

CAINTEGRATOR2 v.1.1

User's Guide



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USING THE caINTEGRATOR2 v.1.1 USER'S GUIDE

This chapter introduces you to the *calIntegrator2 v.1.1 User's Guide* and suggests ways you can maximize its use.

Topics in this chapter include:

- [Introduction to the calIntegrator2 User's Guide](#) on this page
- [Organization of this Guide](#) on this page
- [User's Guide Text Conventions](#) on page 2

Introduction to the caIntegrator2 User's Guide

The *calIntegrator2 v.1.1 User's Guide* is the companion documentation to the calIntegrator2 software application. The user's guide includes information and instructions for the end user about using calIntegrator2.

Organization of this Guide

The *calIntegrator2 v.1.1 User's Guide* contains the following chapters and appendices:

Using the calIntegrator2 User's Guide — This chapter introduces you to the *calIntegrator2 v.1.1 User's Guide* and suggests ways you can maximize its use.

Chapter 1 Getting Started in calIntegrator2 — This chapter introduces general calIntegrator2 procedures and how to obtain help to use calIntegrator2.

Chapter 2 Creating a Study — This chapter describes the processes for creating and managing studies in calIntegrator2.

Chapter 3 Searching a calIntegrator2 Study — This chapter describes the processes for searching studies within calIntegrator2 using the search and browse tools.

Chapter 4 Viewing Search Results — This chapter describes search results that calIntegrator2 returns after queries.

Chapter 5 Analyzing Studies — This chapter describes how to use calIntegrator2 tools to analyze data in clinical or genomic studies that have been deployed in calIntegrator2.

Chapter 6 Administering User Accounts —This chapter describes the process for creating and managing user accounts in calIntegrator2.

Appendix A Data Import Configurations — This appendix describes how MAGE-TAB documents are parsed, validated and imported into calIntegrator2. It also provides examples of the types of MAGE-TAB documents that are expected by calIntegrator2.

Index—This section of the guide provides a complete index.

User's Guide Text Conventions

Table 2.1 illustrates how text conventions are represented in this guide. The various typefaces differentiate between regular text and menu commands, keyboard keys, toolbar buttons, dialog box options and text that you type.


Convention	Description	Example
Bold & Capitalized Command Capitalized command > Capitalized command	Indicates a Menu command Indicates Sequential Menu commands	Admin > Refresh
TEXT IN SMALL CAPS	Keyboard key that you press	Press ENTER
TEXT IN SMALL CAPS + TEXT IN SMALL CAPS	Keyboard keys that you press simultaneously	Press SHIFT + CTRL and then release both.
Monospace type	Used for filenames, directory names, commands, file listings, and anything that would appear in a Java program, such as methods, variables, and classes.	URL_definition ::= url_string
Icon	A toolbar button that you click	Click the Paste button () to paste the copied text.
Boldface type	Options that you select in dialog boxes or drop-down menus. Buttons or icons that you click.	In the Open dialog box, select the file and click the Open button.
<i>Italics</i>	Used to reference other documents, sections, figures, and tables.	<i>caCORE Software Development Kit 1.0 Programmer's Guide</i>
<i>Italic boldface monospace type</i>	Text that you type	In the New Subset text box, enter <i>Proprietary Proteins.</i>
Note:	Highlights a concept of particular interest	Note: This concept is used throughout the installation manual.

Table 2.1 calIntegrator2 User's Guide Text Conventions

Convention	Description	Example
Warning!	Highlights information of which you should be particularly aware.	Warning! Deleting an object will permanently delete it from the database.
{ }	Curly brackets are used for replaceable items.	Replace {root directory} with its proper value, such as c:\cabio

Table 2.1 caIntegrator2 User's Guide Text Conventions (Continued)

CHAPTER

1

GETTING STARTED WITH CAINTEGRATOR2

This chapter introduces general calIntegrator2 procedures and how to obtain help to use calIntegrator2.

Topics in this chapter include:

- [About calIntegrator2](#) on this page
- [Registering as a New calIntegrator2 User](#) on page 6
- [Welcome to calIntegrator2 Workspace](#) on page 8
- [Using Online Help](#) on page 12
- [Logging Out](#) on page 12
- [Application Support](#) on page 13

About calIntegrator2

NCI, Center for Biomedical informatics and Information Technology (CBIIT) is developing a novel translational informatics platform called calIntegrator2 that allows researchers and bioinformaticians to access and analyze subject annotation and experimental data across multiple subject annotation trials and studies. The calIntegrator2 framework provides a mechanism for integrating and aggregating biomedical research data and provides access to a variety of data types (e.g. Immunohistochemistry (IHC), microarray-based gene expression, SNPs, subject annotation trials data, etc.) in a cohesive fashion.

calIntegrator2 is a web based or locally installed portal that allows researchers and study managers to access the biomedical informatics infrastructure and data analysis tools established by calIntegrator2 from one common software platform. As a calIntegrator2 user, you can perform the following tasks:

- Integrate subject annotation data with genomic and/or imaging data
- Import data of various types in a predefined flat format, and create new studies with multiple study data
- Update an existing study to add new attributes or to add/modify data
- Perform analyses on study data

Registering as a New caIntegrator2 User

To request a caIntegrator2 user account, you must register as a new user, completing the following steps:

1. Go to the CBIIT caIntegrator2 login page <http://caintegrator2.nci.nih.gov> or use the URL provided by your System Administrator for the caIntegrator2 instance at your institution.
2. Click the **Register Now** hypertext link, under the caIntegrator2 login section in the upper left of the page. This opens the account registration form (*Figure 1.1*).

Register

Figure 1.1 New user account registration form

3. In the Register form, enter the appropriate information¹.
 - **Security Information**
 - **Do you have an LDAP account** [a user profile with your institution] at [NCICB or your institution]?

¹ Items with an asterisk or highlight are required.

If **Yes**, enter your username and case-sensitive password for the purposes of verifying that it is correct. After you submit your request, you can continue to use calIntegrator2 without an account to browse and search available experiments and download data while your account is verified and activated.

–**Username***

–**LDAP Password***

–**Requested role(s)*** – Select one or more of the roles. Roles are described in [Table 1.1](#).

If your LDAP profile is not validated, calIntegrator2 indicates that the LDAP credentials do not check out. You are asked to reenter them, but you can choose to answer no, and the System Administrator will manually ensure you don't get a duplicate LDAP account during provisioning. You can **Cancel** or talk with your System Administrator about the problem.

If you select **No** [you do not have an LDAP account], the text boxes for entering the LDAP account information disappear. You must indicate the role you would like to be assigned in calIntegrator2, and continue entering the appropriate information in the **Account Details** section.

Role	Description	Permissible 1.0 Actions
Study Manager	Creates, owns and manages studies	Create studies Assign annotations to studies Edit studies Search studies Perform analyses on study data
Study Investigator	Investigates and queries the study data	Query study data Save queries Analyze using K-M Plot Analyze using Gene Expression Plots Analyze using GenePattern

Table 1.1 calIntegrator2 role descriptions

◦ **Account Details**

— **First Name***

— **Last Name***

— **Email [address]***

— **Organization***

— **Address [Lines 1* and 2]**

— **City***

- **State***
- **Country***
- **Postal [or Zip] Code***
- **Phone***
- **Fax**

4. Click **Submit Registration Request** to execute the request, or click **Cancel** to abort the registration.

After registration is sent, the screen displays a confirmation message.

At this point, an email containing all of the information you specified in the new user request form is sent to the caIntegrator2 system administrator and an account request confirmation email is also sent to you, the prospective user, at your specified email address. In response, the caIntegrator2 system administrator uses UPT to create your user account and assign the requested roles (in predefined groups like Study Investigator). When your account is created, the system administrator sends you an email to alert you, after which you can login to caIntegrator2.

When your account is registered, the user ID and password you are assigned determine your access rights for the software.

Welcome to caIntegrator2 Workspace

The caIntegrator2 Welcome workspace enables quick access to all caIntegrator2 functions and information before you login. The Welcome page also displays after you log in, before you open any studies (*Figure 1.2*).

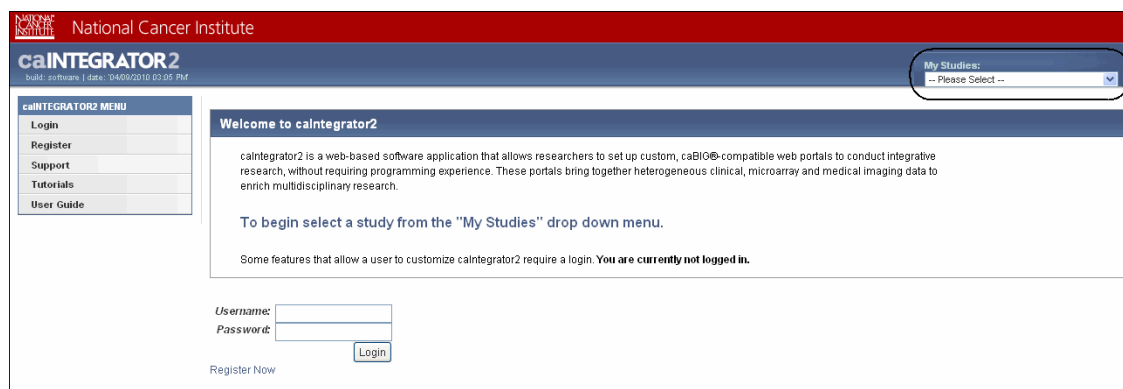


Figure 1.2 Welcome page that displays before and after login

Without logging in, you can browse any public studies. To do so, select from the drop-down list of public studies in the upper right-hand corner of your browser (*Figure 1.2*).

To log into caIntegrator2, follow these steps:

1. On the login page, enter your **username** and **password**.
2. Click the **Login** button. If your login is successful, the Welcome to Browse/ Study page appears

To access caIntegrator2 functions, use the options listed on the left sidebar of the workspace.

caIntegrator2 Functions

When you log into caIntegrator2, before any studies have been created the workspace opens with a Welcome page, as shown in (Figure 1.2). Once a study is created, its name is listed at the top of the left sidebar.

Table 1.2 describes each caIntegrator2 option in the workspace (Figure 1.2).

Sidebar Option	Function
[Study Name]	When you log in, one study displays in the left sidebar by default. Any study that you select in the My Studies drop-down list in the upper right of the page replaces this default selection.
Home	Click this to return to the home page for the selected study.
Search [Study Name]	Click this option to open the Search [Study Name] page from which you can launch queries into your selected study. For more information, see Searching a caIntegrator2 Study .
Study Data	<p>Click Saved Queries > My Queries to open the list of previous queries you saved. Click any item in the list to open the saved query, which displays on the Criteria, Columns and Sorting tabs. From those tabs, you can modify criteria and/or launch the query again. For more information, see Saving a Query on page 54.</p> <p>Click Saved Lists > Global Lists or > My Lists to open gene lists that have been saved for a study. From any page in caIntegrator2 that shows such a group, you can save a such a list of genes to be used for searches or analyses. See Creating a Gene List on page 58.</p>
Analysis Tools	<p>Click any of the listed options to open a page where you can launch an analysis of the data in the selected study.</p> <ul style="list-style-type: none"> • Generate a K-M Plot. See Creating Kaplan-Meier Plots on page 70. • Generate a Gene Expression Plots. See Creating Gene Expression Plots on page 77. • Launch GenePattern Analysis. Analyzing Data with GenePattern on page 91.
Study Management	<p>Click either of the listed options to manage the selected study through editing or deleting it or by creating a new study.</p> <ul style="list-style-type: none"> • Click Manage Studies. See Managing a Study on page 41. • Click Create a New Study. See Configuring and Deploying a Study on page 16.
Application Management	Click Manage Platforms to identify, add or remove platforms that caIntegrator2 supports. For more information, see Managing Platforms on page 42.
caIntegrator2 Menu	<ul style="list-style-type: none"> • Click Support to view contact information for Application Support. • Click Tutorials to view a tutorial to help you get started using caIntegrator2. • Click User Guide to open the caIntegrator2 v.1.0 User's Guide in PDF format.

Table 1.2 caIntegrator2 tabs

In the **My Studies** drop-down list in the upper right of the page, select the study you want to use for your current session. (The list includes all studies to which you are subscribed.) As you do so, the following left sidebar contents change to reflect options relevant to your study selection:

- the logo for the selected study (if it exists)
- the name for the selected study
- the list of saved queries and/or saved lists for that study

Viewing Existing Studies

If you have not logged into caIntegrator2, you can view any public studies in your browser. After logging in, you can view existing studies for which you have been granted permission. In the upper right corner of the page, in the My Studies drop down list, select the study you want to review or work in (*Figure 1.3*).

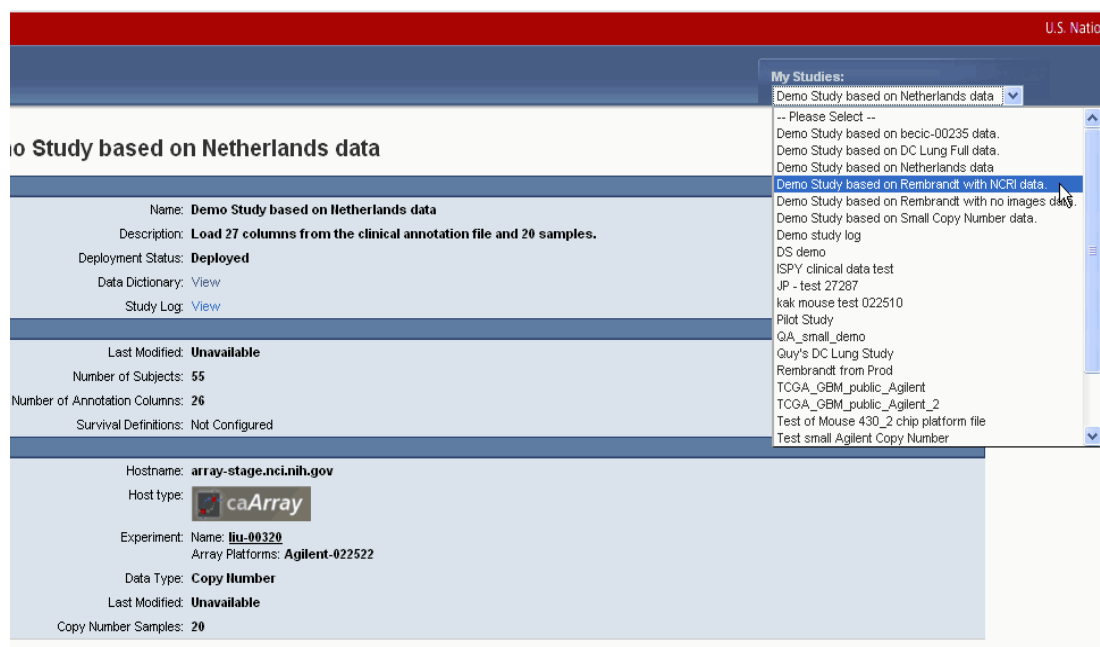


Figure 1.3 Drop-down list for selecting existing studies

The study you select opens in the browser. You can review the study data for which you have been granted permission.

After selecting the study name, in the **My Studies** drop-down list, a study summary should appear, including a status field. If the status is not deployed, or if the study summary does not appear, then the study is not deployed and available for analysis.

When the annotations are uploaded during the creation of the study each field is defined by the study manager.

- Because in looking at the study, you may not know the meaning of all the annotations, you can open a reference page with a summary of the annotations. Click the **Data Dictionary: View** link on the study home page (*Figure 1.4*).

- From the study summary page, you can also open a log for the study. Click the **Study Log: View** link on the page to see all log entries with descriptions.

Data Dictionary

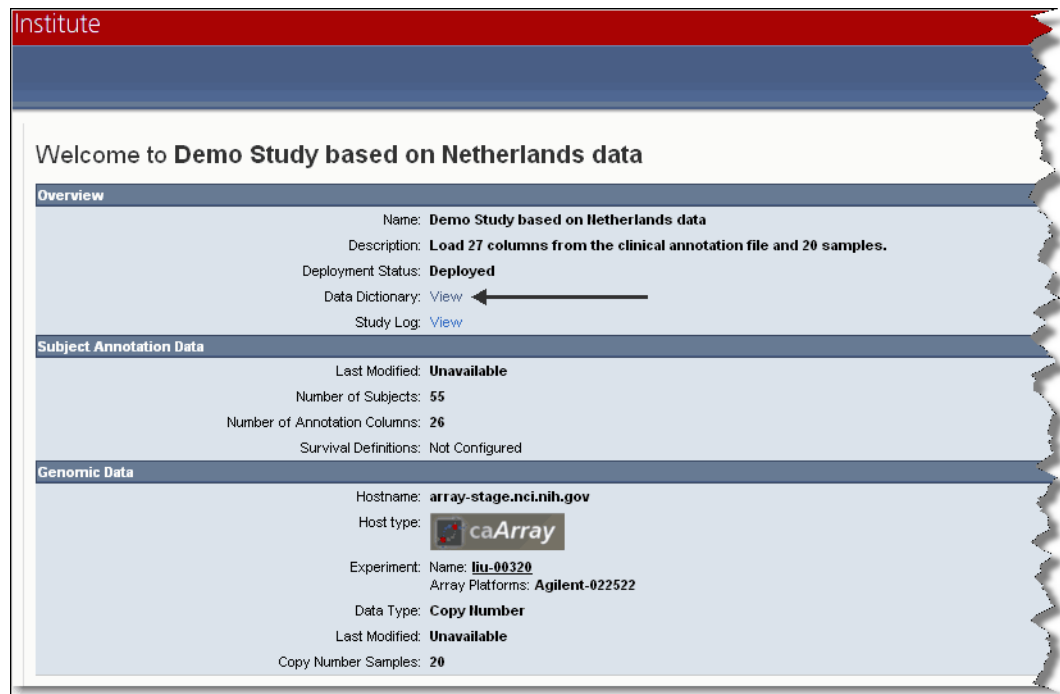


Figure 1.4 A link on a study home page opens a data dictionary summary

The Data Dictionary consists of a table that clarifies all annotations used in the study. It displays their field descriptors, descriptions, caDSR identifiers (if used), caDSR IDs and definitions, data type, and permissible settings (Figure 1.5). The **Restrictions** column indicates whether or not masks have been applied to numeric data in the study. For more information, see *Assigning An Identifier or Annotation* on page 23..

View Data Dictionary

Group Name: Annotations - Default
Description: Default annotation group

Annotation Field Descriptor	Source	Data Type	Description	caDSR ID	Permissible
Age	subject	string	Created via selenium for Rembrandt with NCRI on 12/03/09 09:09:08.		Show/Hide
Block ID	subject	string			Show/Hide
Class	subject	string			Show/Hide
Gender	subject	string	Created via selenium for Rembrandt with NCRI on 12/03/09 09:09:08.		Show/Hide
Grade	subject	string	Grade - created for Netherlands data.		Show/Hide
KRAS	subject	string			Show/Hide

Figure 1.5 Page for viewing data dictionary details

For more information about study details, see *Creating/Editing a Study* on page 17.

Study Log

The study log which you can open by clicking the **Study Log > View** link on the study summary page lists step used to create a study. For more information, see *Viewing/Editing a Log* on page 18.

Using Online Help

The online help explains how to use all of the features.

To access online help, click the help icon at the top of each page to open a context-sensitive topic. Context-sensitive help displays information that corresponds to the page from which help was opened.

When you open online help, the table of contents displays in the left panel.

Once you are in online help, several buttons and/or options help you locate topics of interest.


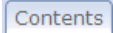
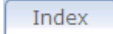
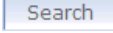
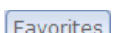



<i>Icon or Button</i>	<i>Description</i>
	Locates and highlights your current topic in the table of contents.
	Select a topic from the complete online help table of contents.
	Select a topic from the online help index.
	Perform word searches of Help by entering query text in the search text box.
	Create a list of your frequently-accessed topics.
 Related Topics 	Opens other closely related topics.
	Prints the current topic.
Topic Name > Topic Name	The breadcrumb trail shows the relative location of the current help topic relative to neighboring topics. Click a breadcrumb link to display that help topic.
Back Forward	Navigates through previously viewed topics.

Table 1.3 Online help tips

Logging Out

To log out of calIntegrator2, click the **logout** link in the upper right-hand corner of the page.

Application Support

For any general information about the application, application support or to report a bug, contact NCICB Application Support.

Email: ncicb@pop.nci.nih.gov	When submitting support requests via email, please include: <ul style="list-style-type: none">• Your contact information, including your telephone number.• The name of the application/tool you are using• The URL if it is a Web-based application• A description of the problem and steps to recreate it.• The text of any error messages you have received
Application Support URL	http://ncicb.nci.nih.gov/NCICB/support
Telephone: 301-451-4384 Toll free: 888-478-4423	Telephone support is available: Monday to Friday, 8 am – 8 pm Eastern Time, excluding government holidays.

CHAPTER 2

CREATING A NEW STUDY

This chapter describes the processes for creating and managing studies in calIntegrator2.

Topics in this chapter include:

- [*Creating a Study – Overview*](#) on this page
- [*Configuring and Deploying a Study*](#) on page 16
- [*Managing a Study*](#) on page 41

Creating a Study – Overview

You can create a calIntegrator2 study by importing subject annotation study data, genomics data and imaging data, using a combination of spreadsheet/files and existing caGrid applications as source data. Each instance of calIntegrator2 can support multiple studies. As the manager creating a study, it is important that you understand the study well and that the data you wish to aggregate has been submitted to the applications whose data can be integrated in calIntegrator2.

- **Subject Annotation** – Subject annotation data refers to pre-subject annotation, phenotypic, subject annotation, pathology or any other annotations associated with a subject. The subject annotation data should be available in CSV files, with a unique patient identifier in one column, one patient per row. Other relevant data can be supplied in other columns to be identified as annotations in the file from within calIntegrator2. You, as the study creator, must have access to the subject annotation data file, as the file does not come from a caBIG® repository.
- **Genomic** – To use calIntegrator2 to integrate array data, the data should be imported into caArray, either locally or the CBIIT installation, using that system's data file import functionality. You must also have a mapping file in CSV format. This file indicates correlations between array files and the subjects in the subject annotation data files. A mapping file consists of two columns: one with the patient ID, and one with the sample ID.

- **Imaging** – Imaging data should have been submitted to the NBIA grid node as public data, either locally or as part of the CBIIT installation. Image annotations, which includes information about images provided by radiologists or other researchers can include such information as tumor size, tumor location, etc. It must be in CSV format, with unique image series IDs in one column and annotation IDs in the second column. You must also have an image mapping file in CSV format. This file indicates correlations between subject annotation subjects or images in NBIA and subjects in the subject annotation data files. A mapping file consists of two columns: one with the patient ID, and one with the NBIA image series ID in the other column.

As you create the study, you define its structure in the process, identifying the data sources and mapping the data between different source data. After the study has been created and deployed, the study can then be used to perform analyses.

Configuring and Deploying a Study

Note: Only a user with a Study Manager role can create a study.

When you create a study, you must specify different data-types (subject annotation, array, image, etc), data sources (caGrid applications – caArray and NBIA) and map the data, (patient to sample, image series, etc.).

To create a new study, follow these steps:

1. In the Study Management section of the left sidebar, click **Create New Study**.
2. In the Create New Study dialog box that opens, provide a name and description for the study you are creating ([Figure 2.1](#)).

