster than hierarchical clustering (if K is small).
- K-Means may produce tighter clusters than hierarchical clustering, especially if the clusters are globular.

Disadvantages to Using this Technique

- Difficulty in comparing quality of the clusters produced (e.g. for different initial partitions or values of K affect outcome).
- Fixed number of clusters can make it difficult to predict what K should be.
- Does not work well with non-globular clusters.
- Different initial partitions can result in different final clusters. It is helpful to rerun the
program using the same as well as different K values, to compare the results achieved.

**Note** the **Warning** in Pearson Correlation and Pearson Squared Distance Metric on use of K-Means clustering.

**Related Topics:**
- Performing K-Means Clustering
- Clustering Overview
- Distance Metrics Overview

**Performing K-Means Clustering**

**Overview**
K-Means clustering generates a specific number of disjoint, flat (non-hierarchical) clusters. It is well suited to generating globular clusters.
For further details, see Overview of K-Means Clustering.

**Actions**
1. Click a complete dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Partitional Clustering** toolbar icon, or select **Partitional Clustering** from the **Clustering** menu, or right-click the item and select **Partitional Clustering** from the shortcut menu. The **Partitional Clustering** parameters dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Clustering Orientation</strong></td>
<td>Cluster by <strong>Genes</strong> or by <strong>Samples</strong>.</td>
</tr>
<tr>
<td><strong>Distance Measurement</strong></td>
<td>Type of distance measurement to use to determine how close two data points are to each other.</td>
</tr>
<tr>
<td><strong>Distance Measurement</strong></td>
<td>Type of distance measurement to use to determine how close two clusters are to each other.</td>
</tr>
</tbody>
</table>

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Type

Set this parameter to **K-Means**.

**Number of Means**

This value specifies the number of clusters the algorithm forms. The value must be **greater than or equal to 2**, and **less than or equal to the number of clusterable items** (genes or samples) in the selected dataset.

**Random Seed**

The seed value for the random number generator. In normal use, setting the random seed is neither necessary nor recommended. On occasion, you may need to determine whether a certain variation in results is due to the random element, or some other cause. For this reason, you are able to set the random seed to a fixed value, thus controlling that source of variation.

4. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the K-Means Clustering operation is performed. To cancel the K-Means Clustering operation, click the **Cancel** button.

Upon successful completion, a new item is added under the original item in the **Experiments** navigator.

**Related Topics:**
- Distance Metrics Overview
- Clustering Overview
- Export Partitional Cluster

**Export Partitional Cluster**

**Overview**

You can export a comma-delimited file (.csv) that contains the genes or samples from a partitional clustering experiment with their associated cluster identifiers.

**Actions**

1. Click on a Partitional Clustering experiment in the **Experiments** navigator. The item is highlighted.

2. Select **Export Partitional Cluster** from the **Clustering** menu, or right-click the item and select **Export Partitional Cluster** from the shortcut menu. The **Save As** dialog is displayed.
3. If necessary, navigate to the folder where the file is to be saved.
4. GeneLinker™ supplies a default file name based on the name of the item in the navigator and a file type extension (.csv). You can use the default file name or you can type over it.
5. Click **Save** to save the file, or click **Cancel** to quit the operation without saving the file.

**Related Topics**
- Overview of Jarvis-Patrick Clustering
- Overview of K-Means Clustering

**Jarvis-Patrick**

**Jarvis-Patrick Clustering Overview**

**Overview**
Jarvis-Patrick clustering is a clustering method based on similarity between neighbors. Similarity (or closeness) is determined by using a distance metric. One or more Neighbors in Common are used to judge the cluster membership of the objects under study. The function is deterministic and non-iterative.

**Algorithm Properties**
- The algorithm chooses the number of clusters.
- There is always at least one item in each cluster.
- The algorithm partitions the input into non-hierarchical clusters.
- The clusters do not overlap.
- If two different items from the input dataset share enough mutual nearest neighbors, then those two items are in the same cluster.

**Parameters**
General clustering parameters, distance measurements between data points, and distance measurements between clusters are used to perform this procedure. In addition to these general clustering parameters, there are two parameters specific to the
Jarvis-Patrick algorithm:

- the number of **Neighbors to Examine**
- the minimum required number of **Neighbors in Common**.

The first parameter, **Neighbors to Examine**, specifies how many of each item’s neighbors to consider when counting the number of mutual neighbors shared with another item. This value must be at least 2. Lower values cause the algorithm to finish faster, but the final set of clusters will have many small clusters. Higher values cause the algorithm to take longer to finish, but may result in fewer clusters and clusters that form longer chains.

The second parameter, **Neighbors in Common**, specifies the minimum number of mutual nearest neighbors two items must have for them to be in the same cluster. This value must be at least 1, and must not exceed the value of the **Neighbors to Examine** parameter. Lower values result in clusters that are compact. Higher values result in clusters that are more dispersed.

**Basic Procedure**

- For each object, find its J-nearest neighbors where ‘J’ corresponds to the **Neighbors to Examine** parameter on the **Partitional Clustering** dialog.
- Two items cluster together if they are in each other’s list of J-nearest neighbors and K of their J-nearest neighbors are in common, where the K value corresponds to the **Neighbors in Common** parameter on the **Partitional Clustering** dialog.

In **GeneLinker™**, input provided to the algorithm is as follows:

- The dataset.
- A distance metric.
- The number of nearest **Neighbors to Examine**.
- The number of nearest neighbors two data points must share to be in the same cluster (**Neighbors in Common**).

**When to Use The Jarvis-Patrick Algorithm**

Use this algorithm when you need to work with non-globular clusters, when tight clusters might be discovered in larger loose clusters, when a deterministic partitional clustering result is desired, or when clustering speed is an issue since the algorithm is not iterative.

**Related Topics:**

- Performing Jarvis-Patrick Clustering
- Clustering Overview
- Tutorial 3: Jarvis-Patrick Clustering

**Performing Jarvis-Patrick Clustering**
**Overview**

The Jarvis-Patrick clustering algorithm is good for detecting chain-like or non-globular clusters. It partitions data into clusters, generating a set of non-overlapping clusters. For further details, see Overview of Jarvis-Patrick Clustering.

**Actions**

1. Click a complete dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Partitional Clustering** toolbar icon, or select **Partitional Clustering** from the **Clustering** menu, or right-click the item and select **Partitional Clustering** from the shortcut menu. The **Partitional Clustering** parameters dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clustering Orientation</td>
<td>Cluster by <strong>Genes</strong> or <strong>Samples</strong>.</td>
</tr>
<tr>
<td>Distance Measurements</td>
<td>Type of distance measurement to use to determine how close two data points are to each other.</td>
</tr>
<tr>
<td>Between Data Points</td>
<td><strong>Type</strong></td>
</tr>
<tr>
<td></td>
<td>Set this parameter to <strong>Jarvis-Patrick</strong>.</td>
</tr>
<tr>
<td>Neighbors to Examine</td>
<td>This value must be at least 2.</td>
</tr>
<tr>
<td>Neighbors in Common</td>
<td>This value must be at least 1, and must not exceed the value of <strong>Neighbors to Examine</strong>.</td>
</tr>
</tbody>
</table>

4. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Jarvis-Patrick Clustering operation is performed. To cancel the Jarvis-Patrick operation, click the **Cancel** button.

Upon successful completion, a new item is added under the original item in the **Experiments** navigator.

**Related Topics:**

Distance Metrics Overview
Agglomerative Hierarchical

Overview
Agglomerative hierarchical clustering is a bottom-up clustering method where clusters have sub-clusters, which in turn have sub-clusters, etc. The classic example of this is species taxonomy. Gene expression data might also exhibit this hierarchical quality (e.g. neurotransmitter gene families). Agglomerative hierarchical clustering starts with every single object (gene or sample) in a single cluster. Then, in each successive iteration, it agglomerates (merges) the closest pair of clusters by satisfying some similarity criteria, until all of the data is in one cluster.
The hierarchy within the final cluster has the following properties:

- Clusters generated in early stages are nested in those generated in later stages.
- Clusters with different sizes in the tree can be valuable for discovery.

A Matrix Tree Plot visually demonstrates the hierarchy within the final cluster, where each merger is represented by a binary tree.

Process
- Assign each object to a separate cluster.
- Evaluate all pair-wise distances between clusters (distance metrics are described in Distance Metrics Overview).
- Construct a distance matrix using the distance values.
- Look for the pair of clusters with the shortest distance.
- Remove the pair from the matrix and merge them.
- Evaluate all distances from this new cluster to all other clusters, and update the matrix.
- Repeat until the distance matrix is reduced to a single element.

Advantages
- It can produce an ordering of the objects, which may be informative for data display.
- Smaller clusters are generated, which may be helpful for discovery.

Disadvantages
- No provision can be made for a relocation of objects that may have been 'incorrectly' grouped at an early stage. The result should be examined closely to ensure it makes sense.
• Use of different distance metrics for measuring distances between clusters may generate different results. Performing multiple experiments and comparing the results is recommended to support the veracity of the original results.

**Divisive Hierarchical Clustering**

• A top-down clustering method and is less commonly used. It works in a similar way to agglomerative clustering but in the opposite direction. This method starts with a single cluster containing all objects, and then successively splits resulting clusters until only clusters of individual objects remain. **GeneLinker™ does not support divisive hierarchical clustering.**

**Related Topics:**
- Clustering Overview
- Performing Agglomerative Hierarchical Clustering

**Performing Agglomerative Hierarchical Clustering**

**Overview**

Agglomerative hierarchical clustering starts with each gene or sample as a single cluster, then in each successive iteration, it merges two clusters together until all genes or samples are in one big cluster.

For further details, see Overview of Agglomerative Hierarchical Clustering.

**Actions**

1. Click a complete dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Hierarchical Clustering** toolbar icon
3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Clustering Orientation</strong></td>
<td>Cluster by <strong>Genes</strong> or by <strong>Samples</strong>.</td>
</tr>
</tbody>
</table>
Distance Measurement Between Data Points Type of distance measurement to use to determine how close two data points are to each other.

Distance Measurement Between Clusters Type of distance measurement to use to determine how close two clusters are to each other.

4. Click OK. The Experiment Progress dialog is displayed. It is dynamically updated as the Agglomerative Hierarchical Clustering operation is performed. To cancel the Agglomerative Hierarchical Clustering operation, click the Cancel button.

Upon successful completion, a new item is added under the original item in the Experiments navigator.

Related Topics:
- Distance Metrics Overview
- Clustering Overview

Self-Organizing Maps (SOMs)

Self Organizing Maps Overview

Overview
The Self-Organizing Map (SOM) is a clustering algorithm that is used to map a multi-dimensional dataset onto a (typically) two-dimensional surface. This surface (a map) is an ordered interpretation of the probability distribution of the available genes/samples of the input dataset. SOMs have been used extensively in many domains, including the exploratory data analysis of gene expression patterns.

There are two particularly useful purposes for this: visualization and cluster analysis. Visualization has typically been a difficult matter for high-dimensional data. SOMs can be used to explore the groupings and relations within such data by projecting the data on to a two-dimensional image that clearly indicates regions of similarity. Even if visualization is not the goal of applying SOM to a dataset, the clustering ability of the SOM is very useful.

Related Topics:
- Performing a SOM Experiment
- Creating a SOM Plot
- Tutorial 4: Self-Organizing Maps
Performing a SOM Experiment

Overview
This procedure explains how to create a SOM experiment for a dataset. The results of this experiment can be visualized in various types of plots to provide you with additional data mining information.

Actions
1. Click a dataset in the Experiments navigator. The item is highlighted.
2. Click the Self-Organizing Map toolbar icon, or select Self-Organizing Map from the Clustering menu, or right-click the item and select Self-Organizing Map from the shortcut menu. The Self-Organizing Map parameters dialog is displayed.

3. Set the dialog parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Orientation</td>
<td>This indicates whether to cluster samples or genes. The default is Genes.</td>
</tr>
<tr>
<td>Distance Metric</td>
<td>This indicates which metric to use to determine distances. The default is Euclidean. Other options are Manhattan, Pearson Correlation, Pearson Squared, Euclidean Squared, and Chebychev.</td>
</tr>
<tr>
<td>Height</td>
<td>This indicates how many nodes high the map shall be. The default height is 4.</td>
</tr>
<tr>
<td>Width</td>
<td>This indicates how many nodes wide the map shall be. The default width is 4.</td>
</tr>
<tr>
<td>Initialization</td>
<td>Method to initialize the reference vectors of the nodes. It can be set to Random Sample (default) or Random Value. Random sample refers to the assignment of randomly selected items</td>
</tr>
</tbody>
</table>
genes/samples) from the dataset to be the initial reference vectors of the map.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Range</strong></td>
<td>If the reference vectors are initialized by Random Values, then <strong>Range</strong> sets the bounds on random values, where values are chosen from the real number range [-value_range, value_range]. The default is 1.</td>
</tr>
<tr>
<td><strong>Number of iterations</strong></td>
<td>Indicates the number of iterations to perform on the SOM. During each iteration, the SOM learns from one item (sample or gene). This must be an integer greater than zero. A good rule-of-thumb is to use the number of cluster items or 500 times the number of nodes, whichever is greater. The default is 8000 to match the default map size (4<em>4</em>500 =8000).</td>
</tr>
<tr>
<td><strong>Radius length</strong></td>
<td>This is an integer that indicates the initial area on the map that can be affected during an iteration of learning (i.e. the bubble neighborhood). The unit of measure is the number of nodes. The default is 3.</td>
</tr>
<tr>
<td><strong>Random seed</strong></td>
<td>This is an integer value that indicates the random seed used by the SOM algorithm, and allows you to perform repeatable experiments. The default is a random number.</td>
</tr>
</tbody>
</table>

4. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the SOM operation is performed. To cancel the SOM operation, click the **Cancel** button.

Upon successful completion, a new SOM item is added under the original item in the **Experiments** navigator.

**Plotting a SOM Experiment:**
- SOM Plot
- Centroid Plot
- Cluster Plot
- Matrix Tree Plot

**Related Topics:**
- Overview of Self-Organizing Maps (SOMs)
- Tutorial 4: Self-Organizing Maps

**Principal Components Analysis (PCA)**

**Overview of Principal Component Analysis (PCA) Functionality**
Overview

Component Analysis is an unsupervised or class-free approach to finding the most informative or explanatory features in data. In particular, Principal Component Analysis (PCA) substantially reduces the complexity of data in which a large number of variables (e.g. thousands) are interrelated, such as in large-scale gene expression data obtained across a variety of different samples or conditions. PCA accomplishes this by computing a new, much smaller set of uncorrelated variables which best represent the original data. PCA is a powerful, well-established technique for data reduction and visualization. 2D and 3D PCA plots often place objects with similar patterns near each other.

GeneLinker™ provides one option for PCA analysis: Orientation by Genes or Orientation by Samples. In brief, PCA oriented by genes is useful for distinguishing sample classes or sample clusters, while PCA oriented by samples is useful for distinguishing gene classes or gene sets.

Mathematical Details and Examples of Orientation

To understand the difference and interpretive implications between the two different orientations - PCA by Genes or PCA by Samples - it is helpful to conceptualize the data analysis from the point of view of covariance matrices. A dataset can be thought of as comprising distinct mathematical or statistical variables (e.g. columns) for which there are statistical samples (e.g. rows).

a) Genes vs. Genes (Orientation by Genes)

- Typically, genes are considered the mathematical or statistical variables and samples are considered the statistical samples. The corresponding covariance matrix (if it were computed) would carry the covariance of one gene vs. another gene, assessed over the samples, and recorded for each pairwise combination of genes (i.e., pairwise combinations of the statistical variables). Thus, if there are n genes and m samples, the corresponding covariance matrix would comprise n by n entries, each entry being the covariance of the ith gene vs. the jth gene, i and j running from 1 through n. The ith element along the diagonal of this covariance matrix is simply the conventional variance of the ith variable, in this case, the variance of the ith gene over all the m samples.

b) Samples vs. Samples (Orientation by Samples)

- However, if the samples are considered to be the mathematical or statistical variables, then the genes would play the role of the statistical samples. This case is less typical, but is still useful for biological interpretation in some situations (e.g., when the samples are different specific times of the cell cycle). In this case, the corresponding covariance matrix (if we were to compute it) would comprise m by m entries, each entry being the covariance of the ith sample vs. the jth sample from the data matrix. However, this time i and j run from 1 through m. Again, the ith element along the diagonal of this covariance matrix is simply the conventional variance of the ith variable. In this case, it is the variance of the ith sample (i.e., the ith mathematical or statistical variable) over all the n genes (the statistical samples).

In GeneLinker™, a Principal Component (PC) is defined as a mathematical entity (i.e., vector) computed from the data which is equivalent to a characteristic vector (i.e.,
eigenvector) of a covariance matrix derived from the data. This is equivalent to finding the best lower dimensional linear basis set in which to represent the original data under the constraint of minimizing residual variance. The results obtained from the GeneLinker™ implementation are equivalent to a classical PCA of the data's covariance matrix; however, for computational speed and accuracy, covariance matrices are not explicitly computed by GeneLinker™ for PCA. From a covariance point of view, for example, a dataset typically comprises n genes by m samples. One can conceptualize two different kinds of covariance matrices for this data archetype:

a) **Orientation by Genes**: n by n covariance matrix (genes in the role of the math/statistics variables; hence, n genes vs. n genes, aggregated over all samples) OR

b) **Orientation by Samples**: m by m covariance matrix (samples in the role of the math/statistics variables; hence, m samples vs. m samples, aggregated over all genes).

For example, if there are n=1000 genes and m=12 samples (12 different human subjects, for example), the covariance matrix for case (a) would have 1000000 elements (1000 x 1000), but the covariance matrix for case (b) would have only 144 elements (12 x 12).

**Technical Notes**

Whether PCA orientation by genes or by samples, the maximum number of bona fide Principal Components that can be returned is the smaller of the number of genes or the number of samples. This is an inherent mathematical constraint.

PC calculation does not require parameters, and none are set by you beyond selecting the orientation of the calculation. The **PCA Components to Display** setting in the Preferences (accessed from the Edit menu) only affects display and reporting. The default limit on the number of PCs displayed in the Scree and Loadings plots is 15. This setting does not affect the actual calculation of the PCs. It sets an upper limit only on the number of PC's to display in these plots; therefore it does not have to be set before the PCs are calculated.

Whether the user requests PCA of count data, log data, max-min normalized data, missing value-replaced data, etc., GeneLinker™ automatically zero-means the data 'variables' before the PCA calculation, as is required for the results to be mathematically equivalent to the PCA of the covariance matrix.

GeneLinker™ limits the number of PCs by their contribution towards representing fractions of the total variance of the data (i.e., their numerical relevance). Only PCs associated with respective eigenvalues greater than or equal to 10⁻⁸ are included in the calculation result set. But in practice PCs with respective eigenvalues (i.e., fractions of data total variance) less than about 0.1 are rarely of much interpretive use or value.

Note also that a PC's pointing direction (e.g., southeast rather than northwest) along the line co-linear with the PC is irrelevant. Therefore, reversing the algebraic signs of all the constituent values of a PC in, for example, a Loadings Line Plot, is irrelevant.

**Related Topics:**
Performing PCA for a Dataset

Overview

GeneLinker™ has the facility to perform Principal Components Analysis (PCA) on a dataset. For a complete description of PCA, see Overview of Principal Components Analysis.

Actions

1. Click a complete dataset in the Experiments navigator. The item is highlighted.
2. Click the Principal Component Analysis toolbar icon, or select Principal Component Analysis from the PCA menu, or right-click the item and select Principal Component Analysis from the shortcut menu. The PCA parameters dialog is displayed.

3. Select whether to perform PC calculation on either Genes or Samples.
4. Click OK. The Experiment Progress dialog is displayed. It is dynamically updated as the PCA operation is performed.

Upon successful completion, a new Gene or Sample Principal Components Analysis item is added under the original item in the Experiments navigator.

Plotting PCA Results

1. Click a Gene or Sample Principal Component Analysis item in the Experiments navigator. The item is highlighted.
2. Select a plot type from the PCA menu. For a complete description of the plot, please see:
   - Scree Plot
   - Loadings Line Plot
   - Loadings Scatter Plot
   - Loadings Color Matrix Plot
Classification and Prediction

SLAM

ANN Classification and Prediction Overview

Overview
ANN Classification, in GeneLinker™, is the process of learning to separate samples into different classes by finding common features between samples of known classes. For example, a set of samples may be taken from biopsies of two different tumor types, and their gene expression levels measured. GeneLinker™ can use this data to learn to distinguish the two tumor types so that later, GeneLinker™ can diagnose the tumor types of new biopsies. Because making predictions on unknown samples is often used as a means of testing the ANN classifier, we use the terms training samples and test samples to distinguish between the samples of which GeneLinker™ knows the classes (training), and samples of which GeneLinker™ will predict the classes (test).

Types of Learning
ANN Classification is an example of Supervised Learning. Known class labels help indicate whether the system is performing correctly or not. This information can be used to indicate a desired response, validate the accuracy of the system, or be used to help the system learn to behave correctly. The known class labels can be thought of as supervising the learning process; the term is not meant to imply that you have some sort of interventionist role.

Clustering is an example of Unsupervised Learning where the class labels are not presented to the system that is trying to discover the natural classes in a dataset. Clustering often fails to find known classes because the distinction between the classes can be obscured by the large number of features (genes) which are uncorrelated with the classes. A step in ANN classification involves identifying genes which are intimately connected to the known classes. This is called feature selection or feature extraction. Feature selection and ANN classification together have a use even when prediction of unknown samples is not necessary: They can be used to identify key genes which are involved in whatever processes distinguish the classes.

Manual Feature Selection
Manual feature selection is useful if you already have some hypothesis about which
genes are key to a process. You can test that hypothesis by:

i. constructing a gene list of those genes,

ii. running an ANN classifier using those genes as features, and

iii. displaying a plot which shows whether the data can be successfully classified.

**Feature Selection Using the SLAM™ Technology**

The genes that are frequently observed in associations are frequently good features for classification with artificial neural networks. In GeneLinker™, ANN classification is done using a committee of artificial neural networks (ANNs). ANNs are highly adaptable learning machines which can detect non-linear relationships between the features and the sample classes. A committee of ANNs is used because an individual ANN may not be *robust*. That is, it may not make good predictions on new data (test data) despite excellent performance on the training data. Such a neural network is referred to as being overtrained.

Each ANN (component neural network or learner) is by default trained on a different 90% of the training data and then validated on the remaining 10%. (These fractions can be set differently in the Create ANN Classifier dialog by varying the number of component neural networks.) This technique mitigates the risk of overtraining at the level of the individual component neural network.

The committee architecture further enhances robustness by combining the component predictions in a voting scheme. Finally, by examining a chart of the voting results, difficult-to-classify samples can often be identified for re-examination or further study.

**Related Topics:**

- An Introduction to Classification: Feature Selection
- Association Mining Using SLAM™
- Creating an ANN Classifier
- Classify New Data

**An Introduction to Classification: Feature Selection**

This document introduces the topic of classification, presents the concepts of features and feature identification, and ultimately discusses the problem that GeneLinker™ Platinum solves: finding non-linearly predictive features that can be used to classify gene expression data. Many examples, some very simple, are used clarify subtle and sometimes difficult concepts.

**Classification**

There are several types of classification:

<table>
<thead>
<tr>
<th>Type of Classification</th>
<th>Description</th>
<th>Example</th>
</tr>
</thead>
</table>

---

GeneLinker Gold 3.1 / GeneLinker Platinum 2.1
### Categorical Classification

**Nominal**

Classification of entities into particular categories.

| Example | That thing is a dog. | That thing is a car. |

**Ordinal**

Classification of entities in some kind of ordered relationship.

| Example | You are stronger than him. | It is hotter today than yesterday. |

**Adjectival or Predicative**

Classification based on some quality of an entity.

| Example | That car is fast. | She is smart. |

**Cardinal**

Classification based on a numerical value.

| Example | He is six feet tall. | It is 25.3 degrees today. |

Categorical classification is also called nominal classification because it classifies an entity in terms of the name of the class it belongs to. This is the type of classification we focus on in this document.

### Features

If we think for a minute about how we classify common everyday objects such as people and cars, it's pretty clear that we are using **features** of those objects to do the job. People have legs, that's a feature that cars don't have. Cars have wheels, that's a feature that people don't have. By selecting the **appropriate set of features**, we can do a good job of classification.

To make this kind of feature-based classification work, we need to have some knowledge of what features make good predictors of class membership for the classes we are trying to distinguish. For example, having wheels or not distinguishes people from cars, but doesn't distinguish cars from trains. These are two different classification tasks. Depending on the classification task we are facing, different features or sets of features may be important, and knowing how we arrive at our knowledge of which features are useful to which task is essential.

### Learning

The general process by which we gain knowledge of which features matter in a given discrimination task is called **learning**. For those of us who are parents, one example of this type of learning (feature selection) involves teaching our children about types of animals. We (endlessly) point to animals and say words like *dog* or *cat* or *horse*. We don't generally give our children a feature list that a biologist might use to define *Canis familiaris* or *Felis catus*. Instead, we present examples and expect our children to figure out for themselves what the important features are. And when they make a correct guess about an animal (a correct classification or prediction), we give copious amounts of positive feedback.

This procedure is called **supervised learning**. We present our children or our computer programs with examples and tell them what category each example belongs to, so they learn under our **supervision**. This is in contrast to **unsupervised learning**. In unsupervised learning objects are grouped together based on perceptions of similarity (or more properly, relative lack of difference) without anything more to go on. While unsupervised learning is indispensable, supervised learning has a substantial
advantage over unsupervised learning.

In particular, supervised learning allows us to take advantage of our own knowledge about the classification problem we are trying to solve. Instead of just letting the algorithm work out for itself what the classes should be, we can tell it what we know about the classes: how many there are and what examples of each one look like. The supervised learning algorithm’s job is then to find the features in the examples that are most useful in predicting the classes.

The clustering algorithms in GeneLinker™ Gold are examples of unsupervised learning algorithms. The classification workflows of GeneLinker™ Platinum are examples of supervised learning algorithms. They are more complex than clustering, and sometimes more frustrating due to their additional complexity, but they have considerable advantages.

The classification process with supervised learning always involves two steps:

1. **Training** (with assessment) – this is where we discover what features are useful for classification by looking at many pre-classified examples.
2. **Classification** (with assessment) – this is where we look at new examples and assign them to classes based on the features we have learned about during training.

This process isn’t perfect, particularly if the number of examples used in training is small. A difficult problem is how to handle objects that don’t fall into any of the classes we know about. There is a tendency to categorize them as belonging to one of the classes we do know about, even if the fit is rather poor. For example, upon seeing a horse for the first time, my son announced, 'Look! Big dog!' GeneLinker™ Platinum's classification algorithms are capable of making this kind of error.

### The Problem

The problem that GeneLinker™ Platinum is the solution to is the classification problem, which is:

1. How do we find a set of features that is a good predictor of what class a sample belongs to?
2. Having found a good set of features, how do we use it to predict what classes new samples belong to?

The first part of the classification problem, which is by far the hardest, is solved by the Sub-Linear Association Mining (SLAM™) and other Molecular Mining Corporation proprietary data mining algorithms. The second is solved by our committee of artificial neural networks (ANNs).

### Feature Selection

#### Features in Data

Before getting into feature selection in more detail, it's worth making concrete what is meant by a feature in gene expression data. The figure below shows two genes with 100 samples each. One gene, call it Gene A, clearly has an enhanced expression value around sample 50. This expression level ‘bump’ is a feature. If every gene expression
profile from tissues of the same class showed the same bump, this feature would be a good predictor of what tissue class a new sample of tissue belonged to.

Suppose we observed the following data:

<table>
<thead>
<tr>
<th>Tissue Class</th>
<th>Average Expression Level</th>
<th>Gene A, Sample 50</th>
<th>Gene B, Sample 50</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td></td>
<td>3.5</td>
<td>2.5</td>
</tr>
<tr>
<td>Cancer</td>
<td></td>
<td>2.5</td>
<td>2.5</td>
</tr>
</tbody>
</table>

In this case, Gene A has a feature, an enhanced expression level for sample 50, that is a good predictor of which class (Normal or Cancer) a tissue belongs to. Gene B has no such feature, its average expression level at sample 50 is independent of tissue class.

**Probability**

So far, it may seem as though a nice clean distinction between features that distinguish classes clearly and those that don't always exists. In fact, this is rarely the case. Most of the time all we see is an enhanced correlation between a feature and a class.