The screenshot shows a 'Create New Study' dialog box. The 'Study Overview' section includes a 'Study Name' text field and a 'Study Description' text area. Below these is a checkbox labeled 'Allow public to browse this study:' which is currently unchecked. Further down, the following information is displayed: 'Status: Not Deployed', 'Status Description:', 'Owner:', 'Last Modified By:', and 'Last Modified Date: Unavailable'. At the bottom right of the dialog are two buttons: 'Save' and 'Cancel'.

Figure 2.1 Create Study page

3. Click **Save**.

This opens an Edit Study page where you can add identify data files for your study.

Creating/Editing a Study

The Edit Study page displays the Name and Description that you entered for a new study, or for an existing study that you are editing (*Figure 2.2*).

Edit Study
Configure your study, and click the **Save** or **Deploy Study** button at the bottom of the page when complete.

Study Overview

Study Name: Demo Study based on Rembrandt with NCRI data

Study Description: Rembrandt with NCRI. Study created via selenium.

Allow public to browse this study: ☐

Status: Deployed

Status Description: Minutes for deployment (approx): 3

Owner: marplej

Last Modified By: manager

Last Modified Date: 03/25/2010 08:45:43

Study Log: [View Log](#) [Edit Log](#)

Study Logo: None [Browse...](#)

JPEG/GIF, 200x72 maximum [Upload Now](#)

Annotation Groups [Add New](#)

Group Name	Description	Number of Annotations	Action
Annotations - Default	Default annotation group	28	

Subject Annotation Data Sources [Add New](#) [Edit Survival Values](#)

Type	Description	Status	Last Modified	Action
DELIMITED_TEXT	rembrandt_clinical_Aug08_subset_mod_for_NCRI.csv	Loaded	Unavailable	Edit Annotations Reload

Genomic Data Sources [Add New](#)

Host Name	Experiment Identifier	File Description	Data Type	Status	Last Modified	Action
array.nci.nih.gov	jagla-00034	Mapping File: None Configured Control Sample Mapping File(s): None Configured	Expression	Loaded	Unavailable	Edit Delete

Imaging Data Sources [Add New](#)

Host Name	Collection Name	File Description	Status	Last Modified	Action
imaging.nci.nih.gov	NCRI	Annotation File: ncri_image_annotations.csv Mapping File: ncri_image_mapping.csv	Loaded	Unavailable	Edit Edit Annotations

External Links [Add New](#)

Name	Description	File Name	Number of Links
------	-------------	-----------	-----------------

Figure 2.2 Edit Study page

To continue creating a study or to modify a study, on the Edit Study page complete these steps:

1. Enter or change(if editing) the name and/or description, if you choose.
2. Check the checkbox to make the study publicly available, if appropriate.
3. For the study log feature, click **View Log** or **Edit Log**. See [Viewing/Editing a Log](#) for details about the log.
4. Click **Save**.

Note: You can save the study at any point in the process of creating it. You can resume the definition and deployment process later.

5. If you choose to add a logo for the study, click the **Browse** button corresponding to **Logo File**. Navigate for the file, then click **Upload Now**. Once you save the

study (or its edit), the logo displays in the center of the page ([Figure 2.3](#)). On the home page for the study, the logo displays in the upper left, above the sidebar.



Figure 2.3 Example of a logo added to the caIntegrator2 browser on the Edit Study page

To continue, you can add subject annotation data sources, genomic data or imaging data.

Viewing/Editing a Log

On the Edit Study page, as a study manager you can open a detailed log for the study.

1. Click **View Log** on the Edit Study page to simply review an existing log. The log records all steps comprising activity in the study, with the most recent displaying at the top of the log.
2. To edit a log, click **Edit Log** on the Edit Study page.
Add an appropriate description/annotations to the individual log entries.
3. Check the **Update** box next to the description, then click **Save** to save the edits. The descriptions will now be available when any user views the log.

Working with Annotations – An Overview

One of the most important factors in creating a study in caIntegrator2 is in properly annotating the data. Because the process can be relatively complex, you might want to review the steps for working with annotations.

Annotation workflow summary:

1. Add an annotation group. This optional step is for users who have a rigid data dictionary of all annotations relevant to the study. This step can also be helpful in cases where a study has many annotations. For more information, see [Adding An Annotation Group](#) on page 19.
2. Add subject annotation data. This consists of multiple sub-steps.
 - a. Add a new subject annotation data sources file. This step uploads the file and starts the workflow for assigning uploaded data definitions. See [Editing an Annotation Group](#) on page 20, step 1.
 - b. Edit the annotations. This step opens the Define Fields for Subject Data page. See [Editing an Annotation Group](#) on page 20, step 2.
 - c. In the Define Fields for Subject Data page, review possible definitions in the annotation group associated with this study.

- d. Assign the visibility of each annotation definition. See [Editing an Annotation Group](#) on page 20, step 3.
- e. Locate and verify the assignment as “identifier” for one annotation. See [Assigning An Identifier or Annotation](#) on page 23.
- f. Review, verify and assign definitions for each annotation. You can do this in one of four ways:
 - Accept existing default definitions as described in the associated annotation group. See [Assigning An Identifier or Annotation](#) on page 23.
 - Create or manage definitions manually. See [Assigning An Identifier or Annotation](#) on page 23.
 - Search for and use definitions existing in other calIntegrator2 studies. see [Searching for Annotation Definitions](#) on page 26.
 - Search for and use definitions from caDSR. see [Searching for Annotation Definitions](#) on page 26.
3. Load the Subject Annotation Source. Up until this point, you can periodically save your work with the annotations, but before you can deploy the study, you must complete this step.
4. Deploy the study. See [Deploying the Study](#) on page 41.

Adding An Annotation Group

An annotation group is a group of annotation definitions configured in a CSV file. This feature is primarily meant for the Study Manager who knows that they have tightly restricted vocabulary definitions that are relevant to a study. In this optional step, you can review the uploaded Group Definition Source file before assigning the appropriate definition for your study.

To add an annotation group, follow these steps:

1. On the Edit Study page for a study, Annotation Groups section, click the **Add New** button.
2. On the Edit Annotation Group page that opens, enter a name for the annotation group.
3. Enter a description (optional).
4. Browse for the Group Definition Source CSV file.

The CSV file must include columns with these column headers in the first row: File Column Name, Field Type, Entity Type, CDE ID, CDE Version, Annotation Def Name, Data Type, Permissible, and Visible. Subsequent rows in the file define each subject annotation column in the subject annotation file.

- a. If a subject annotation is defined by a CDE Public ID, values for the following columns are required: File Column Name, Field Type, Entity Type, CDE ID, and Visible; a value for CDE Version is optional.

– OR –

- b. If a subject annotation definition is not defined by a CDE Public ID, values for the following columns are required: File Column Name, Field Type, Entity Type, Annotation Def Name, Data Type (String, Date, Numeric), Permissible (Yes or No), and Visible (Yes or No).
5. Click **Save**. This uploads the file, whose name now displays on the Edit Study page under Annotation Groups.

When you open the Define Fields for Subject Data page, the annotation definitions in the file you uploaded display on the page, available for assignment in the study. Additionally, you can view the definitions by viewing the annotation group listed in the first column of the matrix.

Note: Annotation definitions by default are visible only to the Study Manager's group. They are not visible to all caIntegrator2 users, unless you change the visibility for each.

Editing an Annotation Group

To edit an annotation group, on the Edit Study page for a study with an existing annotation group, click the **Edit Group** button.

1. You can change the Name and Description for the group.
2. A list of annotation definitions applied to the original annotation group displays on the Edit Annotation Group page.
 - In the drop-down list, you can select a different annotation group for the annotation definition.
 - You can change the visibility for the annotation definition.
 - Click **Change Assignment** to modify the properties of the annotation definition.
3. Click **Update Annotations** to confirm your edits for the group.

Adding Subject Annotation Data

The Edit Study page opens after you save a new study or click to edit an existing study.

Note: To edit information for an existing study, follow the same basic directions in this section. Instead of entering new information, you will modify existing information.

To add or edit subject annotation metadata in this page, follow these steps:

1. On the Edit Study page, click the **Add New** button in the Subject Annotation Data Sources section. Navigate to locate a subject annotation data file which is required for a study. Files must be in CSV file format.

- In the same section of the page, if a file has already been uploaded, its information displays in the varying fields. Click **Edit Annotations**. This opens the Define Fields for Subject Data page (*Figure 2.4*).

Define Fields for Subject Data

Assign annotation definitions to data fields and click **Done**.

Annotation Group	Visible	Annotation Definition	Annotation Header from File	Data from File	
Annotations - Default	<input checked="" type="checkbox"/>	Assign Annotation Definition	Subject	ASP221	ASP308
Annotations - Default	<input checked="" type="checkbox"/>	Age Change Assignment	Age	50-54	50-54
Annotations - Default	<input checked="" type="checkbox"/>	Gender Change Assignment	Gender	M	M
Annotations - Default	<input checked="" type="checkbox"/>	Survival Change Assignment	Survival		
Annotations - Default	<input checked="" type="checkbox"/>	Disease Change Assignment	Disease	ASTROCYTOMA	GEM
Annotations - Default	<input checked="" type="checkbox"/>	Grade Change Assignment	Grade		
Annotations - Default	<input checked="" type="checkbox"/>	Race Change Assignment	Race	WHITE	WHITE
Annotations - Default	<input checked="" type="checkbox"/>	Institution Change Assignment	Institution	NIH NEURO-ONCOLOGY BRANCH	NIH NEURO-ONCOLOGY BRANCH

Figure 2.4 Define Fields for Subject Data page

The first column of the table displays the annotation groups that have been created for this study. For more information, see [Adding An Annotation Group](#) on page 19.

- You can specify visibility of specified annotation data in the **Visible** column.
 - Select a checkbox for a row to make the corresponding data visible to all subscribers of the study or anonymous users if the study is made available to the public.
 - Clear a checkbox to hide the corresponding annotation from any subscriber or anonymous user of the study. Data continues to exist but does not show up in query fields nor query results.
- The Annotation Header from File column on the Define Fields for Subject Data page displays column headers taken from the source CSV file. The page also displays data values in the file you have designated. You must map each column name to an existing column name in the calIntegrator2 database or in

caDSR. If it doesn't yet exist, you can create a custom column name (Figure 2.5).

	A	B	C	D	E	F	G	H
1	Pa	Age	Gender	Survival	Disease	Grade	Race	
2	ASP221	50-54	M		ASTROCYTOMA		WHITE	
3	ASP308	50-54	M		GBM		WHITE	
4	FPH113	20-24	M		UNKNOWN		WHITE	
5	FPH114	40-44	M		UNKNOWN		WHITE	
6	FPH118	55-59	M		GBM		WHITE	
7	FPH309	50-54	M		GBM		WHITE	
8	E09238	45-49	M	18-24M	GBM		WHITE	
9	E09239	25-29	M		UNKNOWN		WHITE	
10	E09262	35-39	M		ASTROCYTOMA		WHITE	
11	E09278	30-34	M		UNKNOWN		WHITE	
12	E09331	35-39	M		UNKNOWN		ASIAN NOS	
13	E09332	55-59	M		GBM		WHITE	
14	E09336	30-34	M		GBM		WHITE	
15	E09348	60-64	M		GBM		WHITE	
16	E09378	45-49	M		UNKNOWN		WHITE	
17	E09449	50-54	M		UNKNOWN		OTHER	
18	E09454	0-4	M		UNKNOWN		WHITE	
19	E09489	55-59	M		GBM		WHITE	
20	E09515	35-39	M		UNKNOWN		WHITE	
21	E09569	45-49	M		UNKNOWN		WHITE	
22	E09587	35-39	M		UNKNOWN		OTHER	
23	E09601	40-44	M		GBM		WHITE	
24	E09610	55-59	M		GBM		WHITE	
25	E09611	60-64	M		UNKNOWN		ASIAN NOS	
26	E09615	45-49	M		UNKNOWN		WHITE	
27	E09624	35-39	M		GBM		WHITE	
28	E09645	45-49	M		UNKNOWN		WHITE	
29	E09657	50-54	M		UNKNOWN		WHITE	
30	E09730	40-44	M		UNKNOWN		WHITE	

Figure 2.5 Example of a source CSV file whose data you are mapping in caIntegrator2

The MOST important steps in creating a new study in caIntegrator2:

- You MUST designate one column in the file as a unique “identifier” column type.
- You MUST review and define column annotation definitions for each column header in the file.

Note the following regarding the list of annotations on this page:

- If caIntegrator2 “recognizes” the same column header in other files already in the system, a term, for example “age” or “survival”, which is the current definition appears in the **Annotation Definition** column above the blue **Change Assignment** link.
- When the annotation definition has not been assigned, and the area above the blue **Assign Annotation Definition** link is blank, no correlating term exists in the database. In this case, you must specify the field type, and then the term will populate the space. See [Assigning An Identifier or Annotation](#) for more information.
- A field name that displays in red indicates an error in the annotation. Click the **Change Assignment** button for more information about the error.

5. To indicate the unique identifier of choice, on the row showing the column header (PatientID in the figure, but other examples are subject identifier, sample identifier, etc), click **Change Assignment** in the **Field Definition** column.

Assigning An Identifier or Annotation

When you click **Change Assignment** on the Define Fields... page, the Assign Annotation Definition for Field Descriptor dialog box opens ([Figure 2.6](#)). On this page you can change the column type and the field definition for the specific data field you selected.

Note: When you change an assignment, you must make sure the data types match--numeric, etc.

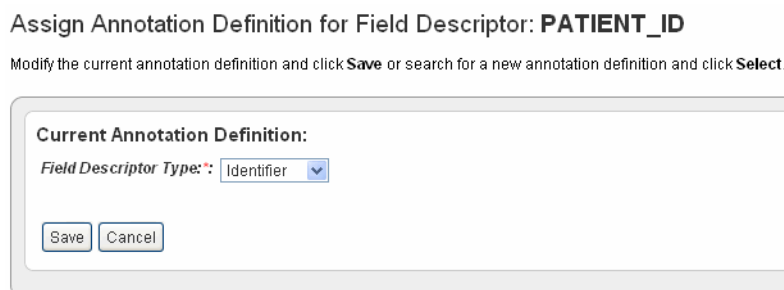


Figure 2.6 The Assign Annotation Definition dialog box

1. For the column (PatientID) that you choose to be the one and only Identifier column, in the **Column Type** drop-down list, select **Identifier**.
2. Click **Save** to save the identifier. This returns you to the Define Fields for Subject Data page where the Identifier is noted in the Field Definition column.
3. After you have defined which field is the Identifier, you must ensure that ALL other fields also have a field definition assignment. For those fields without a Field Definition assignment or for those whose Annotation Definition you want to review, click **Change Assignment**.
4. In the Assign Annotation Definition for Field Descriptor dialog box, select **Annotation** in the drop-down list.

As you select the column type, you can work with column headers in one of four ways in this dialog box.

- You can accept existing default definitions (those that are inherent in the data file you selected). See [Step 5](#).
 - You can create and/or manage your own definitions manually. See [Step 6](#).
 - You can search for and use definitions in other caIntegrator2 studies. See [Searching for Annotation Definitions](#) on page 26.
 - You can search for and use definitions found in caDSR. See [Searching for Annotation Definitions](#) on page 26.
5. Review the current annotation definition in the Assign Definition page, Current Annotation Definition section. Click **Cancel** to return to the Define Fields... page.

You can still initiate a search for another annotation definition in the Search for an Annotation Definition section if you choose to change the definition (Figure 2.7). See [Searching for Annotation Definitions](#) on page 26. Click **Save** to retain any changes.

Assign Annotation Definition for Field Descriptor: **MICROARRAY**

Modify the current annotation definition and click **Save** or search for a new annotation definition and click **Select**.

Current Annotation Definition:

Field Descriptor Type: Annotation

Name: MICROARRAY - JP

Definition:

Keywords: MICROARRAY

Data Type: string

Apply Max Number Mask: ☐ (Max Number)

Apply Numeric Range Mask: ☐ (Numeric Range)

Non-Permissible

- Moff 3354B
- NCI_Lung234_U133A
- NCI_U133A_61L
- CL20041110102AA
- NCI_Lung310_U133
- NCI_Lung_81_U133A
- NCI_Lung270_U133A
- NCI_Lung228_U133A

Permissible

Permissible Values:

Add >

< Remove

New Save Cancel

Search for an Annotation Definition:

Search Search existing studies and caDSR for definitions.

Figure 2.7 Current Annotation Definition

- To enter a new name annotation, or any other information about the annotation definition, click the **New** button and enter the information described in [Table 2.1](#)

Annotation Field	Field Description
Name	Enter the name for the annotation.
Definition	Enter the term(s) that define the annotation.
Keywords	Insert keyword(s) that can be used to find the annotation in a search, separated by commas.
Data Type	Select a string (default), numeric, or date.

Table 2.1 Annotation fields for new definitions

Annotation Field	Field Description
Apply Max Number Mask	<p>This field is available only for numeric-type annotations, or when a new definition is created. This feature is unavailable when permissible values are present.</p> <p>Select the box and enter a maximum number for the mask, such as “80” for age. When you query results above the value of the mask, then the system displays the mask and not the actual age.</p> <p>Note: If you enter masks of both “max number” and “range”, calIntegrator2 applies both masks at the same time.</p> <p>The Data Dictionary page now has a Restrictions column that shows restrictions whenever a mask has been applied.</p>
Apply Numeric Range Mask	<p>This field is available only for numeric-type annotations, or when a new definition is created. This feature is unavailable when permissible values are present.</p> <p>Select the box and enter a width of range for the mask, such as “5” representing blocks of 5 years. For example, if you enter a width of 5, the query only allows age blocks of 0-5, 6-10, 11-15, etc. When you query results above the value of the mask, then the system displays the mask and not the actual age ranges.</p> <p>Note: If you enter masks of both “max number” and “range”, calIntegrator2 applies both masks at the same time.</p> <p>The Data Dictionary page now has a Restrictions column that shows restrictions whenever a mask has been applied.</p>

Table 2.1 Annotation fields for new definitions

Annotation Field	Field Description
Permissible/Non-permissible Values	<p>Note: The first time you load a file, before you assign annotation definitions (step 5 on page 22), these panels may be blank. If the column header for the data is already “recognizable” by calIntegrator2, the system makes a “guess” about the data type and assigns the values to the data type in the newly uploaded file. They will display in the Non-permissible values sections initially. Use the Add and Remove buttons to move the values shown from one list to the other, as appropriate.</p> <p>When you select or change annotation definitions by selecting matching definitions (described in Searching for Annotation Definitions on page 26), this may add (or change) the list of non-permissible values in this section.</p> <p>If you leave all values for a field in the Non-permissible panel, then when you do a study search, you can enter free text in the query criteria for this field.</p> <p>If there are items in the Permissible values list, then the values for this annotation are restricted to only those values. When you perform a study search, you will select from a list of these values when querying this field. If there are no items in the permissible values list then the field is considered free to contain any value.</p> <p>To edit a field's permissible values, you must change the annotation definition. You can do this even after a study has been deployed.</p> <p>Note: You cannot edit permissible values in an existing annotation definition. To change permissible values, you must create a new annotation.</p>

Table 2.1 Annotation fields for new definitions

Searching for Annotation Definitions

An alternative to creating a new definition is to search for annotation definitions already present in calIntegrator2 studies or in caDSR.

1. Enter search keyword(s) in the **Search** text box on the Assign Annotation Definition page. Click **Search** or click **Enter** to launch the search. After a few moments, the search results display on the page (*Figure 2.8*).

Search for an Annotation Definition:

microarray Search existing studies and caDSR for definitions.

Matching Annotation Definitions from caintegrator2				
Name	Actions	CDE Public ID	Data Type	Definition
MICROARRAY	Select		string	Created via selenium for DC Lung Full on C
MICROARRAY	Select		string	Created via selenium for DC Lung Full on C

Matching Annotation Definitions from caDSR					
Name	Actions	CDE Public ID	Context	Status	Definition
Microarray Microarray Analysis Data float One Dimensional Array	Select View	2658378	caBIG	RELEASED	A microarray is a piece of glass or plastic on which different samples have been affix are usually DNA fragments but may also be antibodies, other proteins, or tissues. _Ana microarrays to profile the pattern of proteins).:A collection or single item of factual info drawn. _Generic value domain for a single dimensional array with floating numbers as i
Microarray Identifier java.lang.Long	Select View	2223905	caCORE	RELEASED	A microarray is a piece of glass or plastic on which different samples have been affix are usually DNA fragments but may also be antibodies, other proteins, or tissues. _One

Figure 2.8 Results for annotation definition search

2. To view the definitions corresponding to any of the “Matching Annotation Definitions”, which are those currently found in other caIntegrator2 studies, click the [term], such as “age”, hypertext link. The definition then appears in the Current Annotation Definition segment of the page just above.

In summary, when you click the link, that assigns the definition to the Define Fields for Subject Data page, and it also closes the Annotation Definition page.

You can modify any portion of the definition, as described in [step 6](#) on page 24.

3. The matches from caDSR display some of the details of the search results. To view more details of a match, such as permissible values, click **View**, which opens caDSR to the term. If you click **Select**, the caDSR definition automatically replaces the annotation definition for this field with which you are working.

Caution: Take care before you add a caDSR definition that it says exactly what you want. caDSR definitions can have minor nuances that require specific and limited applications of their use.

4. Once you have settled on an appropriate field definition for the annotation, click **Save**. This returns you to the Define Fields for Subject Data page.

Note: If you have not clicked **Select** for alternate definitions in this dialog box, then click **Save** to return to the Define Field...dialog box without making any definition changes.

5. From the Define Fields for Subject Data page, be sure and designate the data types for each field in the file. Click **Save** on each page to save your entries or click **New** to clear the fields and start again. You will not be able to proceed until every field definition entry on the Fields for Subject Data screen has an entry, one as the unique Identifier and the remainder as annotations.

The Data From File columns on the page display the column header values of the first three rows you designated as “annotations”.

Note: Saving your entries in this way saves the study by name and description, but does not deploy the study. See [Deploying the Study](#) on page 41.

The Edit Study page now displays a “Not Loaded” status for the file whose annotations (column headers) you have defined ([Figure 2.9](#)).

Study Overview

Study Name:

Study Description:

Allow public to browse this study: ☐

Status: Not Deployed

Status Description:

Owner: manager

Last Modified By: manager

Last Modified Date: 05/13/2010 13:43:55

Study Log: [View Log](#) [Edit Log](#)

Study Logo: None [Browse...](#)

JPEG/GIF, 200x72 maximum

[Upload Now](#)

Annotation Groups [Add New](#)

Group Name	Description	Number of Annotations	Action
Annotations - Default	Default annotation group	8	
test2		6	

Subject Annotation Data Sources [Add New](#) [Edit Survival Values](#)

Type	Description	Status	Last Modified	Action
DELIMITED_TEXT	clinical_test_for_doc.csv	Not Loaded	Unavailable	Edit Annotations

Figure 2.9 Example file whose annotations have been defined but not yet loaded

- Click the **Load Subject Annotation Source** button in the Action section to load the data file you have configured, The **Deploy Study** button, to this point has been unavailable, but this step activates the button.