For example, tall people tend to be stronger than short people. There are several reasons for this: tall people have longer arms and legs, which gives their muscles more mechanical advantage; tall people tend to have bigger muscles, simply because they are bigger people; and tall people tend to be men, who have higher testosterone levels, which helps them build more muscle. The fact remains, however, that some short women can lift more weight than some tall men. So if we were to try to classify people into two groups, 'strong' and 'weak', without actually measuring how much they can lift, height might be one feature we would use as a predictor. But, it couldn't be the only one if we wanted our classification to be highly reliable.

If a single feature is not a good class predictor on its own, the alternative is to look for one or more sets of features that together make a good predictor of what class an object falls into. For example, neither height nor weight are particularly good predictors of obesity; but taken together, they predict it fairly well.
Linearly Predictive Features
The tissue data above is an example of a linearly predictive feature. That is, when the expression level of gene A goes up at sample 50, the probability that the tissue is normal goes up too. This can be expressed mathematically by the linear equation:

\[ P(\text{normal}) = k \cdot X_A \]

where \( P(\text{normal}) \) is the probability the tissue is Normal, \( X_A \) is the expression level of gene A, and \( k \) is a constant that depends on the specifics of the data. The expression level of gene A at sample 50 is also a linear predictor of the probability that the tissue is a cancer:

\[ P(\text{cancer}) = 1 - P(\text{normal}) = 1 - k \cdot X_A \]

In this case, the linear relationship is inverted: the higher the expression level of gene A at sample 50, the lower the probability of the tissue being in the cancer class.

Combinations of Linearly Predictive Features
The wonderful thing about linearly predictive features is that they combine linearly. This means that they obey the familiar laws of arithmetic when they are combined: literally, \( 2 + 2 = 4 \) for linearly predictive features. This is not the case for non-linearly predictive features. Not only does this make linearly predictive features easy to understand, it makes the algorithmic mathematical problem of finding them tractable.

For example, consider the example of height and weight as predictors of obesity. Although not strictly linear, they are approximately so. They are in fact an example of monotonic predictors - as they increase or decrease, the probability of a sample being in a particular class increases or decreases as well. It is never the case, for example, that a light person is more likely to be obese than a heavy person. The heavier you are, the more likely you are to be obese, no matter how tall you are. Monotonic predictors can usually be approximated by linear predictors, at least over a limited range, as shown in the following figure.

![Graph showing linear and non-linear combinations of height and weight](image)

Biologically, saturation is a common cause of non-linear but still monotonic behavior. For example, if a given enzyme binds to a particular receptor, more enzyme will result in...
a larger effect up to the point where all the receptors are already bound to enzyme molecules. At that point, the system is saturated and the effect won't increase no matter how much more of the enzyme is added.

The figure below shows body mass index (BMI) as a function of height and weight. A BMI of greater than 25 indicates a person who is overweight, and greater than 29 indicates a person who is obese. The dark gray surface is BMI, the light gray surface is a linear approximation to BMI:

$$\text{BMI} = 40 - 0.29 \times \text{Height} + 0.46 \times \text{Weight}$$

As can be seen from the size of the coefficients, height has a smaller influence on BMI than weight does, but neither of them has such a dramatic influence that it would be possible to ignore the other. The linear combination of features, high weight and low height, or very high weight and high height, is required to classify a person as obese.

Mathematically, combinations of linearly predictive features are easy to extract from even fairly small sets of examples. This is because of the fact that linearly mathematical problems are invertible: in one-dimensional terms, if we know $Y = k \times X$, then we also know $k = Y/X$, which gives us the constant that relates the feature to the probability of being in a given class. This process can be generalized to combinations of features as well, ultimately meaning that there are tedious but straightforward deterministic mathematical algorithms for extracting linear combinations of features that have good predictive power. One such algorithm is principal component analysis (PCA).

**Non-linearly Predictive Features**

Not all classes have linearly predictive features: that is, the probability of an object belonging to a given class cannot be written as a linear function of some set of features. For example, consider weight as a predictor of vehicle class. In particular, consider distinguishing cars from aircraft by weight. If a vehicle is very light, it is probably an aircraft. Most small planes weigh quite a bit less than a car. However, if a vehicle is in the range of one to two thousand pounds, it's probably a car, and if it's much heavier than that it's probably a light jet or larger.

In this case, unlike the monotonic, non-linear case, it is practically impossible to approximate the non-linear features with a linear function over a small range. The probability that a vehicle is a car as a function of weight looks something like this:
As this is not a straight line, linear approximations don’t apply.

**Combinations of Non-linearly Predictive Features**

Combinations of non-linearly predictive features are the most general case a feature selector has to handle. Many biological classification problems can only be solved by such combinations, and unfortunately, the problem of finding a good set of non-linearly predictive features is very nearly intractable.

The reason for this is that unlike linear problems, non-linear problems cannot be inverted. There is no way of turning the equation around and extracting the parameters (the equivalents of the linear constants) that will give us good predictions. This means that the only way we have of finding the combinations of features that give us good predictive power is to search for them, checking combinations of features one by one and trying to figure out what combination gives us the best ability to classify objects into different categories of interest.

**The Combinatoric Explosion**

The simplest way to search for combinations of features that give us good predictive power is to start by looking at features one at a time, and trying to find ones that are predictive of the classes we are interested in. But we’ve already seen that sometimes features that have little or no predictive power on their own, like height for obesity, but are very powerful predictors when combined with other features. Therefore, we have to search not only individual features, but also combinations.

If we have ten genes and look at all pairs, we have $10^2 = 100$ possible combinations. If we look at all possible triples we have $10^3 = 1000$ possible combinations, and so on for quads and quintuplets. For a typical 10,000 gene Affymetrix chip, the number of pairs we have to search through is a hundred million, the number of triples is a trillion, and the number of quads and quintuplets is astronomical.

This dramatic increase in the number of possible combinations as the number of samples goes up is known as the *combinatoric explosion*, and it is the source of intractability in non-linear combinatoric feature selection. Non-linearity forces us to use a
search technique to find the features that give us the best classification of our objects of interest, and the combinatoric explosion makes simple exhaustive search impossible on all but the smallest datasets.

An example of a non-linear combinatoric problem we're all familiar with is time management. At any given time there are dozens of things we might plausibly be doing. Time management is essentially a problem of task categorization. There are two classes of task: 'critical', which is the one we should be doing right now; and 'non-critical', which is everything else.

Each task that faces us has many possible features we might use to categorize it as 'critical': how important is it to our long-term goals, to our short-term goals? How much fun would it be? How important is it to our boss or our spouse or our children or our friends? How long have we been putting it off? Do we need to do it to fulfill some condition on another task we need to get done? And so on. Even selecting a few good features out of this short list to let us classify tasks is a hard problem.

People often focus on the single feature that seems to have the most predictive power on its own. They may use features such as ‘serves my short-term goals’ or ‘makes my spouse or boss happy’ to identify ‘critical’ tasks. They forget that even if we are highly focused on productivity, it's still the case that sometimes the most important task is to go lie on the beach and relax. This is a highly non-linear effect. By itself, ‘makes me feel good’ is not a good predictor of whether or not a task is ‘critical’, but taken in combination with other task features, it becomes a valuable member of the most predictive feature set.

The classification problem is hard because features have non-linear effects, and combine together in non-linear ways. This means that there is no way to select features that have good classifying power without doing some kind of search through combinations of features. Because the number of possible combinations of features is impossibly large, simply searching through all feature combinations is not practical.

The Platinum Solution

In the gene expression analysis arena, the solution to this problem is the SLAM™ algorithm embodied in GeneLinker™ Platinum. This algorithm uses intelligent heuristics to guide the search for combinations of features with high predictive value toward a small subset of combinations that have a good chance of correctly classifying all the examples presented to the algorithm. Once a feature set has been identified by SLAM, it can be used to train a committee of artificial neural networks that can be used to classify new examples. This combined workflow of feature selection, neural network training, and applying the trained classifier to new samples is the core of GeneLinker™ Platinum’s powerful classification solution.

Platinum

Discretization

Overview

Discretization is the process of converting real gene expression data into a typically small number of finite values (e.g. high, medium, low). The variation in the original data is maintained in the discretized dataset. Discretization is a necessary precursor to using
association mining algorithms such as SLAM™ to find associations. Discretization is accomplished by assigning each value in a dataset to a bin. The data ranges (bin boundaries) and number of bins are set on the **Discretization** parameters dialog.

**Quantile Discretization**
- In quantile discretization each bin receives an equal number of data values. The data range of each bin varies according to the data values it contains.

**Range Discretization**
- In range discretization the data range of each bin is equal. The number of data values in each bin varies according to the bin range.

**Discretization Target**
Discretization can be based on the genes, samples or all of the data in a dataset.
- **Per Gene**: each gene is divided up into appropriate ranges.
- **Per Sample**: each sample is divided up into appropriate ranges.
- **All Data**: all values in the dataset are used to determine the bin ranges.

**Actions**
1. Click a dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Discretize Data** toolbar icon or select **Discretize Data** from the **Predict** menu, or right-click the item and select **Discretize Data** from the shortcut menu. The **Discretization** dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Operation</strong></td>
<td>Type of discretization: <strong>Quantile</strong> or <strong>Range</strong>.</td>
</tr>
<tr>
<td><strong>Target</strong></td>
<td>Discretize <strong>Per Gene</strong>, <strong>Per Sample</strong> or <strong>All Data</strong>.</td>
</tr>
<tr>
<td><strong>Number of Bins</strong></td>
<td>The number of discrete groups (bins) to put the values into.</td>
</tr>
</tbody>
</table>

4. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Discretization operation is performed. To cancel the Discretization operation, click [image of dialog].
the **Cancel** button.

Upon successful completion, a new dataset is added under the original dataset in the **Experiments** navigator.

**Related Topics:**
- ANN Classification and Prediction Overview
- SLAM™

**Platinum SLAM™**

**Overview**

SLAM™ (Sub-Linear Association Mining) is a proprietary data mining algorithm of Molecular Mining Corporation (MMC) that is used to find correlations between discretized variables or to predict the outcome of a categorical variable. As an aid to supervised learning, SLAM™ is used to find associations in gene expression data so that a list of interesting genes (features) can be created.

**Association Mining Overview**

Association mining is a machine learning technique which detects when sets of variables have certain values occurring together at a rate greater than would happen by chance. In GeneLinker™, the variables are genes. SLAM™ finds sets of gene expression values which co-occur frequently within each dataset. Such sets are called *associations*. For instance, it may happen that in kidney tissue, repression of gene A results in the up-regulation of genes B and C, and down-regulation of gene Q. In this case, we would expect to find an association like this in the dataset:


**Note:** this says nothing about how B, C, and Q are regulated when A is not repressed, or when a different tissue is being considered.

Such an association can be used in GeneLinker™ to find genes which are connected to certain sample classes. Genes which occur in many such associations, or in associations with very high **support** (see below), are likely to be good predictors – that is to say, good candidates for classification features.

**Association Statistics**

- **Support**: the support statistic of an association is the number of samples in the dataset in which that association appears.

- **Matthews correlation**: a measure of the predictive power of an association: How well those gene values predict that particular class. **(Note** that this is not
related in any simple fashion to the ability of those same genes to predict other classes.)

**Actions**

1. Click a Discretization item in the **Experiments** navigator. The item is highlighted.
2. Click the SLAM toolbar icon, or select SLAM from the **Predict** menu, or right-click the item and select SLAM from the shortcut menu. The SLAM parameters dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Representative Variable</strong></td>
<td>The training variables to be used for prediction</td>
</tr>
<tr>
<td><strong>Number of Iterations</strong></td>
<td>The number of SLAM™ iterations.</td>
</tr>
<tr>
<td><strong>Support Lower Bound</strong></td>
<td>Minimum support threshold for SLAM™.</td>
</tr>
<tr>
<td><strong>Matthews Number Lower Bound</strong></td>
<td>Minimum Matthews threshold for SLAM™.</td>
</tr>
<tr>
<td><strong>Results</strong></td>
<td>If the Matthews and Support bounds settings result in a large number of valid associations being discovered, this setting can be used to limit the results returned to the best 100 or 1000 associations.</td>
</tr>
<tr>
<td><strong>Random Seed</strong></td>
<td>The seed value for the random number generator. In normal use, setting the random seed is neither necessary nor recommended. On occasion, you may need to determine whether a certain variation in results is due to the random element, or some other cause. For this reason, you are able to set the random seed to a fixed value, thus controlling that source of variation. In SLAM™, the random seed can be thought of as prescribing the starting point for the search for associations. If SLAM™ is allowed to run long enough, it will find all of an enormous set of associations which inhabit any given dataset, but the smaller you set the number of iterations, the</td>
</tr>
</tbody>
</table>
greater will be the effect of the random seed. Conversely, the random seed matters less and less as the number of iterations grows greater. It is usually better to set the iteration number high and let SLAM™ run overnight than to do repeated runs with different random seeds.

4. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the SLAM operation is performed. To cancel the SLAM operation, click the **Cancel** button.

Upon successful completion, a new item (SLAM) is added under the Discretization item in the **Experiments** navigator.

If automatic visualizations are enabled in your user preferences, the SLAM Association Viewer is displayed upon completion of the SLAM run.

**Related Topics:**
- Discretization
- SLAM™ Association Viewer
- ANN Classification and Prediction Overview

---

**Platinum**

**Creating an ANN Classifier**

**Overview**

In GeneLinker™, an ANN Classifier is actually a committee of artificial neural networks (ANNs).

**Note:** The terms Learner, Component Classifier, and Artificial Neural Network (ANN) are interchangeable. The term Classifier refers to an ensemble (committee) of learners.

**Classify Parameter Descriptions**

**Learners**

- The number of learners to train. The samples are divided into N subsets. Each learner is trained on a different (N-1)/N samples and validated on the remaining 1/N samples. The default number is 10, corresponding to a conventional 10-fold cross-validation scheme. The number can be made as high as the number of samples (corresponding to leave-one-out cross-validation) or as low as 3. For most problems the default of 10 is fine.

**Learner Votes Required**

- This is the number of learners which must vote for the same class in order for the
Classifier to make a call (prediction) on a given sample. If fewer learners than this number agree, then the Classifier will make a class prediction of 'Unknown'. Raising this number may result in fewer misclassifications. Lowering it may lead to fewer 'Unknown' calls.

**Hidden Units**

- This is the number of nodes in the hidden layer of each ANN. All ANNs have the same three-layer architecture: input nodes, hidden nodes, and output nodes. You can think of each node as corresponding to a neuron, and the interconnections between them as synapses, but this model should not be taken too literally.

- There are as many nodes in the input layer as there are input features (genes) in the training dataset. There are as many nodes in the output layer as there are output classes. The number of hidden nodes in the middle layer is typically between these two numbers.

- Setting the number of hidden nodes higher will usually result in overtraining, leading to poor results on test data. Setting the number of hidden nodes too low might result in an inability to learn even the training data, but this is easily detected by examining the results of the Create Classifier experiment. If the default number of hidden nodes yields good training results but poor test results, reduce the number of hidden nodes. If the default yields poor training results, try increasing the number of hidden nodes.

**Conjugate Gradient Method**

- Polak-Ribiere and Fletcher-Reeves are two variants of the conjugate gradient algorithm used to optimize the neural network internal parameters during training. They differ in the formula used to update the search direction in internal parameter space. For details see, for example, C.M. Bishop, 'Neural Networks for Pattern Recognition', Clarendon Press, Oxford, 1995.

**Steps**

- This is the number of conjugate gradient steps which the learner takes between evaluations of the stopping criteria.

**Stopping Criteria: MSE Fractional Change**

- Training of each ANN is stopped when the MSE (mean squared error) drops by less than this fraction between two successive iterations. The MSE is computed on the validation samples -- see 'Learners' above.

**Stopping Criteria: Maximum Iterations**

- The maximum number of times to evaluate the MSE for any individual ANN. An ANN may occasionally fail to reach the Stopping Criterion Threshold (above) even after running for a long time. This parameter limits the number of training cycles and prevents infinite loops.

**Random Seed**

- Randomization is used to select out the validation data for each learner, and to seed the internal parameters of each learner. Setting the random seed to a constant
value is sometimes useful to determine exact sources of variation between different classifiers.

**Actions**

1. Click a dataset that has variable information associated with it in the **Experiments** navigator. The item is highlighted.

2. Click the **Create ANN Classifier** toolbar icon 🔄, or select **Create ANN Classifier** from the **Predict** menu, or right-click the item and select **Create ANN Classifier** from the shortcut menu. The **Create ANN Classifier** parameters dialog is displayed.

3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Representative Variable</strong></td>
<td>A list of all the variables associated with this dataset are shown in the listbox. Select the one that specifies the correct class values that the classifier is to be trained to predict.</td>
</tr>
<tr>
<td><strong>Learners</strong></td>
<td>The number of component learners in the classifier.</td>
</tr>
<tr>
<td><strong>Learner Votes Required</strong></td>
<td>The threshold at which the classifier will make a prediction.</td>
</tr>
<tr>
<td><strong>Hidden Units</strong></td>
<td>The number of nodes in the hidden layer of the learner.</td>
</tr>
<tr>
<td><strong>Conjugate Gradient Method</strong></td>
<td>Specifies the variant of the method to use.</td>
</tr>
<tr>
<td><strong>Steps</strong></td>
<td>The number of conjugate gradient steps between evaluations of the stopping criteria.</td>
</tr>
<tr>
<td><strong>MSE Fractional Change</strong></td>
<td>Learner training stops when the MSE drops less than this threshold between two successive iterations.</td>
</tr>
<tr>
<td><strong>Maximum Iterations</strong></td>
<td>The maximum number of times to evaluate the MSE for a learner.</td>
</tr>
<tr>
<td><strong>Random Seed</strong></td>
<td>Seed value for the random number generator.</td>
</tr>
</tbody>
</table>

4. Click **OK**. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Create Classifier operation is performed. To cancel the Create Classifier...
operation, click the **Cancel** button.

Upon successful completion, a new item (Trained Classifier) is added under the original item in the **Experiments** navigator.

**Related Topics:**
- ANN Classification and Prediction Overview
- Classify New Data
- Classification Plot
- MSE Plot

**IBIS**

**Platinum**

**IBIS Overview**

**Overview**

IBIS (Integrated Bayesian Inference System) is a system that is able to predict class membership for a gene expression dataset containing measurements for the same phenomenon as the dataset used to train the IBIS classifier. One of the major strengths of the IBIS method is its ability to reveal nonlinear and non-monotonic associations between pairs of genes and their concerted response to a particular stimulus such as a drug. Three types of classifiers are available in GeneLinker™: Linear Discriminant Analysis (LDA), Quadratic Discriminant Analysis (QDA), and Uniform/Gaussian Discriminant Analysis (UGDA). Different classifiers predict different responses to a stimulus for a gene or pair of genes. Each prediction has an associated accuracy percentage and an MSE value.

The concept that gene expression levels for a single gene can be used to predict stimulus response in every case is quite primitive. Although LDA classifiers are able to capture this relationship, there are certainly associations in which the response to a particular stimulus fluctuates as a function of the products of multiple genes. QDA and UGDA classifiers are able to uncover such associations.

**Dataset Requirements**

IBIS requires a complete dataset with an associated variable. The variable must contain more than one class value with at least three observations each (meaning the dataset must have at least six samples). Also, the variable cannot include the class 'unknown'. Generating IBIS classifiers can be time and resource intensive, so filtering to remove genes of no interest first is recommended.
**LDA** can be used to discover linear association between pairs of genes.
**QDA** can be used to discover non-linear associations between pairs of genes.
**UGDA** can be used to discover nonlinear, non-monotonic associations between pairs of genes.

In general, it is best to start by creating classifiers using LDA and single genes. Only if the accuracy and MSE values are unsatisfactory should you try QDA/UGDA as well as gene pairs.

**IBIS Workflow**

If you do not have a specific gene or gene pair in mind, the first step is to search the dataset for a gene or gene pair that would act as a good classifier. The IBIS Search process does this generating a set of proto-classifiers with accuracy and MSE statistics. The results of this process can be viewed in the IBIS Search Results Viewer and in the Classifier Gradient Plot.

Next, create a classifier from one of the proto-classifiers or using the gene or gene pair that is of particular interest to you. The results of this step can be visualized in the Classifier Gradient Plot.

Finally a dataset can be classified using the IBIS classifier and the results of that classification can be visualized in the Classification Plot or in the Classifier Gradient Plot.

**Related Topics:**
- IBIS Search
- Create IBIS Classifier From IBIS Search Results
- Create IBIS Classifier Using a Gene or Gene Pair

**Platinum**

**IBIS Search**

**Overview**

The IBIS search examines all of the genes (or gene pairs) in a dataset as predictors for a target variable. If you already know which gene or gene pair you would like to use to create an IBIS classifier, you do not need to perform an IBIS search. Please see Create IBIS Classifier Using a Gene or Gene Pair.

The IBIS search process creates proto-classifiers using the specified parameters and generates accuracy and MSE statistics for each. An item is added to the Experiments navigator which contains a list of the proto-classifiers and their associated statistics. At the end of the search process, no true classifiers exist, only the information about them and how to produce them, hence the term proto-classifier.

There are three models available for creating classifiers: Linear Discriminant Analysis (LDA), Quadratic Discriminant Analysis (QDA), and Uniform/Gaussian Discriminant Analysis (UGDA). In general, it is best to start by creating classifiers using LDA and single genes. Only if the accuracy and MSE values are unsatisfactory should you try
QDA/UGDA as well as gene pairs.

For a single gene search, one proto-classifier is created for each gene in the dataset (to a maximum of 10000). For a gene pair search, one proto-classifier is created for each pair of genes in the dataset (to a maximum of 1000). Generating a list of IBIS proto-classifiers for gene pairs takes much longer than for single genes. It is recommended that you filter your dataset before performing the search to remove any genes that are not relevant to the search.

**Actions**

1. Click a complete dataset with variable information item (dataset name) in the **Experiments** navigator. The item is highlighted.
2. Select **IBIS Classifier Search** from the **Predict** menu, or right-click the item and select **IBIS Classifier Search** from the shortcut menu. The **IBIS Classifier Search** dialog is displayed.

3. Set parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Representative Variable</strong></td>
<td>It cannot contain the class 'unknown' and it must have at least two classes with a minimum of three observations (samples) for each class.</td>
</tr>
<tr>
<td><strong>Background Class</strong> (UGDA only)</td>
<td>Representative variable class to be used as the background reference. Suggestion: select the variable value with the highest frequency in the training data.</td>
</tr>
<tr>
<td><strong>Classifier Type</strong></td>
<td>Select Linear, Quadratic, or Uniform/Gaussian.</td>
</tr>
<tr>
<td><strong>Classifier Dimension</strong></td>
<td>Dimension of the resultant classifier (1 or 2 genes).</td>
</tr>
<tr>
<td><strong>Minimum Standard Deviation</strong></td>
<td>Use this value to capture your estimate of the error in the data measurements. If the value is too small, degenerate non-useful patterns may be created. If the value is too large, you may miss important patterns due to over-smoothing the classifier. As the name suggests, an</td>
</tr>
</tbody>
</table>
appropriate value would be the smallest standard deviation of the expression of any gene/sample pair over a number of replicate measurements. For full details on this parameter, see Tutorial 7: Appendix.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Committee Size</td>
<td>Number of individual classifiers in the classifier.</td>
</tr>
<tr>
<td>Committee Votes Required</td>
<td>Threshold for classifier to make a prediction.</td>
</tr>
<tr>
<td>Random Seed</td>
<td>An initial value for the random number generator. In IBIS, randomization is only used in cross-validation and the committee structure, that is, in designating training and internal validation samples.</td>
</tr>
</tbody>
</table>

4. Click **OK**. The IBIS search is performed and upon successful completion, a new IBIS Search item is added under the original dataset in the **Experiments** navigator.

**Visualization**

The IBIS Search Results Viewer can be used to examine the results of the IBIS search operation.

**Related Topics:**
- IBIS Overview
- Create IBIS Classifier From IBIS Search Results

**Platinum**

**Create IBIS Classifier From IBIS Search Results**

**Overview**

An IBIS classifier can be created from a proto-classifier created by the IBIS search process. It is created using the parameters that were specified for the search. A proto-classifier has a better chance at being a good classifier if it shows high accuracy and low error. Another path from this point is to create a gene list of genes that show up multiple times in higher ranking gene pair classifiers.

**Actions**

1. Double-click an IBIS Search Results item in the **Experiments** navigator. The item is highlighted and the **IBIS Search Results Viewer** is displayed.
2. Click on the gene/gene pair name of one of the listed proto-classifiers. The item is highlighted.

3. Click **Create IBIS Classifier**. The create classifier operation is performed recycling the parameters used to perform the IBIS search. Upon successful completion, a new IBIS Classifier item is added under original dataset in the **Experiments** navigator.

**Visualization**

A Classifier Gradient Plot can be used to examine the results of the Create IBIS
Classifier operation.

Related Topics:
IBIS Overview
IBIS Search

Platinum

Create IBIS Classifier Using a Gene or Gene Pair

Overview
An IBIS classifier can be created from a specified gene or gene pair. There are three models available for creating classifiers: Linear Discriminant Analysis (LDA), Quadratic Discriminant Analysis (QDA), and Uniform/Gaussian Discriminant Analysis (UGDA). In general, it is best to start by creating classifiers using LDA and single genes. Only if the accuracy and MSE values are unsatisfactory should you try QDA or UGDA as well as gene pairs.

Actions
1. Display a table or plot and select a gene or pair of genes.
2. Select Create IBIS Classifier from the Predict menu. The Create IBIS Classifier dialog is displayed.

3. Set parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Representative Variable</td>
<td>It cannot contain the class 'unknown' and it must</td>
</tr>
<tr>
<td><strong>Background Class (UGDA only)</strong></td>
<td>Representative variable class to be used as the background reference. Suggestion: select the variable value with the highest frequency in the training data.</td>
</tr>
<tr>
<td>-------------------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>Classifier Type</strong></td>
<td>Select Linear, Quadratic, or Uniform/Gaussian.</td>
</tr>
<tr>
<td><strong>Classifier Dimension</strong></td>
<td>This will be set to 1 for a single gene, or set to 2 for a gene pair. You cannot change this setting.</td>
</tr>
<tr>
<td><strong>Minimum Standard Deviation</strong></td>
<td>Minimum standard deviation.</td>
</tr>
<tr>
<td><strong>Committee Size</strong></td>
<td>Number of individual classifiers in the classifier.</td>
</tr>
<tr>
<td><strong>Committee Votes Required</strong></td>
<td>Threshold for classifier to make a prediction.</td>
</tr>
<tr>
<td><strong>Random Seed</strong></td>
<td>Initial random seed value.</td>
</tr>
</tbody>
</table>

At the bottom of the dialog, the gene/gene pair that will be used to create the IBIS classifier is listed.

4. Click **OK**. The create classifier operation is performed. Upon successful completion, a new IBIS Classifier item is added to the **Experiments** navigator under the original dataset.

**Visualization**

An Classifier Gradient Plot can be used to examine the results of this operation.

**Related Topics:**
- IBIS Overview
- IBIS Search

**Platinum**

**Classify New Data**

**Overview**

Classification is the process of using a trained classifier to predict the classes of the items in a dataset.

- **If you use an ANN Classifier**, the dataset to be classified must have the same genes as the training dataset, in the same order and without any extra genes.
- **If you use an IBIS classifier**, the dataset must contain the gene or gene pair used to create the IBIS classifier.

**Actions**

1. Click a raw or filtered dataset in the **Experiments** navigator. The item is highlighted.
2. Click the **Classify** toolbar icon 📊, or select **Classify** from the **Predict** menu. The **Classify** dialog is displayed.
3. Set the parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Name</td>
<td>The name of the new item which will be seen in the Experiments navigator.</td>
</tr>
<tr>
<td>Description</td>
<td>An optional description of the item.</td>
</tr>
<tr>
<td>Classifier</td>
<td>The classifier to be used for the class prediction.</td>
</tr>
</tbody>
</table>

4. Click OK. The **Experiment Progress** dialog is displayed. It is dynamically updated as the Classify operation is performed. To cancel the Classify operation, click the **Cancel** button.

Upon successful completion, a new item (Name) is added under the original item in the Experiments navigator.