Note: You can add as many files as are necessary for a study. Patients 1-20 in first file, 21-40 in second file, or many patients in first file and annotations in second file, etc. As long as IDs are defined correctly, it works.

- Click **Deploy Study**. calIntegrator2 now loads data from the file to the calIntegrator2 database, and the file status changes to “Loaded”.

Note: You can change assignments even after the study is deployed, using the Edit feature. For more information, see [Creating/Editing a Study](#) on page 17.

The Manage Studies page opens when the study is deployed. The **Deployed** status is indicated on the Manage Studies page as well as the Edit Study page. For more information, see [Managing a Study](#) on page 41.

You can continue to perform other tasks in calIntegrator2 while deployment is in process.

See also [Deploying the Study](#) on page 41.

Note: You can repeatedly upload additional or updated subject annotations, samples, image data, array data to the study at later intervals. These later imports do not remove any existing data; they instead insert any new subjects or update annotations for existing subjects.

Defining Survival Values

Survival value is the length of time a patient lived. If you plan to analyze your data in calIntegrator2 to create a Kaplan-Meier (K-M) Plot, then during the Annotation Definition process described above, you must make sure that you have defined at least three fields set to the “date” Data Type. These will be matched to the following three properties during Survival Value definition.

- **Survival Start Date**
- **Death Date**
- **Last Followup Date**

Note: Setting survival values is optional if you do not plan to use the K-M plot analysis feature or if you do not have this kind of data (survival values) in the file.

For some applications, such as REMBRANDT and I-SPY, survival values are pre-defined in the databases when you load the data. In calIntegrator2, however, you can review and define survival value ranges in a data set you are uploading to a study. To be able to do so, you need to understand the kind of data that can comprise the survival values.

To set up survival values, follow these steps:

1. On the Edit Study page, click **Edit Survival Values**. This opens the Survival Value Definitions dialog box ([Figure 2.10](#)).

Edit Survival Values

Select an existing definition and click **Edit** or click **New**.

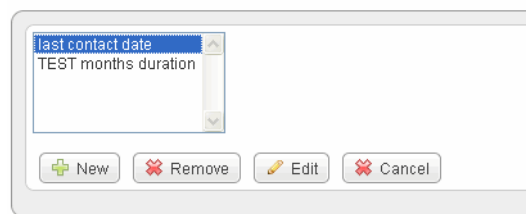


Figure 2.10 Survival Value Definition dialog box

2. Click **New** to enter new survival value definitions.

- OR -

Click **Edit** to edit existing survival value definitions.

3. The dialog box extends, now displaying radio buttons and three drop-down lists that show column headers for date metadata in the spreadsheet you have

uploaded. [Figure 2.11](#) displays survival value ranges that have already been added to a study.

Edit Survival Values

Select an existing definition and click **Edit** or click **New**.

last contact date
TEST months duration

+ New ✖ Remove ✏ Edit ✖ Cancel

Survival Value Definition Properties for 'last contact date'

Survival Definition Type: ☒ By Date ☐ By Length of time in study

* Name: last contact date

* Survival Start Date: ENROLLMENT_DATE

* Death Date: LAST_CONTACT_DATE

* Last Followup Date: LAST_CLINICAL_ASSESSMENT_DATE

Save

Figure 2.11 Survival Definitions example

Survival values can be defined by Date or by Length of time in study. Select the radio button for the category that defines your survival data.

In the drop-down lists, select the appropriate survival value definitions for each field listed. You might want to refer to the column headers in the data file itself. Dates covered by the definitions are already in the data set. You cannot enter specific dates.

- **Survival Definition Type** – Select whether the survival time is defined by dates or length of time subject was in the study.
- **Name** – Enter a unique name that adequately describes the survival values you are defining here. *Example:* Survival from Enrollment Date or Survival from Treatment Start. The name you enter displays later when you are selecting survivals to create the K-M plot.
- **Survival Length Units** – Select the appropriate units for this data.
- **Survival Start Date** – Select the column header for this data.
- **Death Date** – Select the column header for this data.
- **Last Followup Date** – Select the column header for this data.

See also [Creating Kaplan-Meier Plots](#) on page 70.

Updated the Edit Survival Value Definitions page, now has a radio button and 2 different types of ways to define survival values.

Adding/Editing Genomic Data

Note: Genomic data must be parsed and stored in caArray to be able to analyze it in caIntegrator2.

Once you have loaded subject annotation data and identified patient IDs, you can add either one or more sets of array genomic sample data from caArray, which caIntegrator2 maps by sample IDs to the patient IDs in the subject annotation data, covered in this section, or you can load imaging files from NBIA, also mapped by IDs to the patient data, covered in [Adding Imaging Data](#) on page 38. You can also edit genomic data information that you have already added to the study. Genomic sample data and imaging data are independent of each other, so neither is required before loading the other.

It is essential that you are well acquainted with the data you are working with--the subject annotation data, and the corresponding array data in caArray.

caIntegrator2 supports a limited number of array platforms. For more information, see [Managing Platforms](#) on page 42.

To add genomic data to your caIntegrator2 study, follow these steps:

1. On the Edit Study page where you have selected and added the subject annotation data, click the **Add New** button under Genomic Data Sources. You can upload genomic data only from caArray.

This opens the Edit Genomic Data Source dialog box. Enter the appropriate information in the fields ([Figure 2.12](#)). This fields are described below.

Edit Genomic Data Source

Enter data source parameters and click **Save**.

Data Source

caArray Web URL:

caArray Server Hostname:

(Note: caArray v 2.3 or newer is required)

caArray Server JNDI Port:

caArray Username:

caArray Password:

caArray Experiment Id:

Vendor:

Data Type:

Platform:

Use Supplemental Files: ☐

Central Tendency for Technical Replicates:

Indicate if Technical Replicates have high statistical variability: ☒

Standard Deviation Type:

Standard Deviation Threshold:

Figure 2.12 Edit Genomic Source dialog box

- **caArray Web URL** – Enter the URL for the caArray to be used for the genomic data sources. This will enable a user to link to the referenced caArray experiment from the study summary page.
- **caArray Host Name** – Enter the hostname for your local installation or for the CBIIT installation of caArray, array.nci.nih.gov. If you misspell it, you will receive an error message.
- **caArray JNDI Port** – Enter the appropriate server port. See your administrator for more information. *Example:* For the CBIIT installation of caArray, enter **8080**.
- **caArray Username** and **caArray Password** – If the data is private, you must enter your caArray account user name and password; you must have permissions in caArray for the experiment. If the data is public, you can leave these fields blank.
- **caArray Experiment ID** – Enter the caArray Experiment ID which you know corresponds with the subject annotation data you uploaded. *Example:* Public experiment “beer-00196” on the CBIIT installation of caArray (array.nci.nih.gov). If you misspell your entry, you will receive an error message.
- **Vendor** – Select either **Agilent** or **Affymetrix**
- **Data Type** – Select **Expression** or **Copy Number**.
- **Platform (needed only for Agilent)** – If appropriate, select the **Agilent** platform.

Note: Because you can add more than one set of genomic data to a study, a study can also have multiple platforms, one for each set of genomic data.

- **Central Tendency for Technical Replicates** – If more than one hybridization is found for the reporter, the hybridizations will be represented by this method.
- **Indicate if technical replicates have high statistical variability** – If more than one hybridization is found, checking this box will display a ** in the genomic search results when a reporter value has high statistical variability.
- **Standard Deviation Type** - When the checkbox for indicating if technical replicates have high statistical variability is checked, this parameter becomes available. Select in the drop-down the calculation to be used to determine whether or not to display a ** (see previous bullet point).
 - **Relative**, which calculates the Relative Standard Deviation in percentage value
 - **Normal**, which calculates the Standard Deviation in numeric value
- **Standard Deviation Threshold** – When the checkbox for indicating if technical replicates have high statistical variability is checked, this parameter becomes available. This is the threshold at which the Standard Deviation Type is exceeded and the reporter is marked with a **.

2. Click **Save**.

caIntegrator2 goes to caArray, validates the information you have entered here, finds the experiment and retrieves all the sample IDs in the experiment. Once this finishes, the experiment information displays on the Edit Study page under the Genomic Data Sources section ([Figure 2.13](#)).

Host Name	Experiment Identifier	File Description	Data Type	Status	Action
ncias-d227-v.nci.nih.gov	admin-00001	Mapping File(s): nci_sample_mapping.csv Control Sample Mapping File(s): igla_0034_control_samples.csv	Expression	Loaded	Edit Map Samples Delete

Figure 2.13 Genomic Data Sources section of the Edit Study page

3. If you want to redefine the caArray experiment information, you can edit it. Click the **Edit** link corresponding to the Experiment ID. The Edit Genomic Data Source dialog box reopens, allowing you to edit the information.

Note: At any point in the process of working within a study, you can create a gene list. For more information, see [Creating a Gene List](#) on page 58.

Mapping Genomic Data to Subject Annotation Data

Because the goal of caIntegrator2 is to integrate data from subject annotation, genomic and imaging data sources, data from uploaded source files must be mapped to each other.

Note: Supplemental files from caArray for mapping data must be configured appropriately. For information, see [Supplemental Files Configuration](#) on page 119.

To map the samples from the caArray experiment to the patients in the subject annotation data you uploaded, follow these steps:

1. On the Edit Study page, click the **Map Samples** button. This opens the Edit Sample Mappings page ([Figure 2.14](#)).

Edit Sample Mappings
Upload mapping files and click **Map Samples**.

Data Source

caArray Server Hostname: array-stage.nci.nih.gov
caArray Server JNDI Port: 8080
caArray Username:
caArray Experiment Id: jacob-00182
Subject to Sample Mapping File:
Control Sample Set Name:
Control Samples File:

Control Sample Sets

Set Name	Sample Name
Control Set 1	636
	638
	637
	635

Sample Mappings

Unmapped Samples
Sample Name
1
10
100
101
102
103

Figure 2.14 Edit Sample Mappings page showing some already mapped samples

If you have already mapped samples, when you first open this page, they will be listed under Control Sample Sets. If you have not already mapped samples, all of the samples in the caArray experiment you selected are listed as unmapped, because caIntegrator2 does not know how these sample names correlate to the patient data in the subject annotation file until you upload the subject to sample mapping file.

2. At the top of the page, click **Browse** to navigate for the CSV file that identifies the mapping information. This provides caIntegrator2 with the information for mapping patients to caArray samples. Click the **Upload Mapping File** button.

Acceptable mapping file format:

- Affymetrix – The mapping file has only two columns (typically without headers)—one that shows the subject ID (designated in caIntegrator2 as the

“Identifier”) and one that has “Sample name” field from the linked caArray experiment, with one subject per row (*Figure 2.15*).

	A	B	C	D	E	F	G	H
1	E10216	GeneratedSample.UNKNOW	DISEASE_L_E10216_U133P2					
2	E10144	GeneratedSample.UNKNOW	DISEASE_L_E10144_U133P2					
3	E09212	GeneratedSample.UNKNOW	L_20070227_16-22-37-238_E09212_U133P2					
4	E09369	GeneratedSample.UNKNOW	L_20070227_16-22-37-238_E09369_U133P2					
5	E10162	GeneratedSample.UNKNOW	DISEASE_L_E10162_U133P2					
6	E10318	GeneratedSample.UNKNOW	DISEASE_L_E10318_U133P2					
7	E09264	GeneratedSample.OLIGO_L_20070227_11-27-27-881_E09264_U133P2						
8	E10252	GeneratedSample.UNKNOW	DISEASE_L_E10252B_U133P2					

Figure 2.15 An eExample Affymetrix sample mapping file, in CSV format

- Agilent and all other platforms – Raw (level 1) data cannot be mapped; only normalized, processed (level 2) data is acceptable. The 5 column format is as follows

1. Subject ID
2. Sample ID
3. Name of supplemental file (as attached to the experiment in caArray)
4. Name of column header (in the supplemental file) which contains the sample IDs.
5. Name of column header (in the supplemental file) which holds the level 2 data.

Note: When you open the mapping file, make sure that the patient ID is used for mapping.

Unmapped samples continue to show at the top of the caIntegrator2 page. They were loaded from caArray, but they are not in the mapping file. These are not used for integration.

6. Scroll down the page to see samples that are mapped to the patients in the subject annotation data (*Figure 2.16*).

1336	GeneratedSample.OLIGO_L_20070227_11-49-51-876_HF0547_U133P2	
1338	GeneratedSample.UNKNOW_DISEASE_L_E10029_U133P2	
1339	GeneratedSample.GBM_L_20070226_14-05-29-569_HF1356_U133P2	
1340	GeneratedSample.OLIGODENDROGLIOMA_L_HF0599_U133P2	
1342	GeneratedSample.GBM_L_20070226_13-30-40-39_HF0142_U133P2	
1345	GeneratedSample.GBM_L_20070226_14-31-20-427_HF1400_U133P2	
Samples Mapped to Subjects		
Sample ID	Sample Name	Subject Identifier
901	GeneratedSample.UNKNOW_DISEASE_L_E10216_U133P2	E10216
911	GeneratedSample.UNKNOW_DISEASE_L_E10144_U133P2	E10144
914	GeneratedSample.UNKNOW_L_20070227_16-22-37-238_E09212_U133P2	E09212
918	GeneratedSample.UNKNOW_L_20070227_16-22-37-238_E09369_U133P2	E09369
922	GeneratedSample.UNKNOW_DISEASE_L_E10162_U133P2	E10162
925	GeneratedSample.UNKNOW_DISEASE_L_E10318_U133P2	E10318
930	GeneratedSample.OLIGO_L_20070227_11-27-27-881_E09264_U133P2	E09264
940	GeneratedSample.UNKNOW_DISEASE_L_E10252B_U133P2	E10252
954	GeneratedSample.GBM_L_20070226_13-14-06-57_E09624_U133P2	E09624
957	GeneratedSample.ASTROCYTOMA_L_E09137_U133P2	E09137
958	GeneratedSample.UNKNOW_DISEASE_L_E09890_U133P2	E09890
968	GeneratedSample.UNKNOW_L_20070227_16-57-07-283_E09515_U133P2	E09515
1004	GeneratedSample.UNKNOW_L_20070227_17-26-09-910_E09722_U133P2	E09722

Figure 2.16 Example of samples mapped to patients' data

Uploading Control Samples

A Control Samples file is used to calculate fold change data, which compares “tumor” sample gene expression in the caArray experiment to the control samples to identify those that exhibit up or down gene regulation. Control samples can be the “normal” samples, but that is not necessarily the case.

To upload the control samples, follow these steps:

1. On the Edit Sample Mappings page, click the **Map Samples** link.
2. Click **Browse** to navigate for the control samples file, and click the **Upload Control Samples** File button. The control sets display at the top of the page once they have been uploaded (*Figure 2.17*).

The screenshot shows a web interface for mapping samples. At the top, there is a form with the following fields and buttons:

- Array Experiment Id:** A text box containing 'rembr-00037'.
- Subject to Sample Mapping File:** A text box with a 'Browse...' button next to it.
- Control Sample Set Name*:** A text box.
- Control Samples File:** A text box with a 'Browse...' button next to it.
- A 'Map Samples' button is located below the 'Control Samples File' field.

Below the form, there is a section titled 'Control Sample Sets'. It contains a table with two columns: 'Set Name' and 'Sample Name'.

Set Name	Sample Name
Rembrandt controls	GeneratedSample.Normal_L_20070227_14-22-24-128_Normal_5_U133_P2
	GeneratedSample.Normal_L_20070227_14-01-17-731_HF0526_U133P2
	GeneratedSample.Normal_L_20070227_14-01-17-731_HF0131_U133P2
	GeneratedSample.Normal_L_20070227_14-01-17-731_HF0523_U133P2
	GeneratedSample.Normal_L_20070227_14-01-17-731_HF0120_U133P2
	GeneratedSample.Normal_L_20070227_14-01-17-731_HF0137_U133P2
	GeneratedSample.Normal_L_20070227_14-22-24-128_Normal_6_U133_P2
	GeneratedSample.Normal_L_20070227_14-01-17-731_HF0151_U133P2
	GeneratedSample.Normal_L_20070227_14-01-17-731_HF0388_U133P2

Figure 2.17 Example list of control samples

The control samples now display toward the bottom of the page.

3. This information will be used when performing other tasks in caIntegrator2, to be described in other sections.

Note: If a Control Set is to be used in Gene Expression For Annotation, or Gene Expression plots for Annotation Query, then the control set should be composed of only samples which are mapped to subjects.

Configuring Copy Number Data

You can add copy number data for a genomic data source by uploading the mapping file. This allows you to configure parameters to be used when segmentation data is being configured.

The name specified in the third column of the mapping file is specific for each array manufacturer as follows:

- Affymetrix – The third column of the mapping file must contain filenames that end in .cnchp. The corresponding experiment in caArray must have these files and the extensions must match .cnchp.
- Agilent – The third column must name a file which contains level 2 copy number data. Level one copy number will not work. This file name is repeated for each line in the mapping file.

To add copy number data relating to the genomic data you are adding, follow these steps:

1. In the Genomic Data Sources section, for the data you have already added, click **Configure Copy Number Data** button.

Note: This link is available only if you have uploaded copy number data and you are configuring a Copy Number data type (as indicated by the Data Type column on the Edit Study page).

The Edit Copy Number page opens (*Figure 2.18*).

Edit Copy Number Data Configuration

Enter data source parameters and click **Save**.

Figure 2.18 Edit Copy Number page

2. Browse for and enter appropriate information to identify the copy number mapping file. The fields are described in *Table 2.2*. An asterisk* indicates a required field.

Field	Description
Subject and Sample Mapping File	Browse for the appropriate CN mapping file. The file must be a CSV file with 3 column format for mapping single data file and 5 column format for mapping 1 data file per sample.
caDNACopy Service URL*	Control for selecting the URL which hosts the caDNACopy grid service
Change Point Significance Level	Significance levels for the test to accept change-points
Early Stopping Criteria	The sequential boundary used to stop and declare a change

Table 2.2 Fields for retrieving a copy number mapping file.

Field	Description
Permutation Replicates	The number of permutations used for p-value computation
Random Number Seed	The segmentation procedure uses a permutation reference distribution. This should be used if you plan to reproduce the results.

Table 2.2 Fields for retrieving a copy number mapping file.

3. Click **Save Segmentation Data Calculation Configuration** for a genomic data source. On the screen upload a copy number mapping file (format: subject id, sample id, file name) and configure the parameters to be sent when computing segmentation data.

Adding Imaging Data

Once you have loaded subject annotation data and identified patient IDs, you can add either array genomic sample data from caArray which calIntegrator2 maps by sample IDs to the patient IDs in the subject annotation data, or you can load imaging files from NBIA, also mapped by IDs to the patient data, covered in this section. Genomic sample data and imaging data are independent of each other, so neither is required before loading the other.

It is essential that you are well acquainted with the data you are working with--the subject annotation data, and the corresponding imaging data in NBIA.

Any data in NBIA can be uploaded to calIntegrator2. Imaging data consist of images and/or annotations for images.

To add imaging data to the study you are creating or are editing, follow these steps:

1. On the Edit Study page, click the **Add New** button under Imaging Data Sources section. Imaging data can be NBIA images or image annotations, which are uploaded in spreadsheet format.

This opens the Edit Imaging Data Source dialog box. Enter the appropriate information in the fields ([Figure 2.19](#)). Asterisks indicate required fields..

Edit Imaging Data Source

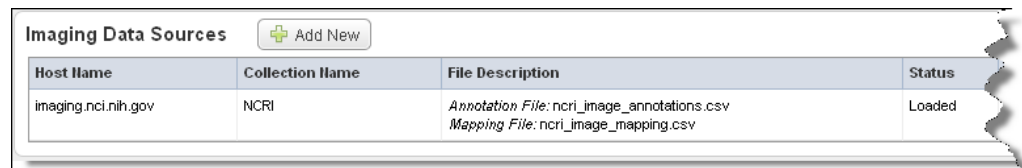
Enter a NBIA Data Source and Image Mapping Data from a file and click **Save**.

Figure 2.19 Edit Image Data Source dialog box

- ° **NBIA Server Grid URL*** – Enter the URL for the grid connection to NBIA

- **NBIA Username and NBIA Password.** This information is not required, as currently all data in the NBIA grid is Public data.
 - **Collection Name*** – Enter the name/source for the collection.
 - **Current Mapping** – If a mapping file has already been uploaded to the study to map imaging data, the file name displays here.
 - **Select Mapping File Type*** – Click to select the file type:
 - **Auto** – No file is required. Selecting this takes all subject annotation subject IDs and attempts to map them to the corresponding ID in the collection in NBIA. If the ID does not exist in NBIA, then no mapping is made for that ID.
 - **By Subject** – Requires a file to be uploaded. The “subject annotation to imaging mapping file” must be in CSV format with two columns that map the caintegrator2 subject annotation subject ID to the NBIA subject ID.
 - **By Image Series** – Requires a file to be uploaded. The subject annotation to imaging mapping file needs to be a two column mapping (CSV) from the caintegrator2 subject annotation subject ID to the NBIA study instance UID.
 - **Subject to Imaging Mapping File** – Click **Browse** to navigate to the appropriate subject annotation to imaging mapping file. See **Select Mapping File Type*** field description.
2. Click **Add** to upload the data to calntegrator2.

The imaging data information displays on the Edit Study page under the Imaging Data Sources section ([Figure 2.20](#)).



Imaging Data Sources + Add New			
Host Name	Collection Name	File Description	Status
imaging.nci.nih.gov	NCRI	Annotation File: ncri_image_annotations.csv Mapping File: ncri_image_mapping.csv	Loaded

Figure 2.20 Imaging Data Sources section of the Edit Study page

3. Once the data is uploaded, you must assign identifiers and annotations to the data in the same way you did with the subject annotation data. For more information, see [Assigning An Identifier or Annotation](#) on page 23 and [Searching for Annotation Definitions](#) on page 26.
4. To deploy the study, see [Deploying the Study](#).

Adding External Links

This feature on the Edit Study page allows you to configure a CSV file with URLs to be used as external links relevant to a study. This allows you to easily share or configure references.

To add an external link, follow these steps:

1. As a study manager, you can configure a CSV file with URLs to be used as external links.
2. On the Edit Study page, click the **Add** button under External Links section. External links can be any URL(s) to resources that are hosted external to calIntegrator2 but are relevant to the study being deployed.
3. Assign a name to the external link.
4. Add a description for the link, if appropriate.
5. Browse for the CSV file containing URLs (HTTP linked) to resources outside of calIntegrator2.
6. Click **Upload Now**. calIntegrator2 does not validate any links in the file being uploaded.

Once you have created external links for a study, when the study is open, an External Links section showing the link(s) displays on the left sidebar of the page (*Figure 2.21*).

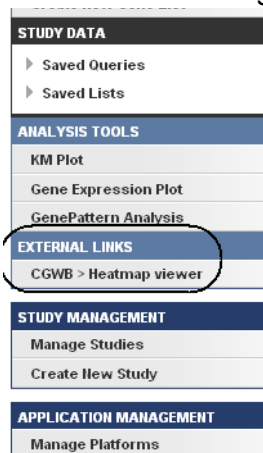


Figure 2.21 Left sidebar displaying external links

Click the link to open a page that displays appropriately formatted web page links (*Figure 2.22*). .