**Reasons For Misclassifications:**

There are often no misclassifications in the training data – artificial neural networks are fairly powerful and adaptable learners. If there are misclassifications, however, it may be for one of several possible reasons:

- We may be using a set of genes which do not discriminate between the sample classes.
- The training set may be unbalanced. That is, it may have too many examples of one class and not enough of another.
- We may have set the number of hidden units in the neural networks too small.
- The data may contain errors such as mislabelled samples or incorrect measurements.
- The voting threshold may be set too low.
- The stopping criteria may have been set too loose (maximum iterations too small).

The above reasons may affect either training or test results. If the training results are excellent but the test results are poor, it may be for one of the following additional reasons:

- We may have set the number of hidden units in the neural networks too large.
- We may have too many features (genes) for the number of samples in the training set.
set.

- The test data may be drawn from a significantly different population than the training data.
- The test data may not have been normalized in a similar fashion to the training data.
- The test dataset may have been filtered with different genes than the training dataset. (GeneLinker™ checks only that the number of genes used in training and prediction is the same, not their identities).
- The stopping criteria may have been set too tight (maximum iterations too large).

Related Topics:

ANN Classification and Prediction Overview
Classifier Viewer
IBIS Overview

Plots

Clustering Plots

Creating a Scatter Plot

Overview
The scatter plot can be used for the pair-wise comparison of either two samples or two genes. This plot can be launched from the table viewer, color matrix, or matrix tree plot by selecting either two samples or two genes.

In the case of samples, this plot can be used to visually determine those genes that show significant induction or repression between the two selected samples since differentially expressed genes will lie either above or below the line of slope=1.

Similarly, if two genes are selected, the plot will visually display the relative proportion of the two selected genes across all samples. This plot could be used in the case where a great deal of information exists about two genes - for example in the case of co-regulated genes. In this case you might expect the genes to maintain a constant proportion across all samples. Such a plot could be used to visually inspect this hypothesis.

Actions
1. Display a table view or color matrix plot of a dataset, or a matrix tree plot of a clustering experiment.
2. Select two rows (to plot sample vs. sample) or two columns (to plot gene vs. gene) in the table by clicking on the row/column names while holding down the <Ctrl> key.
3. Select Scatter Plot from the Explore menu. A scatter plot of the two rows/columns is displayed.
Interacting With the Plot
Selecting Items
Displaying an Expression Value

Plot Functions
Exporting an Image
Lookup Gene
Annotate
Color by Gene Lists or Variables

Customizing the Plot
Configuring Plot Components
Resizing a Plot

Related Topics:
Creating a Table View of Gene Expression Data
Creating a Color Matrix Plot
Creating a Matrix Tree Plot

Creating a Coordinate Plot
Overview
The coordinate plot is used to view the profile of a gene's expression pattern over all samples, or a sample's expression pattern over all genes.
For a large dataset, a coordinate plot of all genes over all samples may be very busy. For more refined behavior, select one or more genes or samples from the table viewer before creating the coordinate plot. In this case only the selected genes or samples are plotted.

Actions
Displaying a Coordinate Plot of All Genes
1. Click a dataset item in the Experiments navigator. The item is highlighted.
2. Select Coordinate Plot from the Explore menu. A coordinate plot of all genes is displayed.

Displaying a Coordinate Plot of Selected Genes or Samples
1. Click a dataset item in the Experiments navigator. The item is highlighted.
2. Click the Table View toolbar icon, or select Table View from the Explore menu, or right-click the item and select Table View from the shortcut menu. A table view of the dataset is displayed.
3. Select one or more genes or samples for display:
   • Selecting a gene or sample: click on a column or row heading. The name is highlighted.
   • Selecting multiple genes or samples: press and hold the <Ctrl> key and click on column or row headings. The names are highlighted.
• **Selecting a series of genes or samples**: press and hold the <Shift> key and click on column or row names. The names are highlighted.

4. Select **Coordinate Plot** from the **Explore** menu. A coordinate plot of the selected gene(s) is displayed.

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**Interacting With the Plot**

Selecting Items
Displaying an Expression Value

**Plot Functions**

Exporting an Image
Lookup Gene
Annotate
Create Gene List from Selection or Cluster

**Customizing the Plot**

Configuring Plot Components
Resizing a Plot

**Related Topics:**

Summary Statistics

**Creating a Centroid Plot**
Overview

A centroid plot can be used to visualize the centroid or exemplar profile for each of the clusters resulting from a partitional clustering experiment. For example, if you select a K-Means clustering experiment where K=5, a centroid plot of it shows 5 profiles. Each profile represents the average value for all of the members of one of the clusters.

- **If genes were clustered**, each of the profiles represents the average expression value for the genes in a cluster over all samples.
- **If samples were clustered**, each of the profiles represents the average expression value for the samples in a cluster over all genes.

By selecting one or more cluster centroids and then launching a cluster plot, it is possible to visually 'drill down' into the clusters to view the individual member profiles.

Actions

1. Click a Partitional Clustering experiment in the Experiments navigator. The item is highlighted.
2. Select **Centroid Plot** from the Clustering menu, or right-click the item and select **Centroid Plot** from the shortcut menu. A plot of all cluster centroids is displayed.

Using the Plot

Selecting Items
Displaying an Expression Value
Shared Selection

Plot Functions

Lookup Gene
Creating a Cluster Plot

Overview
A cluster plot can be used to display the profiles of individual members within a cluster. The cluster plot can be launched from a partitional clustering experiment in the Experiments navigator or from a centroid plot. By selecting one or more cluster centroids and then launching the cluster plot it is possible to visually 'drill down' into the clusters to view the individual member profiles.

Actions
Showing a Cluster Plot of All Clusters
1. Click a Partitional Clustering experiment in the Experiments navigator. The item is highlighted.
2. Select Cluster Plot from the Clustering menu, or right-click the item and select Cluster Plot from the shortcut menu. A cluster plot of the experiment is displayed.
Showing a Cluster Plot of Selected Cluster(s)

1. Click a Partitional Clustering experiment in the **Experiments** navigator. The item is highlighted.

2. Select **Centroid Plot** from the **Clustering** menu, or right-click the item and select **Centroid Plot** from the shortcut menu. A centroid plot of the experiment is displayed.

3. Select one or more clusters:
   - **Selecting a single cluster**: click on a cluster on the plot or click on a name in the legend.
   - **Selecting multiple clusters**: press and hold the <Ctrl> key and click on clusters on
the plot or in the legend.

- **Selecting a series of clusters**: press and hold the <Shift> key and click on the first and last cluster name in the legend.

4. Select **Cluster Plot** from the **Clustering** menu, or right-click on the plot (or a selected legend item) and select **Cluster Plot** from the shortcut menu. A cluster plot of the selected cluster(s) is displayed.

**Using the Plot**

**Selecting Items**

**Plot Functions**

- Lookup Gene
- Annotate
- Create Gene List from Selection
- Exporting an Image
Customizing the Plot
Configuring Plot Components
Resizing a Plot

Related Topic:
Summary Statistics

Creating a Matrix Tree Plot

Overview
Tree plots visually highlight clustering relationships. They are indispensable for hierarchical clusterings, and can also be used to view partitional clusterings (K-Means and Jarvis-Patrick), and SOMs.

The matrix tree plot is a combined display of a tree plot and a color matrix. At the top, the plot legend consists of a color gradient above an expression value scale. The default range for the scale is from the minimum to the maximum value contained within the dataset. The cluster tree appears to the right of the color array when samples are clustered, or below it when genes are clustered.

The tree for a hierarchical clustering is a close reflection of the agglomerative algorithm that produced it. Consider gene clustering: two very similar genes are joined at a 'node', representing a cluster. That line is joined to the next nearest gene or sub-cluster by another line a little lower, and so on. In the end, closely related genes tend to appear beside each other in the diagram. (Note that the converse is not true - genes appearing beside each other in the tree diagram are only closely related if they are also linked by lines).

In the picture above:
• Cluster Node 1 contains A and B
• Cluster Node 2 contains A, B, and C
• Cluster Node 3 contains A, B, C, D and E
• Cluster Node 4 contains D and E
• Cluster Node 1 merged together the 'closest', Cluster Node 4 the next 'closest', and
Cluster Node 2 the next 'closest' after that. Cluster Node 3 contains all the items from the entire dataset, representing the cluster with the largest distance between its members.

For partitional clustering, there is a separate comb for each cluster, and the combs have only one level (hence the alternative name 'flat clustering'). All items (genes or samples) in a cluster appear together but no further ordering is done on the items within a cluster.

**Actions**

1. Double-click a hierarchical or partitional clustering experiment in the **Experiments** navigator. The item is highlighted and a matrix tree plot of the selected item is displayed.

OR

1. Click a hierarchical or partitional clustering, or a SOM experiment item in the **Experiments** navigator. The item is highlighted.

2. Click the **Matrix Tree Plot** toolbar icon, or select **Matrix Tree Plot** from the **Clustering** menu, or right-click and select **Matrix Tree Plot** from the shortcut menu. A matrix tree plot of the selected item is displayed.

**Plot Indicators**

As you move the mouse pointer over a gene or sample name, a gray bounding box is drawn around its column or row so you can easily see which tiles belong to it.

The name of selected genes or samples are highlighted in dark blue with white text. It is not possible to select genes and samples concurrently.

**Interacting With the Plot**

**Selecting Items**
Displaying a Gene Expression Value

**Plot Functions**
Profile Matching
Color by Gene Lists or Variables
Exporting an Image

**Customizing the Plot**
Changing the Gradient Color and Scale
Resizing Cells in a Color Grid
Toggling the Color Grid On or Off

**Related Topic:**
Creating a Summary Statistics Chart

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**Creating a Two Way Matrix Tree Plot**

**Overview**
A two way matrix tree plot is useful for visualizing the results of two clustering experiments simultaneously. One must be based on genes, and the other on samples and both must be derived from the same original dataset.

**Actions**
1. Press and hold the <Ctrl> key, and then click on two clustering experiments under the same original dataset in the Experiments navigator. One must be sample-based; the other gene-based. Both items are highlighted.
2. Click the Two Way Matrix Tree Plot toolbar icon 📉, or select Two Way Matrix Tree Plot from the Clustering menu, or right-click on of the highlighted items and select Two Way Matrix Tree Plot from the shortcut menu. The plot is displayed with the sample clusters on the right side and the gene clusters on the bottom, relative to the color matrix portion of the plot.
Plot Indicators
As you move the mouse pointer over a gene or sample name, a gray bounding box is drawn around its column or row so you can easily see which tiles belong to which gene or column.

As you move the mouse pointer over the dendrogram portion of the plot, the gray bounding box surrounds the genes that are in that node cluster and a tooltip is displayed listing the number of members and a cluster merge distance reference value.

The name of a selected item (genes or samples) is highlighted in dark blue with white text. One or more items can be selected, however, it is not possible to select genes and samples concurrently.

Interacting With the Plot
Selecting Items
Displaying a Gene Expression Value

Plot Functions
Profile Matching
Color by Gene Lists or Variables
Exporting an Image

Customizing the Plot
Changing the Gradient Color and Scale
Resizing Cells in a Color Grid
Toggling the Color Grid On or Off

Related Topic:
Creating a Summary Statistics Chart

SOM Plots

Creating a SOM Plot

Overview
The SOM plot is a composition of a proximity-gradient map, a cluster membership list showing the items (samples/genes) contained in the selected cluster, and a node/cluster profile plot comparing node and cluster profiles.

The Proximity Gradient Map
The main part of the chart is the proximity-gradient map (it appears as the background in the upper-left hand pane of the chart). This proximity-gradient map is a high-level view of the average proximity (or similarity) between the reference vectors of the SOM. One end of the gradient is used to indicate areas of high average similarity, and the other end of the gradient indicates low average similarity.

Each node in the map is depicted as a small, filled-in circle, and each node represents a single cluster. The nodes of the map are numbered first from left to right, then from bottom to top. Nodes are numbered starting at one. You can see the node's number in a tooltip that appears when you hover the mouse pointer over that node in the map.

The dashed circles around the nodes, called cardinality rings, indicate how many items are contained in the cluster represented by the node. Nodes with the largest radius contain the most items. The selected node has a dashed cardinality ring and its items are listed in the cluster membership list.

The proximity grid shows more accurately the similarity between adjacent nodes. One color indicates high similarity and another color indicates low similarity. Shades in between those two specific colors indicate intermediate degrees of similarity.

The Cluster Membership List
The list to the right of the proximity-gradient map is the cluster membership list. This list always shows the items (samples/genes) in the cluster represented by the selected node.

The Node/Cluster Profile
The plot below the proximity-gradient map is the node/cluster profile. This plot provides information about the map node and the cluster that it represents for the selected node. The blue line in the plot is the profile of the reference vector of the selected node. The red line is the profile of the centroid of the cluster represented by that node. Comparing
these two profiles allows you to determine how well the characteristic profile of the cluster matches the profile of the node.

The pink area behind the node and centroid profiles is the area of one standard deviation around the centroid. The size of that area indicates the fitness of the cluster. Large areas indicate low fitness and small areas indicate high fitness.

**Actions**

1. Double-click a SOM experiment in the **Experiments** navigator. The item is highlighted and a SOM plot of the selected item is displayed.

   OR

1. Click a SOM experiment in the **Experiments** navigator. The item is highlighted.

2. Select **SOM Plot** from the **Clustering** menu, or right-click on the SOM experiment and select **SOM Plot** from the shortcut menu. A SOM Plot of the selected item is displayed.

**Selecting a Node**

1. Click on a node (cluster) in the proximity-gradient map (upper left of plot). The node is ringed by a rotating dashed circle. To the right, a list of the members in the cluster is displayed and below, there is a plot of the cluster profile.
Displaying a Cluster Plot of a Node
1. Click on a node to select it.
2. Select Cluster Plot from the Clustering menu, or right-click on the proximity-gradient map or on the profile plot and select Cluster Plot from the shortcut menu. A cluster plot of the selected node is displayed.

Using the Cluster Membership List Shortcut Menu
1. Right-click in the cluster membership list to display the shortcut menu. Select an item on the menu to activate that function.
   - Lookup gene in a database.
   - Annotate a gene.
   - Create a gene list.

Related Topics:
  - Customizing the SOM Plot
  - Resizing the SOM Plot
  - Tutorial 4: Self-Organizing Maps

Creating a SOM Centroid Plot

Overview
A centroid plot from a SOM plot can be used to see the profiles of the values in the dataset that have been associated with a particular node.

Actions
1. Click on a SOM experiment in the Experiments navigator. The item is highlighted.
2. Select Centroid Plot from the Clustering menu, or right-click on the item and select Centroid Plot from the shortcut menu. A centroid plot of the SOM experiment is displayed.
Interpretation

The SOM centroid plot shows the characteristics of the clusters, i.e., the representative profile (the centroid) and the fitness of the cluster in terms of the standard deviation above and below the representative. This provides important abstract information about how the gene expression data relates to the clustering provided by the SOM. It also shows the corresponding node's reference vector, which allows comparison of the representative profile of the cluster with the node's reference vector to determine how well (on average) the points associated with that node actually match that node's characteristics.

Using the Plot

Selecting Items
Displaying an Expression Value
Shared Selection

Plot Functions

Lookup Gene
Annotate
Create Gene List from Selection or Cluster
Exporting a .PNG Image

Customizing the Plot

Configuring Plot Components
Resizing a Plot
Creating a SOM Cluster Plot

Overview
A SOM cluster plot makes it possible to visually 'drill down' into the a SOM cluster to view the individual member profiles.

Actions
1. Click a SOM experiment in the Experiments navigator. The item is highlighted.
2. Select Cluster Plot from the Clustering menu, or right-click on the item and select Cluster Plot from the shortcut menu. A cluster plot of the SOM experiment is displayed.

Using the Plot
Selecting Items

Plot Functions
Lookup Gene
Annotate
Create Gene List from Selection
Exporting a .PNG Image
Customizing the Plot
Configuring Plot Components
Resizing a Plot

Related Topics:
  - Overview of Self-Organizing Maps (SOMs)
  - Tutorial 4: Self-Organizing Maps

Creating a SOM Matrix Tree Plot

Overview
Tree plots are used to visualize clustering relationships. GeneLinker™ displays a tree plot in conjunction with a color matrix display of values, typically gene expression levels. A legend displays a color gradient and the scale from the minimum to maximum expression value range. The cluster tree appears to the right of the color array (when samples are clustered), or below the color matrix plot (when genes are clustered).

Actions
1. Click on a SOM experiment in the Experiments navigator. The item is highlighted.
2. Click the Matrix Tree Plot toolbar icon, or select Matrix Tree Plot from the Clustering menu, or right-click the item and select Matrix Tree Plot from the shortcut menu. A matrix tree plot of the SOM experiment is displayed.

Plot Indicators
As you move the mouse pointer over a gene or sample name, a gray bounding box is drawn around its column or row so you can easily see which tiles belong to it.
The names of one or more selected items (genes or samples) are highlighted in dark blue with white text. It is not possible to select genes and samples concurrently.
Hover the mouse pointer over a colored tile to see the gene name, sample name and value in a tooltip.

**Interacting With the Plot**
- Selecting Items
- Displaying a Gene Expression Value

**Plot Functions**
- Profile Matching
- Color by Gene Lists or Variables
- Exporting a PNG Image

**Customizing the Plot**
- Changing the Gradient Color and Scale
- Resizing Cells in a Color Grid
- Toggling the Color Grid On or Off

**Related Topics:**
- Overview of Self-Organizing Maps (SOMs)
- Tutorial 4: Self-Organizing Maps

**PCA Plots**

**Creating a Scree Plot**

**Overview**
A Scree Plot is a simple line segment plot that shows the fraction of total variance in the data as explained or represented by each PC. The PCs are ordered, and by definition are therefore assigned a number label, by decreasing order of contribution to total variance. The PC with the largest fraction contribution is labeled with the label name from the preferences file. Such a plot when read left-to-right across the abscissa can often show a clear separation in fraction of total variance where the 'most important' components cease and the 'least important' components begin. The point of separation is often called the 'elbow'. (In the PCA literature, the plot is called a 'Scree' Plot because it often looks like a 'scree' slope, where rocks have fallen down and accumulated on the side of a mountain.)

**Note:** the maximum number of Principal Components to display is set in Preferences under the Edit menu. This only applies to what is displayed in the Scree Plot and the Loadings Line Plot. This setting does not affect the actual calculation of the PCs. It solely sets an upper limit on the number of PC's to display in these two plots; therefore it does NOT have to be set before the PCs are calculated.
GeneLinker™ also limits the number of PCs by their contribution towards representing fractions of the total variance of the data (i.e., their numerical relevance). Only PCs associated with respective eigenvalues greater than or equal to $1E-8$ are included in the calculation result set. But in practice, PCs with respective eigenvalues (i.e., fractions of data total variance) less than about 0.1, are rarely of much interpretive use or value.

Note also that a PC's pointing direction (e.g., southeast rather than northwest) along the line co-linear with the PC is irrelevant. Therefore, reversing the algebraic signs of all the constituent values of a PC in, for example, a Loadings Line Plot, is irrelevant.

**Actions**

1. Click a PCA Experiment in the **Experiments** navigator. The item is highlighted.
2. Select **Scree Plot** from the **PCA** menu, or right-click the item and select **Scree Plot** from the shortcut menu. The Scree Plot is displayed:

The x axis contains the Principal Components sorted by decreasing fraction of total variance explained. (The numerical labels assigned to each PC are according to this
ordering, and persist whether or not the Scree Plot is actually displayed.) The y axis contains the fraction of total variance explained. Along the red line, numerical values of each PC can be seen in a tool tip. Note the 'elbow' in the red line at PC3 in this example; hence, PC1 and PC2 are the most important. PC3 through PC7 are interpreted then as unimportant. Sometimes the PC at the 'elbow' can be considered important too if its fraction of the total is substantial (it is not in this example). The cumulative fraction of total variance explained is also shown in yellow-orange. Numerical values can be seen in a tooltip.

**Interpretation:**

The Scree Plot has two lines: the lower line shows the proportion of variance for each principal component, while the upper line shows the cumulative variance explained by the first N components. The principal components are sorted in decreasing order of variance, so the most important principal component is always listed first.

**Using the Plot**

Selecting Items
Displaying an Expression Value

**Customizing the Plot**

Configuring Plot Components
Resizing a Plot

**Plot Functions**

Exporting a PNG Image
Lookup Gene
Annotate

**Related Topics:**

Overview of Principal Component Analysis (PCA) Functionality
Tutorial 5: Principal Component Analysis (PCA)

**Creating a Loadings Color Matrix Plot**

**Overview**

The Loadings Color Matrix Plot is one of three closely related plots (Loadings Line Plot, Loadings Scatter Plot, and Loadings Color Matrix Plot) that displays the individual elements the PCs. (Since a PC is a vector, it has constituent elements which are called the loadings. By mathematical definition of PC adopted by GeneLinker™, the Euclidean norm, i.e., vector length, of each PC is 1.) The loadings of a given PC represent the relative extent to which the original 'variables' (Genes or Samples, depending on the
Orientation selected for the PCA) influence the PC. The Loadings Color Matrix Plot displays these loadings as a tiled grid of colored rectangles such as those typically used to view tables and clustering results.

The PCs are in the columns and the 'variables' are in the rows (e.g., Genes if PCA by Genes). The legend displays a color gradient, and the scale of values from the minimum to maximum coefficient value. Often there are so many variables in gene expression data that this type of visualization makes it easier to gain an overview and to interpret than the Loadings Line Plot and Loadings Scatter Plot.

The loadings (color-coded rectangular tiles) can be interpreted as the derived relative weightings of the original 'variables' in the derived linear combination that constitutes each PC. Thus, the color-coded tiles express the relative weights of association between the original 'variables' and the computed PCs.

The default sort for the Loadings Color Matrix plot is in absolute descending order of the first PC.

**Actions**

1. Click a PCA Experiment in the Experiments navigator. The item is highlighted.
2. Click the Loadings Color Matrix Plot toolbar icon, or select Loadings Color Matrix Plot from the PCA menu, or right-click the item and select Loadings Color Matrix Plot from the shortcut menu. The Loadings Color Matrix Plot is displayed.

**Sorting by Principal Component**

At the top of the plot, under each PC label is a button. Only one of these buttons is active at a time. It indicates the current plot sort and by which PC.

The rows of the plot can be sorted by a single PC in:

- Absolute descending order (highest to lowest value regardless of sign).
• Descending order (highest to lowest value).
• Ascending order (lowest to highest value).

The default sort for the Loadings Color Matrix plot is in absolute descending order of the first PC.

To sort by a PC, click on the button under the PC label. This button operates in a cyclic fashion. The cycle is as follows:

1. Click once, the sort is in absolute descending order.
2. Click the same button again, the sort is in descending order.
3. Click the same button again, the sort is in ascending order.
4. Click the same button again and the cycle begins again (absolute descending order).

Each time a sort button is clicked, the plot is updated to reflect the new sort status. In the example below, the samples are sorted in descending order by the 6th PC.

Launching a Loadings Line Plot
1. Select one or more PCs by clicking on the PC label. Press and hold the <Ctrl> key to select multiple PCs. To select a series of PCs, press and hold the <Shift> key and click on the first and last PC labels in the series.
2. Select Loadings Line Plot from the PCA menu, or right-click on the color grid and select Loadings Line Plot from the shortcut menu.

Launching a Loadings Scatter Plot
1. Press and hold the <Ctrl> key and click on two PC labels.
2. Select Loadings Scatter Plot from the PCA menu, or right-click on the color grid and select Loadings Scatter Plot from the shortcut menu.
Creating a Loadings Line Plot

Overview
The Loadings Line Plot is one of three closely related plots (Loadings Line Plot, Loadings Scatter Plot, and Loadings Color Matrix Plot) that displays the individual elements of the PCs. (Since a PC is a vector, it has constituent elements which are called the coefficients or loadings. By mathematical definition of PC adopted by GeneLinker™, the Euclidean norm, i.e., vector length, of each PC is 1.) The loadings of a given PC represent the relative extent to which the original 'variables' (Genes or Samples, depending on the Orientation selected for the PCA) influence the PC. The Loadings Line Plot displays these loadings of a particular PC as a connected line graph. The coefficients or component loadings can be interpreted as the derived relative weightings of the original 'variables' (Genes or Samples, depending on selected Orientation) in the derived linear combination that constitutes each PC. Thus, the coefficients or component loadings express the relative weights of association between the original 'variables' (Genes or Samples) and the computed PCs.

The Loadings Line Plot x axis shows the original 'variables' (e.g., Genes) in the same order in which they appear in the dataset from which the PCs were derived. The y axis shows the numerical values of the loadings. GeneLinker™ assumes the original measurements reflect gene expression levels; hence, the y axis label is 'Loading' regardless of which normalizations may have been performed in producing the dataset upon which the PCA was performed. The y axis ranges across a continuum restricted between -1 and 1, by mathematical definition of PCs (i.e., PCs form an orthonormal basis).

Note that the maximum number of Principal Components (PCs) to display is set in Preferences under the Edit menu. This only applies to what is displayed in the Scree Plot and the Loadings Line Plot. This setting does not affect the actual calculation of the PCs. It solely sets an upper limit on the number of PC's to display in these two plots; therefore it does not have to be set before the PCs are calculated.

GeneLinker™ also limits the number of PCs by their contribution towards representing fractions of the total variance of the date (i.e., their numerical relevance). Only PCs associated with respective eigenvalues greater than or equal to 1E-8 are included in the
calculation result set. But in practice, PCs with respective eigenvalues (i.e., fractions of data total variance) less than about 0.1, are rarely of much interpretive use or value. Note also that a PC’s pointing direction (e.g., southeast rather than northwest) along the line co-linear with the PC is irrelevant. Therefore, reversing the algebraic signs of all the constituent values of a PC in, for example, a Loadings Line Plot, is irrelevant. If you choose the same principal component for both axes, the points may fall outside the unit circle.

**Actions**
1. Click a PCA Experiment in the **Experiments** navigator. The item is highlighted.
2. Select **Loadings Line Plot** from the **PCA** menu, or right-click the item and select **Loadings Line Plot** from the shortcut menu. The PCA Loadings Line Plot is displayed:

![Loadings Line Plot](image)

A Loadings Line Plot allows you to see the relative influence of **Genes** (if PCA by **Genes**) or **Samples** (if PCA by **Samples**) on the PCs. The numerical values can be
interrogated by selecting individual curves for clarity and viewing tooltips. Because the maximum possible range for loadings is the same for all PCs (-1 to 1), it makes comparisons of loadings commensurable. Thus, you could compare, for example, the loading for a given gene on the x axis across each PC as well as compare different genes among one another in their respective contributions to a given PC. In some contexts where the Genes or Samples have been pre-sorted or clustered into meaningful groups, it is possible to identify which groups are most heavily represented in each PC. This can help to identify good PCs for separating gene or sample classes.

**Plot Operations**
- Selecting Items
- Configuring Plot Components
- Resizing a Plot
- Exporting a PNG Image

**Related Topics:**
- Overview of Principal Component Analysis (PCA) Functionality
- Tutorial 5: Principal Component Analysis (PCA)

**Creating a Loadings Scatter Plot**

**Overview**

The Loadings Scatter Plot is one of three closely related plots (Loadings Line Plot, Loadings Scatter Plot, and Loadings Color Matrix Plot) that displays the individual elements the PCs. (Since a PC is a vector, it has constituent elements which are called the loadings. By mathematical definition of PC adopted by GeneLinker™, the Euclidean norm, i.e., vector length, of each PC is 1.) The loadings of a given PC represent the relative extent to which the original 'variables' (Genes or Samples, depending on the Orientation selected for the PCA) influence the PC. The Loadings Scatter Plot displays these loadings compared to one another in a scatter plot of one selected PC vs. another selected PC.

The component loadings or coefficients can be interpreted as the derived relative weightings of the original 'variables' (Genes or Samples, depending on selected Orientation) in the derived linear combination that constitutes each PC. Thus, the component loadings or coefficients express the relative weights of association between the original 'variables' (Genes or Samples) and the computed PCs.

The x axis contains a user-selected PC, and the y axis contains another user-selected PC. The resulting scatter plot then displays the relative associations the original 'variables' with the user-selected PCs. You can then directly change selection of the PCs in the Loadings Scatter Plot on an axis.

**Note:** Plotting a PC against itself will correctly result in points falling outside the unit circle as expected. This is the only case that will do so. However, you should not plot a PC against itself because this provides no useful information.
Actions
1. Click a PCA Experiment in the **Experiments** navigator. The item is highlighted.
2. Select **Loadings Scatter Plot** from the **PCA** menu, or right-click the item and select **Loadings Scatter Plot** from the shortcut menu. The Loadings Scatter Plot is displayed:

![Loadings Scatter Plot](image)

By default, the Loadings Scatter Plot uses the first two PCs as axes.

Changing the PCs
- To change the PC represented by the x-axis, click on a PC in the x-axis drop-down list in the upper left corner of the plot. The plot is updated using the new x-axis PC.
- To change the PC represented by the y-axis, click on a PC in the y-axis drop-down list in the upper center of the plot. The plot is updated using the new y-axis PC.

Using the Plot
Selecting Items
Displaying an Expression Value

Customizing the Plot
Configuring Plot Components
Resizing a Plot

Plot Functions
Exporting a PNG Image
Creating a Score Plot

Overview
The Score Plot involves the projection of the data onto the PCs in two dimensions. The PCs were computed to provide a new space of uncorrelated 'variables' which best carry the variation in the original data and in which to more succinctly represent the original 'samples'. The typical application of PCA is to find the PCs of the Genes ('variables'), and then project the Samples ('samples') onto those PCs. Since typically there are many fewer PCs than genes, it is often easier to see structure in your data with this projection-based plot than it would be in the original data.

The Score Plot is a scatter plot. The x axis contains a user-selected PC. The y axis contains another user-selected PC. The plot contains points that represent the original 'samples' (e.g., projected Samples if PCA by Genes (the 'variables'), projected Genes if PCA by Samples (the 'variables')) projected onto the user-selected PCs. By default, the Score Plot shows data on the first two PCs.

Actions
1. Click a PCA Experiment in the Experiments navigator. The item is highlighted.
2. Select Score Plot from the PCA menu, or right-click the item and select Score Plot from the shortcut menu. The Score Plot is displayed.
Normalizing the Data

The **Raw Data/Normalize** button in the upper right corner of the plot acts as a switch between two views of the data: raw and normalized. The button 'pressed' state displays the normalized view, the 'unpressed' state shows the raw view. The normalized view is shown below:

The normalized view is strictly analogous to, and presents the same information as, the raw view. The essential difference is that in the normalized view, before the points are plotted, the projected values are divided by the Euclidean norm, i.e., vector length, of the respective row of **Samples** (if PCA by **Genes**) or respective column of **Genes** (if PCA by **Samples**).
In some cases, the PCs can be interpreted biologically. This normalized view allows you to easily identify the genes or samples that share the properties of the PCs selected for axes of the plot.

Values close to 1 (one) for any normalized view indicate that the sample or gene is almost parallel to the principal component; -1 implies anti-parallel. This view provides a relative measure of how closely correlated each Sample (if PCA by Genes) or Gene (if PCA by Samples) is to an axis PC.

**Note:** Plotting a PC against itself may correctly result in points falling outside the unit circle. This is the only case that will do so. Plotting a PC against itself provides no useful information.

**Note:** The term 'normalized' here refers to the re-scaling of projections for the 3D Score Plot. It does not refer to any normalizations of the raw data that may, or may not, have been done prior to performing the PCA.

### Changing the PCs

- To change the PC represented by the x-axis, click on a PC in the x-axis drop-down list in the upper left corner of the plot. The plot is updated using the new x-axis PC.
- To change the PC represented by the y-axis, click on a PC in the y-axis drop-down list in the upper center of the plot. The plot is updated using the new y-axis PC.

### Using the Plot

Selecting Items

Displaying an Expression Value

### Customizing the Plot

Configuring Plot Components

Resizing a Plot

### Plot Functions

Exporting a PNG Image

Lookup Gene

Annotate

### Related Topics:

Overview of Principal Component Analysis (PCA) Functionality

Tutorial 5: Principal Component Analysis (PCA)

### Creating a 3D Score Plot
Overview

The 3D Score Plot is a scatter plot. The x, y and z axes represent individual Principal Components (PCs). The plot contains points that represent the original data (projected Samples if PCA by Genes or projected Genes if PCA by Samples) projected onto the individual PCs. By default, the 3D Score Plot shows data on the first three PCs.

Actions

1. Double-click a PCA experiment in the Experiments navigator. The item is highlighted and a 3D score plot of the selected item is displayed.

OR

1. Click a PCA experiment in the Experiments navigator. The item is highlighted.
2. Click the 3D Score Plot toolbar icon or select 3D Score Plot from the PCA menu, or right-click the item and select 3D Score Plot from the shortcut menu. A 3D score plot of the selected item is displayed.

The text area at the bottom of the plot displays the first three principal component values for the point the mouse cursor is pointing at.

Normalizing the Data

The Raw Data/Normalize button in the upper right corner of the plot acts as a switch between two views of the data: raw and normalized. The button 'pressed' state displays the normalized view, the 'unpressed' state shows the raw view.

1. Click the Raw Data/Normalize button. A normalized view of the data is displayed.
The normalized view is strictly analogous to, and presents the same information as, the raw view. The essential difference is that in the normalized view, before the points are plotted, the projected values are divided by the Euclidean norm, i.e., vector length, of the respective row of Samples (if PCA by Genes) or respective column of Genes (if PCA by Samples).

In some cases, the PCs can be interpreted biologically. This normalized view allows you to easily identify the genes or samples that share the properties of the PCs selected for axes of the plot.

Values close to 1 (one) for any normalized view indicate that the sample or gene is almost parallel to the principal component; -1 implies anti-parallel. This view provides a relative measure of how closely correlated each Sample (if PCA by Genes) or Gene (if PCA by Samples) is to an axis PC.

**Note:** The term 'normalized' here refers to the re-scaling of projections for the 3D Score Plot. It does not refer to any normalizations of the raw data that may, or may not, have been done prior to performing the PCA.

**Note:** Plotting a PC against itself may correctly result in points falling outside the unit circle. This is the only case that will do so. Plotting a PC against itself provides no useful information.

**Home Button**
- The Home button returns the plot to its original orientation.

**Refresh Button**
- The Refresh button refreshes the display after you change the choice of principal
components.

Changing the PCs

- To change the PC represented by the X-axis, click on a PC in the X-axis drop-down list in the upper left corner of the plot. Click the Refresh button to update the plot.
- To change the PC represented by the Y-axis, click on a PC in the Y-axis drop-down list in the upper center of the plot. Click the Refresh button to update the plot.
- To change the PC represented by the Z-axis, click on a PC in the Z-axis drop-down list in the upper center right of the plot. Click the Refresh button to update the plot.

Plot Functions

3D Plot Functions

Related Topics:

- Overview of Principal Component Analysis (PCA) Functionality
- Tutorial 5: Principal Component Analysis (PCA)
- Troubleshooting

Classification Plots

Platinum

SLAM™ Association Viewer

Overview

The SLAM™ association viewer is used to visualize the associations found by SLAM™ and to create gene lists. Associations are patterns of a certain value of the target variable co-occurring with certain values of certain genes. For each association, the viewer displays its Matthews correlation, support statistic (the number of samples in the dataset which contain the pattern), class, number of genes in the association, and the list of the gene identifiers.

The Matthews correlation measures the 'interestingness' of an association. More precisely, it measures how well the association can be used to predict its class. If all the samples in a dataset are labelled as true positive (TP), true negative (TN), false positive (FP) or false negative (FN) depending on whether both the expression pattern and the class match the association, then:

\[ M = \frac{(TP \times TN) - (FP \times FN)}{2[TP \times FP \times (TN + FN) \times (TP + FN) \times (FP + TN)]} \]

This gives a value between 1 (very interesting) and -1 (anti-predictive), with a value of zero representing no useful information. Thus values of the Matthews correlation below about 0.5 are unlikely to be of great interest, and values below zero are unlikely to occur.
Support is easier to understand but less powerful than Matthews correlation. The support is simply the number of instances (samples) in the dataset which match the association pattern. In other words, it is the number of true positives (TP) in the Matthews computation. Because SLAM may identify patterns which only cover part of a certain class -- e.g. previously unrecognized molecular subtypes of a cancer -- it is important to remember that a large support number does not necessarily identify a useful association: There may be very interesting (high Matthews) patterns which characterize only parts of the entire dataset and hence have low support.

**Actions**

1. Double-click a SLAM experiment in the **Experiments** navigator. The item is highlighted and the SLAM association viewer is displayed.

OR

1. Click a SLAM experiment in the **Experiments** navigator. The item is highlighted.
2. Click the **Association Viewer** toolbar icon or select **Association Viewer** from the **Predict** menu, or right-click the SLAM item and select **Association Viewer** from the shortcut menu. The SLAM association viewer is displayed.
Creating a Gene List
The SLAM™ association viewer lists the associations on the left and has a place to create a gene list on the right. To populate the gene list, select associations by clicking on the checkboxes next to them in the associations list.

Sorting
To sort the Association list, click on a column header (except Genes). The association list is sorted by that characteristic in the direction indicated by the arrowhead in the column header. The sorting process behaves in a cumulative multi-level manner. Each successive time you click on a column header to sort the list, that characteristic becomes the primary sort key. Previous sorts are maintained in descending order of importance.

To sort the Gene List, click on a column header.

Using the Association Filter
This filter is a real-time control of what is seen in the association list. Click and drag the Minimum Matthews Number slider to expand or contract the number of associations displayed in the association list. The list is updated when you release the mouse button.

To filter the associations by a gene name characteristic, select the characteristic using the drop down list (choices are: is, starts with, contains, does not contain and ends with) and type the gene name or fragment into the text box. The association list is updated (with a slight delay) as you type.

Related Topics:
Creating Gene Lists
Prediction using SLAM™

Platinum
Classification Plot Training Results

Overview
The Classification plot can be used to display the results of training a classifier.

Description
At the top of the viewer is the legend. Dark green is the color of the predicted class and red is the color of a true class.

Each row (sample) has:
• Sample name;
• Prediction (predicted class);
• Class boxes showing the distribution of the votes for each of the possible classes.
• A box that is highlighted in dark green is the predicted class for that sample.
• A box that is highlighted in red is the true class of that sample as specified in the training classes dataset.

**Actions**

1. Click a Trained Classifier item in the **Experiments** navigator. The item is highlighted.
2. Select **Classification Plot** from the **Predict** menu, or right-click the item and select **Classification Plot** from the shortcut menu. A Classification plot of the training results is displayed.

**Interpretation**

• The class of a training sample (that has a true class) that has a dark green box and no red box has been predicted correctly.

• The class of a training sample that has a dark green box and a red box has been predicted incorrectly.

• If no prediction has been made for a sample, it will have no class listed under prediction and no dark green box.

• If a training sample has no true class, it will not have a red box.

**Related Topics:**

Create ANN Classifier
Classify
MSE Plot

**Platinum**

**Classification Plot Classification Results**

**Overview**

The Classification plot can be used to show the results of classification using a trained
classifier.

Description
At the top of the viewer is the legend. Dark green is the color of the predicted class and red is the color of the comparison class, if one is selected. You may choose as a comparison variable any variable of the same variable type as the classifier associated with the same dataset as you are making predictions for.

Each row (sample) has:
- Sample name;
- Prediction (predicted class);
- Class boxes showing the distribution of the votes for each of the possible classes.
- A box that is highlighted in dark green is the predicted class for that sample.
- A box that is highlighted in red is the true class of that sample as specified in the training classes dataset.

Actions
1. Click a Classified item in the Experiments navigator. The item is highlighted.
2. Select Classification Plot from the Predict menu, or right-click the item and select Classification Plot from the shortcut menu. A Classification plot of the classification results is displayed.
3. In the legend, set the Comparison Variable. The classification plot is updated using the comparison variable information.
Interpretation
• The class of a training sample (that has a true class) that has a dark green box and no red box has been predicted correctly.
• The class of a training sample that has a dark green box and a red box has been predicted incorrectly.
• If no prediction has been made for a sample, it will have no class listed under prediction and no dark green box.
• If a training sample has no true class, it will not have a red box.

If the variable you want does not appear in the Comparison variable drop-down list, it may have been imported as a different variable type. Use the Variable Manager to see all the variables available for a given dataset, and what types are assigned to each.

Related Topics:
Create ANN Classifier
Classify
IBIS Overview

Confusion Matrix

Overview
A confusion matrix is a plot used to evaluate the performance of a classifier during supervised learning. It is a matrix plot of the predicted versus the actual classes of the gene expression data.

Actions
1. Select Variable Manager from the Tools menu. The Variable Manager is displayed.

2. Press and hold the <Ctrl> key and click on the two variables of interest (for example, one predicted and one observed).

3. Click Show Confusion Matrix. The Confusion Matrix is displayed.
Interpretation
A confusion matrix is an array showing relationships between true and predicted classes. Entries on the diagonal of the matrix, in blue, count the correct calls. Entries off the diagonal, in red, count the misclassifications. The totals are shown in light blue.

Note that the unknown class is not included in calculating the accuracy of the classifier.

Related Topics:
- Run Classifier
- Classifier Viewer
- Variable Manager

Platinum
MSE Plot

Overview
The Mean Squared Error plot shows the results for each component learner in a training run.

Actions
1. Click an ANN Classifier in the Experiments navigator. The item is highlighted.
2. Select Mean Squared Error Plot from the Predict menu, or right-click on the item and select Mean Squared Error Plot. The training results are displayed.
Interpretation

The MSE is computed by taking the differences between the target and the actual neural network output, squaring them and averaging over all classes and internal validation samples. Because the neural network outputs are real numbers between 0 and 1, this results in a Mean Squared Error between 0 and 1. As the neural network is iteratively trained, the MSE should drop to some small, stable value. Each neural network (component classifier) has its MSE plotted independently. Some components may stop if they reach stability earlier than others, and hence have MSE plots which do not extend over all iterations.

This plot may be used to diagnose certain types of training problems. If several component classifiers show large MSE values even at the end of training, it may be desirable to adjust the training parameters and try again. For instance, the number of hidden units might be increased, the maximum iterations in the stopping criteria might be increased, or the conjugate gradient method or steps number might be changed. If on the other hand only one or two component classifiers show large MSEs at the end, it may indicate inconsistencies between training samples. Consult the Classification Plot and look for samples which show inconsistent voting or 'untidy' histograms. In this case the voting structure of the classifier might result in reasonable classification despite problems with individual component learners.

Related Topics:
Create ANN Classifier
Classify
Classifier Viewer

Platinum

IBIS Search Results Viewer
Overview
The IBIS search results viewer displays a table view of the proto-classifiers that were generated by the IBIS Search using the specified search parameters. For each proto-classifier, the gene/gene pair name, accuracy, and MSE values are listed.

The information displayed in this viewer can be used to assess the proto-classifiers generated by the IBIS search process as a pretext to creating an IBIS classifier. Interesting genes can also be used to create a gene list.

Actions
1. Double-click on an IBIS Search Results item in the Experiments navigator, or right-click the item and select IBIS Search Results Viewer from the shortcut menu. The item is highlighted and the IBIS Search Results Viewer is displayed.
Sorting the List of Proto-Classifiers

1. Click on a column header to sort the list by that characteristic. The list can be sorted in ascending or descending order of gene/gene pair name, accuracy, or MSE.

• **Note:** sorting by gene name for a list of gene pair proto-classifiers sorts on the name of the first gene in each pair.

Displaying a Classifier Gradient Plot
A classifier gradient plot of a single selected proto-classifier can be displayed. A selected proto-classifier is highlighted in blue (whether or not its box is checked).

1. Click on a single gene/gene pair name to select the proto-classifier. The line is highlighted.
2. Click Classifier Gradient Plot. A classifier gradient plot of the selected proto-classifier is displayed.

Creating an IBIS Classifier

An IBIS classifier can be made from a single selected proto-classifier. A selected proto-classifier is highlighted in blue (whether or not its checkbox is checked).

1. Click on the gene/gene pair name of a single proto-classifier to select it. The line is highlighted.
2. Click Create IBIS Classifier. The IBIS classifier is created recycling the parameter settings from the IBIS search. An IBIS Classifier item is added under the training dataset in the Experiments navigator.

Creating a Gene List

1. For Single Gene Proto-Classifiers: Check one or more proto-classifier checkboxes. You can use the <Ctrl> key to check multiple checkboxes, or the <Shift> key to check a series.

   For Gene Pair Proto-Classifiers: Check one or more proto-classifier checkboxes to add their genes to the Genes list box. If the gene is already in the Genes box, then the count for that gene is incremented instead. Check the gene checkboxes in the Genes list box.

2. Click Create Gene List. The Create Gene List dialog is displayed.

3. Provide a Name for the gene list.
4. Optionally provide a Description for the gene list.
5. Click OK. The gene list is created and is added to the Gene Lists navigator.

Related Topics:
   IBIS Overview
   IBIS Gradient Plot
   Create IBIS Classifier From IBIS Search Results
Classifier Gradient Plot

Overview
A classifier gradient plot can be used to visualize the results of creating an IBIS classifier, an IBIS search operation, or classification of a dataset using an IBIS classifier.

Plot Description

**Data points:** The points on the plot represent the gene expression values for the samples in the displayed dataset. By default, the points are colored by the training variable. They may be colored by any associated variable, not just the training variable, to show how well the classifier predicts the other variable. You may display the data points from a compatible dataset or no data points at all.

**Background Gradient:** The plot grid coordinates are run through the classifier to create a background gradient. The color of each pixel in the background represents the classifier's class prediction for that coordinate location. For example, if you represent class x with bright red, then any spot on the background that is red is in a region that the classifier would predict that a sample belongs in class x.

In cases where the classifier is not able to make a certain prediction. (For instance in regions where the predictions shift from class x to y), you may notice that the background blends from one color to the next. The actual color does not change with the strength of the classifier vote; its transparency does. At a point where the committee is 80% sure that the point is blue and 20% sure that the point is red, the final color will be a translucent blue which is 20% transparent and a red which is 80% transparent. (0% being opaque and 100% being invisible). In many cases, the IBIS classifiers are quite certain with their predictions, so a tight boundary usually exists between classes. If you de-select the dominant color, then the other colors become visible.

If you look at the bottom right corner of the left plot, you will notice the color is neither red, green, nor blue. If you uncheck all of the colors and enable them one at a time, you will see that the corner is a combination of red and blue, indicating that the committee of IBIS classifiers was unsure about the class in that region. Some of the committee members voted for red and others voted for blue. The relative intensity of the color tells you if one is more probable than the other.
The blending of colors is much more obvious in the rainbow plot on the right. This plot is of the same data, but the classifier used on it was created with a smaller committee size. With a smaller committee, the chances of it settling on a prediction at a boundary decreases, resulting in much larger shifts in the predictions. You can see regions where the classifier thought there was a chance of the prediction being red, blue and green all at the same time. So although this is a good example on how to interpret the coloring scheme, in general, this exemplifies the value of having a larger committee size (at least 10 or the number of samples in the dataset, whichever is smaller).

**Plot Size:** The X and Y axis ranges are determined by the gene expression values for the data that was used to create the classifier (the training dataset). If you drag a compatible dataset (a dataset that contains the classifier gene or gene pair) onto the viewer, the data points on the plot are replaced with the expression values from the new dataset. If the range of the new data is larger than that of the training data, the scales of the X and/or Y axes are increased to accommodate the new data values. If this happens, a new gradient is produced. The original plot area (training data value ranges) is highlighted by a rectangle on the new plot.

- **Note:** the classifier will not necessarily make informative decisions about a prediction if the data to be predicted is well outside the range that was used to create the classifier.

**Actions**
1. Click an IBIS item in the **Experiments** navigator. The item is highlighted.
2. Select **Classifier Gradient Plot** from the **Predict** menu, or right-click the item and select **Classifier Gradient Plot** from the shortcut menu. A classifier gradient plot of the item is displayed.

---

**Scatter Plot Data Series**

<table>
<thead>
<tr>
<th>Setting</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>None</strong></td>
<td>Turns off the display of the data points from the plot leaving the background gradient.</td>
</tr>
<tr>
<td><strong>Training Data</strong></td>
<td>This is the default setting. The data points are the expression values for the classifier gene or gene pair in the training dataset.</td>
</tr>
<tr>
<td>-------------------</td>
<td>--------------------------------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td><strong>Other Dataset</strong></td>
<td>A dataset that contains the classifier gene or gene pair, with or without associated variables. Drag a dataset from the navigator and drop it on the box. The points on the plot are replaced with the values from the new dataset. <strong>Note:</strong> only one set of data points can be displayed at one time.</td>
</tr>
</tbody>
</table>

**Color by Variable**

Click the **Color by Variable** icon to turn the coloring of the displayed data points on or off.

The variable drop-down list is used to select the variable for coloring the data points. The default setting is coloring by the classes of the training variable.

**Gradient Legend**

This is a list of the classes in the training variable. Each class has a checkbox next to it. If the checkbox is checked, that background gradient color is displayed. To turn off the display of a background class color (e.g. to show a less dominant color as in the example), click the checkbox next to it to uncheck it.

To display the Color Manager, double-click in the **Gradient Legend** box on the dialog, or select **Color Manager** from the **Tools** menu. Use the **Color Manager** to customize the colors used for the plot points and the gradient legend. In the example above, the dominant colors in the background gradient have been turned off.

**Samples**

To the right of the plot is a list of the samples in the currently displayed dataset.

- **To highlight a point and its sample name**, click on a sample in the **Samples** list or a point on the plot.
• To highlight multiple points and their sample names, press and hold the <Ctrl> key and click on the sample names in the Samples list or on points on the plot.

• To highlight a series of points and their sample names, press and hold the <Shift> key and click on the first and last sample names in the Samples list.

Interpretation
This plot could be useful in creating general cause and effect rules. For example, you might be able to tell that there is a correlation between gene expression levels and variable class.

Related Topics:
IBIS Overview
IBIS Search
Classification Plot - Classification Results

Plot Functions

Selecting Items

Overview
You can select one or more genes, samples or clusters on a plot. This can be done on the plot itself or on the plot legend.

Actions
Selecting a Single Gene or Sample
Click on the gene or sample name. The gene is highlighted in the legend and on the plot where appropriate.

Selecting Multiple Genes or Samples
Press and hold the <Ctrl> key and click on the item names.

Selecting a Series of Genes or Samples
Press and hold <Shift> and click on the first item in the series. This becomes the anchor point until the <Shift> key is released. Keep holding the <Shift> key and click another item name. All item from the first clicked to the last clicked (inclusive) are selected.
If you click on another item name, the selected series is de-selected and a new series from the anchor item to the last item clicked is selected.

De-selecting Individual Items in a Series
Release the <Shift> key and press and hold the <Ctrl> key and click on the selected item(s) to be de-selected.
Selecting a Node
Click on the dendrogram when the gray bounding box surrounds the items in the node. The names of the items are highlighted.

- To display a coordinate plot of the selected node, right-click on the plot and select Coordinate Plot from the shortcut menu.
- To display a summary statistics chart of the selected node, right-click on the plot and select Summary Statistics from the shortcut menu.

Select All
To select all of the items in the plot legend and their corresponding items on the plot, right-click on the plot and select Select All from the shortcut menu.

Select None
To de-select all of the items in the plot legend and their corresponding items on the plot, right-click on the plot and select Select None from the shortcut menu.

Related Topics:
- Changing the Gradient Color and Scale
- Resizing Cells in a Color Grid
- Toggling the Color Grid On or Off

Displaying an Expression Value

Overview

Actions
Hover the mouse pointer over the cell in the color grid for which you want to know the value. A tooltip appears displaying the column name, row name and expression value. The tooltip disappears when you move the mouse pointer off that tile. If the expression value is missing then 'N/A' is displayed.

Related Topics:
- Changing Your User Preferences
- Color by Gene Lists or Variables

Shared Selection
Overview

When studying a dataset, it is common practice to examine it from many perspectives. In GeneLinker™, this is done by displaying the dataset values in a table or color matrix plot, or by performing experiments (such as clustering) on the data and displaying the results in different types of plots.

Shared selection is the process by which selecting one or more elements of the same type (such as genes, samples, or clusters) in one table or plot, selects the same element or elements in all other applicable tables or plots instantaneously. This powerful facility makes the features you want to study distinct in all locations concurrently.

For example, if you have a table view and a color matrix plot of a dataset, and a matrix tree and cluster plot of a clustering experiment based on that dataset, selecting a gene in the table viewer instantly selects the same gene in all the other plots.

Element Scope

- A gene has global scope. This means that if a gene is present in more than one dataset, selecting it in a table or plot of one dataset selects it in the tables or plots of the other dataset.
- A sample is relevant to all datasets and experiments derived from a single source dataset. In the Experiments navigator, this means the scope of a sample is a single branch of the tree.
- A cluster is relevant only to the experiment it was created within.
- A principal component is relevant only to the experiment it was created within.

If you have a gene selected, and you display another table or plot that contains that gene, the gene will be highlighted when the new table or plot is displayed.

Actions

**Highlight a gene** on any table or plot or in the Genes, or Gene Lists navigator. The gene is highlighted wherever it exists (tables, plots, navigators).

**Highlight a sample** in a table or plot. The sample and all samples related by sample merging are highlighted on all other tables or plots of datasets or experiments derived from the same dataset.

**Highlight a cluster (or node)** on a centroid or SOM plot (either in the legend or on the plot). One or more of the genes or samples in that cluster are highlighted on any other plots derived from the same source dataset.

Related Topics:

- Selecting Items
- Creating a Table View of Gene Expression Data
- Creating a Color Matrix Plot

Configuring Plot Components
Overview

Several plots - the centroid, cluster, scatter, coordinate, scree, score, loadings line, and loadings scatter plots can be configured using this function to highlight certain features or otherwise enhance the plot. For example, you may find it helpful to customize one or more of the following properties:

- foreground/background colors
- line styles and colors
- axis properties (e.g. logarithmic scale)
- titles

*All customizations made to the appearance of a plot using this function are lost once the plot or GeneLinker™ is closed.*

Actions

1. Right-click on an appropriate type of plot.
2. Select **Customize** from the shortcut menu. The **Properties** dialog is displayed.
3. Click the item you wish to change, and edit the values accordingly. The plot is updated using the new values.
4. Click the **x** icon in the upper right corner of the dialog to close it.

**Related Topic:**

Exporting Images

Resizing a Plot

Overview

The graph portion of a plot can be resized.

Actions

1. Right-click on a plot to display a shortcut menu.
2. Select **Resize** from the shortcut menu. The **Resize** dialog is displayed.
3. Set the **New width** and/or **New height**.
4. Click **OK**. The plot is re-drawn at the specified size.

**Related Topics:**
- Selecting Items
- Configuring Plot Components
- Displaying an Expression Value

**Color By Gene Lists or Variables**

**Overview**
The color matrix, matrix tree, two way matrix tree, scatter, and 3D score plots can be colored by gene list membership and/or by variable. The loadings color matrix plot can be colored by variable only.

- **When color by gene list is enabled**, the color indicator box just below each gene name label is colored according to the color plan specified in the color manager.
- **When color by variable is enabled**, the color indicator box just beside each sample name label is colored according to the color plan specified in the color manager.
- **When both color by gene list and variable are enabled**, the gene list and variable color indicator boxes are colored according to the color plan specified in the color manager.

**Actions**

**Color Matrix, Loadings Color Matrix (Color by Variable only), Matrix Tree, Scatter, or Two Way Matrix Tree Plot**

- **Coloring by Variable**
  1. Select a variable item from the **Color Scheme** list box at the top of the plot in the **Color by** group. **Note:** the **Color by** group is on the plot only if there are variables associated with the displayed dataset or experiment.
  2. Click the **Color Variable** button at the top of the plot (pressed = on). The indicator boxes are colored according to the selected class variable item using the color scheme defined in the Color Manager.

- **Coloring by Gene List**
  1. Select **Color Manager** from the **Tools** menu. The **Color Manager** dialog is displayed.
2. Click the **Gene Lists** tab.

3. Check the boxes to the left of the gene lists to select them.

4. Click the **Coloring by Gene List** button to turn on this feature (‘is on’ is appended to the button name when it is on). The gene names are colored according to list membership in order of priority.

**Note:** for the color indicator boxes to be drawn for genes and/or samples, the color tiles must be at least 10 pixels in width and/or height.

**3D Score Plot**

- **Coloring by Gene List**
  1. Select **Color Manager** from the **Tools** menu. The **Color Manager** dialog is displayed.
2. Check the boxes to the left of the gene lists to select them.

3. Click the **Coloring by Gene List** button to turn on this feature ('is on' is appended to the button name when it is on). The gene names and corresponding points on the plot are colored according to list membership in order of priority.