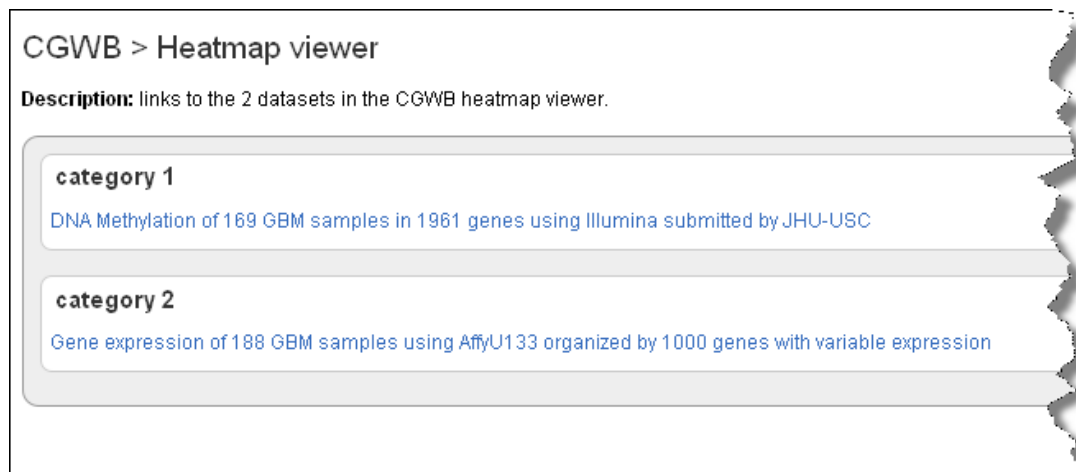


Figure 2.22 An example of external links

Deploying the Study

When you are ready to deploy the study, click the **Deploy Study** button on the Edit Study page. caIntegrator2 retrieves the selected data from the data service(s) you defined and makes the study available to a study manager or to anyone else who may want to analyze the study's data. Using the Manage Studies feature, you can then configure and share data queries and data lists with all investigators who access the study.

Note that you can continue to work in caIntegrator2 while study is being deployed.

Managing a Study

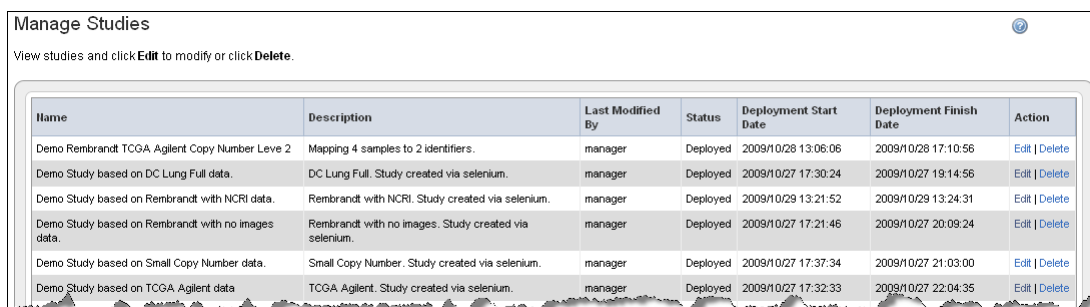
Note: A user without management privileges has no access to this section of caIntegrator2.

Once you have started to create a study or have deployed it, you can update an existing study in the following ways:

- Add new attributes (annotations) and upload relevant data to an existing study.
- Delete a study
- Modify existing annotation definitions
- Reload subset of study data and re-deploy the study and perform new analyses
- Re-deploy the entire study with new set of data and mappings.

To update, edit or delete a study, follow these steps:

1. On the left sidebar, click **Manage Studies**. The Manage Studies page appears (*Figure 2.23*).



Manage Studies

View studies and click **Edit** to modify or click **Delete**.

Name	Description	Last Modified By	Status	Deployment Start Date	Deployment Finish Date	Action
Demo Rembrandt TCGA Agilent Copy Number Leve 2	Mapping 4 samples to 2 identifiers.	manager	Deployed	2009/10/28 13:06:06	2009/10/28 17:10:56	Edit Delete
Demo Study based on DC Lung Full data.	DC Lung Full. Study created via selenium.	manager	Deployed	2009/10/27 17:30:24	2009/10/27 19:14:56	Edit Delete
Demo Study based on Rembrandt with NCRI data.	Rembrandt with NCRI. Study created via selenium.	manager	Deployed	2009/10/29 13:21:52	2009/10/29 13:24:31	Edit Delete
Demo Study based on Rembrandt with no images data.	Rembrandt with no images. Study created via selenium.	manager	Deployed	2009/10/27 17:21:46	2009/10/27 20:09:24	Edit Delete
Demo Study based on Small Copy Number data.	Small Copy Number. Study created via selenium.	manager	Deployed	2009/10/27 17:37:34	2009/10/27 21:03:00	Edit Delete
Demo Study based on TCGA Agilent data	TCGA Agilent. Study created via selenium.	manager	Deployed	2009/10/27 17:32:33	2009/10/27 22:04:35	Edit Delete

Figure 2.23 Manage Studies page

All of the “in process” or “completed” studies display on this page, with associated metadata. Note that whoever edited or updated the study last is shown in the Last Modified Column, indicated as the Study Manager.

- Click the **Edit** link corresponding to your study of choice to open the Edit Studies page ([Figure 2.24](#)).

Edit Study
Configure your study, and click the **Save** or **Deploy Study** button at the bottom of the page when complete.

Study Overview

Study Name: Demo Study based on Rembrandt with NCRI data

Study Description: Rembrandt with NCRI. Study created via selenium.

Allow public to browse this study: ☐

Status: Deployed

Status Description: Minutes for deployment (approx): 3

Owner: marplej

Last Modified By: manager

Last Modified Date: 03/25/2010 08:45:43

Study Log: [View Log](#) [Edit Log](#)

Study Logo: None [Browse...](#)
JPG/GIF, 200x72 maximum
[Upload Now](#)

Annotation Groups [Add New](#)

Group Name	Description	Number of Annotations	Action
Annotations - Default	Default annotation group	28	

Subject Annotation Data Sources [Add New](#) [Edit Survival Values](#)

Type	Description	Status	Last Modified	Action
DELIMITED_TEXT	rembrandt_clinical_Aug08_subset_mod_for_NCRI.csv	Loaded	Unavailable	Edit Annotations Reload

Genomic Data Sources [Add New](#)

Host Name	Experiment Identifier	File Description	Data Type	Status	Last Modified	Action
array.nci.nih.gov	jagla-00034	Mapping File: None Configured Control Sample Mapping File(s): None Configured	Expression	Loaded	Unavailable	Edit Add

Imaging Data Sources [Add New](#)

Host Name	Collection Name	File Description	Status	Last Modified	Action
imaging.nci.nih.gov	NCRI	Annotation File: ncri_image_annotations.csv Mapping File: ncri_image_mapping.csv	Loaded	Unavailable	Edit Edit Annotations

External Links [Add New](#)

Name	Description	File Name	Number of Links
------	-------------	-----------	-----------------

Figure 2.24 Edit Studies page where you can edit any details for an existing study

On this page you can edit any details such as adding or deleting files, survival values, and so forth. For information about working with the Edit Study feature, see [Creating/Editing a Study](#) on page 17.

- Click the **Delete** link to delete the corresponding study.

Managing Platforms

calIntegrator2 supports a limited number of array platforms, all of which originate from Agilent or Affymetrix. While they do not represent all of the platforms supported by caArray, calIntegrator2 must have array definitions loaded for the platforms it supports, and be able to properly load the data from caArray and parse it.

You can create a study without genomic data, but you cannot add genomic data to a calIntegrator2 study without a corresponding supported array platform. If you add more than one set of genomic data to the study, you can specify more than one platform for the study.

On the Manage Platforms page, you can identify, add or remove supported platforms.

To manage platforms in calIntegrator2, follow these steps:

1. Click **Manage Platforms** on the left sidebar.

The Manage Platforms page that opens lists the platforms calIntegrator2 currently supports, those that the system can pull from caArray ([Figure 2.25](#)). You can also add a new platform by entering information in the fields in the Create a New Platform section.

Platform Name	Platform Type	Platform Channel Type	Vendor	Array Name(s)
Agilent-022522	Agilent Copy Number	Two-Color	Agilent	Agilent-022522
AgilentG4502A_07_01	Agilent Gene Expression	Two-Color	Agilent	AgilentG4502A_07_01
AgilentG4502A_07_3	Agilent Gene Expression	Two-Color	Agilent	AgilentG4502A_07_3
GeneChip Human Mapping 100K Set	Affymetrix Copy Number	One-Color	Affymetrix	Mapping50K_Hind240, Mapping50K_Hind240
GenomeWideSNP_6.Full	Affymetrix SNP	Two-Color	Affymetrix	GenomeWideSNP_6
HG-U133 Plus 2	Affymetrix Gene Expression	One-Color	Affymetrix	HG-U133 Plus 2

Figure 2.25 Manage Platforms page

2. To add a platform, in the Platform Type field, select the appropriate platform type from the drop down list.
3. Click **Browse** to navigate for the Affymetrix or Agilent file you want to add.
4. Enter a **Platform Name** if the file is a NON-GEML.xml file.

Depending on the Platform Type you select, there may be other parameters to provide here as well, such as **Platform Channel Type** for an Agilent platform.

5. Click the **Browse** button to browse for the appropriate annotation file. When you have located it, click **Open** in the Upload File dialog box. The system displays the annotation file you select in the Annotation File box.
6. Once all parameters have been entered, click **Create Platform**.

The platform deployment can be time-consuming. If the platform takes more than 12 hours to deploy, calIntegrator2 displays a “timed out” message. At that point, you can delete the platform, even if it has not loaded to the system.

Note: Platform loading can fail if the manufacturer’s platform annotation file is missing data.

CHAPTER 3

SEARCHING A CAINTEGRATOR2 STUDY

This chapter describes the processes for searching studies within calIntegrator2.

Topics in this chapter include:

- [Search Overview](#) on this page
- [Searching a Study](#) on page 46
- [Managing Queries](#) on page 54

Search Overview

The search and browse functions in calIntegrator2 allow you to search for subject annotation data, genomic data or imaging data that were uploaded into the application as part of a study. When gene expression and imaging data are uploaded into a calIntegrator2 study, mapping files that correlate the data in those files to patient IDs in the subject annotation data file must also be uploaded. When you launch a search, calIntegrator2 finds and integrates the subject annotation, genomic and imaging data based on the mapping files and the criteria that you define in the search query.

In a search query, you can specify criteria for just one of the data types, or configure complex search criteria that join two or three data types. The available criteria for the query were defined when the study was deployed.

The basic workflow for a study search follows these steps:

1. Select the study to be searched.
2. Select one data type:
 - **Annotations** – Searches one or more uploaded CSV files for data identifiers or annotations (column headers) specified when the study was created

- **Genomic** – Searches caArray experiments samples uploaded in the study for gene expression data by gene name or reporter ID.
 - **Image Series** – Searches NBIA files uploaded in the study for image annotations or links to images, identified by subject identifiers or image series IDs.
3. Define criteria for the search in the selected data type and run the search.
 4. For a more complex search, select multiple criteria from more than one data type.
 5. Specify whether you want subject/imaging annotations to display or genomic data to display.
 6. Review search results.
 7. Configure results column and sorting display settings. You can do this before or after you run a search. If you choose to do it after, you must re-run the search.
 8. Download annotation search results as a CSV file. The CSV file contains only the data you specified in the annotation and display configurations.
 9. Follow links to NBIA in the search results to view or download images located in the search.

Searching a Study

To initiate a search of all annotations and/or other data in a study, follow these steps:

1. In caIntegrator2, in the upper right hand corner, select the study you want to browse or perform a simple search.
2. On the left sidebar, under the first section that displays the study name, click **Search [Study Name]**. This opens a simple search query page with five tabs (*Figure 3.1*).

Search Demo Study based on Netherlands data

Criteria Results Type Sorting Query Results Save query as...

Define Query Criteria for: Unsaved Query

Annotations - Default Add

No criteria added. Please select criteria from the pulldown box.

☐ or ☒ and

Run Query

Figure 3.1 Search page

3. On the Criteria tab, in the drop-down list, select the type of data you want to search. The listed options reflect the type of data that have been uploaded to the study.

Note: You can perform a search using one or more criteria you set in one of the data types, or you can define criteria in more than one data type per query, creating a more complex search.

- **Annotations**
- **Gene Expression**
- **Image Series**

4. Click **Add** to define annotation elements for the search.

Continue with:

Annotation and Image Data Searches on page 47

Gene Expression Data Searches on page 49

5. To add additional criteria for the search, repeat steps 3 and 4, as appropriate. You can set more than one data type or more than one criterion for a data type. The criteria become cumulative, thus refining the search.
6. Once you have configured the query criteria, select the Boolean **Or** or **And** search operator at the bottom of the page.
 - **Or** finds a data subset with at least one of the search criteria
 - **And** finds a data subset with both/or all search criteria.
7. Click the **Remove** button to clear any data elements you have defined.
8. You can launch the search from this tab. Click the **Run Search** button. For information about the search results, see *Chapter 4 Viewing Query Results*. You may want to run the search first to see what kind of results you get before you configure the data display, described in step 9.

– or –

9. On the Results Type tab, you can specify the columns you want to display in the search results data. On the Sorting tab, you can specify how the data is to be sorted. For more information, see *Results Type Tab* on page 51 and *Sorting Tab* on page 53.

Note: As long as you are still in the current query session, you can return to the Criteria, Columns and Sorting tabs to add, modify or remove data and display criteria and re-run the search. If you configure another query without saving the first, the first query will be lost. If you save the query, your current search criteria are saved.

Annotation and Image Data Searches

- If you select the Annotation data type which is the default selection, an additional drop-down list displays data elements that are annotation definitions specified when the data was uploaded into the study (*Figure 3.2*). Select a search criterion from among the options. You can make only one selection at a time.

Note: If the study includes imaging data, imaging annotations should be available in the Annotations list.

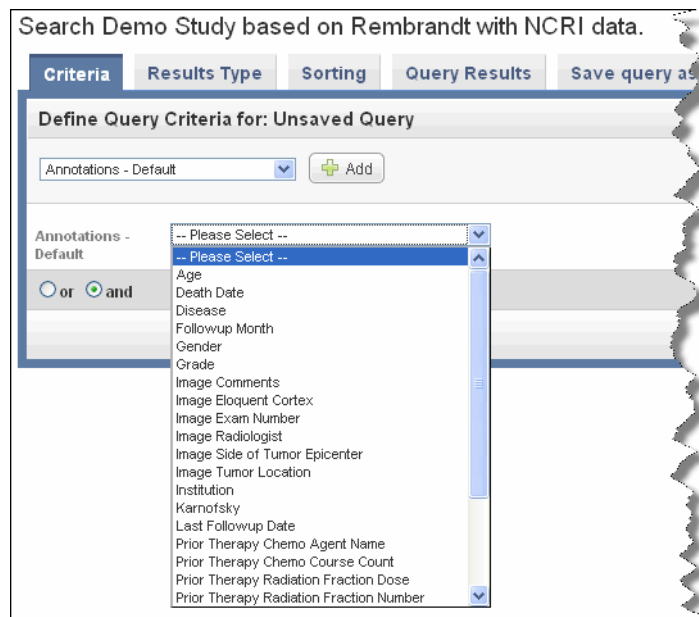


Figure 3.2 Annotation search criteria, including criteria for imaging

- Each choice opens other fields relevant to the selection where you can further define your search query.
 - If permissible values were added when the annotation was defined, you must select among the values in a drop-list that displays on the right side of the page.
 - If no permissible values were defined as part of the annotation, you have the option to enter descriptive text in a text box on the right side of the page (Figure 3.3).

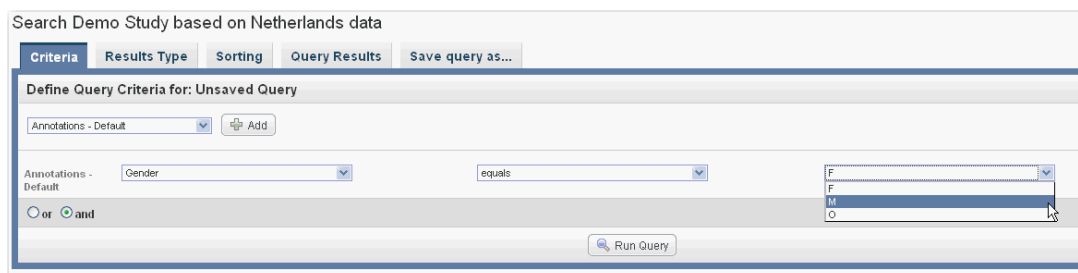


Figure 3.3 You may be able to further define search criteria when you select a specific subject annotation or imaging annotation element

Note: When working with image data, if only an Imaging Mapping file was uploaded when the study was created and not an Image Series Annotation file, you cannot enter image search criteria. The search results will, however, display a link that allows you to view the associated images in NBIA.


Continue with step 5 in [Searching a Study](#) on page 46.

Gene Expression Data Searches

1. For the Gene Expression selection, select **Gene Name** or **Fold Change**. If the study includes multiple platforms, a **Platform** option is also visible.
2. **Gene Name** or **Fold Change** – Enter one or more gene symbols in the text box or click the icons to locate genes in the following databases. If you enter more than one gene in the text box, separate the entries by commas. If multiple platforms are part of the study, your platform selection in the Fold Change query criteria determines the control samples that are available.

Note: If you leave the gene symbols field blank, calIntegrator2 searches all gene symbols for a match to the other criteria you specify.

calIntegrator2 provides three methods whereby you can obtain gene names for a gene expression search.

- **caBIO** – This link searches caBIO, then pulls identified genes into calIntegrator2 for analysis.
 - a. Click the **caBIO** icon ().
 - b. Enter **Search Terms**. Note that calIntegrator2 can perform a search on a partial HUGO symbol. For example, as search using *nicotin* would find matches with “nicotinic” and “nicotinamide”.
 - c. Select if you want to search in **Gene Keywords**, **Gene Symbols** or **Pathways** (from the drop-down list).
 - Selecting **Gene Keywords** searches only the Full Name field in caBIO.
 - Selecting **Gene Symbols** searches only the Unigene and HUGO gene symbols in caBIO.
 - Selecting **Pathways** searches only the pathway names in caBIO. Note that searching in Pathways is a two step process. First, the initial Pathway search produces search results which are pathways. Second, from the pathway search results screen, you must select pathways of interest, then click **Search Pathways for Genes** to obtain a list of genes related to the selected pathways.
 - d. Select the **Any** or **All** choice to determine how your search terms will be matched. **Any** finds any match for any search term you entered. **All** finds only results that match all of the search terms.

- e. Choose the **Taxon** from the drop-down list and click **Search**. The search results display ([Figure 3.4](#)).

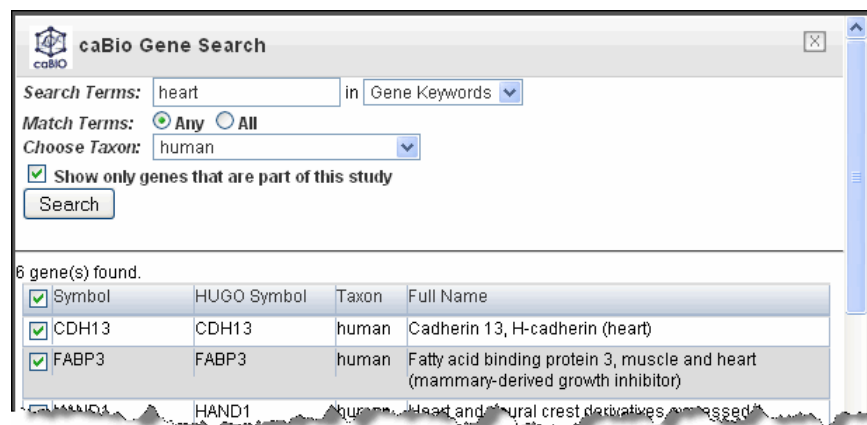


Figure 3.4 Example caBIO gene search criteria and search results

- f. In the search results, use the check boxes to identify the genes whose symbols you want to use in the gene expression analysis.
- g. Click **Use Genes** at the bottom of the page. This pulls the checked genes into the Criteria tab ([Figure 3.5](#)).

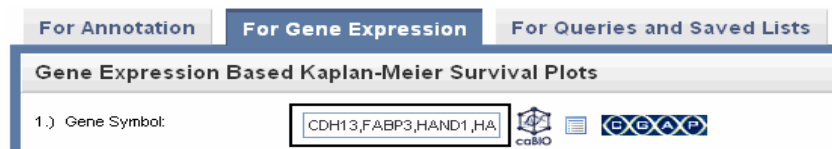



Figure 3.5 Genes pulled in from caBIO display on the Criteria tab

- **Gene List** – This link locates gene lists saved in caIntegrator2.
 - a. Click the Genes List icon () to open a small dialog that lists previously saved gene lists. For more information, see [Creating a Gene List](#) on page 58.
 - b. In the drop-down menu, select a gene list. In the list that appears, use the check boxes to identify the genes whose symbols you want to use in the gene expression analysis.
 - c. Click **Use Genes** at the bottom of the dialog. This pulls the checked genes into the Search Criteria tab.
- **CGAP** – Use this directory to identify genes. Before clicking this link you must enter gene symbols in the text box. This link does not pull anything into caIntegrator2 but does provide information about the gene(s) whose names you entered.

Additional fields display for the Fold Change selection.

The fold change option appears only if genomic control samples have been uploaded to the study. Fold change identifies genes with expression differences compared to control samples, as defined when the study was deployed in

calIntegrator2. You can enter query values in greater/lesser-than-or-equal-to arguments.

3. Select or enter data for the Fold change fields shown in [Figure 3.6](#):

The screenshot shows a web form for calIntegrator2. On the left, there's a label 'Gene Expression' followed by a dropdown menu currently showing 'Fold Change'. To the right of this is a text input field for 'Gene Symbol(s) (comma separated list) or blank for all genes'. Below these are two more dropdowns: 'Control Sample Set' (showing 'Control Set 1') and 'Regulation Type' (showing 'Up'). At the bottom right is a text input field for 'Up-regulation folds' containing the value '2.0'. There are some small icons on the right side of the form.

Figure 3.6 Fields for identifying fold change search criteria

- **Control Sample Set** – Select from the drop down list the name of the uploaded control sample set to serve as the fold change reference.
- **Regulation Type** – Select the term that describes the gene expression in comparison with the control samples: **Up** is increased expression; **Down** is decreased expression; **Up or Down** is increased or decreased; **Unchanged** means no change in expression.
- **Up-Regulation Folds** – Enter a numerical value representing fold change. The number you enter here is dependent upon the Regulation Type you selected.
 - **Up** = Up Regulation Folds – Samples with a fold change greater than this value, when compared to the control samples, will be returned.
 - **Down** = Down Regulation Folds – Samples with a fold change less than this value, when compared to the control samples, will be returned.
 - **Up or Down** = Down Regulations Folds, Up Regulation Folds – Samples with a fold change either up or down, when compared to the control samples, will be returned.
 - **Unchanged** = Samples with a fold change between the two specified values will be returned.

For example, if you enter 2.0 in this field, after selecting **Up** in the previous field, the search will locate genes whose expression is 2 times (2-fold up regulation) the base value.

Continue with step 5 in [Searching a Study](#) on page 46.

Results Type Tab

You can specify columns for the way you want the search results to display either before or after you run the search. If you run the search directly from the Criteria tab before setting the results type/sorting features, by default only the Subject Identifiers display on the Search Results tab. You can then come back to the Results Type tab and [Sorting Tab](#) to expand the display options and re-run the search, having set the display parameters.

The selection you make on the Results Type tab determines whether calIntegrator2 displays search results for subject annotation or genomic data. It filters the search based on the criteria you set on the Criteria tab, whether it is annotation, gene expression or image series data type(s). In other words, if you select annotation criteria

on the Criteria tab, but select Genomic on the Results Type tab, the data subset that displays on the Search Results tab is genomic data that is filtered by the annotation criteria you defined on the Criteria tab.

1. On the Results Type tab, select the **Annotation** or **Genomic** radio button to search annotation data (*Figure 3.7*).

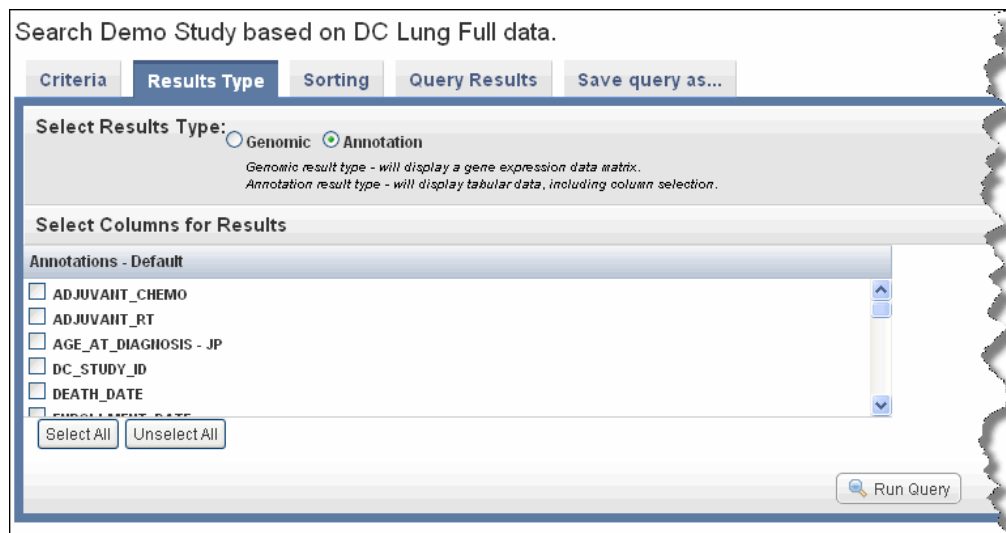


Figure 3.7 Results Type tab, annotation options

Annotation – Select the annotation elements that you want to display in the search results. All elements listed are column headers in the data uploaded to the study. You can make multiple selections on this list.

Note: For subject annotations, the Patient or Subject Identifier displays by default in the search results.

Results display as tabular data.

Genomic – Select the Reporter Type and Results Orientation:

- **Gene Name** – Finds and summarizes at the gene level all reporters that match criteria for the gene you defined on the Criteria tab
- **Reporter ID** – Finds all reporters that map to the gene(s) you identified on the Criteria tab
- **Genes in rows/Subjects in columns** or **Genes in columns/Subjects in rows** – Determines query results matrix format

Results display in a gene expression data matrix. For more information, see [Genomic Data](#) on page 56.

Imaging – If imaging annotations have been added to the study, annotation elements also display on the lower right section of this page when you select **Annotation**. All elements listed are column headers in the image annotation data uploaded to the study. You can make multiple selections on this list.

Note: If you select even one Image Annotation on the Results Type tab, the Image Series IDs display by default in the search results. If you select no

Image Annotations on the Results Type tab, however, even if you have selected image series criteria on the Criteria tab, no image series IDs display in the search results. The fact that images can be located, however, in NBIA is indicated by two image-related buttons at the bottom of the Query Results page. You can open the images in NBIA, but they will be at StudyInstance UID level. See [Relationship of Patient to Study to Series to Images](#) on page 66.

Results display as tabular data. For more information, see [Subject Annotation and Imaging Data](#) on page 56.

2. Use the **Select All** or **Unselect All** buttons to aid you in making your selections.

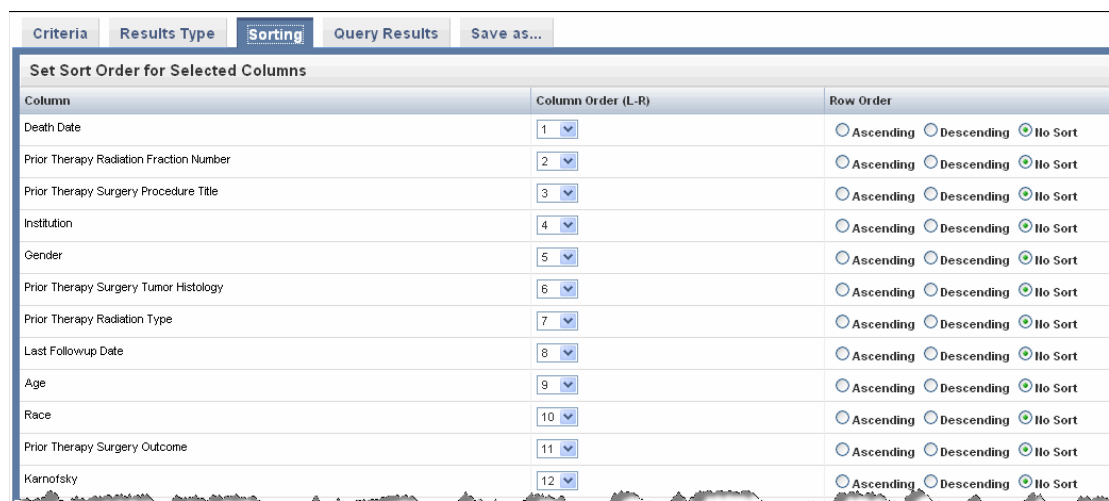
The column selection is saved as part of the query if you save it. See [Saving a Query](#) on page 54.

Sorting Tab

On the Sorting tab, you can set the sort order for data columns in the query results. You can also indicate whether column contents are sorted in ascending or descending order.

The columns that display on the Sorting tab are those criteria that you selected on the [Results Type Tab](#) for an Annotation Results type search.

1. Select the Sorting tab and indicate the left to right column order of the Search Results by changing one or more numbers in the Column Order column in this table ([Figure 3.8](#)).



Column	Column Order (L-R)	Row Order
Death Date	1	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Prior Therapy Radiation Fraction Number	2	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Prior Therapy Surgery Procedure Title	3	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Institution	4	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Gender	5	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Prior Therapy Surgery Tumor Histology	6	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Prior Therapy Radiation Type	7	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Last Followup Date	8	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Age	9	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Race	10	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Prior Therapy Surgery Outcome	11	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort
Karnofsky	12	<input type="radio"/> Ascending <input type="radio"/> Descending <input checked="" type="radio"/> No Sort

Figure 3.8 Sorting tab

2. In the **Row Order** column, indicate how you want columns sorted, **Ascending** or **Descending**, or leave the default, **No Sort**, if you choose.
3. Click **Run Query** at the bottom of the page to execute your sorting changes in the search results. When you do so, the change in column order is visible on the Query Results tab, as well as on the Sorting tab. For example, any column that you have indicated to be number “1” now appears in Query Results immediately

after the Subject Identifier column and at the top of the Set Sort Order table on the Sorting tab.

Sorting parameters are saved as part of the query if you choose to save it using the Save Query feature. See [Saving a Query](#) on page 54.

4. If you click the **Reset** button before running the query from the Sorting tab, the original column settings are restored.

For information about the search results, see [Chapter 4 Viewing Query Results](#).

Managing Queries

When you create a search query in calIntegrator2, you can save the query for later use or edit it.

[Exporting Query Results](#)

Saving a Query


To save a query, follow these steps:

1. Click the **Save As** tab and enter a **Search Name** and **Search Description**, unique to the search. *Example: **Batch ID 6 and female***
2. Click **Save**.

Once the query is saved, it is listed by its name under the **Study Data > Queries > My Queries** in the left sidebar, whenever the study to which the query applies is selected. Click on the saved query in this list to either edit or re-run the query. Click on the query name to retrieve query results. If you hover over the Name text for the query, a pop-up displays the query description.

Editing a Query

To edit a query, follow these steps:

1. To edit a query, select it in the left sidebar under the **Study Data > Queries > My Queries**.
2. Click the **Edit** icon () corresponding to the study.
3. Change the query and display criteria on the Criteria, Columns and Sorting tabs.
4. On the Save As tab, check the appropriate options and click **Save As**. You can use the same name as the original query or modify the name as needed.

Exporting Query Results

After running a search, you can export the result set or a subset as a tab-delimited text file. For more information, see [Exporting Data](#) on page 67.

CHAPTER 4

VIEWING QUERY RESULTS

This chapter describes search results that calIntegrator2 returns after queries.

Topics in this chapter include the following:

- [Query Results Overview](#) on this page
- [Browsing Query Results](#) on page 56

Query Results Overview

After you launch a search of a calIntegrator2 study, the system automatically opens the Query Results tab showing the results of your search.

If you have not configured the column and sort display parameters before launching the search, by default the tab shows only the subject identifiers and a column that allows you to select each row of the data subset.

To display and/or sort additional data, you must return to the Columns and/or Sorting tabs to set display parameters, then re-run the search. The new search results will display the additional information, with the columns and data sorted as you specified. See [Results Type Tab](#) on page 51.

calIntegrator2 paginates search results into pages of configurable size (default 20) with standard paginated navigation controls. To sort columns by ascending or descending parameters for on any displayed field, click on the underlined column header.

You can download search results as a CSV file. The file contains the annotations, columns and data sort configurations you specified in the search query. See [Exporting Query Results](#) on page 54.

Browsing Query Results

The query results that can display depend upon the criteria you established for the search. Follow the links below for more information about the category of data you searched.

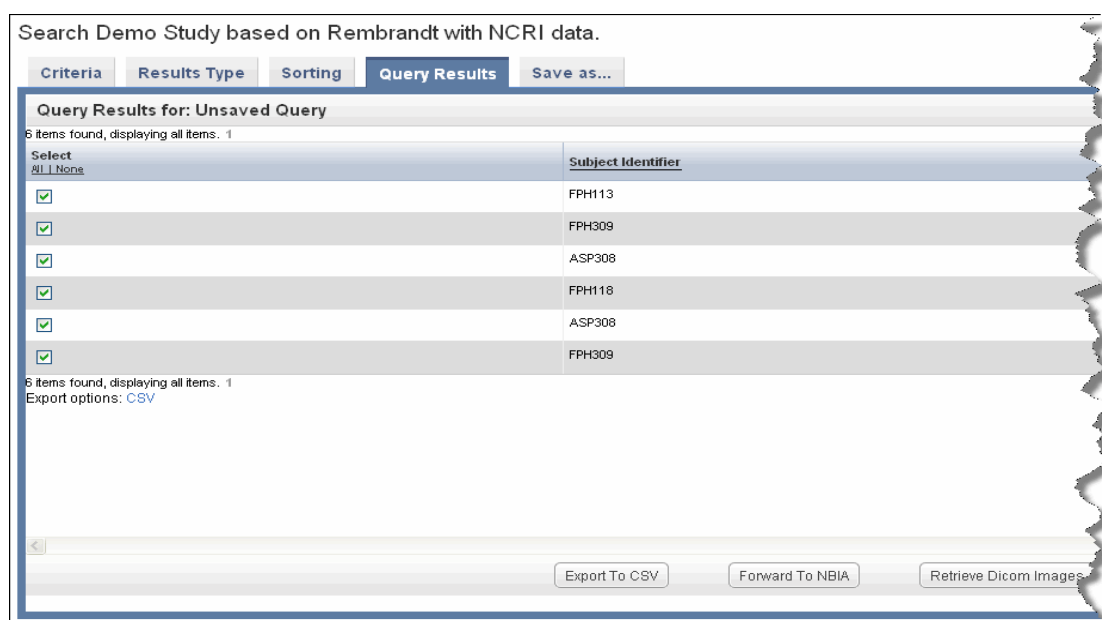
[Subject Annotation and Imaging Data](#) on page 56

[Genomic Data](#) on page 56

[Expanding Imaging Data Results](#) on page 62

Subject Annotation and Imaging Data

If you run the search before configuring column and sort display parameters, only the [subject] ID that meet the criteria and a column allowing you to select each row appear on the table ([Figure 4.1](#)). .



Search Demo Study based on Rembrandt with NCRI data.

Criteria Results Type Sorting **Query Results** Save as...

Query Results for: Unsaved Query

6 items found, displaying all items. 1

Select	Subject Identifier
<input checked="" type="checkbox"/>	FPH113
<input checked="" type="checkbox"/>	FPH309
<input checked="" type="checkbox"/>	ASP308
<input checked="" type="checkbox"/>	FPH118
<input checked="" type="checkbox"/>	ASP308
<input checked="" type="checkbox"/>	FPH309

6 items found, displaying all items. 1
Export options: [CSV](#)

Export To CSV Forward To NBIA Retrieve Dicom Images


Figure 4.1 Query Results page

You can add details for one or more subjects by configuring them on the Results Type tab. Annotations listed there are the column headers in the CSV file(s) that were uploaded to the study. For information about using the Results Type tab, see [Results Type Tab](#) on page 51.

Genomic Data

If after defining gene expression criteria on the Criteria tab, you select the **Genomic** result type on the Results Type tab, genomic data search results display in a gene expression data matrix. Because the data was downloaded from caArray, the data permissions granted there still apply. In other words, if you have been given access to the data in caArray, you can see it in caIntegrator2.

You can select on the Results Type tab a preferred orientation for displaying the results: genes in rows and subjects in columns, or genes in columns and subjects in rows.

For Gene criteria, the cells display the median gene expression value for each gene. By each gene symbol, calIntegrator2 displays an icon () which you can click to open the Cancer Genome Anatomy Project (CGAP) showing data for the gene (Figure 4.2).

Search Demo Study based on DC Lung Full data.

Criteria

Results Type

Sorting

Query Results

Save query as...

Query Results for: JP - two genes

	Subject ID	134	48	231	150	320	230	109	348	183	459	614	232	538
	Sample ID	134	48	231	150	320	230	109	348	183	459	614	232	538
Gene	Reporter ID													
AKT1	207163_s_at	1318.12	1164.96	1520.83	630.27	1491.69	1169.84	858.65	986.98	1113.31	851.28	888.96	1051.76	1117.1
TERT	207199_at	7.7	15.92	42.03	57.93	50.94	29.93	16.03	27.4	35.17	25.54	47.79	27.09	25.1

Figure 4.2 Genomic query result matrix after gene criterion has been specified

If you have selected “Genomic” on the Results Type tab, then the column headers are a clickable label which sorts the entire table on that column. If you selected Reporter ID on the Results Type tab, the Reporter ID is clickable (and the gene is not clickable).

For fold-change criteria, the cells display the normalized signal-based value for a given reporter for a given sample. In the results matrix, calIntegrator2 highlights matrix values for fold change results that meet fold change criteria. Red represents upregulated values and blue indicates downregulated values (Figure 4.3, Figure 4.4).

Criteria	Results Type	Sorting	Query Results	Save query as...																
Query Results for: Unsaved Query																				
	Subject ID	45	153	472	35	486	467	497	80	82	360	237	347	83	469	375	178	456	544	221
	Sample ID	45	153	472	35	486	467	497	80	82	360	237	347	83	469	375	178	456	544	221
Gene Name																				
CDH13		2.45	-1.9	2.24	2.06	-1.76	-1.05	2.6	-1.88	-2.69	-1.85	-1.03	-3.18	-1.39	-2.36	-1.48	-3.38	-6.18	3.64	-2.02
FABP3		1.09	2.06	1.77	1.21	2.55	1.5	1.39	-1.1	2.03	1.01	-1.02	-1.29	4.57	-1.15	1.33	1.09	4.22	1.06	1.67
HAND1		-1.07	1.48	1.65	1.39	2.2	2.85	1.22	1.42	1.58	2.2	1.53	7.25	-1.11	1.78	1.26	2.1	2.11	1.77	2.91
HAND2		1.63	1.06	-1.72	-1.59	-2.9	-1.24	-1.28	-1.09	-1.77	-2.99	2.1	1.49	-1.14	-2.06	-1.34	1.66	-1.33	-2.1	-2.1
LBH		1.27	2.21	1.59	-1.54	1.15	1.73	2.75	2.34	-1.78	-1.97	3.6	-1.02	2.34	1.06	1.65	1.03	1.69	-1.44	-1.18
LOC128102		1.2	5.04	1.75	1.39	-1.16	2.49	-2.77	1.88	-1.3	1.64	1.04	2.45	1.3	1.4	2.05	3.33	-1.54	1.69	2.23

Export options: CSV

Export To CSV

Figure 4.3 Gene Name search 6 genes, Reporter Type: Gene. Genes display in rows and subjects appear in columns.

Criteria	Results Type	Sorting	Query Results	Save query as...																
Query Results for: Unsaved Query																				
		Subject ID	134	48	231	150	320	230	109	348	459	614	232	538	207	451	351	375	163	227
		Sample ID	134	48	231	150	320	230	109	348	459	614	232	538	207	451	351	375	163	227
Gene Name	Reporter ID																			
CDH13	204726_at		-1.27	1.78	-2.16	1.99	1.58	-2.09	1.29	-1.51	-1.23	-2.19	-2.43	1.52	1.62	2.67	-1.14	-1.48	-1.37	2.99
FABP3	205738_s_at		1.11	5.06	2.67	-1.62	3.12	1.7	-1.02	-3.3	2.71	1.71	-1.04	-1.42	-1.04	2.11	2.52	1.71	-2.62	1.41
FABP3	214285_at		2.39	3.9	2.86	1.23	3.47	2.61	1.15	1.84	2.67	2.9	-1.56	-1.17	-2.81	2.12	2.4	-1.75	1.18	2.56
HAND1	220138_at		1.86	1.2	5.06	-1.2	2.9	1.47	-1.45	1.03	1.77	1.69	1.24	2.0	2.43	1.07	2.29	1.26	2.63	1.73
HAND2	220480_at		-2.01	-2.46	-1.77	-1.09	3.01	-1.85	-1.67	-1.95	-1.66	3.53	1.09	-1.53	1.19	-2.27	2.74	-1.34	-1.35	1.04
HSD3B2, LOC391081, HSD3B1, LOC128102	215665_at		4.42	-3.0	3.83	-3.97	9.44	5.29	5.71	1.69	1.02	2.51	6.55	1.82	2.2	-2.17	3.93	4.54	5.68	4.8
LBH	221011_s_at		1.57	1.36	-1.33	1.1	-2.68	-1.79	1.82	2.36	-1.0	1.49	-1.2	-2.42	-1.07	-1.78	2.02	1.65	-1.81	1.9
LOC128102	216819_at		-1.07	-3.23	-1.53	1.04	-3.72	-1.13	-2.19	-2.6	-2.06	-1.12	-1.43	-2.18	-1.17	-5.93	-1.7	-1.19	4.77	5.21
<div><div></div><div></div></div>																				
<div>Export To CSV</div>																				

Figure 4.4 Gene Name search 6 genes, Reporter Type: Reporter ID. Genes display in rows and subjects appear in columns.

- Genomic data does not display in tandem with subject annotation and imaging data; it only displays when you select the Genomic result type on the Results Type tab. Genomic data is however, filtered by subject annotation and imaging query criteria configured on the Criteria tab.
- Click the Export Options CSV link to download the CSV file whose data displays on the Search Results tab. When you do so, the CSV file opens automatically in MS Excel or similar applications for working with spreadsheets, showing the columns and sorting as you defined them in calIntegrator2 on the appropriate tabs.
- You can save genes identified in the search results as a gene list.

Creating a Gene List

From any page in calIntegrator2 that shows such a group, you can save a list of genes so you can use it for searches or analyses. To create a gene list, follow these steps:

1. Click the **Create New Gene List** link in the left sidebar. This opens the Manage Gene List page ([Figure 4.5](#)):

Manage Gene List

Click **Create Gene List** to create a new Gene List.


The screenshot shows a web form titled "Create a New Gene List". It contains the following elements:

- A text input field for "Gene List Name" with a red asterisk indicating it is required.
- A larger text area for "Gene List Description".
- A checkbox labeled "Make Visible to Others".
- A text input field for "Gene Symbols".
- An "Upload File:" button with a "Browse..." link below it. A note "(csv file format)" is present.
- Two icons: a blue cube icon labeled "caBIO" and a blue box icon labeled "CXP".
- A "Create Gene List" button at the bottom right.

Figure 4.5 Manage Gene List page

2. Enter a name for the gene list.
3. Enter a description (optional).
4. Select **Make Visible to Others** if you want the list to be visible to anyone who views the study. This selection places the list in the Global List folder in the left sidebar under Saved Lists. In any box where you can select lists, the term '**Global**' will identify any list so identified when the list is created.
5. For **Gene Symbol**, enter one or more gene symbols in the text box or click the icons to locate genes in the following databases. If you enter more than one gene in the text box, separate the entries by commas.

calIntegrator2 provides three methods whereby you can obtain gene names for creating a gene list:

- **caBIO** – This link searches caBIO, then pulls identified genes into calIntegrator2 for analysis.
 - a. Click the **caBIO** icon ().
 - b. Enter **Search Terms**. Note that calIntegrator2 can perform a search on a partial HUGO symbol. For example, as search using *nicotin* would find matches with “nicotinic” and “nicotinamide”.
 - c. Select if you want to search in **Gene Keywords**, **Gene Symbols** or **Pathways** (from the drop-down list).
 - Selecting **Gene Keywords** searches only the Full Name field in caBIO.
 - Selecting **Gene Symbols** searches only the Unigene and HUGO gene symbols in caBIO.
 - Selecting **Pathways** searches only the pathway names in caBIO. Note that searching in Pathways is a two step process. First, the initial

Pathway search produces search results which are pathways. Second, from the pathway search results screen, you must select pathways of interest, then click **Search Pathways for Genes** to obtain a list of genes related to the selected pathways.

- d. Select the **Any** or **All** choice to determine how your search terms will be matched. **Any** finds any match for any search term you entered. **All** finds only results that match all of the search terms.
- e. Choose the **Taxon** from the drop-down list and click **Search**. The search results display in the same dialog box (*Figure 4.6*).