- **Coloring by Variable**
  1. Click the **Color Scheme** button in the upper left of the plot to turn on color by variable (pressed = on). The sample names and corresponding points on the plot are colored according to their class. To edit the color scheme, use the Color Manager (variables tab).
Related Topics:
  Color Manager
  Creating a Color Matrix Plot
  Creating a 3D Score Plot

Color Manager

Overview
The Color Manager is used to set the colors used for coloring the color matrix, matrix tree, two way matrix tree, or 3D score plot items by gene list and/or variable.

The Color Manager is also used to create the color priority hierarchy for gene list coloring. If a gene is in more than one list, the color used for that gene is the color associated with that gene's highest priority list. For example, if gene 'A' is in lists 1, 2, and 3, and the lists are prioritized with 1 as the highest and 3 as the lowest, the color used for gene 'A' is the color for list 1.

The color scheme is saved between GeneLinker™ sessions.

Actions
  1. Select **Color Manager** from the **Tools** menu. The **Color Manager** dialog is displayed.
Coloring by Gene List
1. Click the Gene Lists tab on the Color Manager dialog.

Enabling/Disabling Coloring by Gene List Function
- Click the button at the top of the Gene Lists pane to toggle coloring by gene list on or off. The button state (pressed/unpressed) and label reflect the current state of the button.

Setting the Gene List Color Priority Hierarchy
1. Click a gene list name. The gene list item is highlighted.
- Click the Up button to move the selected gene list up one spot in the hierarchy (top of list = highest priority).
- Click the Down button to move the selected gene list down one spot in the hierarchy (bottom of list = lowest priority).

To Sort the Gene List Color Priority Hierarchy
- Click the blank column header above the check boxes. The list can be sorted in ascending or descending order of inclusion in display.
- Click the blank column header above the colors. The list can be sorted in ascending or descending order of color.
- Click the Name column header. The list can be sorted in ascending or descending alphabetical order.
- Click on the # column header. The list can be sorted in ascending or descending numerical order.

Enabling/Disabling Coloring by Specific Gene Lists
- Check the checkbox beside a gene list to enable coloring by that gene list.
- Un-check the checkbox beside a gene list to disable coloring by that gene list.

Modifying the Color Used for a Gene List
1. Click a gene list. The item is highlighted.
2. Click the Color button. The Pick a Color dialog is displayed.

3. Select a color for the gene list.
4. Click OK. The Color Manager and all applicable plots are updated with the new color.

**Coloring by Variable**
1. Click the Variables tab on the Color Manager dialog.

**To Select the Class Variables for Coloring**
1. Select a variable item from the Variable Type drop-down list.

**To Sort the Class List**
- Click the Class list header. A small upward pointing triangle appears next to the title indicating the list is sorted in ascending alphabetical order.
- Click the Class list header again. A small downward pointing triangle appears next to the title indicating the list is sorted in descending alphabetical order.

**Modifying the Color Used for a Class**
1. Click on a class. The item is highlighted.
2. Click the Color button. The Pick a Color dialog is displayed.
3. Select a color for the class.
4. Click OK. The **Color Manager** and all applicable plots are updated with the new color.

**Related Topic:**
* Color By Gene Lists or Variables

**Exporting an Image**

**Overview**
You can export the image of a plot to a graphics file. The choices for image file type are PNG, SVG and PDF. PNG (Portable Network Graphics) is a raster graphics format, while SVG (Support Vector Graphics) and PDF (Portable Document Format) are vector graphics formats.

**Raster graphics** are pictures made up of pixels. A photo is the perfect example of a raster graphic. One limitation of raster graphics is that clarity is dependent on resolution. The resolution of a raster graphic is the number of dots, pixels, or lines per inch of graphic. The higher the resolution, the crisper the image.

**Vector graphics** are line based art. A vector image can be scaled to any size because the lines themselves have no resolution and the fills are mathematical expressions. Vector graphics have a number of advantages over raster graphics:

- easily scale to different display sizes and resolutions.
- compact.
- can be enlarged without loss in quality.
- can be edited more easily since you can resize or alter the components that make up the image (extracting features like this from raster images is difficult).
- provide efficient color support for geometrical shapes.
- support advanced interactive content.
- support metadata and text search.

**PNG File**
The PNG format (.PNG) is supported by all major browsers and image processing...
applications.
If you require very high resolution graphics (e.g. for magazine publications), the SVG and PDF formats are recommended.

**SVG File**
The SVG format (.SVG) is a language for describing two-dimensional vector graphics in XML. SVG 1.0 is a Web standard (a W3C Recommendation). SVG images can be edited using the latest versions of Corel Draw and Adobe Illustrator.

**PDF File**
PDF (.PDF) is a file format that was specified by Adobe Systems Inc. to be portable across many platforms. Adobe Acrobat and Adobe Illustrator are examples of applications that support PDF.

**Actions**
1. Click the plot you wish to export to make it the active window.
2. Select Export Image from the File menu, or right-click on the plot and select Export Image from the shortcut menu. The Save dialog is displayed.

![Save dialog]

3. To the right of the file list area is a group entitled Files to Export. All of the components of the plot (if there are more than one) that can be exported are listed here. You have the option to choose which components of the plot you want to export. Check the checkbox next to each of the components you want to export to an image file. By default, the main plot is selected for you.
4. Navigate to the folder where the file is to be saved.
5. Type in a File name.
6. Select a graphics file type from the Files of type drop-down list.
7. Click Save. The image is saved to the specified file(s). If you selected multiple components, each one will be exported to a separate file (using the same file name prefix).

**Related Topics:**
Exporting Data
Find

Overview
The Find function highlights the first gene (or cluster that contains the gene) which matches or contains the search string. This function applies to most plots and table views.

Actions
1. Display a dataset in a table or color matrix plot or display a plot of an experiment.
2. Click the Find toolbar icon, or press <Ctrl> F, or select Find from the Edit menu. The Find dialog is displayed.

3. Set the Find parameters.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Find what</td>
<td>Type the search string into this text box.</td>
</tr>
<tr>
<td>Match Case</td>
<td>Check this box to search in a case-sensitive manner.</td>
</tr>
<tr>
<td>Find whole words only</td>
<td>Check this box to find only whole words that match the search string.</td>
</tr>
</tbody>
</table>

4. Click Find. The Find operation is performed and the name of the first gene that matches the search string (or cluster containing the gene) is highlighted in the table or plot. The search string and the gene containing it are listed in the status bar.
   - If no gene matches the search string, a message is displayed in the status bar.

Related Topics:
Find Next
Find Previous

Find Next
Overview
The Find Next function highlights the next gene (or cluster containing the gene) which matches or contains the search string. The Find Next function is active immediately after the Find, Find Next, or Find Previous function has been used.

- This function wraps around. Searching begins at the gene after the highlighted gene and continues to the end of the list. If no match is found, searching continues from the start of the list.

Actions
1. Press <F3>, or select Find Next from the Edit menu. The Find Next operation is performed and the name of the next gene that matches the search string (or cluster containing the gene) is highlighted in the table or plot. The search string and the gene containing it are listed in the status bar.

Related Topics:
Find
Find Previous

Find Previous

Overview
The Find Previous function highlights the previous gene (or cluster containing the gene) which matches or contains the search string. The Find Previous function is active immediately after the Find, Find Next, or Find Previous function has been used.

- This function wraps around. Searching begins at the gene before the highlighted gene and continues to the start of the list. If no match is found, searching continues from the end of the list.

Actions
1. Press <Shift><F3>, or select Find Previous from the Edit menu. The Find Previous operation is performed and the name of the previous gene that matches the search string (or cluster containing the gene) is highlighted in the table or plot. The search string and the gene containing it are listed in the status bar.

Related Topics:
Find
Find Next
Profile Matching

Overview
The Profile Matching function is used to reorder the display in a Color Matrix, Matrix Tree, or Two Way Matrix Tree plot based on the profile of one or more selected genes. Profile Matching can be applied to complete datasets only. If you have an incomplete dataset, you could apply missing value estimation or a filtering operation to create a complete dataset from your original one. Use the new complete dataset for profile matching operations.

Actions
1. Display a Color Matrix Plot of a complete dataset, or a Matrix Tree Plot of a clustered dataset, or a Two Way Matrix Tree Plot of two appropriate clustered datasets.
2. Select a reference.
   • To select a single gene, click on the name of the gene on the plot. The gene name is highlighted.
   • To select multiple genes, press and hold the <Ctrl> key and click on the names of the genes on the plot. The selected genes are highlighted.
3. Click the Profile Matching toolbar icon, or select Profile Matching from the Tools menu, or right-click on the plot and select Profile Matching from the shortcut menu. The Profile Matching dialog is displayed.

4. Set the Distance Metric for the profile matching calculations.
   Note: If you try to perform profile matching using less than the necessary number items, a message is displayed, then the dialog is displayed again so you can select more items.
   • Pearson Correlation or Pearson Squared, at least two items must be checked.
   • Spearman, at least three items must be checked.
   • All others, at least one item must be checked.
5. Under the Include heading, a sample with a checkmark is included in the profile matching calculations. The default is all samples included.
   • Click an included sample to exclude it.
6. For a single gene profile match, the values listed under the Profile heading are the actual values for those samples. For a multiple gene profile matching, the values listed under the Profile heading are the average value for those samples for the selected genes. These are the values used in the profile matching calculations. Double-click on a value to edit it. The value you enter is used in place of the original value in the profile matching calculations.

7. Click OK. The Experiment Progress dialog is displayed. It is dynamically updated as the Profile Matching operation is performed. To cancel the Profile Matching operation, click the Cancel button.

The genes in the plot are rearranged with the genes sorted from the best match at the left to the worst match at the right. **Note:** on a Matrix Tree or Two Way Matrix Tree plot, the tree portion is no longer displayed.

**Saving a Profile**

a) To save a profile, right-click on the plot and select Save Profile from the shortcut menu or close the plot, and then click Yes on the Save Profile dialog. The Profile Matching item is added to the Experiments navigator pane under the original dataset.

**Note** that if you exit the application without saving a profile, you will be prompted to do so.

**Related Topics:**
- Creating Color Matrix Plots
- Creating Matrix Tree Plots
- Creating a Two Way Matrix Tree Plot

**Matrix Tree Plot Node Selection**

**Overview**

The node selection feature gives you a quick way to select all the genes (or samples) in one or more nodes on a Matrix Tree Plot. The selected genes or samples can then be displayed in a plot, or used to create a gene list (genes only), or apply Profile Matching to the Matrix Tree Plot (genes only).

**Actions**

1. Display a matrix tree plot of a Hierarchical Clustering experiment.
2. Move the mouse pointer over the dendrogram portion of the plot (below the color tiles for gene clustering; to the right of the color tiles for sample clustering). A rectangle outlines the genes belonging to the current node.

3. Click the mouse while the rectangle outlines the genes (or samples) you are interested in. The genes (or samples) in the node are highlighted.

- **Selecting multiple nodes**: press and hold the <Ctrl> key and click on each node.
- **Selecting a series of nodes**: press and hold the <Shift> key and click on the first and last node in the series.

4. Use the selected genes to:

Create a Gene List
- Click the Create Gene List toolbar icon 🌐.
Display a Plot or Perform Profile Matching

• Right-click on the plot to display the shortcut menu.

<table>
<thead>
<tr>
<th>Menu Option</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scatter Plot</td>
<td>Display a Scatter Plot of the two selected genes or samples.</td>
</tr>
<tr>
<td>Coordinate Plot</td>
<td>Display a Coordinate Plot of the selected genes or samples.</td>
</tr>
<tr>
<td>Summary Statistics</td>
<td>Display a Summary Statistics chart of the selected genes or samples.</td>
</tr>
<tr>
<td>Profile Matching</td>
<td>Apply Profile Matching to the Matrix Tree Plot using the selected genes as</td>
</tr>
<tr>
<td></td>
<td>the reference.</td>
</tr>
</tbody>
</table>

Related Topics:
Creating a Matrix Tree Plot
Hierarchical Clustering

Changing the Gradient Color and Scale

Overview
At the top of the color matrix, matrix tree and two way matrix tree plots is a legend. The legend consists of a color gradient and a corresponding expression level scale. The scale shows the minimum, middle and maximum expression values mapped on the plot. Each colored tile on the plot represents the expression level of that gene (column name) for that sample (row name). The color of a tile is determined by the color gradient at that expression level.

Actions
Changing the Scale of the Gradient
1. Right-click on the plot and select Customize from the shortcut menu. The Customize dialog is displayed.

   ![Customize dialog]

   • Type a new value into the Minimum and/or the Maximum field and press <Enter> or use the scroll arrows to set the value(s).
• Click the **Use actual range** button to set the minimum and maximum for the display from the actual minimum and maximum values in the dataset. As the new values are entered or set, the plot is re-drawn using the new values giving you a chance to preview your changes.

4. Click **OK** to keep the new values, or click **Cancel** to revert to the previous ones.

**Changing the Color of the Gradient**

1. Right-click on the plot, and select **Customize** from the shortcut menu. The **Customize** dialog is displayed.

2. Click a new color scheme from the **Palette** drop-down list. The plot is re-drawn using the new values giving you a chance to preview your changes.
3. Click **OK** to keep the new color scheme, or click **Cancel** to revert to the previous color scheme.

**Note** that the color scheme is universal. All matrix tree, color matrix and two way matrix tree plots displayed will use the selected color scheme.

**Related Topics:**
- Selecting Items
- Resizing Cells in a Color Grid

**Resizing Cells in a Color Grid**
Overview
The size of the color tiles on the color matrix, matrix tree and two way matrix tree plots can be changed by using the resize function. The size of the dendrogram or partitional comb height on a matrix tree or two way matrix tree plot can be changed using the same function.

Actions
1. Right-click on a color matrix, matrix tree, or two way matrix tree plot, and select Resize from the shortcut menu. The Resize dialog is displayed.
   
   For a Color Matrix Plot:

   ![Color Matrix Plot Resize Dialog](image1)

   For a Matrix Tree Plot (hierarchical clustering / partitional clustering):

   ![Matrix Tree Plot Resize Dialog](image2)

   For a Two Way Matrix Tree Plot:

   ![Two Way Matrix Tree Plot Resize Dialog](image3)

2. Type in or use the scroll arrows to set the Cell width and/or Cell height of the color tiles.

   Note: if you choose a value for the width or height that designates less space than is required to display the row or column names, the names are not displayed.

3. For the matrix tree or two way matrix tree plots, type in or use the scroll arrows to set the Dendrogram or Partitional Comb height.

4. Click OK to display the plot using the new values, or click Cancel to revert to the previous ones.

Related Topics:
- Changing the Gradient Color and Scale
- Toggling the Color Grid On or Off
- Selecting Items
Toggling the Color Grid On or Off

Overview
Turning off the color grid makes it easier to discern cluster membership as this action will place the cluster lines adjacent to their associated labels. In the shortcut menu there is an item that toggles the color grid on and off.

Actions

Toggling the Color Grid Off
• When the color grid is visible, right-click and select *Hide Color Matrix* to turn the color grid off.

Toggling the Color Grid On
• When the color grid is not visible, right-click and select *Show Color Matrix* to turn the color grid on.
Related Topics:
Changing the Gradient Color and Scale
Resizing Cells in a Color Grid
Selecting Items

SOM Plot Functions

Customizing the SOM Plot

Overview
The appearance of the SOM plot proximity-gradient map can be customized. The color gradient used in the background to indicate areas of similarity and several other
characteristics can be changed. For complete details about the SOM plot, see Creating a SOM Plot.

**Actions**
1. Right-click on the proximity-gradient map to display a shortcut menu.
2. Select **Customize**. The **SOM Properties** dialog is displayed.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Similarity</td>
<td>The color gradient to use for the proximity-gradient map.</td>
</tr>
<tr>
<td>Show Cardinality</td>
<td>Toggle on (checked) or off (unchecked) to show and hide</td>
</tr>
<tr>
<td>Rings</td>
<td>the cardinality rings.</td>
</tr>
<tr>
<td>Ring Color</td>
<td>The color of the cardinality rings.</td>
</tr>
<tr>
<td>Show Nodes</td>
<td>Toggle on (checked) or off (unchecked) to show and hide</td>
</tr>
<tr>
<td></td>
<td>the nodes on the map.</td>
</tr>
<tr>
<td>Node Color</td>
<td>The color of the nodes on the map.</td>
</tr>
<tr>
<td>Show Proximity</td>
<td>Toggle on (checked) or off (unchecked) to show and hide</td>
</tr>
<tr>
<td>Grid</td>
<td>the proximity grid.</td>
</tr>
<tr>
<td>Strong Connection</td>
<td>The color associated with high similarity in the proximity grid.</td>
</tr>
<tr>
<td>Weak Connection</td>
<td>The color associated with low similarity in the proximity grid.</td>
</tr>
<tr>
<td>Show Profile</td>
<td>Toggle on (checked) or off (unchecked) to show/hide the profile.</td>
</tr>
</tbody>
</table>

3. Set the parameters.
4. Click **OK** to apply the changes, or click **Cancel** to keep the previous plot settings.

**Related Topics:**
- Performing a SOM Experiment
- Creating a SOM Plot
- Resizing the SOM Plot

**Resizing the SOM Plot**

**Overview**
Both the proximity-gradient map and the node/cluster profile can be resized.

**Actions**

**Zooming the Proximity-Gradient Map**
1. Select **Zoom** from the **View** menu, or right-click the proximity-gradient map and select **Zoom** from the shortcut menu. The **Resize** dialog is displayed.

![Resize dialog](image)

2. Set the **Zoom percentage**.
3. Click **OK**. The map is zoomed to the specified percentage.

**Resizing the Node/Cluster Profile**
1. Right-click on the node/cluster profile (displayed in the lower pane of the window).
2. Select **Resize** from the shortcut menu. The **Resize** dialog is displayed.

<table>
<thead>
<tr>
<th>Element</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Width (in pixels)</td>
<td>The width of the profile plot.</td>
</tr>
<tr>
<td>Height (in pixels)</td>
<td>The height of the profile plot.</td>
</tr>
<tr>
<td>Maximum</td>
<td>The maximum value of the y-axis.</td>
</tr>
<tr>
<td>Minimum</td>
<td>The minimum value of the y-axis.</td>
</tr>
<tr>
<td>Set to cluster range</td>
<td>Automatically adjust the y-scale to fit the cluster.</td>
</tr>
<tr>
<td>Set to dataset range</td>
<td>Automatically adjust the y-scale to fit the entire dataset.</td>
</tr>
</tbody>
</table>

3. Set the parameters.
4. Click **OK** to apply the changes, or click **Cancel** to keep the previous plot settings.

**Related Topics:**
- Performing a SOM Experiment
- Creating a SOM Plot
- Customizing the SOM Plot

3D Plot Functions
3D Plot Functions

Overview
This describes the various techniques available for interacting with 3D plots.

Actions
Displaying the Coordinates of a Point
Hover the mouse pointer over the point. The coordinates show in the area below the plot.

Selecting a Point
Click a point on the plot or click on an item in the legend. The selection is highlighted on the plot and in the legend.

Selecting Multiple Items
Press and hold the <Ctrl> key and click on items in the legend or points on the plot. The items are highlighted in the legend and on the plot.

Selecting a Series of Items
Press and hold the <Shift> key and click on the first and last item in the series on the legend. The items are highlighted in the legend and on the plot. (<Shift>-click has the same behavior as <Ctrl>-click on the plot).

Rotating the Plot
Click on the plot and drag. The plot rotates in the direction the mouse moves.

Zooming the Plot
Press the <Alt> key and then click and drag up or down on the plot.
- Drag up to shrink.
- Drag down to enlarge.

Panning the Plot
Right-click and drag on the plot.

Displaying the Plot Shortcut Menu
Right-click on the legend to display a shortcut menu:
Select an enabled function item.

<table>
<thead>
<tr>
<th>Element</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Select All</td>
<td>Select all items on a plot.</td>
</tr>
<tr>
<td>Select None</td>
<td>De-select all items on a plot.</td>
</tr>
<tr>
<td>Color</td>
<td>Select a color from the color context menu. The selected item is re-drawn</td>
</tr>
<tr>
<td></td>
<td>using the new color.</td>
</tr>
<tr>
<td>Export Image</td>
<td>Export an image of the plot.</td>
</tr>
</tbody>
</table>

Using Plot Buttons

- Click **Home** on the upper part of the plot to return the plot to its original state.
- Click **Normalize/Raw Data** on the upper part of the plot to switch between viewing a plot of the raw data and a plot of the data after it has been normalized.

Related Topics:

- Color By Gene Lists or Variables
- Troubleshooting

Exporting a Dataset

Exporting Data

Overview

Gene expression data can be exported to a .csv file (comma separated values). If your dataset has variable information associated with it, you are given the option to embed the variable data within the exported file.

Actions

1. Click a dataset in the **Experiments** navigator. The dataset is highlighted.
2. Select **Export Data** from the **File** menu. If the dataset has variable information, the **Export Gene Expression Values** dialog is displayed.
• Select GeneLinker Tabular to export data to a file without embedding variable data, or select GeneLinker Tabular with Variables to export data to a file with embedded variables.

3. Click Export. The Save As dialog is displayed.

4. If necessary, navigate to the folder where the file is to be saved.
5. GeneLinker™ supplies a default file name based on the name of the item in the navigator and a file type extension (.csv). You can use the default file name or you can type over it.
6. Click Save. The data is saved to the specified file.

Note on Embedded Variable Data
GeneLinker™ imports data and variable information from separate files. Some programs, such as Spotfire®'s DecisionSite™, import data and variable information from a single combined source file.

Related Topics:
   Exporting Images
   Generating Reports
   Exporting to DecisionSite

Exporting to DecisionSite

Overview
Gene expression data can be exported directly into Spotfire®'s DecisionSite™ application.
Enabling Export to DecisionSite™

You must have Spotfire®’s DecisionSite™ installed to use this feature, so install it if necessary.

The second thing you must do is edit your GeneLinker.conf file to tell GeneLinker™ where DecisionSite™ lives. This file is created in the GeneLinker™ install directory (default Program Files\MMC\GeneLinker Platinum or Gold) the first time you run GeneLinker™, so if you haven’t run GeneLinker™ since installing it, please start GeneLinker™ and then exit the program.

If GeneLinker™ is running, please exit the program. The GeneLinker.conf file must be edited while GeneLinker™ is not running.

If you edit the GeneLinker.conf file while GeneLinker™ is running, GeneLinker™ will wipe out your changes when you restart it.

The following two entries must be edited with the correct directory paths from your DecisionSite™ install. The two lines below show the default directories for each.

```
mmc.genelinker.decisionsite.workingdirectory=C:\Program Files\Spotfire\DecisionSite\Data
mmc.genelinker.decisionsite.location=C:\Program Files\Spotfire\DecisionSite\Program
```

If these preferences are not set, the Export to DecisionSite™ menu item is not visible in the GeneLinker™ File menu.

Actions

1. Click a dataset in the Experiments navigator. The item is highlighted.

2. Select Export to DecisionSite from the File menu.

3. Select whether to write each gene as a DecisionSite record, or each sample as a DecisionSite record.

4. Click OK.

- If DecisionSite™ is installed properly and the preferences have been properly set, the dataset is exported to a .csv file in the DecisionSite™ working directory using the dataset name from the Experiments navigator. The DecisionSite™ application is then launched and automatically loads the dataset which GeneLinker™ has just exported.

- If you chose to export the data with Samples as Records, and if there are variables associated with the selected experiment, then they will also be included in the exported file and will appear in DecisionSite.

Once the dataset is in DecisionSite™, it can be saved to a DecisionSite™ format file.
**Genes: Structures and Functions**

**Genes Overview**

**Overview**
A gene, in the context of GeneLinker™, consists of an identifier of a specific type, an optional short name, optional description, and an associated lookup URL.

Please note that gene identifiers have a length restriction of 25 characters. This means that on import of a dataset or a gene list, identifiers that are longer than 25 characters are truncated.

Genes are imported into your GeneLinker™ database when you import a dataset or a gene list. All of the genes in your database are listed in an alphabetical list in the Genes navigator. Genes can be annotated, looked up in an external database, or included in a gene list.

**Related Topics:**
Changing Your User Preferences

**Lookup Gene**

**Overview**
You have the option of looking up gene information in a database on the World Wide Web from the Genes or Gene Lists navigators, the table viewer, and many of the plots. The results of a lookup gene operation are displayed using the HTML browser specified in your user preferences. See Disclaimer.

**Actions**
1. On a plot or in a table view, click on one or more genes (the Find function can be used to locate a gene). Alternatively, you can click on the Genes tab in the navigator and click on one or more genes, or click on the Gene Lists tab, and click on a gene list. The items are highlighted.
2. Click the Lookup Gene toolbar icon, or select Lookup Gene from the Tools menu, or right-click a selected item and select Lookup Gene from the shortcut menu.
3. Your HTML browser is launched displaying the available information for those genes.
If you selected more than one gene, the gene names are displayed in the left frame, and the information about the selected gene is displayed in the right frame.

- The database accessed for gene information is dependent on which Gene ID the genes have. For example, if the genes you are looking up have GenBank Gene IDs, GeneLinker™ will use the GenBank URL specified in the user preferences when it launches the HTML browser.

Related Topics:
- GenBank Identifiers
- UniGene Identifiers
- Affymetrix Identifiers

Predefined Identifier Types

Affymetrix Identifiers

Overview
Affymetrix identifiers are also known as Affymetrix probe set identifiers. They are used by Affymetrix to identify the probe included on their GeneChips®. They resemble GenBank identifiers, but usually also contain a suffix or prefix. These identifiers can be used in conjunction with the NetAffx™ website to provide information and links to gene specific information. See Disclaimer.

Actions
To set the Affymetrix URL to the NetAffx website:
1. Select Preferences from the Tools menu. The User Preferences dialog is displayed.
2. Click the **Gene Database** tab. The **Gene Database** pane is displayed.

3. Set the **Lookup Gene Database URL** for Affymetrix to:

   https://www.netaffx.com/LinkServlet?probeset=MMC_ID
GenBank Identifiers

Overview
GenBank identifiers are used to index GenBank sequence entries, and thus can be used to retrieve information about a particular gene or DNA sequence from the GenBank database. This information also includes links to similar sequence entries and other public databases.

GenBank is the National Institute of Health (NIH) genetic sequence database, an annotated collection of all publicly available DNA sequences. It is maintained by the National Center for Biotechnology Information (NCBI) within the National Institute of Health (NIH). It is part of the International Nucleotide Sequence Database Collaboration, which also includes the DNA DataBank of Japan (DDBJ) and the European Molecular Biology Laboratory (EMBL).

The GenBank database and related resources can be freely accessed via the National Center for Biotechnology Information (NCBI) home page at the following URL (see Disclaimer):


UniGene Identifiers

Overview
UniGene is a database of non-redundant sequence clusters where each entry represents a unique gene. UniGene identifiers contain both an organism tag as well as a unique numerical index. These identifiers can be used to query UniGene to retrieve gene specific information, which includes the chromosomal map location in addition to tissue specific expression information.

UniGene is produced and maintained by the National Center for Biotechnology Information (NCBI) within the National Institute of Health (NIH).
Gene Lists: Structures and Functions

Gene Lists Overview

Overview
A Gene List is a set of gene identifiers that has a name and optionally a description. Gene lists can be created within GeneLinker™ from one of its many plot or experiments, or gene lists can be imported. Importing a gene list imports any genes that are not already in the database. Importing a gene list can also be used to add descriptive information to genes that already exist in the GeneLinker™ database.

Gene lists can be used to reduce the number of features (genes) in a dataset under study or to specify the features for a supervised learning experiment.

Gene Lists Navigator
All gene lists are listed alphabetically in the Gene Lists navigator.

- Click on a gene list to display information about it (name, description, creation date, etc.) in the description pane located below the navigator.
- Double-click on a gene list to expand the list of genes under the gene list name in the Gene Lists navigator.
- Click on a gene list name or genes within a gene list to lookup gene information in a database.

Related Topics:
- Modifying or Deleting a Gene List
- Gene List Filtering
- Exporting a Gene List

GeneLinker™ Gene List Native File Format

Overview
Features
- Text following a comment character ';' is ignored if the ';' is at the beginning of the line or is immediately preceded by a whitespace (a blank or a tab).
- Blank lines are ignored.
- Text enclosed in '[' and ']' on a single line marks the beginning of a list. Genes listed thereafter belong to this list
- Text between the '[' and ']' is the name, and optionally a description of the list. If
the description appears, it must follow the name separated by a '|' (pipe).

- The name of the first list in the file is optional, and if absent then the name of the first list is assumed to be that of the file being imported (minus the extension).
- Genes are listed with one gene entry per line. Each entry has 1 to 3 fields separated by commas (if commas appear in the text of the gene entry, then that text must be quoted).
- The first field is required, and is the database identifier of the gene.
- The second field is optional and is the gene name.
- The third field is optional and is a short description for the gene.

Example 1: Two gene lists in the same import file.

; Simple Gene List Example
[Simple Gene List]
Hs.178452
Hs.48876
Hs.99910
; Second Simple List in the same file
[Second Simple List]
Hs.289271
Hs.75593
Hs.91379

Example 2: Single, more complex list in a file.

; More Complex Gene List Example
[More Complex Gene List | This list adds names and some descriptions.]
Hs.178452,Gene 1
Hs.48876,Gene 2
Hs.99910,Gene 3,I particularly like this gene.
Hs.289271,Gene 4
Hs.75593,Gene 5,"This description, unlike the other, contains commas."
Hs.91379,Gene 6

Example 3: The simplest example. The name of this list is assumed to be the name of the file that contains it (minus the extension).

Hs.178452
Importing a Gene List

Overview
GeneLinker™ can import gene lists from files in two different formats. The acceptable formats are:

- A file containing a simple list of gene identifiers, or
- A file containing one or more lists of gene identifiers, a header for each list giving the list name, and optionally a short and a long name or description for each gene.

Gene identifiers may be one of the following: GenBank, Affymetrix, UniGene, or custom. Please note that gene identifiers have a length restriction of 25 characters. This means that on import of a dataset or a gene list, identifiers that are longer than 25 characters are truncated.

If you are importing a file with multiple gene lists, all gene identifiers in the file should be from the same database, e.g. all GenBank, or all UniGene - not some of each. If you want to associate both identifiers with a single gene, choose one to be the gene identifier and incorporate the other into the description. If you are using the gene list import feature to update short names or descriptions for your genes, it is best to do all the genes from a given database at once, rather than one gene list at a time. The short names and descriptions only need be updated once per gene, not once per gene list in which that gene appears.

File Formats
A file in the first format (simple list) looks like the following:

```
Hs.48876
Hs.99910
Hs.289271
Hs.75593
Hs.91379
```

A file in the second format (containing headers) looks like the following:
A gene list can be imported to bring new genes into the database, or to update the information for genes that are already in the database.

**Actions**

**Importing a Gene List File**

1. Select **Import** from the **File** menu and **Gene List** from the sub menu. The **Open** dialog is displayed.

2. Navigate to the correct folder and click on the file to be imported. The file name is highlighted.

3. Click **Open**. The **Import Gene List** dialog is displayed.

4. Select the **Gene Database** from the drop-down list. This should match the type of identifier the genes being imported have. For example, if the gene list contains genes that have GenBank identifiers, select GenBank.

5. Click **OK**. If the name of the gene list being imported is the same as an existing gene list, the **Edit Gene List Information** dialog is displayed for you to enter a new, unique gene list name and optionally a description. Click **Save**.
If the gene list being imported contains genes that are not yet in the database, they are imported.

If it contains genes that are already in the database, a conflict arises if a gene's name or description in the gene list file differs from the corresponding entry in the GeneLinker™ database. See Conflict Resolution for details on how to resolve conflicts.

6. The gene list(s) are imported and the new item(s) are added to the Gene Lists navigator.

Related Topics:
- Gene Lists Overview
- Creating a Gene List

Conflict Resolution

Overview

When importing a gene list, a conflict arises if a gene's name or description in the gene list file differs from the corresponding entry in the GeneLinker™ database. When a conflict arises, the Conflict Resolution dialog is displayed.

The dialog lists information about the gene that is in conflict:

Data File: The name of the gene list file.
**Gene Database:** The type of gene identifiers the genes in the gene list file have.

**Gene Identifier:** The identifier of the gene that is in conflict.

The mid portion of the dialog displays the gene **Name** and **Description** from both sources - the gene list file and in the database. Please note that if the **Description** is longer than 40 characters, it is displayed on the dialog in truncated form.

**Actions**

1. Read the gene information displayed on the dialog.
2. Select the gene information **Source** that is correct (the gene list file or the database) by clicking the radio button next to it.
3. You have the option to set the source to resolve any subsequent conflicts for the remainder of the current gene list import operation. If you do not check the checkbox in the **Don't ask again** group, you will have to resolve conflicts on a gene by gene basis.
4. Click **OK**.

Once all the conflicts are resolved, the gene list import completes.

**Related Topic:**

Importing a Gene List

**Creating a Gene List Within GeneLinker™**

**Overview**

A gene list can be created from a selection in a table view or plot.

**Actions**

1. Display a table view of a dataset or a plot of an experiment.
2. Select the genes to be included in the gene list.
   - **Selecting a single gene:** click on the gene name in the table or plot.
   - **Selecting multiple genes:** press and hold down the <Ctrl> key and click on the gene names.
   - In a SOM Plot, click on a plot cluster, or select one or more genes in the legend.
3. Click the **Create Gene List from Selection** toolbar icon, or select **Create Gene List from Selection** from the **Edit** menu. The **Create a Gene List** dialog is displayed.
4. Type in a unique **Name** and optional **Description** for the gene list. The gene list name *must be* unique. If it is not, a message is displayed (the **Save** button is disabled until a unique name is entered). Click **OK** and enter a unique name.

5. Click **Save**. A new item is added to the list under the **Gene Lists** tab in the navigator.
   - Click the **Gene Lists** tab to see the list of gene lists.
   - Click the **Experiments** tab to return to the **Experiments** navigator.

**Related Topics:**
- Gene Lists Overview
- Importing a Gene List

**Platinum**

**Creating a Gene List from the SLAM™ Association Viewer**

**Overview**
A gene list can be created from the SLAM™ Association Viewer.

**Actions**
1. Click on a SLAM item in the **Experiments** navigator. The item is highlighted.
2. Select **SLAM Results** from the **Predict** menu. The **SLAM™ Association Viewer** is displayed.
3. Click the checkbox to the left of the desired associations in the Associations list. Their genes are added into the Genes list box displayed to the right of the Associations list.

- As genes are added to the Genes list box, their include check boxes are checked. Only checked genes are included when you save a gene list.
- Note that only one copy of a gene name is listed in the Genes list box. The Count column in the Genes list box indicates the number of associations the gene occurs within.

4. Click the Save As button. The Create a Gene List dialog is displayed.

5. Type in a unique name and optional description for the gene list.
6. Click OK. A new item is added to list under the Gene Lists tab in the navigator.
   a) Click the Gene Lists tab to see the list of gene lists.
   b) Click the Experiments tab to return to the Experiments navigator.

Related Topics:
Gene Lists Overview
Importing a Gene List

Modifying or Deleting Gene Lists

Overview
You can rename a gene list or edit its description. Gene lists can be deleted.

Actions
Modifying a Gene List
1. Right-click a gene list in the Gene Lists navigator. The item is highlighted and the shortcut menu is displayed.
2. Select Edit Gene List from the shortcut menu. The Edit Gene List Information dialog is displayed.

3. Enter a new name for the gene list.
4. Optionally enter, edit, or delete the existing description.
5. Click OK to update the gene list information, or click Cancel to keep the original information.

Deleting a Gene List
1. Right-click a gene list in the Gene Lists navigator. The gene list is highlighted and a shortcut menu is displayed.
2. Select Delete Gene List from the shortcut menu. A confirmation dialog is displayed.
3. Click **Delete** to delete the gene list, or click **Cancel** to keep the gene list.

**Related Topic:**
Gene Lists Overview

### Exporting a Gene List

#### Overview
Gene list files can be used to share gene information between users. The two formats for exporting gene lists are: **Include GeneLinker Header**, which creates a text file containing the header information described in Importing a Gene List, or **Gene Identifiers Only** which creates a file containing a bare list of genes.

#### Actions
1. Click the **Gene Lists** tab in the navigator.
2. Right-click an item in the **Gene Lists** navigator. The item is highlighted and a shortcut menu is displayed.
3. Select **Export Gene List** from the shortcut menu. The **Export Gene List** dialog is displayed.

4. Select **Include GeneLinker Header** to export to a GeneLinker™ native file format gene list (with headers). Select **Gene Identifiers Only** to export to a gene list file without headers. The **Save As** dialog is displayed.
5. Navigate to the destination folder, type in a name for the file, and click **Save**. The gene list is exported (saved) to the file.

**Note on File Formats:**

The first format, **Include GeneLinker Header**, creates a file that looks like the following:

```plaintext
[Exported on 02-29-2002 16:36:38
[Gene List 1]
<table>
<thead>
<tr>
<th>Affymetrix Gene Name</th>
<th>Gene Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>AFFX-HSAC07_000251_3_at</td>
<td></td>
</tr>
<tr>
<td>AFFX-HSAC07_000251_1_at</td>
<td></td>
</tr>
<tr>
<td>D48974_s_at</td>
<td></td>
</tr>
<tr>
<td>B36974_at</td>
<td></td>
</tr>
<tr>
<td>U08499_at</td>
<td></td>
</tr>
<tr>
<td>M25079_s_at</td>
<td></td>
</tr>
<tr>
<td>M06802_at</td>
<td></td>
</tr>
<tr>
<td>Z70759_at</td>
<td></td>
</tr>
<tr>
<td>284721_cds2_at</td>
<td></td>
</tr>
<tr>
<td>ntn_a4_at</td>
<td></td>
</tr>
</tbody>
</table>
```

For full details on this format, please see GeneLinker™ Gene List Native File Format.

The second format, **Gene Identifiers Only**, creates a .lst file that looks like the following:

```plaintext
AFFX-HSAC07_000251_3_at
AFFX-HSAC07_000251_1_at
D48974_s_at
B36974_at
U08499_at
M25079_s_at
M06802_at
Z70759_at
284721_cds2_at
ntn_a4_at
```

**Note:** If you select multiple gene lists for simultaneous export, and choose **Gene Identifiers Only**, the resulting file contains the concatenation of all the selected gene lists.

**Related Topics:**
- Gene Lists Overview
- Importing a Gene List
- GeneLinker™ Gene List Native File Format

**Annotations and Report Generation**
Annotations Overview

Overview
An annotation is a text note that can be associated with a gene, sample, dataset or experiment. Annotations can be added, viewed, edited, output in a report, or deleted. Annotations can be used to record your intentions and discoveries at each step of an analysis run from the initial raw dataset, through preprocessing of the data, to a final clustering, classification, or other experiment. These annotations can then be output as part of a workflow report.

Annotation Components
- user identification,
- date and time (time created, last modified),
- subject heading,
- body text.

Gene Annotations
The scope of a gene is global, so the scope of a gene annotation is global. Wherever you view a gene (Genes navigator, gene list, dataset, or experiment), you can view its annotations.

Sample Annotations
The scope of a sample is local to a dataset and its descendent experiments (but not derived datasets). For example, if you annotate the first sample in a dataset and then you cluster it, the first sample in the clustered experiment has the annotation. If, however, instead of clustering, you normalized the dataset, the first sample in the normalized dataset will not have the annotation.

Dataset/Experiment Annotations
Any dataset or experiment listed in the Experiments navigator can be annotated.

Related Topics:
Annotations Editor/Viewer
Generating Reports

Annotations Viewer/Editor

Overview
The annotations viewer/editor is used to view, add, edit, or delete annotations for an item. An item can be a gene in the Genes or Gene Lists navigator, a gene or sample in a
table or plot, or a dataset or experiment listed in the **Experiments** navigator.

### Actions

1. Click an item. The item is highlighted.
2. Click the **Annotate** toolbar icon ➔, or select **Annotate** from the **Edit** menu, or right-click the item and select **Annotate** from the shortcut menu. The **Annotations for** editor dialog is displayed.

3. Click an annotation (blank to add) in the upper list box. The annotation is highlighted and the details of that annotation appear in the **Subject** and text boxes in the lower part of the dialog.

   **Adding/Editing an Annotation**
   
   - **To change the subject information**, click in the **Subject** field, and then type in the new information.
   
   - **To change the text content**, click in that area, and then type in the new information.

4. Click **OK** to apply the changes, or **Cancel** to discard changes made since the editor was opened.

### Related Topic:

Annotations Overview

### Generating Reports

#### Overview

GeneLinker™ can generate two types of reports:

- **A Single experiment report** is a report for the experiment selected in the **Experiments** navigator.
For example, generating a single experiment report for a clustering experiment produces a report that includes information just about that clustering experiment.

- **A Workflow report** is a report for the workflow leading up to and including the experiment selected in the **Experiments** navigator.

For example, generating a workflow report for the same clustering experiment produces a report that includes information about the original dataset, any intermediate elimination or estimation of missing values, any normalization and/or filtering steps, and the clustering experiment.

**Information provided in the reports includes (where applicable):**

- Dimensions of the dataset,
- Experiment parameters,
- Experiment results,
- Experiment annotations,
- Sample annotations,
- List of genes,
- Gene annotations.

**Actions**

1. Click an item in the **Experiments** navigator. The item is highlighted.
2. Select **Generate Report** or **Generate Workflow Report** from the **File** menu. The **Save As** dialog is displayed.

3. Navigate to the folder where the file is to be saved.
4. GeneLinker™ provides a default file name, based on the selected item’s name, with an extension of .html. You may rename the default path and file name by typing over them.
5. Click **Save**. The report is saved as an HTML file in the specified folder. When the report generation is finished, GeneLinker™ automatically spawns your browser displaying the report. The browser is specified in your user preferences.
Gene Lookup

If the report includes a list of genes (such as the cluster membership list on a partitional clustering experiment), click on one or more gene names to look them up in an external database.

Related Topics:
- Exporting Data
- Exporting Images
- Lookup Gene

Reference

Cancelling an Operation or Experiment

Overview

An operation or experiment can be cancelled while it is running. Cancelling an operation or experiment returns the database to the state it was just before the operation/experiment was started.

Actions

1. While an operation or experiment is running, the Experiment Progress dialog is
displayed. It is dynamically updated as the operation or experiment progresses.

2. To cancel the running operation or experiment, click the **Cancel** button, or press <Esc>. A confirmation dialog is displayed.

- If you click **No**, the operation/experiment proceeds.
- If you click **Yes**, the operation/experiment is cancelled (even if it completed after you clicked **Cancel**). The **Experiment Progress** dialog is updated indicating the cancel process is in progress.

The dialog disappears once the cleanup of the database is complete.

**Related Topic:**
GeneLinker™ Functions List

**Keyboard Shortcuts**

<table>
<thead>
<tr>
<th>Shortcut</th>
<th>Function Invoked</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;Ctrl&gt;+D</td>
<td></td>
</tr>
<tr>
<td></td>
<td><code>&lt;Alt&gt;+F4</code></td>
</tr>
<tr>
<td>---</td>
<td>-----------</td>
</tr>
<tr>
<td></td>
<td></td>
</tr>
<tr>
<td>GeneLinker Gold 3.1</td>
<td>GeneLinker Platinum 2.1</td>
</tr>
<tr>
<td>---------------------</td>
<td>------------------------</td>
</tr>
<tr>
<td>F2</td>
<td>Ctrl+E</td>
</tr>
<tr>
<td></td>
<td>Shift+F3</td>
</tr>
</tbody>
</table>

Note: The table is incomplete and may require additional context to understand fully.
<table>
<thead>
<tr>
<th>Key</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt;Ctrl&gt; + &lt;Shift&gt; + T</td>
<td>Create a Table View of Reliability Measures</td>
</tr>
<tr>
<td>&lt;Ctrl&gt; + M</td>
<td>Create Item</td>
</tr>
</tbody>
</table>
<Ctrl>+B

Variable Viewer

<Ctrl>+U

<Ctrl>+B

Variable Viewer

<Ctrl>+U
<table>
<thead>
<tr>
<th>Ctrl</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>Tree Plot</td>
</tr>
<tr>
<td>3</td>
<td>3D Scope</td>
</tr>
<tr>
<td>V</td>
<td>Ctrl+L</td>
</tr>
<tr>
<td>---</td>
<td>-------</td>
</tr>
<tr>
<td>Alt+Enter</td>
<td>Look up</td>
</tr>
</tbody>
</table>
## Glossary of Terms/Acronym List

Clicking the **Index** tab in the left pane of the online help may find additional information on terms not listed below.

<table>
<thead>
<tr>
<th>A</th>
<th>Annotations</th>
<th>Comments or suggested links to additional information. Annotations are associated with items such as genes, samples, or datasets.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Annotations editor</td>
<td>The window that allows annotations to be viewed, added, modified and/or deleted.</td>
</tr>
<tr>
<td></td>
<td>ANOVA or Analysis of Variance</td>
<td>A statistical procedure to estimate the significance of differential expression between two or more groups of samples. The test involves comparing the variance of the whole sample set to the variances within the groups – hence the name. In GeneLinker the term ANOVA is used generically to describe both the F-test and the Kruskal-Wallis test. (Some statistical texts use the term ANOVA for the F-test but not for the Kruskal-Wallis test.)</td>
</tr>
<tr>
<td></td>
<td>Application</td>
<td>The GeneLinker™ software.</td>
</tr>
<tr>
<td></td>
<td>Apriori</td>
<td>An association mining algorithm.</td>
</tr>
<tr>
<td></td>
<td>Artificial Neural Network (ANN)</td>
<td>A type of classifier (learner) loosely inspired by the interconnected nature of biological neurons. There are numerous excellent texts which discuss ANNs. Two are: Christopher M. Bishop, Neural Networks for Pattern Recognition (Oxford: Clarendon/Oxford University Press, 1995), and Simon Haykin, Neural Networks: A Comprehensive Foundation (New York: MacMillan, 1994).</td>
</tr>
<tr>
<td></td>
<td>Association</td>
<td>A pattern of feature values which occurs in a dataset more often than would be expected randomly. In GeneLinker™, a set of genes and their expression levels which co-occur with a certain sample class more often than would be expected randomly.</td>
</tr>
<tr>
<td></td>
<td>Association mining</td>
<td>The process of searching a dataset for associations. The algorithm used in GeneLinker™ Platinum is SLAM™.</td>
</tr>
<tr>
<td></td>
<td>Attribute</td>
<td>A single property of the dataset.</td>
</tr>
<tr>
<td>B</td>
<td>Bubble neighborhood</td>
<td>A rectangular neighborhood around a node, where the bounds are based on the current...</td>
</tr>
</tbody>
</table>
radius. The left boundary is radius nodes to the left of the node (including the node itself). Similarly, the top, right and bottom boundaries are radius nodes up, to the right and down from the node respectively. A neighborhood with a radius of one contains only a single node.

C
Centroid Plot
Useful for visualizing the centroid or exemplar points for each of the resulting clusters of a non-hierarchical experiment.

Chebychev distance metric
The maximum distance between two points \(X=(X_1, X_2, \text{etc.})\) and \(Y=(Y_1, Y_2, \text{etc.})\) along a single dimension.

Classification
(1) A division of a set of samples into classes; a discrete categorical variable. (2) The process of assigning or predicting the class of a sample.

Classifier
A device which assigns or predicts classes based on the pattern of features shown by a sample. For example, a classifier might be trained to predict whether a gene expression pattern arises from one cancer type or another. GeneLinker™ Platinum uses a committee of neural networks as a classifier.

Clustering
Also referred to as Cluster Analysis, this is a technique for sorting cases (genes, samples, etc.) into groups, or clusters, so that the degree of association is strong between members of the same cluster and weak between members of different clusters. Data subsets of genes or samples get grouped together (clustered) based on their similarities. Clustering techniques include Agglomerative Hierarchical, K-Means, Jarvis-Patrick and SOM.

Cluster Plot
Used to display the profiles of the individual members within a cluster.

Color Matrix Plot
A color plot used to visualize a dataset of values (e.g. gene expression levels). The display consists of a tiled grid of colored squares, samples in the rows, genes (note that gene names are case-sensitive) in the columns, and a legend. It can also be used to view a results of Principal Component Analysis.

Comb
A comb is a structure used in a Matrix Tree or Two Way Matrix Tree plot of a dataset that has a flat (non-hierarchical) cluster structure.
<table>
<thead>
<tr>
<th>Committee of neural networks</th>
<th>The comb is analogous to the dendrogram which is used to show hierarchical structure. An ensemble of neural networks, each one of which is trained slightly differently, that together makes predictions.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Component classifier</td>
<td>A member of a committee of neural networks (see above). Also known as a learner.</td>
</tr>
<tr>
<td>Continuous data, continuous variable</td>
<td>A trait or variable which can assume any of a range of numerical values. For instance, gene expression data is continuous. Contrast 'discrete'.</td>
</tr>
<tr>
<td>CSV file</td>
<td>A Comma Separated Value file is a typical file type used for storing data. Each record is stored as text, a comma delimiter separates each field, and a line feed and a return character mark the end of the record.</td>
</tr>
<tr>
<td>Cy5/Cy3</td>
<td>The ratio of two fluorescent intensities (Cy5 dye and Cy3 dye) on a spotted array.</td>
</tr>
<tr>
<td>D</td>
<td></td>
</tr>
<tr>
<td>Data mining</td>
<td>Also known as Knowledge Discovery and Data mining (KDD). Data mining is an automated analysis process used for gleaning valid, previously unknown, potentially useful information from stored data.</td>
</tr>
<tr>
<td>Data point</td>
<td>A single item in a dataset. Each item has one value for each attribute (or feature) of the data space in which the dataset exists.</td>
</tr>
<tr>
<td>Delimiter</td>
<td>A separator between data values (see CSV File).</td>
</tr>
<tr>
<td>Dendrograms</td>
<td>A pictorial description of the hierarchy created through hierarchical clustering. It shows at a glance which clusters are strongly or weakly joined by indicating the distance between them when they were joined. See also Matrix Tree Plots and Partitional Clustering Plots. Contrast 'comb'.</td>
</tr>
<tr>
<td>Discrete data, discrete variable</td>
<td>A trait or variable which can only assume a small number of distinct values is said to be discrete. For instance, 'gender' is a discrete variable which can typically assume one of two values in humans. Contrast 'continuous'.</td>
</tr>
<tr>
<td>Distance metrics</td>
<td>Quantitative measurements of similarity between two data points under study.</td>
</tr>
<tr>
<td>E</td>
<td></td>
</tr>
<tr>
<td>EST</td>
<td>1. Eastern Standard Time</td>
</tr>
<tr>
<td></td>
<td>2. Expressed Sequence Tags, short segments of cDNA used to uniquely identify a gene.</td>
</tr>
</tbody>
</table>
Euclidean distance metric
The straight line distance between any two points.

Exemplar
A model attribute value derived from example of that attribute. This can be done statistically or by selecting a representative example.

Exemplar point
A data point with attribute values such that its attribute signature represents the attribute signature of the collection or data points it represents.

Experiments navigator pane
The hierarchical tree control for datasets and experiments. It is the upper left pane of the GeneLinker™ main window. The pane has three tabs (Experiments, Genes and Gene Lists). Experiments is the default.

Expression level
mRNA abundance, commonly measured by fluorescent intensities on gene chips.

Feature
In machine learning, a trait used as input to supervised or unsupervised learning experiment. In GeneLinker™, genes are features.

Feature Selection
The process of deciding which available features a classifier will use as inputs.

Filtering
Methods that allow the exclusion of some genes from further analysis.

Flat Classification Structure
A classification structure in which no cluster contains any other cluster. See also Partitional Clustering.

F-Test
A parametric ANOVA intended to estimate the significance of differential expression between two or more groups of samples. The F-test is designed for normally-distributed data and can give misleading results if applied to severely non-normal data.

GenBank
A public repository of DNA, maintained by the NCBI (Website: http://www.ncbi.nlm.nih.gov/GenBank see Disclaimer).

Gene Chip
See Microarray.

Gene expression
The relative abundance of all mRNA species in a cell or tissue as they vary with environmental or biological factors or conditions.

Gene Expression Profile
Line plot showing how gene properties vary with environmental or biological factors or conditions.
Globular Cluster

A cluster which is very roughly spherical or elliptical is referred to as globular. A more precise mathematical term is convex, which roughly means that any line you can draw between two cluster members stays inside the boundaries of the cluster. Contrast ‘non-globular cluster’ - it may have a very complicated, convoluted boundary. Members of globular clusters typically bear some resemblance to the mean of the cluster. The mean of a non-globular cluster is often irrelevant, and can even lie outside the cluster.

Green dye intensity

The sample of interest, or denominator, in a spotted array relative gene expression ratio experiment. Also described as a Cy5/Cy3, test/background experiment, where in this case it represents Cy3 or background.

Hierarchical clustering

A method of cluster analysis in which data is organized into a tree-like graph based on similarity. **Agglomerative Hierarchical Clustering** is a bottom up clustering method in which all data points start in individual clusters, and at each step of the clustering process the two closest clusters are merged until only one cluster remains. **Divisive Hierarchical Clustering** is a top-down clustering method and is essentially the reverse of agglomerative hierarchical clustering. GeneLinker™ does not support divisive hierarchical clustering.

Housekeeping genes

A housekeeping gene is a gene that is assumed to be constitutively expressed at a constant level. Common examples include beta-actin and GAPDH. Although they are assumed to be constitutive, they are often expressed at different levels and hence need to be normalized.

Hybridization array

An array where hybridization occurs between the pre-attached genetic materials (DNA, RNA etc.) and relevant complementary genetic materials (DNA, RNA etc.) under study.

Iteration (SOM)

(SOM) A single step within which the map 'learns' a single item from the input dataset.

Jarvis-Patrick clustering

A clustering method; see Overview of Jarvis-
Patrick Clustering for detailed information.

K-Means clustering
An algorithm that generates fixed-sized, flat classifications and clusters based on distance metrics for similarity. The specified K value will determine the number of clusters that are created. See Overview of K-Means Clustering for detailed information.

Kruskal-Wallis
A non-parametric ANOVA intended to estimate the significance of differential expression between two or more groups of samples. The Kruskal-Wallis test is applicable to any sort of data, whether normally-distributed or not, but is less powerful than the analogous F-test.

Linear Discriminant Analysis (LDA)
A probabilistic classification model that produces linear boundaries between samples from different classes.

Loadings Line Plot
The Loadings Line Plot is one of three closely related plots (Loadings Line Plot, Loadings Scatter Plot, and Loadings Color Matrix Plot) that displays the individual elements of the PCs in Principal Component Analysis, allowing you too see the relative influence of genes or samples on the PCs.

Loadings Scatter Plot
The component loadings are the linear combinations for each principal component, and express the correlation between the original variables and the newly formed components. This type of scatter plot is used for PCA, where the x and y axes represent user-selected principal components. This shows the correlation of the variables with the user-selected principal components.

Loadings Color Matrix Plot
The loadings of a given PC represent the relative extent to which the original variables (genes or samples, depending on the Orientation selected for the PCA) influence the PC. The Loadings Color Matrix Plot displays these loadings as a tiled grid of colored rectangles such as those typically used to view tables and clustering results.
Lowess
Locally Weighted Regression and Smoothing Scatter plots.

Manhattan distance metric
The distance between two points $X=(X_1, X_2, \text{ etc.})$ and $Y=(Y_1, Y_2, \text{ etc.})$ computed as the sum of the distances along every dimension.

Map
(SOM) A collection of interconnected nodes.

Matrix Tree Plot
A tree plot used to visualize clustering relationships for hierarchical clusterings; can also be used to represent partitional clusterings. See Dendrograms and Partitional Clustering.

Matthews correlation
Matthews correlation measures the predictive accuracy of an association for its class. If all samples in the dataset are labeled true positive, true negative, false positive or false negative, and their frequencies represented by TP, TN, FP, FN then the Matthews correlation $= (TP\times TN- FP\times FN)/\sqrt{(TP+FP)(TN+FN)(TP+FN)(FP+TN)}$.

Microarray
A group of DNA features arranged on a microchip; may be high-density (i.e. more than 2500 features per chip) or low-density (2500 features or fewer per chip). Some researchers prefer to use high density microarrays which provide more information, some of it not required; others prefer to use customized low-density microarrays that contain only the data of interest.

Microarray process
The process of moving a sample from a source plate to the microarray, hybridizing the microarray with probes, scanning the slide, and evaluation of the spots. Example: collect the mRNA sample, isolate the nucleic acid, purify the products, deposit the DNA to create a microarray, hybridize a fluorescent probe to the microarray, detect the fluorescence using a scanner, and analyze the fluorescent image.