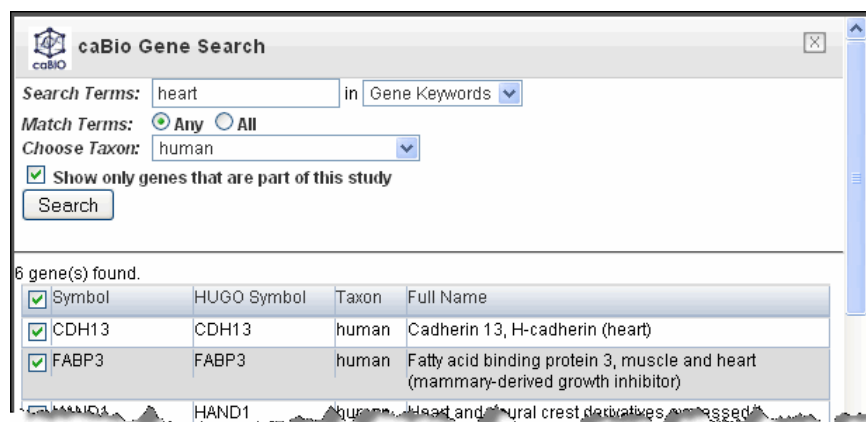



Figure 4.6 Example caBIO gene search criteria and results

- f. In the search results, use the check boxes to identify the genes whose symbols you want to include in the gene list.
- g. Click **Use Genes** at the bottom of the page. This pulls the checked genes into the Gene Symbols field on the Gene List page.
- **Gene List** – This link locates gene lists saved in calIntegrator2.
 - a. Click the Gene List icon () to open a small dialog that lists prior-saved gene lists in calIntegrator2.
 - b. In the drop-down menu, select a gene list. In the list that appears, use the check boxes to identify the genes whose symbols you want to use in the plot analysis.
 - c. Click **Use Genes** at the bottom of the dialog. This pulls the checked genes into the Gene Symbols field on the Gene List page.
- **CGAP** – Use this directory to identify genes. Before clicking this link you must enter gene symbols in the text box. This link does not pull anything into calIntegrator2 but does provide information about the gene(s) whose names you enter.
6. If you so choose, you can upload a gene list. For the Upload File field, click the **Browse** button to navigate to a .csv file made up of gene symbols. calIntegrator2 converts the comma-separated content to a gene list.

- Click **Create Gene List** at the bottom of the page. caIntegrator2 now opens the Edit Gene List page which shows the name and symbols of the newest gene list ([Figure 4.7](#)).

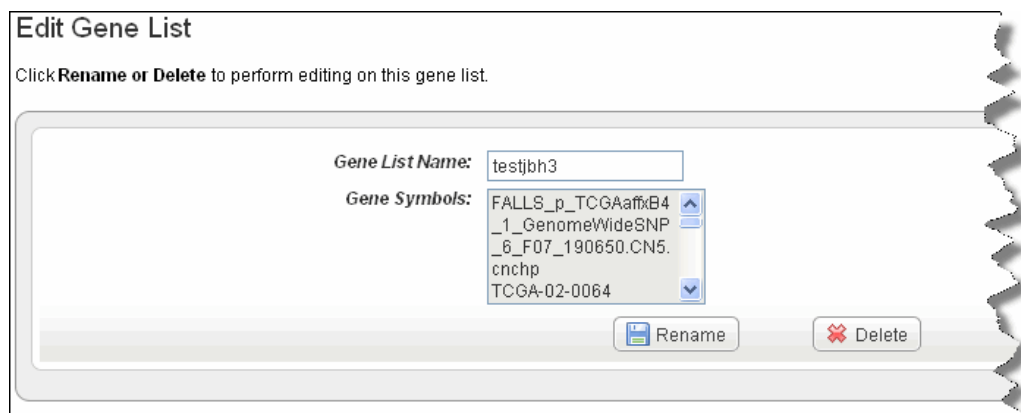


Figure 4.7 The Edit Gene List for reviewing, editing the name or deleting a gene list.

Editing a Gene List

To view a gene list in caIntegrator2, under Study Data in the left sidebar, click **Saved Lists > Global Lists** or **My Lists**. The system displays gene lists that have been saved for the open study ([Figure 4.8](#)).

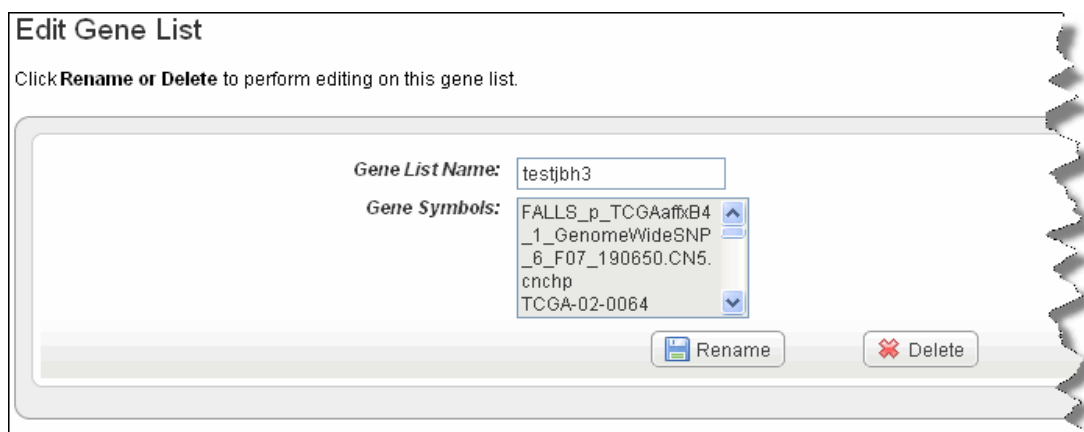


Figure 4.8 Edit Gene List allows you to edit gene lists for a study

You can initiate the following functions on this page:

- Click on any of the gene list names or the gene list icon (📄) to rerun the query from which the gene list was first created. In the query results, you can click on the gene icon (📄) to open the Cancer Genome Anatomy Project (CGAP) showing metadata for the gene.
- Click the edit icon (✏️) to open an Edit Gene List dialog box. On this page you can review the list of gene symbols included in the list.
- To rename the list in the **Gene List Name** text box, click the **Rename** button.
- To delete the study by clicking the **Delete** button.

See also [Creating a Gene List](#).

Expanding Imaging Data Results

In reviewing imaging search results, it is important to understand the hierarchy of submissions in NBIA. For more information, see [Relationship of Patient to Study to Series to Images](#) on page 66.

If you run a search before configuring column and sort display parameters, only the Subject Identifiers for the patients/images that meet the criteria and a column containing one check box per row display by default ([Figure 4.9](#)).

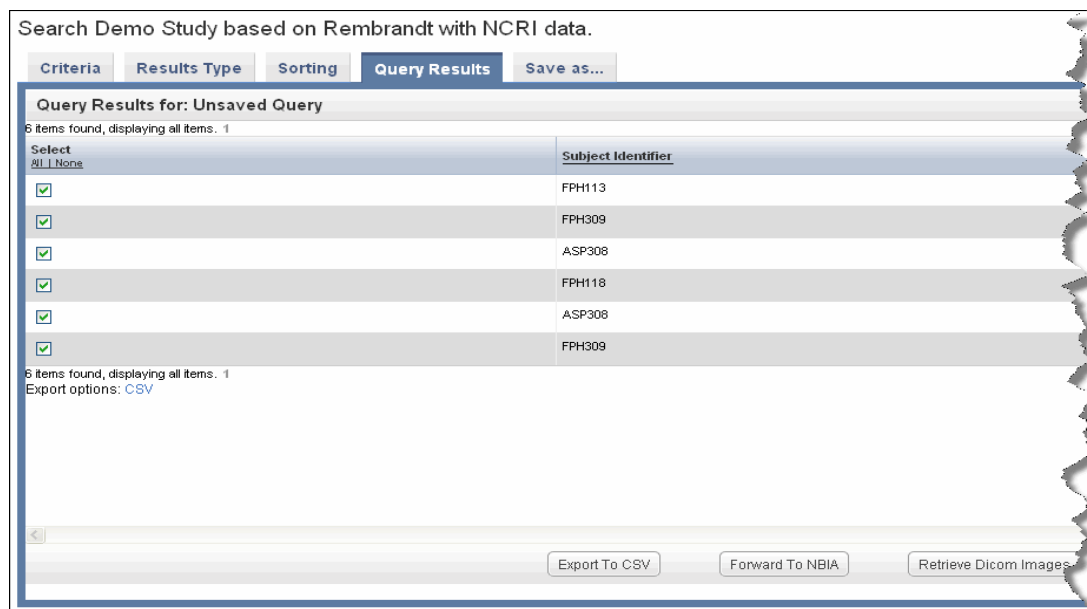


Figure 4.9 With imaging criteria only and no column definition, only Subject IDs display

If your annotation choice on the Columns page identifies annotations such as tumor size or tumor location, the search results display image series subsets that have those

annotations. The check boxes work in conjunction with buttons at the bottom of the results page (*Figure 4.10*).

Search Demo Study based on Rembrandt with NCRI data.

Criteria Results Type Sorting Query Results Save as...

Query Results for: Unsaved Query Results per Page: 20 Apply

6 items found, displaying all items. 1

Select	Subject Identifier	Image Series Identifier	Tumor Location
<input checked="" type="checkbox"/>	ASP308	2.16.124.113543.6003.121591217.13842.19801.1684612788 View in NBIA	Frontal
<input checked="" type="checkbox"/>	ASP308	2.16.124.113543.6003.2317586685.40219.20287.3012655789 View in NBIA	Frontal
<input checked="" type="checkbox"/>	FPH309	2.16.124.113543.6003.549598832.64081.17387.2785982861 View in NBIA	Frontal
<input checked="" type="checkbox"/>	FPH309	2.16.124.113543.6003.2205896078.6864.16978.1740991361 View in NBIA	Frontal
<input checked="" type="checkbox"/>	FPH113	2.16.124.113543.6003.2255697655.34510.18599.2966603150 View in NBIA	Frontal
<input checked="" type="checkbox"/>	FPH118	2.16.124.113543.6003.2241039616.45708.20383.2016653450 View in NBIA	Frontal

6 items found, displaying all items. 1
Export options: [CSV](#)

Figure 4.10 By expanding display parameters, you can view complete details for image search results

You can add more details for images by configuring image annotations on the Results Type tab. Annotations listed there are the column headers in the image series CSV file(s) that were uploaded to the study. Examples of image details include the following:

- All image details (name, size, etc.)
- The series to which the image belongs
- Image feature attributes
- The subject ID. Click the subject ID under Annotations on the Results Type tab to display this.

You can set display parameters for the results on the Columns and Sorting tabs. For more information, see *Results Type Tab* on page 51.

See also *caIntegrator2 and NBIA*, *Retrieving Dicom Images* and *Example of Retrieving Images*:

caIntegrator2 and NBIA

Images can be accessed in NBIA if you see buttons on the Search Results page. See the Imaging Note in *Results Type Tab* on page 51. You can click links on the Search Results tab to view or download image data.

- **View in NBIA** – This link corresponds to each Image Series listed in the results table. If you click the link, NBIA opens to the login page. After you log in, NBIA brings up the first image in the corresponding image series (*Figure 4.11*). You must log into NBIA to see the data. On the NBIA page that opens, you can opt to view the entire series containing this image, or you can display the image as a large JPEG-formatted image. You can also add the image to the NBIA basket.

For more information, see the NBIA online help or user's guide accessible from NBIA.



Figure 4.11 An example of displaying the first image in image series

- Forward to NBIA** – This button is linked to results you have selected by row. Click the button to open NBIA, where the image series you select are loaded in the NBIA image basket. In the event that the calIntegrator2 study was NOT configured with image annotation for an image series, calIntegrator2 sends NBIA a list of Study Instance UIDs, for which NBIA will add all corresponding image series to the basket. In the event that the calIntegrator2 study was configured with annotations for an image series, the system sends NBIA a list of Image Series IDs, for which NBIA adds all corresponding image series to the basket.

Retrieving Dicom Images

On the Imaging data Search Results page, you can click the **Retrieve DICOM Images** button which is linked to results you have selected by row. calIntegrator2 retrieves the corresponding image(s) from NBIA through the grid. NBIA organizes the download file by patient ID, StudyInstance UID, and ImageSeries UID, and compresses it into a zip file. When calIntegrator2 notifies you that the file is retrieved, the DICOM Retrieval page

indicates whether the retrieved files are Study Instance UIDs or Image Series UIDs (Figure 4.12). For more information, see the note below.



Figure 4.12 DICOM Retrieval result

Click the **Download DICOM** link to download and save the file. caIntegrator2 unzips the file and displays the list of images in the file. To open the DICOM images, you must have a DICOM image viewer application installed on your computer. For more information, see <http://dicom.online.fr/fr/download.htm>.

In the search results, not all of the patients in the data subset may be mapped to image series IDs. If you select a mixture of patients that have image annotations as indicated by an image series ID and patients that do not have image annotations (no image series ID), when you click the **Retrieve DICOM Images** button, NBIA retrieves the images for the entire *NBIA study instance UID* that includes the image seriesIDs you checked.

If on the Search Results tab you select only patients that have image annotations as indicated by an image series ID, when you click the **Retrieve DICOM Images** button, NBIA retrieves images for the *NBIA image series* that were matched in the search. If the results are a mixture, but you select one specific row with a valid image annotation, caIntegrator2 aggregates to the *image series*. If results are a mixture and you select multiple rows, caIntegrator2 aggregates to the NBIA study in which multiple image series you have selected in the search results are found.

If your query does not have image annotations and all check boxes are selected, results will go up to image series UID and gives all image series in it. Search results may ultimately depend on how the study was created. For example, if no image series display in query results, it means they were not mapped in the study. In that case, the results “move” up to Study Instance UIDs.

To best understand this, it is important to review the hierarchy of submissions in NBIA. For more information, see [Relationship of Patient to Study to Series to Images](#) on page 66.

Example of Retrieving Images:

If you are searching a study that has image data and image annotation(s) for at least one image series, you would follow these steps:

1. Open a study that has imaging data associated with it that points to the production NBIA server.

2. Make a query that will have image series or patients who are associated to Image Studies and select a few of those patients in the check box.
3. Click the **Retrieve Dicom Images** button.

Note that it aggregates to the image study.

4. Now go back to Results Type tab, select all image annotations and run the query again.
5. Select an image series type column and click the **Retrieve Dicom Images** button.

calIntegrator2 now aggregates to the Image Series that were selected and not the Image Study.

6. Select a row that doesn't have image series data, and a row that does, and push the button.

This should aggregate to the study for the rows selected.

7. Click **Forward to NBIA**. You should see the same types of aggregation for these tests.

When the image Study is in the checked boxes (regardless of image series being there or not), the system aggregates up to the Image Study level.

Relationship of Patient to Study to Series to Images

This flowchart illustrates the relationship of patient to study to series and lastly to images.

subject annotation trial > Patient (Subject) > Study > Series > Images

For example, the Study Instance UID is the set of images resulting from one patient office visit. When you upload a spreadsheet of an image series, the hierarchy of images in an image series might look like this:

Study Instance UID (one office visit):

Brain (image series)

- Brain image 1
- Brain image 2
- Brain image 3

Leg (image series)

- Leg image 1
- Leg image 2
- Leg image 3

You can add details for images by configuring image annotations on the Results Type tab. Annotations listed there are the column headers in the image series CSV file(s) that were uploaded to the study. Examples of image details include the following:

- All image details (name, size, etc.)
- The series that the image belongs to
- Image feature attributes
- The subject ID. Click the subject ID under Annotations on the Results Type tab to display this.

Exporting Data

You can choose to download tabular search results as a CSV file. Click the **Export .csv** link at the bottom of the page. You may need to scroll the page to see it. The file contains the annotations, columns and data sort configurations you specified in the search query.

Note: You will not see the Export option when genomic data displays as query results.

CHAPTER 5

ANALYZING STUDIES

This chapter describes how to use calIntegrator2 tools to analyze data in subject annotation or genomic studies that have been deployed in calIntegrator2.

Topics in this chapter include the following:

- [Data Analysis Overview](#) on this page
- [Creating Kaplan-Meier Plots](#) on page 70
- [Creating Gene Expression Plots](#) on page 77
- [Analyzing Data with GenePattern](#) on page 91

Data Analysis Overview

Once a study has been deployed, you can analyze the data using calIntegrator2 analysis tools.

You can verify that the study has “Deployed” status by selecting the study name in the My Studies dropdown selector. After selecting the study name, click **Home** in the left sidebar of the calIntegrator2 menu. A study summary should appear, including a status field. If the status is not deployed, or if the study summary does not appear, then the study is not deployed nor available for analysis.

If the study is ready for analysis, you will see an **Analysis Tools** menu in the left sidebar with the following options:

- **K-M Plot:** This tool analyzes subject annotation data, generating a Kaplan-Meier (K-M) plot based on survival data sets. See [Creating Kaplan-Meier Plots](#) on page 70.
- **Gene Expression Plot:** This tool analyzes annotation, subject annotation or genomic data based on gene expression values. See [Creating Gene Expression Plots](#) on page 77.

- **GenePattern:** This feature provides an express link to GenePattern where you can perform analyses on selected calIntegrator2 studies, or it enables you to perform several GenePattern analyses on the grid. See [Analyzing Data with GenePattern](#) on page 91 .

After defining or running the analysis on selected data sets, analysis results display on the same page, allowing you to review the analysis method parameters you defined.

Creating Kaplan-Meier Plots

The Kaplan-Meier method analyzes comparative groups of patients or samples. In calIntegrator2, the K-M method compares survival statistics among comparative groups. You can configure the survival data in the application. For example, you might identify a group of patients with smoking history and compare survival rates with a group of non-smoking patients, or compare the survival data for two groups of patients with a specific disease type, based on Karnofsky scores. You could compare groups of patients with varying gene expression levels. You can also identify data sets using the query feature in the application, saving the queries, then configuring the K-M to compare groups identified by the queries.

The key is to first identify subsets of patients or samples that meet criteria you want to establish, thus filtering the data you want to compare. Next, generate a K-M plot based on their survival probability as a function of time. Survival differences are analyzed by the log-rank test.

calIntegrator2 calculates the log-rank p-value for the data, indicating the significance of the difference in survival between any two groups of samples. The log rank p-value is calculated using the Mantel-Haenszel method. The p-values are recalculated every time a new plot is generated.

Note: To perform a K-M plot analysis, survival data must have been identified for the study you want to analyze. For more information, see [Defining Survival Values](#) on page 29.

K-M Plot for Annotations

The groups identified for this K-M plot generation are based on annotations.

1. Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page.
2. Under Analysis Tools on the left sidebar, select **K-M Plot**.
3. Select the **For Annotation** tab at the top of the page ([Figure 5.1](#)).

Kaplan-Meier Survival Plots

For Annotation | For Gene Expression | For Queries and Saved Lists

Annotation Based Kaplan-Meier Survival Plots

Annotation Group	Annotation	Values
1) Patient Groups:	Select Annotation Group	Select Annotation

Survival Value

2) Select Survival Measure: death is last contact date

Reset Create Plot

Figure 5.1 Fields for defining annotation data for a K-M plot

4. The groups to be compared in the K-M plot originate from one patient group. Varying data sets are based upon multiple values corresponding to the selected annotation. Define Patient Groups using these options:
 - **Annotation Type** – Select the annotation type that identifies the patient group. Selections are based on the data in the chosen study.
 - **Annotation** – Select an annotation. Fields are based on the annotation type you select. For example, if you choose **Subject**, then you could select **Gender** or **Radiation Type** or any field that would distinguish the patients into groups based upon their values.
 - **Values** – Using conventional selection techniques, select two or more values which will be the basis for the K-M plot. Permissible (available) values or “No Values” correspond to the selected annotation.
5. **Survival value** is the length of time the patient lived. calIntegrator2 displays valid survival values entered for this study. Select the survival measure which is the unit of measurement for the survival value to be used for the plot.
6. Click the **Create Plot** button.

calIntegrator2 generates the plot which then displays below the plot criteria (*Figure 5.2*).

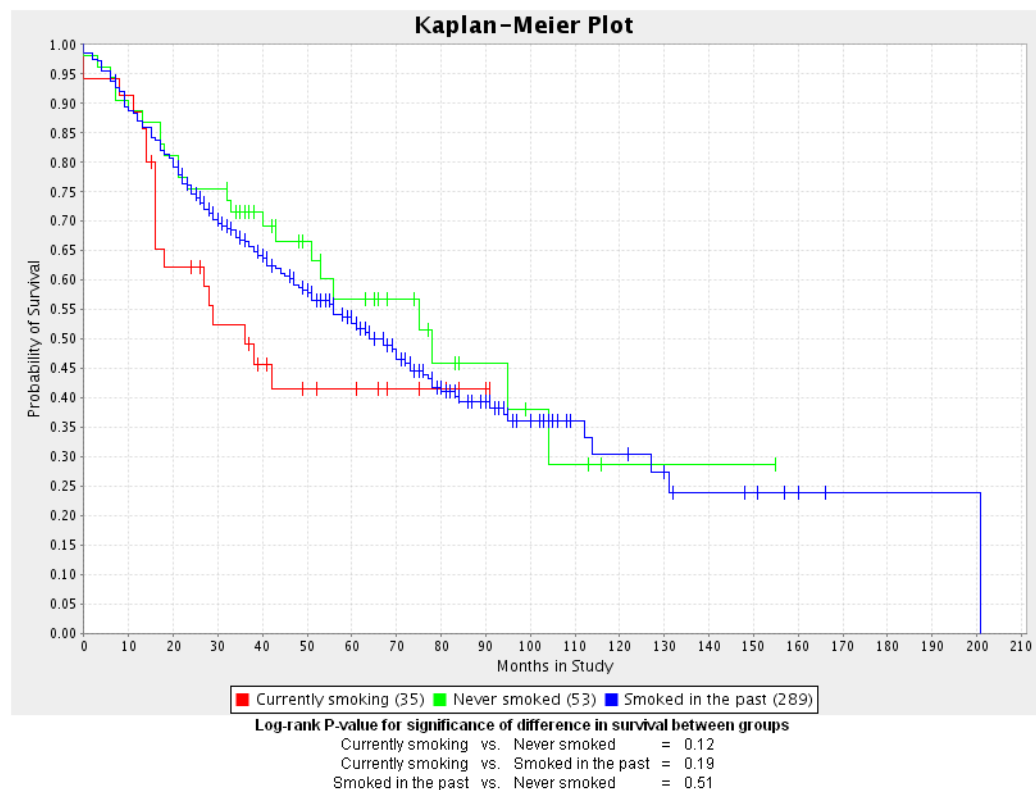


Figure 5.2 A K-M plot generated for groups based on annotations

- The number of subjects for each group appears embedded in the legend of the graph below the plot.

- calIntegrator2 generates a P-value for the selected groups; it displays at the bottom of the page. A low P-value generally has more significance than a high P-value.

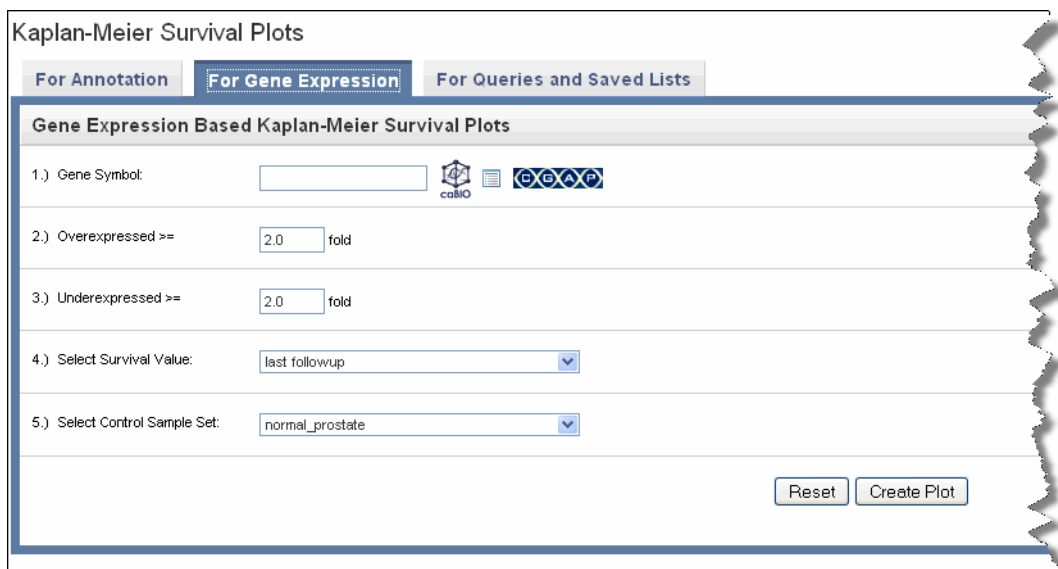
Note: For information regarding the P-value calculation, see [Creating Kaplan-Meier Plots](#) on page 70.

K-M Plot for Gene Expression

calIntegrator2 allows you to compare expression levels for one given gene in different representative groups. The relative expression level is referred to as “fold change”. Fold change is the ratio of the measured gene expression value in an experimental sample as determined by a reporter to a reference value calculated for that reporter against all control samples. The reference value is calculated by taking the mean of the \log_2 of the expression values for all control samples for the reporter in question. The \log_2 mean value (n) is then converted back to a comparable expression signal by returning 2 to the exponent n .

To create a K-M plot illustrating gene expression values, follow these steps:



- Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page. You must select a study with gene expression data.
- Under Analysis Tools on the left sidebar, select **K-M Plot**.
- Select the **For Gene Expression** tab ([Figure 5.3](#)).



Kaplan-Meier Survival Plots

For Annotation | **For Gene Expression** | For Queries and Saved Lists

Gene Expression Based Kaplan-Meier Survival Plots

1.) Gene Symbol:  

2.) Overexpressed >= fold

3.) Underexpressed >= fold


4.) Select Survival Value:

5.) Select Control Sample Set:

Figure 5.3 Fields for defining gene expression data for a K-M plot

- For **Gene Symbol**, enter one or more gene symbols in the text box or click the icons to locate genes in the following databases. If you enter more than one gene in the text box, separate the entries by commas.

calIntegrator2 provides three methods whereby you can obtain gene names for calculating a KM plot:

- **caBIO** – This link searches caBIO, then pulls identified genes into calIntegrator2 for analysis.
 - a. Click the **caBIO** icon ().
 - b. Enter **Search Terms**. Note that calIntegrator2 can perform a search on a partial HUGO symbol. For example, as search using *nicotin* would find matches with “nicotinic” and “nicotinamide”.
 - c. Select if you want to search in **Gene Keywords**, **Gene Symbols** or **Pathways** (from the drop-down list).
 - Selecting **Gene Keywords** searches only the Full Name field in caBIO.
 - Selecting **Gene Symbols** searches only the Unigene and HUGO gene symbols in caBIO.
 - Selecting **Pathways** searches only the pathway names in caBIO. Note that searching in Pathways is a two step process. First, the initial Pathway search produces search results which are pathways. Second, from the pathway search results screen, you must select pathways of interest, then click **Search Pathways for Genes** to obtain a list of genes related to the selected pathways.
 - d. Select the **Any** or **All** choice to determine how your search terms will be matched. **Any** finds any match for any search term you entered. **All** finds only results that match all of the search terms.
 - e. Choose the **Taxon** from the drop-down list and click **Search**. The search results display in the same dialog box (*Figure 5.4*).

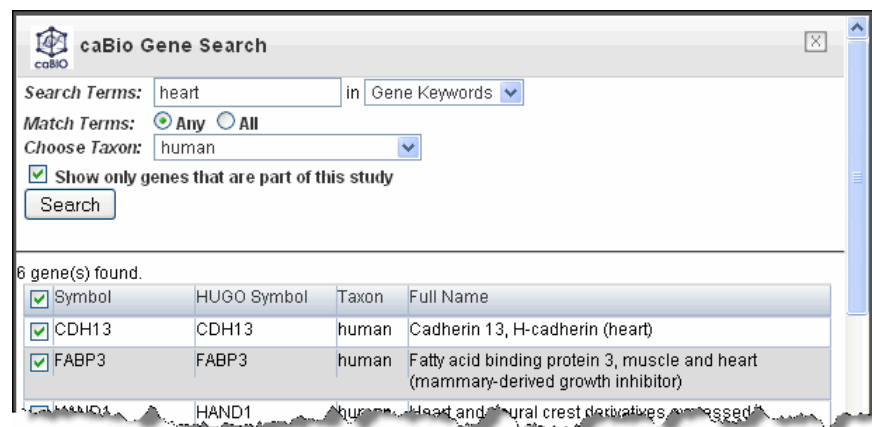


Figure 5.4 Example caBIO gene search criteria and results

- f. In the search results, use the check boxes to identify the genes whose symbols you want to use in the plot calculation.

- g. Click **Use Genes** at the bottom of the page. This pulls the checked genes into the For Gene Expression tab in a comma-separated list ([Figure 5.5](#)).

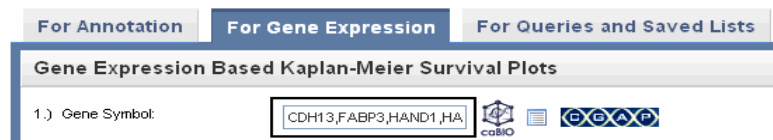



Figure 5.5 Genes pulled in from caBIO display on the selected tab

- **Gene List** – This link locates gene lists saved in calIntegrator2.
 - a. Click the Gene List icon () to open a small dialog that lists prior-saved gene lists in calIntegrator2. See [Creating a Gene List](#) on page 58.
 - b. In the drop-down menu, select a gene list. In the list that appears, use the check boxes to identify the genes whose symbols you want to use in the plot analysis.
 - c. Click **Use Genes** at the bottom of the dialog. This pulls the checked genes into the For Gene Expression tab.
- **CGAP** – Use this directory to identify genes. Before clicking this link you must enter gene symbols in the text box. This link does not pull anything into calIntegrator2 but does provide information about the gene(s) whose names you enter.
- 5. **Over-expressed/Under-expressed** – Define the over- and under-expression criteria, expressed in terms of fold-change. Fold change is the ratio of the measured gene expression value for an experimental sample to the expression value for the control sample.
- 6. **Survival value** – The length of time the patient lived. For **Survival Value**, select the survival measure which is the unit of measurement for the survival value to be used for the plot.
- 7. **Control Sample Sets** – One or more are created by the study manager when a study is deployed. Select the **Control Sample Set** you would like to use to calculate fold-change.

Note: If the study has more than one platform associated with it, the platform is inherently selected when you select the control set. Control sets are comprised of samples from only one platform.
- 8. Click the **Create Plot** button. calIntegrator2 generates the plot which then displays below the plot criteria ([Figure 5.6](#)).

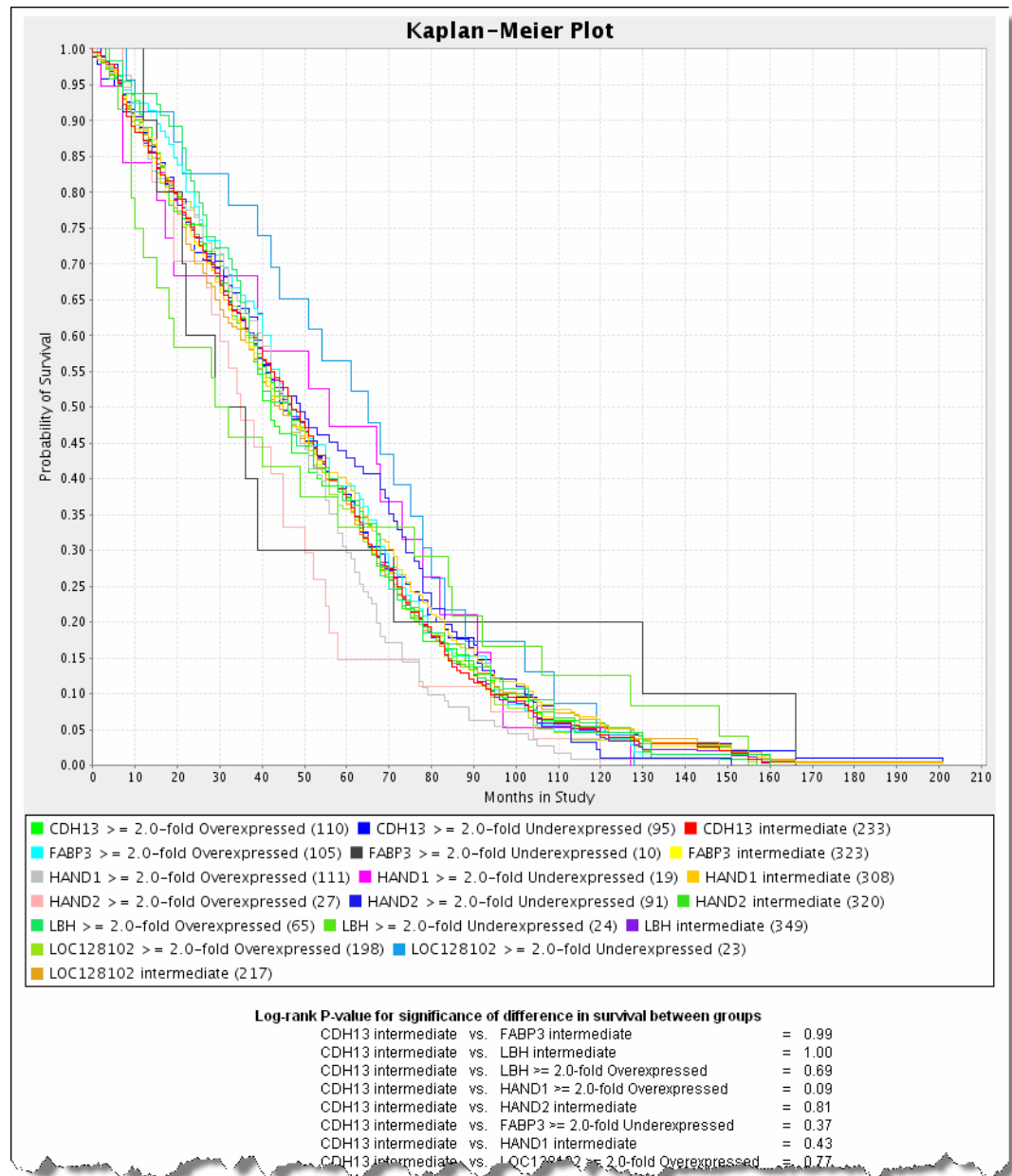


Figure 5.6 K-M plot generated from gene expression data.

- The gene symbol for each group represented in the data appears with its color correlation to the plot embedded in the legend of the graph below the plot. Three lines on the plot represent each gene symbol entered for the plot. Each line of the three represents a subgroup of people carrying the gene—one line for overexpressed values, one line for under expressed values and one line for intermediate values which represents gene values that are not up-regulated nor down-regulated.
- In queries that include a fold change criterion and that are configured to return genomic data, raw expression values are replaced with calculated fold change values.

- A P-value is also generated for the selected groups; it displays at the bottom of the page. A low P-value generally has more significance than a high P-value.

Note: For information regarding the P-value calculation, see [Creating Kaplan-Meier Plots](#) on page 70.

K-M Plot for Queries and Saved Lists

You can identify data sets using the query feature in the application. You can manipulate the queries to find the groups you want to compare, save the queries, then configure the K-M to compare the query groups. This is one method of limiting the data considered in the K-M plot calculation.

1. Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page. The queries you identify for the K-M plot must have been saved previously in calIntegrator2.
2. Under Analysis Tools on the left sidebar, select **K-M Plot**.
3. Select the **For Queries and Saved Lists** tab ([Figure 5.7](#)).

Kaplan-Meier Survival Plots

For Annotation For Gene Expression **For Queries and Saved Lists**

Kaplan-Meier Survival Plots based on Saved Queries and Saved Lists

1.) Select Saved Queries and Lists:

Available Queries and Lists

- [0]-JP - prostate genes over expressed 2X
- [0]-JP - prostate genes up or down 2X
- [0]-JP - test UPMC
- [0]-all genes - up reg 50X plus age 50+
- [0]-all genes - up reg 5X

Add >

< Remove

Selected Queries and Lists

2.) ☐ Exclusive Subjects (Subjects in upper Selected Queries or Lists are removed from subsequent Selected Queries or Lists)

3.) ☐ Add additional group containing all other subjects not found in selected queries and lists.

4.) Select Survival Value: last followup

Reset Create Plot

Figure 5.7 Fields for defining K-M plot parameters based on saved queries in calIntegrator2

4. **Queries** – Select **Queries** whose data you want to analyze from the **All Available Queries** panel and move them to the **Selected Queries** panel using the **Add >>** button.

Note: Genomic queries do not appear in the lists; they cannot be selected for this type of K-M plot.

5. **Exclusive Subject in Queries** – Check the box if you want to exclude any subjects that appear in both (or all) queries selected for the plot, thus eliminating overlap.
6. **Add Additional Group...all other subjects** – Check the box to create an additional group of all other subjects that are not in selected query groups.
7. **Survival value** – The length of time the patient lived. Select the survival measure which is the unit of measurement for the survival value to be used for the plot.

8. Click the **Create Plot** button. calIntegrator2 generates the plot which then displays below the plot criteria ([Figure 5.8](#)).

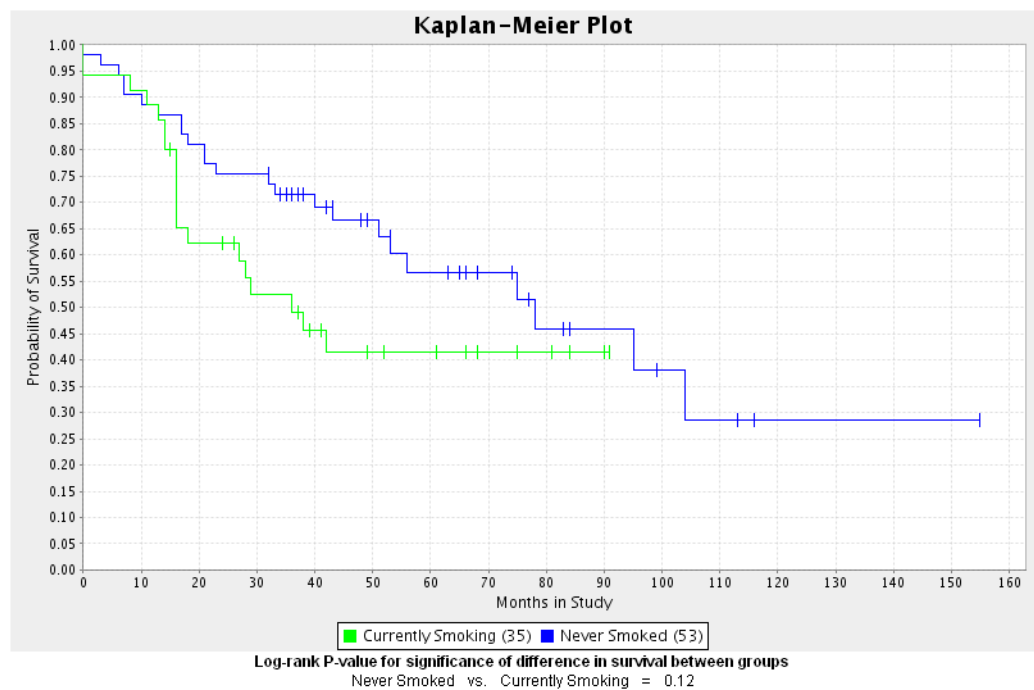


Figure 5.8 K-M Plot comparing statistics between subjects in two queries

- The number of subjects for each group appears embedded in the legend of the graph below the plot.
- A P-value is also generated for the selected groups; it displays at the bottom of the page. A low P-value generally has more significance than a high P-value.

Note: For information regarding the P-value calculation, see [Creating Kaplan-Meier Plots](#) on page 70.

Creating Gene Expression Plots

Gene expression plots compare signal values from reporters or genes. This statistical tool allows you to compare values for multiple genes at a time, but it does not require only two sets of data to be compared. It also allows you to compare expression levels for selected genes against expression levels for a set of control samples designated at the time of study definition.

calIntegrator2 provides three ways to generate meaningful gene expression plots, indicated by tabs on the page. The tabs are independent of each other and allow you to select the genes, reporters and sample groups to be analyzed on the plot.

- [Gene Expression Value Plot for Annotation](#) – You can locate genes in the caBIO directories or calIntegrator2 Gene Lists. You can learn more about the genes in the CGAP directory. You can define criteria for the plot using subject annotation and image annotations.

- **Gene Expression Value Plot for Genomic Queries** – You can select data based on saved genomic queries.
- **Gene Expression Value Plot for Annotation and Saved List Queries** – You can select data based on saved subject annotation queries. You can locate genes in the caBIO directories or calIntegrator2 Gene Lists.

See also [Understanding a Gene Expression Plot](#) on page 87.

Gene Expression Value Plot for Annotation

To generate a gene expression plot, follow these steps:

1. Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page. (You must select a study which has genomic data.)
2. Under Analysis Tools on the left sidebar, select **Gene Expression Plot**. This opens a page with three tabs
3. Select the **For Annotation** tab ([Figure 5.9](#)).

Gene Expression Value Plots

For Annotation | For Genomic Queries | For Annotation Queries and Saved Lists

Annotation Based Gene Expression Plots

1.) Gene Symbol(s) (comma separated list):

2.) Select Reporter Type: ☒ Reporter Id ☐ Gene

Annotation Group	Annotation	Values
Select Annotation Group	Select Annotation	

3.) Sample Groups:

4.) ☐ Add additional group containing all other subjects not found in selected queries.

5.) ☐ Add additional group containing all control samples for this study:

Figure 5.9 Gene expression value tab for configuring gene expression annotation value plot

4. **Gene Symbol** – Enter one or more gene symbols in the text box or click the icons to locate genes in the following databases. If you enter more than one gene in the text box, separate the entries by commas.

calIntegrator2 provides three methods whereby you can obtain gene names for calculating a gene expression plot:

- **caBIO** – This link searches caBIO, then pulls identified genes into calIntegrator2 for analysis.
 - a. Click the **caBIO** icon ().
 - b. Enter **Search Terms**. Note that calIntegrator2 can perform a search on a partial HUGO symbol. For example, as search using **nicotin** would find matches with “nicotinic” and “nicotinamide”.
 - c. Select if you want to search in **Gene Keywords**, **Gene Symbols** or **Pathways** (from the drop-down list).

- Selecting **Gene Keywords** searches only the Full Name field in caBIO.
 - Selecting **Gene Symbols** searches only the Unigene and HUGO gene symbols in caBIO.
 - Selecting **Pathways** searches only the pathway names in caBIO. Note that searching in Pathways is a two step process. First, the initial Pathway search produces search results which are pathways. Second, from the pathway search results screen, you must select pathways of interest, then click **Search Pathways for Genes** to obtain a list of genes related to the selected pathways.
- d. Select the **Any** or **All** choice to determine how your search terms will be matched. **Any** finds any match for any search term you entered. **All** finds only results that match all of the search terms.
- e. Choose the **Taxon** from the drop-down list and click **Search**. The search results display in the same dialog box ([Figure 5.4](#)).

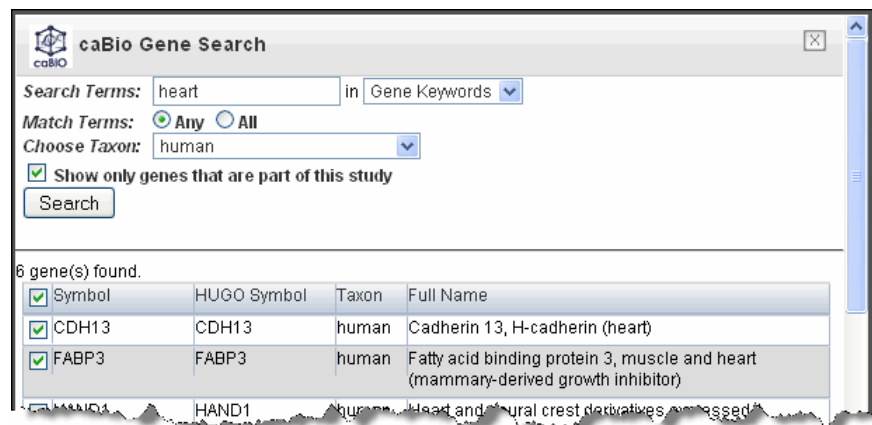


Figure 5.10 Example caBIO gene search results

- f. In the search results, use the check boxes to identify the genes whose symbols you want to use in the gene expression plot calculation.
- g. Click **Use Genes** at the bottom of the page. This pulls the checked genes into the For Annotation tab ([Figure 5.5](#)).

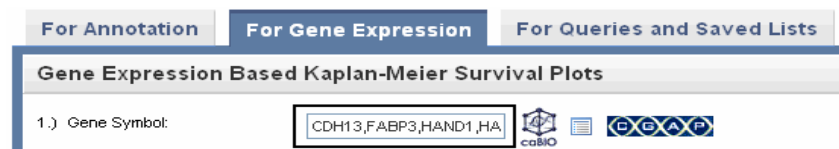



Figure 5.11 Genes pulled in from caBIO display on the tab

- **Gene List** – This link locates gene lists saved in caIntegrator2.
 - a. Click the Gene List icon () to open a small dialog that lists prior-saved gene lists in caIntegrator2. For more information, see [Creating a Gene List](#) on page 58.

- b. In the drop-down menu, select a gene list. In the list that appears, use the check boxes to identify the genes whose symbols you want to use in the plot analysis.
 - c. Click **Use Genes** at the bottom of the dialog. This pulls the checked genes into the For Annotation tab.
- **CGAP** – Use this directory to identify genes. Before clicking this link you must enter gene symbols in the text box. This link does not pull anything into calIntegrator2 but does provide information about the gene(s) whose names you enter.
5. **Reporter Type** – Select the radio button that describes the reporter type:
 - **Reporter ID** – Summarizes expression levels for all reporters you specify.
 - **Gene Name** – Summarizes expression levels at the gene level.
 - **Platform** – This field displays only if the study has multiple platforms. Select the appropriate platform for the plot. The platform you select determines the genes used for the plot.
6. **Sample Groups** – Choose among the following options:
 - **Annotation Type** – Select the annotation type. Selections are based on the data in the chosen study
 - **Annotation** – Select an annotation. Fields are based on the annotation type you select. For example, if you choose Subject, then you could select Gender or Radiation Type or any field that would distinguish the patients into groups based upon study values.
 - **Values** – Using conventional selection techniques, select one or more values which will be the basis for the plot. Permissible (available) values or “No Values” correspond to the selected annotation.
7. **Add Additional Group...** – Define as follows:
 - **...all other subjects** – Check the box to create an additional group of all other subjects that are not in selected query groups.
 - **...control group** – Check the box to display an additional group of control samples for this study. The control set should be composed of only samples which are mapped to subjects. See [Uploading Control Samples](#) on page 36.
8. Click the **Create Plot** button. calIntegrator2 generates the plot which then displays below the plot criteria in bar graph format ([Figure 5.6](#)).