MMC
Molecular Mining Corporation

Navigator
The upper left pane of the GeneLinker™ main window. Referred to as the Experiments, Genes or Gene Lists navigator pane, depending on which of the three tabs is selected. Experiments is the default.

Neighborhood
On a map, a node's neighborhood consists
<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neighbors in Common</td>
<td>Refers to the number of data points in the nearest neighbor list that two data points must have in common for the two data points to be clustered together. The Jarvis-Patrick clustering algorithm clusters two data points together if they are in each other's near neighbor list and have at least a minimum (specified) number of Neighbors in Common.</td>
</tr>
<tr>
<td>Neighbors to Examine</td>
<td>Refers to the minimum required number of near neighbors to examine for a particular data point. The Jarvis-Patrick clustering algorithm clusters two data points together if they are in each other's nearest neighbor list and have at least a minimum (specified) number of nearest Neighbors in Common. This value limits the number of nearest Neighbors to Examine when determining the number of Neighbors in Common.</td>
</tr>
<tr>
<td>Neural network</td>
<td>See Artificial Neural Network.</td>
</tr>
<tr>
<td>N-Fold Culling</td>
<td>A filtering method that allows genes without a large enough relative change to be ignored during analysis.</td>
</tr>
<tr>
<td>Node (SOM)</td>
<td>A single unit within a map.</td>
</tr>
<tr>
<td>Non-globular clusters</td>
<td>In contrast to globular clusters, non-globular clusters do not have well defined centers. Non-globular clusters can have a chainlike shape. Algorithms such as Jarvis-Patrick are good at finding chainlike clusters.</td>
</tr>
<tr>
<td>Normality, normally-distributed</td>
<td>Data which have a histogram with a particular bell-shape, also referred to as a Gaussian distribution, are normally-distributed. See any basic statistical text for a detailed discussion. You can examine a histogram of your data in GeneLinker using the Summary Statistics function.</td>
</tr>
<tr>
<td>Normalization</td>
<td>A family of techniques intended to ensure that all variables have equivalent status and all samples have equivalent status during analysis. This may involve adjustments to remove non-biological sources of variability, or to remove biological sources of variability which are known to be irrelevant to the scientific question at hand.</td>
</tr>
<tr>
<td>Outlier</td>
<td>An outlier refers to a data point that exists outside the main grouping of data points. Outliers can be the result of experimental error or other environmental causes.</td>
</tr>
<tr>
<td>Overtraining</td>
<td>A common problem in supervised learning in which increasing accuracy on training data results, paradoxically, in decreasing accuracy on test data.</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Partitional clustering</td>
<td>Partitional clustering shows cluster membership by drawing a set of 'comb' structures, where each 'comb' connects entries in the same cluster. These plots visualize the results of partitional clustering algorithms (e.g. K-Means, Jarvis-Patrick). See also Dendrograms and Matrix Tree Plots.</td>
</tr>
<tr>
<td>Principal Component</td>
<td>Principal Component Analysis, a method of projecting data onto a lower-dimensional subspace in a way that is optimal in a sum-squared error sense.</td>
</tr>
<tr>
<td>Pearson Correlation</td>
<td>A measurement of the linear dependencies between two variables.</td>
</tr>
<tr>
<td>Preprocessing</td>
<td>The act of arranging data so that it is in an acceptable format for optimal use in a software application.</td>
</tr>
<tr>
<td>P-Value</td>
<td>The probability that a given effect is due to random chance as opposed to a systematic influence. More precisely, the p-value is the probability of observing the data or observing the effect when a null hypothesis is true, the null hypothesis asserting that there is no systematic influence. The observed effect, for example, might be the difference between the expression of a certain gene under a treatment condition and its expression under a different condition. A p-value must fall between 1 and zero. A p-value near one implies an observed effect that can easily occur by chance (i.e., an insignificant effect). Whereas, a p-value near zero (e.g., 0.01 or smaller) implies little role for chance to account for the observed effect (i.e., a statistically significant effect due to some kind of systematic influence).</td>
</tr>
</tbody>
</table>
| Quadratic Discriminant Analysis (QDA) | }
A probabilistic classification model that produces non-linear, curved boundaries between samples from different classes.

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>R</td>
<td>(SOM) The distance, counted in nodes, over which a new cluster item's influence is felt during learning.</td>
</tr>
<tr>
<td>Radius length</td>
<td>The random seed allows you to always get identical results when you repeat any type of analysis that uses a random number generator (e.g. the initial random assignment of points in K-means clustering, or the random sampling of rows in SLAM). Since computers are deterministic, they don't really generate random numbers. They use <em>pseudo random number generators</em> to mimic random numbers. A pseudo random number generator is essentially a function that produces a sequence of numbers that appear random. The actual pseudo random number generator takes the current number in a sequence and produces the next number in the sequence. The random seed is essentially a way of specifying exactly where to start in this sequence. If you specify the same random seed, you will always get the same behaviour if you try to repeat an analysis. If you specify a different random seed, you will probably get slightly different results. You might be able to get a sense of how robust your results are if you tend to see the same results with different random seeds.</td>
</tr>
<tr>
<td>Random Seed</td>
<td>In a comma-delimited file (.csv) a record is a row of data. A record generally refers to a sample as samples are usually in the rows of a dataset.</td>
</tr>
<tr>
<td>Record</td>
<td>The sample of interest, or numerator, in a spotted array relative gene expression ratio experiment. Also described as a Cy5/Cy3, test/background experiment, where in this case it represents Cy5 or test.</td>
</tr>
<tr>
<td>Red dye intensity</td>
<td>(SOM) A sequence of feature values. The reference vector is comparable to (i.e. has the same dimensions as) items to be clustered.</td>
</tr>
<tr>
<td>Reference vector</td>
<td>The designated key variable in training a classifier or running SLAM™. Typically this will be the variable which you are trying to</td>
</tr>
<tr>
<td>Representative variable</td>
<td>The designated key variable in training a classifier or running SLAM™. Typically this will be the variable which you are trying to</td>
</tr>
</tbody>
</table>
predict, e.g. tissue type or disease class. Contrast 'feature'.

**Robust**
A classifier which makes accurate predictions on test data is said to be robust.

**Sample**
All gene expression measurements from a single hybridization or chip or microarray experiment. A single row in GeneLinker (usually).

**Scaling**
Adjusting the values across samples (gene chips) so that the slope of each sample is equivalent.

**Scatter Plot**
A summary of the data showing the relationship between two variables (represented by X and Y axes).

**Score Plot**
The component scores are the data on the principal components. They project the original individuals onto the newly formed components, and currently support 2D and 3D score plots. The Score Plot is a scatter plot used for PCA, where the axes represent user-selected principal components. The plot contains the individuals projected onto those principal components.

**Scree Plot**
A simple line or bar plot for PCA; shows the ordered percentage of variance explained by each principal component. It resembles a scree slope (where rocks have fallen down the side of a mountain).

**Session**
The time span between starting (opening) and stopping (closing, exiting) the GeneLinker™ application.

**SLAM™**
An acronym for Sub-Linear Association Mining, SLAM™ is MMC’s proprietary fast stochastic method for association mining in discrete data.

**SOM (Self Organizing Map)**
A SOM is an algorithm that forms a topologically ordered mapping from the input signal space onto a neural network. It can be thought of as a non-linear projection of the probability density function of the input signal space onto a two-dimensional map. It organizes a set of samples on a map such that their distribution indicates their relative similarities. SOMs can be used for preprocessing patterns for their recognition, or, if the neural network is a regular two-dimensional array, to project and visualize high-dimensional signal spaces on such a
Spearman Correlation is a measure that identifies certain linear and non-linear correlations between sequences. Spearman Correlation ranks the values of two sequences and finds the linear correlation of the ranks.

A spotted array is a microarray of genes (printed by a robot, usually spot cDNA) containing many features (spots), where each spot corresponds to a specific gene. Therefore, the intensity of the spots on the array indicates where more information is present for a specific gene.

Spotted array scaling is the process of taking the multiple measurements taken for each gene and reducing them to a single value less biased or more representative than the constituent measurements if taken alone. The most common case will involve measuring Cy5 and Cy3 fluorescent intensity values and calculating their ratio. The process can also include background measurements for Cy5 and Cy3, subtracting their values before calculating the ratio.

A statistic is used to rank associations (all and within a class) in terms of their relevance to the target variable (Matthews column, phenotype, potential consequent).

A status bar is the bar that appears in the lower right corner of the application used to display information to the user.

Stochastic describes any algorithm which employs random sampling and therefore may show some variation in results when run over and over again on the same data.

A sub-experiment is an experiment derived from another experiment.

Supervised analysis, supervised learning, is a method that finds patterns in high-dimensional data by initially relying upon some assumptions of particular categories or relationships in the data. Commonly used techniques include classifiers such as linear discriminants, artificial neural networks, and support vector machines. These have been successfully applied to many different kinds of data. For gene expression data, these methods are often used to assign an observed expression profile to a predetermined class.

Support in association mining is the number of
samples in a dataset in which a given association appears.

**SVM**
Support Vector Machine. Algorithm used to identify patterns in datasets.

**T**
Tab-delimited
A data file which uses the tab character (ASCII character 9) to separate entries within a row.

Tabular
A data file in the form of a regular table is described as tabular. Each line of a tabular data file has the same number of fields (or columns, or delimiters). Each row corresponds to a sample and each column to a gene, or vice versa.

**Target node**
(SOM) The node in the map that is most similar to the selected item from the input dataset.

**Target variable**
See Representative variable.

**Test data**
Data held back from a classifier until after it is trained. The classifier is then used to make predictions about the test data. The accuracy of those predictions is a fair measure of the accuracy that the classifier can be expected to make on any similar data in the future.

**Training**
A classifier must be exposed to known samples before it can be used to make predictions on unknown samples. This process of optimizing the classifier's internal parameters is called training.

**Training data**
Data used as examples to train a classifier. Training samples must have known classes associated with them. These known classes comprise the representative variable for training.

**Transformation**
A technique to achieve a different dataset by applying some user-defined functions to the original data.

**U**
Uniform/Gaussian Discriminant Analysis (UGDA)
A probabilistic classification model that treats one class as a diffuse 'background' class, and the other classes as 'hot spots', defined by elliptical boundaries.

Unsupervised analysis, Unsupervised
Unsupervised analysis finds patterns in high-
learning dimensional data without relying upon a priori assumptions of particular categories or relationships in the data. Techniques include hierarchical clustering, K-Means clustering, and Self-Organizing Maps (SOM). These have been successfully applied to a wide variety of complex data including microarrays.

<table>
<thead>
<tr>
<th>V</th>
<th>Validation data</th>
</tr>
</thead>
<tbody>
<tr>
<td>V</td>
<td>Data used to validate or control the training of a classifier.</td>
</tr>
</tbody>
</table>

**Variable**

In GeneLinker™, a set of observations associated with samples. For instance, if a pathologist determined a tumor type for each sample in a dataset those observations might comprise a variable named 'known tumor type'. Such a variable could be compared against other variables of the same type (see below), e.g. 'predicted tumor type'.

**Variable type**

Variables which comprise distinct measurements of the same phenomenon are grouped together in GeneLinker™ into variable types. An example of a variable type is 'tumor type', and two variables of that type might be 'known' and 'predicted by model #4'.

**Vector**

Mathematically, this is a sequence of numbers; biologically, this is an agent that transfers material (usually DNA).

**Visualization**

A method used to view gene expression data profiles using tables or graphs (e.g. Scatter Plots, Matrix Tree Plots, Color Matrix Plots, etc.).

**W**

**X**

**XML**

eXtensible Markup Language

**Y**

**Z**

---

**Default Experiment Naming Convention**

**Legend**

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>v</td>
<td>value, short for gene expression value</td>
</tr>
<tr>
<td>rel</td>
<td>reliability measure</td>
</tr>
</tbody>
</table>

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<table>
<thead>
<tr>
<th>p</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>#mv</td>
<td>number of missing values</td>
</tr>
<tr>
<td>max</td>
<td>maximum</td>
</tr>
<tr>
<td>NN</td>
<td>nearest neighbors</td>
</tr>
<tr>
<td>Euclid</td>
<td>Euclidean</td>
</tr>
<tr>
<td>Pearson</td>
<td>Pearson Correlation</td>
</tr>
<tr>
<td>Chebych</td>
<td>Chebychev</td>
</tr>
<tr>
<td>Eucl Sq</td>
<td>Euclidean Squared</td>
</tr>
<tr>
<td>Manhatn</td>
<td>Manhattan</td>
</tr>
<tr>
<td>Pear Sq</td>
<td>Pearson Squared</td>
</tr>
<tr>
<td>Spear</td>
<td>Spearman</td>
</tr>
<tr>
<td>avg</td>
<td>average</td>
</tr>
<tr>
<td>#</td>
<td>number of genes to keep</td>
</tr>
<tr>
<td>{}</td>
<td>enclose a gene's or gene list's name</td>
</tr>
<tr>
<td>[]</td>
<td>enclose a sample's name</td>
</tr>
<tr>
<td>()</td>
<td>enclose a list of parameters</td>
</tr>
<tr>
<td>&quot; &quot;</td>
<td>enclose a variable's name</td>
</tr>
<tr>
<td>N</td>
<td>committee size</td>
</tr>
<tr>
<td></td>
<td>separate independent parameters; use</td>
</tr>
<tr>
<td>or</td>
<td>contextual clues or a longish list</td>
</tr>
<tr>
<td>,</td>
<td>join closely related parameters</td>
</tr>
<tr>
<td>/</td>
<td>per</td>
</tr>
</tbody>
</table>

**Default Names for Experiments**

**Remove Values**

=> table

- by Expression
  - {<=, =, =>} numeric_value
    >Removed: v <= 7.6
    >Removed: v = 10.2
    >Removed: v >= 33.3

- by Reliability Measure
  - pvalue-ish thing (numerically high values are removed)
    >Removed: p > 0.65

**Estimate Missing Values**

=> table

+ min_number_of_missing_values required for gene removal
+ replace with
  - central tendency: mean or median
> Estimated: #mv < 5 | mean
> Estimated: #mv < 2 | median
- nearest neighbours: number_of_neighbours, {euclidean, pearson correlation}
> Estimated: #mv < 8 | nn=2 | euclid
> Estimated: #mv < 1 | nn=4 | pear sq

- arbitrary value: the_value
> Estimated: #mv < 5 | v=17.078

filter genes
=> table
- gene list: name_of_gene_list (keep or remove)
  > Filtered: keep {myGeneList}
  > Filtered: remove {your Favourite Gene List}
- maximum culling: number_of_genes_to_keep
  > Filtered: max #=25
- N-Fold Culling with N: minimum_n_fold_min/max_ratio
  > Filtered: n-fold with n >= 2.5
- N-Fold Culling with number of genes: number_of_genes_to_keep
  > Filtered: n-fold #=100
- range culling: number_of_genes_to_keep
  > Filtered: range #=256
- spotted array n-fold culling: induction/repression_threshold
  > Filtered: spotted array n >= 1.26

Normalize
=> table
- logarithmic
  - base {2, e, 10}
  > Norm: log2
  > Norm: ln
  > Norm: log10
- sample scaling
  - Central Tendency
    - divide by {mean, median} user-specified arbitrary_new_{mean, median}
  > Norm: Sample scaling: divide, mean=6.7
  > Norm: Sample scaling: divide, median=150
    - subtract {mean, median}
  > Norm: Sample scaling: subtract mean
> Norm: Sample scaling: subtract median
- Linear Regression
  + baseline sample: sample_name
  + control genes: \{all, gene_list\}
> Norm: LinReg: \[16-ALL B\] | \{likelyC56\}
- Lowess
  - window_width = \{0..1\}
> Norm: Lowess: window=0.25
- positive and negative control genes
  + gene list: gene_list
  + control: \{negatives, positives\}
  + value: \{mean, median\}
  + range: \{within each sample, across all samples\}
> Norm: Neg ctrls: \{u14-P inhibitors\} | median | all samples
> Norm: Pos ctrls: \{some other gene list\} | mean | each sample
- other transformations
  - divide by maximum
> Norm: Divided by max
  - scaling between 0 and 1
> Norm: Scaled min to max
  - standardize
> Norm: Standardized

**F-Test**
=> F-Test results
- grouping variable
  > F-test: "my Variable name here"

**Kruskal-Wallis Test**
=> K-W Test results
- grouping variable
  > K-W test: "my Variable name here"

**Hierarchical Clustering**
=> Hierarchical Clustering results
+ cluster orientation: \{Genes, Samples\}
+ distance metric (points): \{Chebychev, Euclidean, Euclidean Squared, Manhattan, Pearson Correlation, Pearson Squared, Spearman\}
+ dm between clusters: \{average linkage, single linkage, complete linkage\}
+ algorithm properties: {agglomerative}
  > Hier: genes | Euclid | single
  > Hier: samples | Chebych | complete
} avg, single, complete

**Partitional Clustering**

=> Partitional Clustering results
+ cluster orientation: {Genes, Samples}
+ distance metric (points): {Chebychev, Euclidean, Euclidean Squared, Manhattan, Pearson Correlation, Pearson Squared, Spearman}
+ dm between clusters: {average linkage, single linkage, complete linkage}
+ algorithm properties:
  + type: {K-Means, Jarvis-Patrick}
    K-Means:
    + number of means: number_of_clusters = {2...}
    + random seed: random_integer
    Jarvis-Patrick:
    + neighbours to examine: int_check
    + neighbours in common: int_required
  > K-means, k=4: samples | Chebych | complete
  > J-P (4, 2): samples | Manhatn | avg
} avg, single, complete

**Self-Organizing Map**

=> SOM results
- orientation: {genes, samples}
- distance metric: {Chebychev, Euclidean, Euclidean Squared, Manhattan, Pearson Correlation, Pearson Squared, Spearman}

- map dimension
  - height = {1...}
  - width = {1...}
- reference vector
  - initialization: {random sample, random value}
  - range: float_range
- Algorithm Properties
  - number of iterations:
  - radius length: rlength = {1...}
  - random seed: int_random
  > SOM: genes | 3x4 | Euc Sq
> SOM: samples | 5x4 | Spear
* widthxheight

**Principal Component Analysis**
=> PCA results
- PCA orientation: {Genes, Samples}
  > PCA: genes
  > PCA: samples

**Discretize Data**
=> Discritization results
+ operation: {Quantile Discretization, Range Discretization}
+ target: {per gene, per sample, all data}
+ number of bins: number_of_bins = {2...}
  > Discretized: 3 bins/sample | quantile
  > Discretized: 6 bins/gene | quantile
  > Discretized: 4 bins/all data | range

**SLAM**
=> SLAM results
+ representative variable: variable
+ number of iterations: number_of_iterations
+ minimum support: = minimum_support = {2...}
+ minimum Matthews Number: min_Matthews
+ random seed: random_seed
  > SLAM: "my Rep Variable #2" | 10,000 | 2 | 0.6

**Create ANN Classifier**
=> ANN classifier
+ representative variable: variable
+ committee size: committee_size
+ committee votes required: committee_votes_required
+ hidden units: hidden_units
+ Conjugate Gradient Method
  - Polak-Ribiere
  - Fletcher-Reeves
    + steps: number_of_steps
+ MSE Fractional Change: minimum_mean_squared_error_fractional_change_to_stop
+ max iterations: maximum_iterations_before_stopping
+ random seed: random_seed
  > ANN: "leukemia-Dr D" | 16-5-3 | N=10 | 0.001 | 15
* where the last g-h-c bit is
  g # of genes in training dataset
  h # of hidden units
  c # of classes in representative variable

**IBIS Classifier Search**
  => IBIS search results
+ representative variable: variable
+ classifier type: {linear, quadratic, uniform/gaussian}
  - background class: {n/a, a_class_from_variable}
+ dimension {1 gene, 2 genes}
+ minimum standard deviation: min_std_dev
+ committee size: committee_size
+ committee votes required: committee_votes_required
+ random seed: random_seed
  > IBIS search: "Awl or AML test" | LDA | 1D

  } LDA, QDA, UGDA
* IBIS search: "varName" xDA nD

**Create IBIS Classifier**
  => IBIS classifier
+ representative variable: variable
+ classifier type: {linear, quadratic, uniform/gaussian}
  - background class: {n/a, a_class_from_variable}
+ gene or genes
+ minimum standard deviation: min_std_dev
+ committee size: committee_size
+ committee votes required: committee_votes_required
+ random seed: random_seed
  > IBIS: "leukemia-Dr B" | LDA | 1D | N=10
  > IBIS: "leukemia-Dr C" | QDA | 2D | N=10
  > IBIS: "leukemia-Dr A" | UGDA, ALL | 1D | N=10

**Classify**
  => classification/variable
+ variable name
+ classifier used to produce
  > myNewVariableName

  * no change from today; the output is the variable name as specified

**Profile Matching**

=> Profile Matching results
+ Distance Metric = {Chebychev, Euclidean, Euclidean Squared, Manhattan, Pearson Correlation, Pearson Squared, Spearman}
+ starting profile: gene or average of selected genes
+ gene expression values per sample
  > Profile: {avg custom} | Spear
  > Profile: {custom} | Chebych
  > Profile: {D86874_at} | Pearson

  * today: Profile Matching: Average of Selected Genes
    Profile Matching: Artificial Profile 1
    Profile Matching: D86974_at // single gene, no changes

**Changing Your License Information**

**License Overview**

**Overview**
When you start GeneLinker™, your license is checked for validity in accordance with your license agreement before the application can run.

**License Types**

<table>
<thead>
<tr>
<th>Type</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demo</td>
<td>A demo license is a temporary, time-limited license for running GeneLinker™ on a single computer.</td>
</tr>
<tr>
<td>Licensed Client</td>
<td>A licensed client is a single license for running a single copy of GeneLinker™ on a single computer.</td>
</tr>
<tr>
<td>(Node-locked)</td>
<td></td>
</tr>
<tr>
<td>Floating Client</td>
<td>A floating client is part of a network solution for multiple users of GeneLinker™. A floating client requests a license from the license server.</td>
</tr>
<tr>
<td><strong>License Server</strong></td>
<td>A license server is part of a network solution for multiple users of GeneLinker™. The license server has a fixed number of licenses available to assign to floating clients.</td>
</tr>
</tbody>
</table>

**Floating Licenses**
Floating licenses are a network solution for multiple users of GeneLinker™. On one
network computer, GeneLinker™ runs as a license server. On all other network computers that have GeneLinker™ installed, GeneLinker™ runs as a floating client. We recommend that license servers (for floating licenses) be installed on machines that are running the Windows® NT or Windows® 2000 operating system.

When a floating client GeneLinker™ starts up, it requests a license from the license server. The floating client must receive a license back from the license server before GeneLinker™ can run. If there are more network computers that have GeneLinker™ installed than there are floating licenses supported by the license server, then the floating clients must compete for the available licenses.

- **If the license server has a license available**, it assigns it to the floating client that requests it. When the floating client receives the license from the license server, GeneLinker™ can start.

- **If the license server has no license available** (that is, they are all in use by other floating client GeneLinker™ users), the license server will deny a license to the requesting floating client. In this case, the requesting floating client GeneLinker™ will not start and the user is informed of the situation.

**Actions**

**Changing Your License Type**

If your license changes, you will have to update the license information within GeneLinker™. Please follow the instructions appropriate to the type of change you are making.

<table>
<thead>
<tr>
<th>From:</th>
<th>To:</th>
<th>Instructions:</th>
</tr>
</thead>
<tbody>
<tr>
<td>Demo</td>
<td>Licensed Client (Node-locked)</td>
<td>Updating Demo License to Licensed Client</td>
</tr>
<tr>
<td>Demo</td>
<td>License Server</td>
<td>Updating Demo License to License Server</td>
</tr>
<tr>
<td>Licensed Client (Node-locked)</td>
<td>License Server</td>
<td>Changing from Licensed Client to License Server</td>
</tr>
</tbody>
</table>

**Licensed Client: System Changes**

For GeneLinker™ Platinum, if your machine name has been changed, on startup, a dialog is displayed indicating that your license information has been updated and that you need to reboot the computer.

If you have a licensed client (node-locked) GeneLinker™ and your computer configuration changes (such as a new motherboard or hard drive), follow the instructions in Licensed Client: Configuration Change to update the GeneLinker™ license information.

To move a licensed client (node-locked) GeneLinker™ from one computer to another computer, follow the instructions in Licensed Client: Moving from One Computer to Another to update the GeneLinker™ license information on the new computer.

**License Server: System Changes**

To move a GeneLinker™ license server from one computer to another, follow the instructions in License Server: Moving from One Computer to Another.
If you have a license server GeneLinker™, and your computer configuration changes (such as a new motherboard or hard drive), follow the instructions in License Server: Configuration Change.

**Floating Client: Server Change**
To update floating clients after a license server move, follow the instructions in Updating Floating Client after Server Move.

**Demonstration Client: Time Extension**
If you need a bit more time running the GeneLinker™ demo version before purchasing, follow the instructions in Demo License Time Extension.

**Additional Information on the License Product**
For information on the licence product FLEXlm, please visit the Macrovision and Globetrotter Software website at: [http://www.globetrotter.com/flexlm/flexlm.shtml](http://www.globetrotter.com/flexlm/flexlm.shtml).

**Related Topic:**
Starting the Program

**Demo License Time Extension**

**Demo License Time Extension**

**Overview**
When your demo license expires, GeneLinker™ will no longer run. Please contact Molecular Mining Corporation (MMC) sales for purchase information.
If you need additional time using the demo version before purchasing, follow the instructions below.

**Actions**
1. Start the demo version of GeneLinker™. Since the old license has expired, the program will not run. Instead, a message is displayed.

![Message](image)

2. Click **Edit License Information**. The **License Information** dialog is displayed.
3. If you have not already received your new demo license key and expiry date, call MMC technical support. The support representative will need the following information from the License Information dialog:
   • Your machine name.
   • Your volume serial number.

   Using this information, the support representative will provide you with:
   • A new demo license key.
   • An expiry date.

4. On the License Information dialog, ensure Demonstration Client is selected in the Installation Type list.

5. Enter the new Expiry Date (Year, Month, Day - mixed case permitted).

6. Enter the new 12-digit demo License Key. Please note that the license key is case sensitive. Be sure that all letters are typed in upper case.

7. Click Save. The dialog closes and the update license information operation is performed. A message is displayed.

8. Click OK.

9. Re-boot the computer. This step is necessary to activate the new license information.

Related Topics:
   License Overview
   Starting the Program
   Contacting Molecular Mining Corporation

License Changes
Changing from Licensed Client to License Server

Overview
Use this procedure to convert GeneLinker™ from a licensed client (node-locked) to a floating license server.

Actions
1. Start GeneLinker™ on your computer.
2. Select License Information from the Tools menu. The License Information dialog is displayed.

3. If you have not already received your new extended license key, expiry date, and number of floating licenses to support, call Molecular Mining Corporation (MMC) technical support. The support representative will need the following information:
   - Your machine name (on the License Information dialog)
   - Your computer MAC address. If your computer has the Windows® operating system, this information can be found by typing ipconfig/all at a command prompt. The MAC address is listed as the Physical Address. For other operating systems, the support representative will direct you on how to find this information and if necessary, on how to manually create the license file.

Using this information, the support representative will provide you with:
   - A new extended license key.
   - An expiry date.
   - The number of floating licenses to support.
4. Select License Server from the Installation Type list. The License Information dialog is updated.
5. Enter the new **Expiry Date** (Year, Month, Day - mixed case permitted).
6. Enter the new 24-digit **License Key. Please note that the license keys are case sensitive. Be sure that all letters are typed in upper case.**
7. Enter the **Number of Licenses** (floating) the license server is to support.
8. Click **Save**. The dialog closes and the update license information operation is performed. A message is displayed.
9. Click **OK**.
10. Re-boot the computer. This step is necessary to activate the new license information.

**Related Topics:**
- License Overview
- Starting the Program
- Contacting Molecular Mining Corporation

**Updating Demo License to Licensed Client**

**Overview**
This procedure is used to change the license information when installing a Licensed Client GeneLinker™, or this procedure is used to convert GeneLinker™ from a Demonstration Client to a Licensed Client.

**Actions**
1. Start GeneLinker™ on your computer.
   If your demo license has expired, the program will not run. Instead, a message is displayed.

   • Click **Edit License Information**.
   If your demo license is still valid or if you are installing a Licensed Client, after the program has started, select **License Information** from the **Tools** menu.

   The **License Information** dialog is displayed.

2. If you have not already received your new license key and expiry date, call Molecular Mining Corporation (MMC) technical support. The support representative will need the following information from the **License Information** dialog:
   - Your machine name.
   - Your volume serial number.

   Using this information, the support representative will provide you with:
   - A new license key.
   - An expiry date.

3. Select **Licensed Client** from the **Installation Type** list. The **License Information** dialog is updated.
4. Enter the new Expiry Date (Year, Month, Day - mixed case permitted).
5. Enter the new 12-digit License Key. Please note that the license key is case
   sensitive. Be sure that all letters are typed in upper case.
6. Click Save. The dialog closes and the update license information operation is
   performed. A message is displayed.

7. Click OK.
8. Re-boot the computer. This step is necessary to activate the new license information.

Related Topics:
   License Overview
   Starting the Program
   Contacting Molecular Mining Corporation

Updating Demo License to License Server

Overview
This procedure is used to change the license information when installing a floating
License Server GeneLinker™, or this procedure is used to convert GeneLinker™ from a
Demonstration Client to a floating License Server.

Actions
1. Start GeneLinker™ on your computer.
   If your demo license has expired, the program will not run. Instead, a message is
   displayed.
• Click **Edit License Information**.
If your demo license is still valid or if you are installing a floating License Server, after the program has started, select **License Information** from the **Tools** menu.

The **License Information** dialog is displayed.

2. If you have not already received your new extended license key, expiry date, and number of floating licenses to support, call Molecular Mining Corporation (MMC) technical support. The support representative will need the following information from the **License Information** dialog:

- Your machine name.
- Your computer MAC address. If your computer has the Windows® operating system, this information can be found by typing `ipconfig/all` at a command prompt. The MAC address is listed as the Physical Address. For other operating systems, the support representative will direct you on how to find this information and if necessary, on how to manually create the license file.

Using this information, the support representative will provide you with:

- A new extended license key.
- An expiry date.
- The number of floating licenses to support.

3. Select **License Server** from the **Installation Type** list. The **License Information** dialog is updated.
4. Enter the new **Expiry Date** (Year, Month, Day - mixed case permitted).
5. Enter the new 24-digit **License Key**. *Please note that the license keys are case sensitive. Be sure that all letters are typed in upper case.*
6. Enter the **Number of Licenses** (floating) the license server is to support.
7. Click **Save**. The dialog closes and the update license information operation is performed. A message is displayed.

8. Click **OK**.
9. Re-boot the computer. This step is necessary to activate the new license information.

**Related Topics:**
- License Overview
- Starting the Program
- Contacting Molecular Mining Corporation

**Computer or Network Changes**

**Licensed Client: Configuration Change**

**Overview**
Use this procedure to update the GeneLinker™ license information after a configuration change (such as a new motherboard or hard drive) on your computer.

**Actions**
1. Start GeneLinker™. Since the license information is no longer correct, the application
will not run. Instead, a message is displayed.

2. Click **Edit License Information**. The License Information dialog is displayed.

3. If you have not already received your new license key and expiry date, call Molecular Mining Corporation (MMC) technical support. The support representative will need the following information from the License Information dialog:
   - Your machine name.
   - Your volume serial number.
   Using this information, the support representative will provide you with:
     - A new license key.
     - An expiry date.
4. On the License Information dialog, ensure **Licensed Client** is selected in the Installation Type list.
5. Enter the new Expiry Date (Year, Month, Day - mixed case permitted).
6. Enter the new 12-digit License Key. **Please note that the license key is case sensitive. Be sure that all letters are typed in upper case.**
7. Click **Save**. The dialog closes and the update license information operation is performed. A message is displayed.

8. Click **OK**.
9. Re-boot the computer. This step is necessary to activate the new license information.

Related Topics:
License Overview
Starting the Program
Contacting Molecular Mining Corporation

Licensed Client: Moving From One Computer to Another

Overview
Use this procedure to move a licensed client GeneLinker™ from one computer to another.

Repository
To preserve your data, you will have to move your repository from the old computer to the new one. The repository is located in the Repository folder under the GeneLinker™ main directory (the default main directory is MMC in Program Files).

Actions
1. If desired, copy your repository from the old computer to a temporary location on the new computer or to a disk or CD-ROM.
2. Uninstall GeneLinker™ from the old computer.
4. Start GeneLinker™. Since the license information is not valid, the program will not start. A message is displayed.

5. Click Edit License Information. The License Information dialog is displayed.
6. If you have not already received your new license key and expiry date, call Molecular Mining Corporation (MMC) technical support. The support representative will need the following information from the License Information dialog:

- Your machine name.
- Your volume serial number.

Using this information, the support representative will provide you with:

- A new license key.
- An expiry date.

7. On the License Information dialog, ensure Licensed Client is selected in the Installation Type list.

8. Type in the new Expiry Date (Year, Month, Day - mixed case permitted).

9. Type in the new License Key. Please note that the license key is case sensitive. Be sure that all letters are typed in upper case.

10. Click Save. The dialog closes and the update license operation is performed. A message is displayed.

11. Click OK.

12. If you saved a copy of your repository, copy the files to the Repository folder under the GeneLinker™ main directory overwriting the files that were installed. Note: if you copy the Repository folder (instead of its files), be sure that you do not end up with a Repository folder inside the GeneLinker™ Repository folder.

13. Re-boot the computer. This step is necessary to activate the new license information.

Related Topics:
- License Overview
- Starting the Program
License Server: Configuration Change

Overview
Use this procedure to update the GeneLinker™ license information after a configuration change (such as a new motherboard or hard drive) on the license server computer.

Actions
1. Start GeneLinker™. Since the license information is no longer correct, the application will not run. Instead, a message is displayed.

   ![GeneLinker™ error message]

2. Click **Edit License Information**. The **License Information** dialog is displayed.

   ![License Information dialog]

3. If you have not already received your new extended license key, expiry date, and number of floating licenses to support, call Molecular Mining Corporation (MMC) technical support. The support representative will need the following information from the **License Information** dialog:
   - Your machine name.
   - Your computer MAC address. If your computer has the Windows® operating system, this information can be found by typing `ipconfig/all` at a command prompt. The MAC address is listed as the Physical Address.
For other operating systems, the support representative will direct you on how to find this information and if necessary, on how to manually create the license file.

Using this information, the support representative will provide you with:

- A new extended license key.
- An expiry date.
- The number of floating licenses to support.

4. On the License Information dialog, ensure License Server is selected in the Installation Type list.
5. Enter the new Expiry Date (Year, Month, Day - mixed case permitted).
6. Enter the new 24-digit License Key. Please note that the license keys are case sensitive. Be sure that all letters are typed in upper case.
7. Enter the number of floating licenses to support.
8. Click Save. The dialog closes and the update license information operation is performed. A message is displayed.

9. Click OK.
10. Re-boot the computer. This step is necessary to activate the new license information.

Related Topics:

License Overview
Starting the Program
Contacting Molecular Mining Corporation

License Server: Moving from One Computer to Another

Overview
Use this procedure to move the GeneLinker™ license server from one computer to another.

Repository
To preserve your data, you will have to move your repository from the old computer to the new one. The repository is located in the Repository folder under the GeneLinker™ main directory (the default main directory name is MMC).

Actions
1. If desired, copy your repository from the old computer to a temporary location on the new computer or to a disk or CD-ROM.
2. Uninstall GeneLinker™ from the old computer.
3. Install GeneLinker™ to the new computer as a Floating License Server. See GeneLinker™ Installation for detailed instructions on how to install GeneLinker™.
4. Start GeneLinker™. Since the license information is not valid, the program will not start. A message is displayed.

5. Click Edit License Information. The License Information dialog is displayed.

6. If you have not already received your new extended license key, expiry date, and number of floating licenses to support, call Molecular Mining Corporation (MMC) technical support. The support representative will need the following information from the dialog:
   • Your machine name.
   • Your computer MAC address. If your computer has the Windows® operating system, this information can be found by typing `ipconfig/all` at a command prompt. The MAC address is listed as the Physical Address. For other operating systems, the support representative will direct you on how to find this information and if necessary, on how to manually create the license file.

   Using this information, the support representative will provide you with:
   • A new extended license key.
   • An expiry date.
   • The number of floating licenses to support.
7. On the License Information dialog, ensure License Server is selected in the Installation Type list.
8. Type in the new Expiry Date (Year, Month, Day - mixed case permitted).
9. Enter the new 24-digit License Key. Please note that the license keys are case sensitive. Be sure that all letters are typed in upper case.
10. Enter the number of floating licenses to support.
11. Click Save. The dialog closes and the update license information operation is performed. A message is displayed.

12. Click OK.
13. If you saved a copy of your repository, copy the files to the Repository folder under the GeneLinker™ main directory overwriting the files that were installed. Note: if you copy the Repository folder (instead of its files), be sure that you do not end up with a Repository folder inside the GeneLinker™ Repository folder.
14. Re-boot the computer. This step is necessary to activate the new license information.
15. Inform the users of the floating client computers of the new license server name so they can update their license information.

Related Topics:
- License Overview
- Starting the Program
- Contacting Molecular Mining Corporation

Updating Floating Client after Server Move

Overview
Use this procedure to update the license information for GeneLinker™ floating clients when the GeneLinker™ license server moves from one computer to another.

Required Information
You will need the following information from your system administrator:
- The new server name.

Actions
GeneLinker™ Floating Client Running When License Server Changes
1. A message is displayed indicating that GeneLinker™ has lost contact with the license
server.

• **Note**: this message can occur for other reasons, so please check with your system administrator to determine the cause of the message. See Troubleshooting for further information.

2. Select **License Information** from the **Tools** menu. The **License Information** dialog is displayed.

3. Enter the new **Server Name** (mixed case permitted).
4. Click **Save**. The dialog closes and the update license information operation is performed.
5. Exit GeneLinker™. This step is necessary to activate the new GeneLinker™ license information.
6. Restart GeneLinker™. Rebooting the computer is not necessary.

**GeneLinker™ Floating Client Not Running When License Server Changes**

1. Start the GeneLinker™ floating client. The application will not start because it does not know the name of the new license server. Instead, a message is displayed.

2. Click **Edit License Information**. The **License Information** dialog is displayed.

3. Enter the new **Server Name** (mixed case permitted).
4. Click **Save**. The dialog closes and the update license information operation is performed.
5. Start GeneLinker™.

**Related Topics:**

*GeneLinker Gold 3.1 / GeneLinker Platinum 2.1* 483
Troubleshooting/Technical Support

Troubleshooting

Overview

License Issues

- If you are running the demo version of GeneLinker™ and your temporary license expires, contact Molecular Mining Corporation (MMC) sales to purchase a license.
- If you move GeneLinker™ from one machine to another or if your license server changes, you will need to update GeneLinker™. See the Maintenance section for full details.

Floating Client Lost Contact With the License Server

It is possible for a floating client to lose contact with the license server. Some possible causes for this could be:

- The network card in the floating client computer has become unplugged.
- The license server has crashed.
- The license server has been moved to another computer. See Updating Floating Client after Server Move for instructions on how to update the floating client license information.

If the problem is resolved and contact is reestablished with the license server, the floating client GeneLinker™ will not terminate (a message is displayed).

If the problem is not resolved within ten minutes, the floating client GeneLinker™ will terminate. Please note: any running experiment will complete even if it takes more than ten minutes and all data is saved.

3D Plots are Black

The PCA color plots can appear black if the color for the monitor is set to 256 colors. Sometimes games change the color setting but forget to set it back.

To check your current color settings:

1. Click **Start**.
2. Select **Settings**.
3. Select **Control Panel**.
4. Double-click on **Display**.
5. Click the **Settings** tab.
6. If **Colors** is set to 256 Colors, change it to the highest setting appropriate for your system.
3D Plots Crashing

The most common cause for crashes when displaying 3D plots is having older video drivers. To determine what video card and driver you have, and to update to the latest driver:

1. Click **Start**.
2. Select **Settings**.
3. Select **Control Panel**.
4. Double-click the **System** icon.
5. Click the **Hardware** tab (Windows 2000).
6. Click **Device Manager**.
7. Click the plus next to **Display Adapters**. This shows the name and type of video card on your system.
8. Click on the video card entry to highlight it.
9. Click the **Properties** button, or right-click on the video card name and select **Properties**.
10. Click the **Driver** tab. The driver version number is listed.
   - Go to the video card manufacturer website (e.g. www.ati.com) to find out what the latest driver is for your video card and download it. This process transfers the new driver to your system so it can be installed.
   - Most video card manufacturer websites have a *find a driver* or *download driver* option or page. For example, on the ATI site, the option is at the left of the main page in the Customer Service column. Be sure to download the correct driver for your operating system and video card.
11. To update the driver on your system, click **Update Driver** button on the **Properties** dialog. Follow the instructions in the **Update Device Driver** wizard.
12. Re-boot your computer to activate the new video driver.
13. Display a 3D plot.

In rare instances, the above procedure will not resolve the problem. In this case, you need to turn off hardware acceleration. This solves the problem by slowing things down a bit.

*To turn off hardware (video) acceleration in Windows 95/98/ME:*

1. Click **Start**.
2. Select **Settings**.
3. Select **Control Panel**.
4. Double-click the **System** icon.
5. Click the **Performance** tab.
6. Click the **Graphics** button.
7. Move the slider for **Hardware acceleration** to the left (None).
8. Click **OK**.
9. Close all the dialogs and all programs.
10. Reboot the computer.

**To turn off hardware (video) acceleration in Windows 2000:**
1. Click **Start**.
2. Select **Settings**.
3. Select **Control Panel**.
4. Double-click the **Display** icon. The **Display Properties** dialog is displayed.

![Display Properties dialog]

5. Click the **Settings** tab.
6. Click the **Advanced** button.

![Advanced settings and Troubleshooting tab]

7. Click the **Troubleshooting** tab.
8. Move the slider for **Hardware acceleration** to the left (None).
9. Click **OK**.

10. Close all the dialogs and all programs.

11. Reboot the computer.

**Note About Power Saving**

If you intend to run long experiments, we recommend not enabling your computer's power save features.

**Related Topics:**
- List of System Messages
- Handling a System Crash or Hang

**Handling a System Crash or Hang**

**Overview**

**Program Operation Indicators**

Check the **molecule spinner** in the upper right corner of the window. While GeneLinker™ is busy performing a function (such as preparing to display a plot), this indicator is active. It may be that the experiment you are performing is complex and hence taking a long time to finish. In this situation, wait for the experiment to complete.

The **Experiment Progress** dialog reflects the progress of the running experiment. To cancel an experiment while it is running, click the **Cancel** button on the **Experiment Progress** dialog. When an experiment is cancelled, the data repository is returned to the state it was in as the experiment was started.

**Program Hang**

One indication that the application is hung is if the mouse cursor indicates that the application is busy, but it never returns from this busy state. Alternatively, the system may be hung if the mouse pointer appears normal but there is no response to input.

If the application crashes, GeneLinker™ may simply disappear, or the operating system may crash. Alternatively, the operating system may report that GeneLinker™ or Java has caused a problem and GeneLinker™ is going to be terminated.

While inconvenient, a hang or a crash may also cause data to be lost. GeneLinker™ uses a data caching mechanism as a means to recover smoothly from hangs or crashes. When GeneLinker™ is restarted, it attempts to recover as much data as possible from its cached files.

**Actions**

If GeneLinker™ appears to be hung, on Windows® NT or 2000 it may be possible to see if it is still working by checking the Windows® Task Manager, as follows:
• Right-click on an empty section of the Windows® Taskbar and select Task Manager. This launches the Task Manager applet.

• Display the programs currently running by selecting the Processes tab. GeneLinker™ appears in this list as 'java.exe' or 'javaw.exe'. The number under the CPU column header indicates the percentage of processor power that 'java.exe' or 'javaw.exe' is using.

  • If this number is zero, then GeneLinker™ is probably hung.
  • If it is not zero, then GeneLinker™ may be busy completing some task and you may wish to wait for it to complete.
  • If it stays at a high value (95+) for an inordinate length of time, GeneLinker™ may be hung. **Note:** the SLAM™ operation can take a very long time to complete its data processing. If you are running SLAM™, wait for the operation to complete.

  **Warning:** Closing GeneLinker™ by ending the process from the Task Manager may lose recent changes to the data.

  • If GeneLinker™ is hung, you can try to close the application by clicking the close icon in the top right corner of the window. Closing GeneLinker™ in this way preserves changes to the data.
  • If GeneLinker™ crashes, restart the application. If the operating system crashes, reboot the computer.

**Related Topic:**
Contact Information for Molecular Mining Corporation

**List of System Messages**

**Initialization Messages**

'Warning: GeneLinker™ has failed to initialize correctly - Perhaps there is another instance already running.'

  • One common reason for this is that you may have clicked too many times and started more than one instance of GeneLinker™. After this message is displayed GeneLinker™ exits. To fix this problem, ensure GeneLinker™ is not already running, then restart the application.

'Warning: GeneLinker™ will expire on: Expiry Date.'

'Preference file missing a mmc.genelinker.license.filename entry. GeneLinker™ cannot start.'

'Could not find license manager file. GeneLinker™ cannot start.'

'License for GeneLinker™ has expired. GeneLinker™ cannot start.'

'Couldn't get license for GeneLinker™. GeneLinker™ cannot start.'

  • Ensure the files listed as 'missing' or 'not found' are present in the license folder in the GeneLinker™ directory, or obtain a new license if required, then restart the application. Alternatively, call Technical Support.
Messages on Startup

'Thank you for evaluating GeneLinker. Its free demonstration period has expired. To purchase a license, contact sales at Molecular Mining Corporation.'

'The GeneLinker license for this computer has expired. To renew your license, please contact sales at Molecular Mining Corporation. If you have an up-to-date GeneLinker license key for this computer, click 'Edit License Information'.

'The GeneLinker license for the license server 'Your Server Name' has expired. To renew your license, please contact sales at Molecular Mining Corporation. If you have an up-to-date license key for this computer, click 'Edit License Information'.

'The GeneLinker license for this computer is invalid. To obtain a license, please contact sales at Molecular Mining Corporation. If you have a GeneLinker license key for this computer, click 'Edit License Information'.

'The GeneLinker license for the license server 'Your Server Name' is invalid. To obtain a license, please contact sales at Molecular Mining Corporation. If you have a GeneLinker license key for this computer, click 'Edit License Information'.

'The GeneLinker license server 'Your Server Name' was not found on your network. If the name or address of your GeneLinker license server has changed, click 'Edit License Information'.

'GeneLinker requires the GeneLinker License Manager Service, but it isn't currently running on this computer. Restarting the computer should restart the service. Failing that, reinstalling GeneLinker may help. If problems persist, contact technical support at Molecular Mining Corporation.'

'GeneLinker could not connect to the license server on the network computer 'Your Server Name'. If the name or address of your GeneLinker license server has changed, click 'Edit License Information'.

'The GeneLinkerPlatinum.conf file is missing an entry for the license file name. The application can not start.'

• No license file name entry in the configuration file.

'Could not find the license.dat file at the location specified within GeneLinkerPlatinum.conf. The application can not start.'

• No license file in specified location.

'Could not connect to the FlexLM license manager. The application can not start.'

• The server (lmgrd) has not been started yet, or the wrong port@host or license file is being used, or the port or host name in the license file has been changed.

'GeneLinker Platinum could not obtain license from server. All available licenses are checked out.'

• Licensed number of users already reached.

'The feature requested could not be found in the license file for GeneLinker Platinum. The application can not start.'

• The feature could not be found in the license file.
'GeneLinker Platinum's license server does not support the feature requested. The feature may have expired or the version number is not supported.'
  
  - The feature has expired (on the server), or has not yet started, or the version is greater than the highest supported version.

'GeneLinker Platinum's license server has detected invalid license keys. Please see your system administrator to obtain valid license keys.'

  - The code in the license file line does not match the other data in the license file.

**Messages after Startup**

'GeneLinker has lost communication with its licence manager service running on the network computer '<server name>'. GeneLinker is now trying to re-establish contact, but will automatically shut itself down if it fails to do so before <current time + 10 minutes>. Any experiments in progress at that time will run to completion and will be saved automatically before GeneLinker quits.'

  - Three possible reasons. Connectivity problems (physical), the server has crashed, or the license manager is not running.

'Connection has been re-established with the license manager. GeneLinker will not shut itself down.'

  - The problem that caused the lost communication with the license manager has been resolved within the time out period (10 minutes).

'There has been no connection to License Manager for the past 10 minutes. Application is being shut down.'

  - All attempts to reconnect to the license manager have failed during the last 10 minutes.

**License Messages**

'A problem was encountered while initializing the dialogue needed to update your license file. The application will exit after this dialog is closed. Please check the log files for the problem details.'

'The licensing information for GeneLinker has been updated. You must restart this computer for these changes to take affect.'

'The server name for this GeneLinker floating client has been updated. You must restart GeneLinker for this change to take affect.'

**Upgrade Messages**

'Welcome to GeneLinker! GeneLinker is upgrading your data repository to the latest format.'

'This should take less than a minute or two.'

'This may take a few minutes.'

**Data Import Messages**

'Could not open <filename> for reading.'

  - This means that the file <filename> is either not present on the system or the user
does not have permission to read it.

'Could not open <filename> for writing.'

• This means that the user does not have permission to open the file <filename>,
  which will generally be a temporary output file opened by a script.

'Could not find header in file: <filename>.'

• This means the file is corrupt or has the wrong format, and the script cannot detect
  the data header.

'Could not find data in file: <filename>.'

• This means the file is corrupt or has the wrong format, and the script could not
  detect the start of the numeric data in the file.

'Could not understand expression column: <column name>.'

• This means the script could not find a column of the given name in the file. The
  header is probably corrupt, or the file is of the wrong format.

'Could not understand confidence column: <column name>.'

• This means the script could not find a column of the given name in the file. The
  header is probably corrupt or the file is of the wrong format.

'Script did not get any input files!'

• The script has been run without any input files selected.

'Script did not get any expression output file!'

• The script was not passed a temporary filename for the preprocessed results.

'Incorrect file format.'

• The GenePix header string (ATF) was not detected in a GenePix Axon Text File.

'The name 'dataset' is already taken. Enter a unique name for this dataset.'

Variable Import Messages

'A variable named 'variable name' already exists. To create a new variable type, you
must use another name.'

'A variable named 'variable' already exists in this dataset. To import a new variable, you
must use another name.'

Navigator Messages

'Are you sure you want to delete 'your experiment' experiment? This action cannot be
undone.'

'Are you sure you want to delete 'your experiment' and all of its derived experiments?
This action cannot be undone.'

'Are you sure you want to delete these experiments? This action cannot be undone.'

'Are you sure you want to delete these experiments and all of their derived
experiments? This action cannot be undone.'

Filtering Messages

For N Fold Culling With N:
The user specified value can not be less than or equal to zero.'

For N Fold Culling With Number of Genes, Range Culling, Maximum Culling, Spotted Array N Fold Culling:
'The user specified value can not be less than or equal to zero.'
'The user specified value cannot be larger than or equal to the number of genes.'

Normalization Messages
'The Gene List just created cannot be used in this experiment. You have selected 'one gene (for example). Please see the Help topics on using gene lists in Normalization. This new Gene List will still be available for other experiments.'
'The Gene List just created cannot be used in this experiment. You have selected all the genes. Please see the Help topics on using gene lists in Normalization. This new Gene List will still be available for other experiments.'

Clustering Messages
For K Means For the Number of Means:
'The number of clusters must not be less than 2.'
'The number of clusters must not exceed the number of clusterable items: #.'

For Jarvis-Patrick for the Neighbors to Examine:
'The number of Neighbors to Examine must not be less than 2.'
'The number of Neighbors to Examine must not exceed the number of clusterable items: #.'

For the Neighbors in Common:
'The required number of Neighbors in Common must not be less than 1.'
'The required number of Neighbors in Common must not be greater than or equal to the number of Neighbors to Examine.'

- Make the required changes to the clustering parameter Gold(s) and try again. If that is unsuccessful, call Technical Support.

Message when Launching Summary Statistics
'Summary Statistics requires a selection that contains at least two data values. Change your selection and try again.'

- Select a dataset or gene/sample with more than one value to view summary statistics.

Messages when Exporting Images
'Error encoding PNG file: <filename>'
'Ran out of memory making PNG file: <filename>'
'Error writing out file: <filename>'

- If any other applications are running, close them to free up some memory. Try the export operation again. If that is unsuccessful, exit GeneLinker™, restart the application (and possibly reboot the computer), and try the export again. If that fails, call Technical Support.

**Experiment Messages**

'The experiment couldn't be completed. Check that the operation and its parameters are appropriate to the data.'

- The most common cause of this message is GeneLinker™ attempting to carry out an impossible mathematical operation, such as dividing by zero or taking the logarithm of a negative number.
- Create a table view of your data and inspect it for negative numbers, genes with zero expression or other features that might invalidate the operation you requested. Once you have determined the source of the problem, try filtering or preprocessing the data then run the operation that previously failed.

'Are you sure you want to cancel the experiment?'

**Gene Lists Messages**

'Are you sure you want to delete gene list 'Your Gene List'? This action cannot be undone.'

'Are you sure you want to delete these 'Your Gene Lists' gene lists? This action cannot be undone.'

**Create Classifier Messages**

'The number of learners must be between 2 and the number of samples in the dataset (inclusive).'

'The number of hidden units must be between 1 and four times the number of genes in the dataset (inclusive). In general, the number of hidden units should be much smaller than the number of genes.'

'The number of conjugate gradient steps must be between 2 and 2147483647 (inclusive). In general, the number of steps should be much less than 1,000.'

'The maximum number of iterations must be between 1 and 2147483647 (inclusive). In general, the maximum number of iterations should be less than 10,000.'

**SLAM Messages**

'This value must be at least zero.'

'The number of iterations must be greater than zero.'

'The range for Matthews numbers is -1 through 1, inclusive. In general, associations with Matthews numbers that are less than 0.5 or so are not of interest.'

'The minimum support measure must be between one and the number of samples (inclusive).'
Related Topics:
Handling a System Crash or Hang
Troubleshooting/Technical Support

Contact Information for Molecular Mining Corporation

Sales
To purchase a GeneLinker™ product license or for a free onsite, in-depth presentation on the GeneLinker™ application suite, please call the Molecular Mining Corporation sales team at:
1 - 877 - 454 - 8570
or send an email to:
sales@molecularmining.com

Customer Technical Support
A Help Desk representative will make every effort to get back to you within one business day.
Toll-free within North America, call
1 - 877 - 454 - 8570 Monday-Friday, 9:00am - 5:00pm EST.
International callers, call
1 - 613 - 547 - 9752 Monday-Friday, 9:00am - 5:00pm EST.
or send an email to:
support@molecularmining.com

Suggestions
We are very interested in your feedback and suggestions on our GeneLinker™ family of products. Please send an email to:
suggestions@molecularmining.com

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GeneLinker™ Tour - Importing, Viewing, and Preprocessing Data

Importing a Dataset and its Genes
The import process copies a dataset of expression values and all of its genes from your files into the GeneLinker™ database. This process consists of three major steps:
1. Select a template (such as Affymetrix MAS 5.0 or GenePix Green/Red). This template tells GeneLinker™ how to interpret the contents of your data files.
2. Select the file or folder where your data file or files are located.
3. Select how to orient your data (genes in columns is the default for GeneLinker).
Once imported, the dataset is listed in the Experiments navigator and the genes are listed in the Genes navigator.

Importing a Gene List
Genes can be imported separately from expression data by importing a gene list. This can be done to add new genes to the database, or to update the information associated with genes already in the database.

Viewing a Gene Expression Dataset
A dataset can be viewed in two different ways: the table viewer (left half of image) shows a spreadsheet-like view of the values in the dataset, and the color matrix plot (right half of image) shows a color grid with its cells colored along a gradient representing the data values.

Preprocessing Your Data
GeneLinker™ offers a variety of preprocessing options which can be applied one or more times to a dataset. You can then view the preprocessed data as you would raw data.

Eliminate or estimate missing values
- If your dataset contains missing (null) values, you can apply techniques for
estimating them. You can also eliminate genes that have too many missing values.

**Filtering**
- Filtering operations can be applied to your data to create a new dataset containing a reduced number of genes.

**Normalization**
- Normalization is used to minimize uninteresting sources of variation. GeneLinker™ provides multiple techniques for normalizing your data.

**Remove values**
- Data values can be eliminated from a dataset by value or by reliability measure.
GeneLinker(TM) Tour - Statistical Functions

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