Legends below the plot indicate the plot input. By default, the plot shows the mean of the data. [Figure 5.12](#) displays a plot with gene expression median calculation summaries.

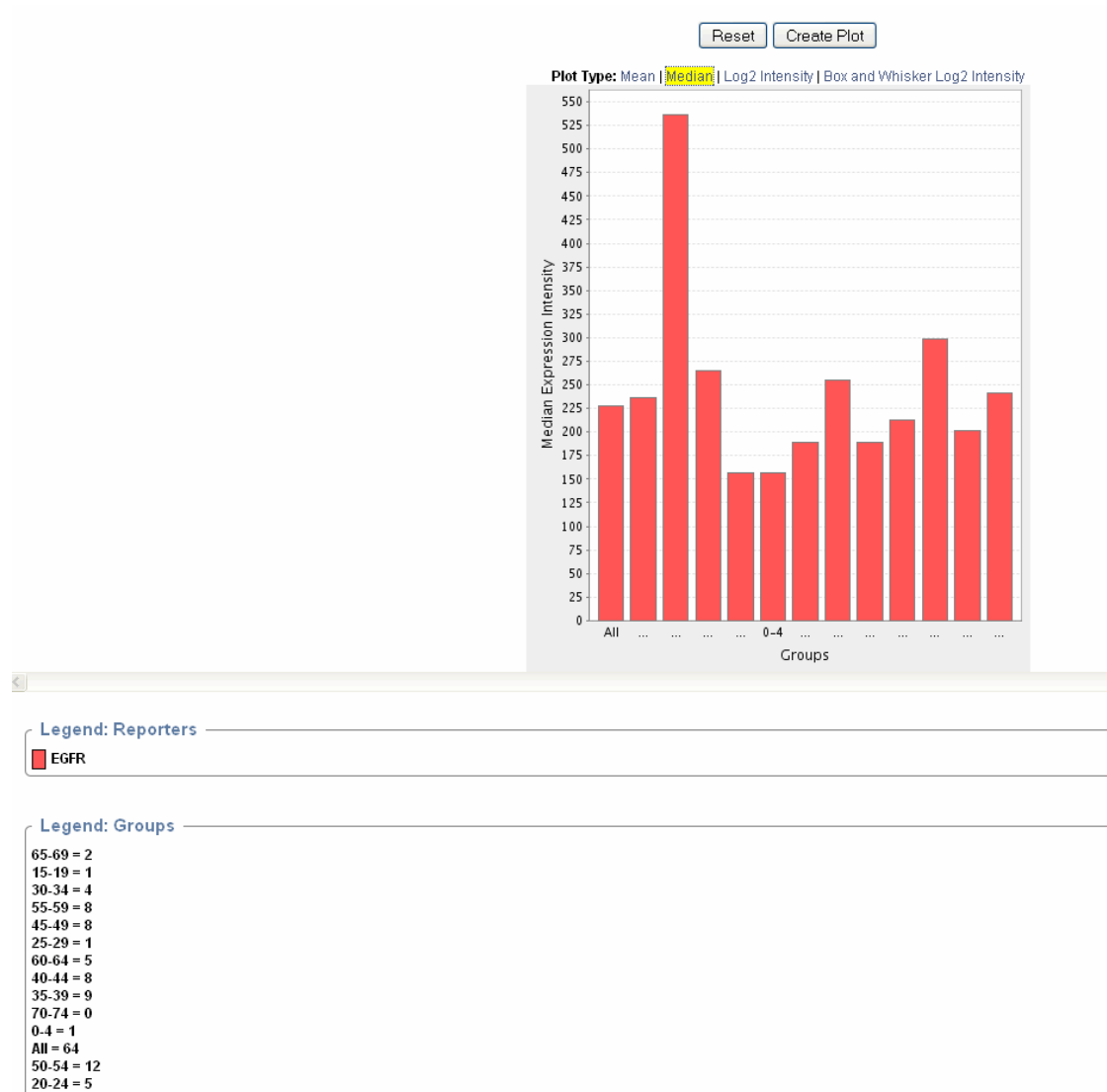


Figure 5.12 Gene expression plot based on selected annotations

- You can recalculate the data display by clicking the **Plot Type** above the graph. See [Understanding a Gene Expression Plot](#) on page 87.
- You can modify the plot parameters and click the **Reset** button to recalculate the plot.

Gene Expression Value Plot for Genomic Queries

Data to be analyzed on this tab must have been saved as a genomic query. For more information, see [Saving a Query](#) on page 54.

To generate a gene expression plot using a genomic query, follow these steps:

- Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page. (You must select a study which has genomic data.)

2. Under Analysis Tools on the left sidebar, select **Gene Expression Plot**.
3. Select the **For Genomic Queries** tab (*Figure 5.13*).

The screenshot shows the 'Gene Expression Value Plots' window with the 'For Genomic Queries' tab selected. The window has three tabs: 'For Annotation', 'For Genomic Queries' (active), and 'For Annotation Queries and Saved Lists'. Below the tabs is a section titled 'Genomic Query Based Gene Expression Plots'. It contains two main steps:

- 1.) Select a Genomic Query: A dropdown menu is open, showing the following options:
 - all genes - up reg 5X
 - JP - prostate genes overexpressed 2X
 - JP - prostate genes up or down 2X
 - all genes - up reg 50X plus age 50+
- 2.) Select Reporter Type: Two radio buttons are present: 'Reporter Id' (selected) and 'Gene'.

At the bottom right of the window are two buttons: 'Reset' and 'Create Plot'.

Figure 5.13 Gene expression value tab for configuring gene expression genomic queries plot

4. **Genomic Query** – Click on the genomic query upon which the plot is to be based.
5. **Reporter Type** – Select the radio button that describes the reporter type:
 - **Reporter ID** – Summarizes expression levels for all reporters you specify.
 - **Gene Name** – Summarizes expression levels at the gene level.

- Click the **Create Plot** button. calIntegrator2 generates the plot which then displays below the plot criteria. Legends below the plot indicate the plot input ([Figure 5.14](#)).



Figure 5.14 A gene expression plot (Mean) based on a genomic query.

- You can recalculate the data display by clicking the **Plot Type** above the graph. See [Understanding a Gene Expression Plot](#) on page 87.
- You can modify the plot parameters and click the **Reset** button to recalculate the plot.

Gene Expression Value Plot for Annotation and Saved List Queries

Data to be analyzed on this tab must have been saved as a subject annotation query, but it must have genomic data identified in the query. For more information, see [Adding/Editing Genomic Data](#) on page 31. For the genomic data, you must identify genes whose expression values are used to calculate the plot.

To generate a gene expression plot using an annotation query, follow these steps:



- Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page. You must select a study saved as a subject annotation study, but which has genomic data.
- Under Analysis Tools on the left sidebar, select **Gene Expression Plot**.

3. Select the **For Annotation Queries and Saved Lists** tab (Figure 5.15).

Gene Expression Value Plots

For Annotation For Genomic Queries **For Annotation Queries and Saved Lists**

Gene Expression Plots based on Saved Queries and Saved Lists

1.) Gene Symbol(s) (comma separated list):  

2.) Select Reporter Type: ☒ Reporter Id ☐ Gene

3.) Select Saved Queries and Lists:

Available Queries and Lists

Selected Ou

4.) ☐ Exclusive Subjects (Subjects in upper Selected Queries or Lists are removed from subsequent Selected Queries or Lists)


5.) ☐ Add additional group containing all other subjects not found in selected queries and lists.

6.) ☐ Add additional group containing all control samples for this study:

Figure 5.15 Gene expression value tab for configuring gene expression annotation queries plot

4. **Gene Symbol** – Enter one or more gene symbols in the text box or click the icons to locate genes in the following databases. If you enter more than one gene in the text box, separate the entries by commas.

calIntegrator2 provides three methods whereby you can obtain gene names for calculating a gene expression plot:

- **caBIO** – This link searches caBIO, then pulls identified genes into calIntegrator2 for analysis.
 - a. Click the caBIO icon ().
 - b. Enter **Search Terms**. Note that calIntegrator2 can perform a search on a partial HUGO symbol. For example, as search using **nicotin** would find matches with “nicotinic” and “nicotinamide”.
 - c. Select if you want to search in **Gene Keywords**, **Gene Symbols** or **Pathways** (from the drop-down list).
 - Selecting **Gene Keywords** searches only the Full Name field in caBIO.
 - Selecting **Gene Symbols** searches only the Unigene and HUGO gene symbols in caBIO.
 - Selecting **Pathways** searches only the pathway names in caBIO. Note that searching in Pathways is a two step process. First, the initial Pathway search produces search results which are pathways. Second, from the pathway search results screen, you must select pathways of interest, then click **Search Pathways for Genes** to obtain a list of genes related to the selected pathways.

- d. Select the **Any** or **All** choice to determine how your search terms will be matched. **Any** finds any match for any search term you entered. **All** finds only results that match all of the search terms.
- e. Choose the **Taxon** from the drop-down list and click **Search**. The search results display in the same dialog box ([Figure 5.4](#)).

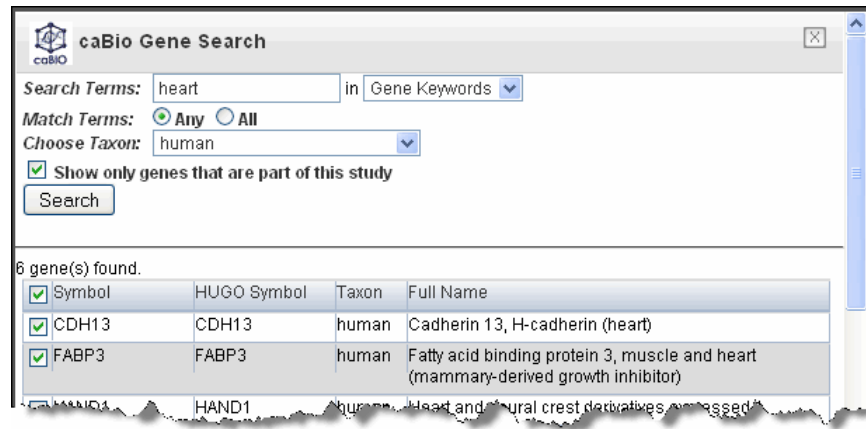


Figure 5.16 Example caBIO gene search results

- f. In the search results, use the check boxes to identify the genes whose symbols you want to use in the plot calculation.
- g. Click **Use Genes** at the bottom of the page. This pulls the checked genes into the tab ([Figure 5.5](#)).

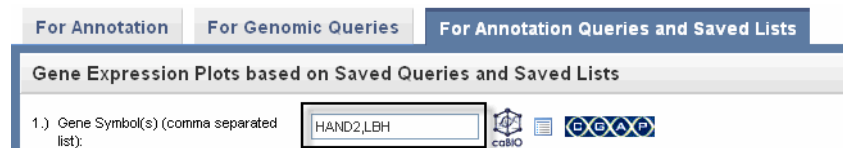



Figure 5.17 Genes pulled in from caBIO display on the tab

- **Gene List** – This link locates gene lists saved in calIntegrator2.
 - a. Click the Gene List icon () to open a small dialog that lists prior-saved gene lists in calIntegrator2. For more information, see [Creating a Gene List](#) on page 58.
 - b. In the drop-down menu, select a gene list. In the list that appears, use the check boxes to identify the genes whose symbols you want to use in the gene expression analysis.
 - c. Click **Use Genes** at the bottom of the dialog. This pulls the checked genes into the tab.
 - **CGAP** – Use this directory to identify genes. Before clicking this link you must enter gene symbols in the text box. This link does not pull anything into calIntegrator2 but does provide information about the gene(s) whose names you enter.
5. For **Reporter Type**, select the radio button that describes the reporter type:

- **Reporter ID** – Summarizes expression levels for all reporters you specify.
 - **Gene Name** – Summarizes expression levels at the gene level.
 - **Platform** – This field displays only if the study has multiple platforms. Select the appropriate platform for the plot. The platform you select determines the genes used for the plot.
6. For **Saved Queries**, choose among the available saved queries and lists. Build your selections in the right panel by using the **Add >** and **Remove <** buttons.

Note: The [SL] and [Q] prefixes to list names indicate “Subject Lists” or “Saved Queries”. A “G” in the prefix indicates the list is Global. For more information, see [Creating a Gene List](#) on page 58.
 7. Check the **Exclusive Subjects...** option to remove subjects in your queries and lists selection from queries or lists you use subsequently for analysis, using them exclusively for the current analysis.
 8. For the **Add Additional Group...** options, define as follows:
 - **...all other subjects** – Check the box to create an additional group of all other subjects that are not in selected query groups.
 - **...control group** – Check the box to display an additional group of control samples for this study. The control set should be composed of only samples which are mapped to subjects. See [Uploading Control Samples](#) on page 36.
 9. Click the **Create Plot** button. caIntegrator2 generates the plot which then displays below the plot criteria in bar graph format ([Figure 5.6](#)).

By default, calIntegrator2 displays the mean of the data below the plot criteria. Legends below the plot indicate the plot input.

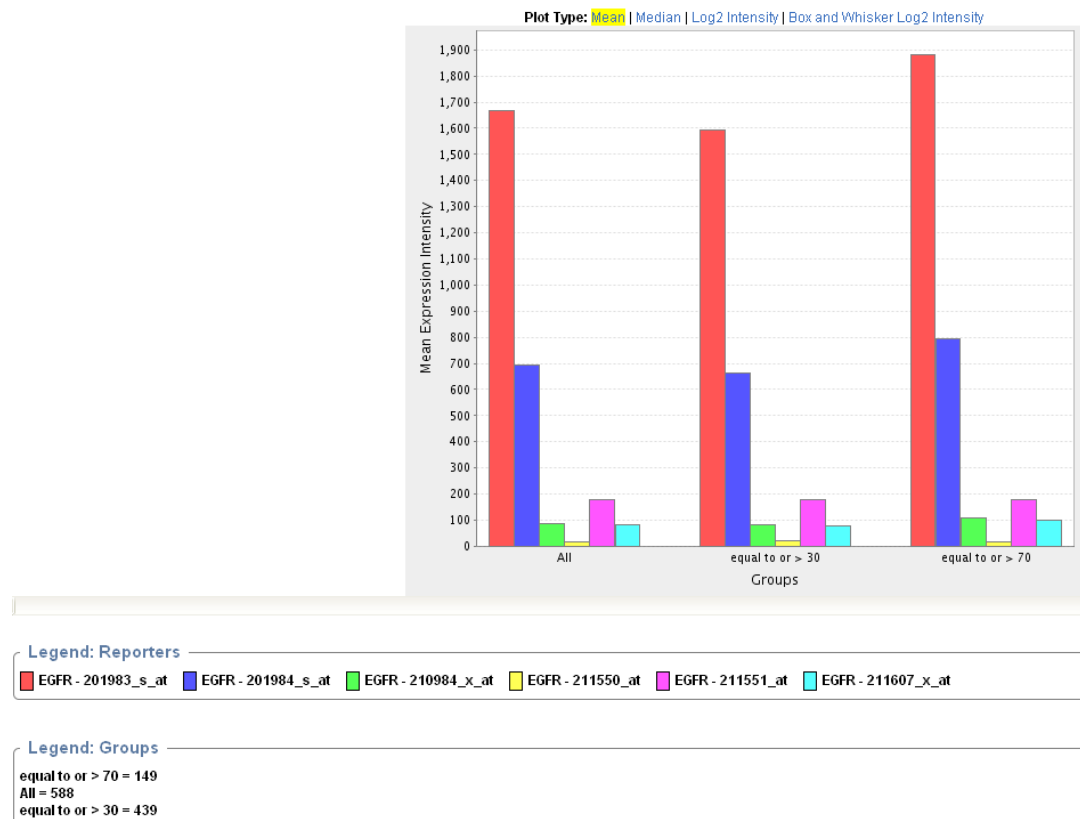


Figure 5.18 Gene expression plot based on annotation queries gene expression values

- You can recalculate the data display by clicking the **Plot Type** above the graph. See [Understanding a Gene Expression Plot](#) on page 87.
- You can modify the plot parameters and click the **Reset** button to recalculate the plot.

Understanding a Gene Expression Plot

Above the plot, you can select various plot types. When you do so, the plot is recalculated. Although all of the plots in this section appear similar, note the differences in calculation results and legends between the Y axis on each of the plots.

When you perform a Gene Expression simple search, by default the **Mean** Gene Expression Plot ([Figure 5.19](#)) appears.

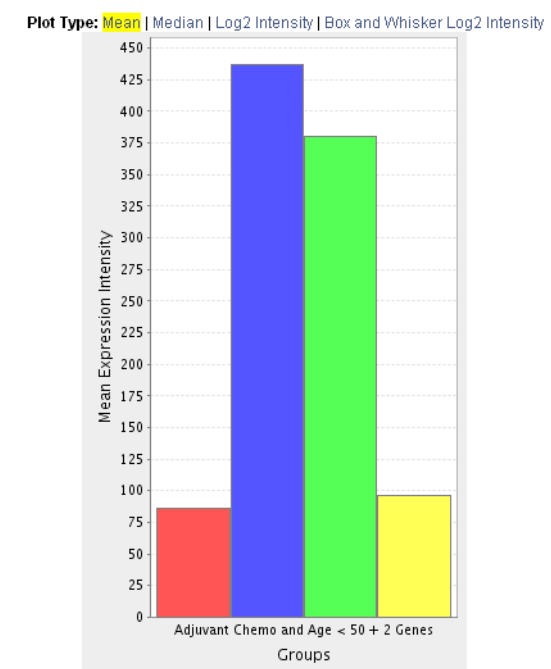


Figure 5.19 Gene expression plot calculating the mean

The **Mean** Gene Expression Plot ([Figure 5.19](#)) displays mean expression intensity (Geometric mean) versus Groups.

The **Median** Gene Expression Plot ([Figure 5.20](#)) displays the median expression intensity versus Groups.

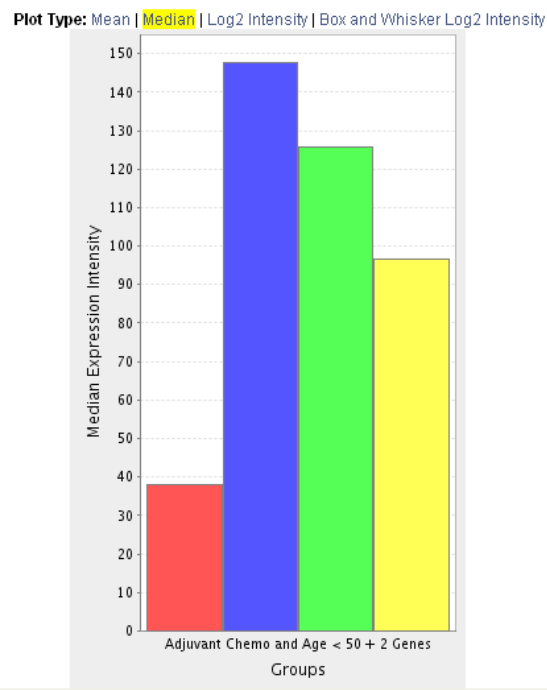


Figure 5.20 Gene expression plot calculating the median

The **Log2 Intensity** Gene Expression Plot ([Figure 5.21](#)) displays average expression intensities for the gene of interest based on Affymetrix GeneChip arrays (U133 Plus 2.0 arrays).

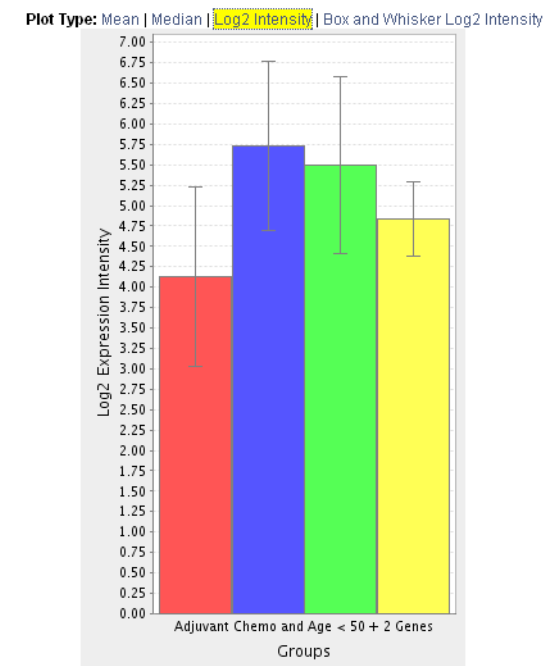


Figure 5.21 Gene expression plot displaying log2 intensity values

The box and whisker log2 expression intensity plot displays a box plot ([Figure 5.22](#), [Figure 5.23](#)). Example uses of box and whisker plots include the following:

- Indicate whether a distribution is skewed and whether there are potential unusual observations (outliers) in the data set.
- Perform a large number of observations.
- Compare two or more data sets.
- Compare distributions because the center, spread, and overall range are immediately apparent.

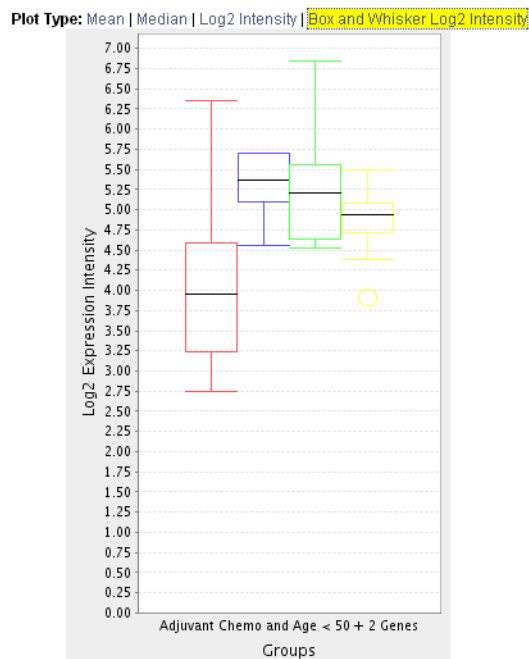


Figure 5.22 Box and whisker plot based on the same data set as represented in [Figure 5.19](#), [Figure 5.20](#), [Figure 5.21](#)

In descriptive statistics, a box plot or boxplot, also known as a box-and-whisker diagram or plot, is a convenient way of graphically depicting groups of numerical data through their five-number summaries (the smallest observation excluding outliers, lower quartile [Q1], median [Q2], upper quartile [Q3], and largest observation excluding outliers).

The box is defined by Q1 and Q3 with a line in the middle for Q2. The interquartile range, or IQR, is defined as Q3-Q1. The lines above and below the box, or 'whiskers', are at the largest and smallest non-outliers. Outliers are defined as values that are

more than $1.5 \times \text{IQR}$ greater than Q3 and less than $1.5 \times \text{IQR}$ than Q1. Outliers, if present, are shown as open circles (*Figure 5.23*).

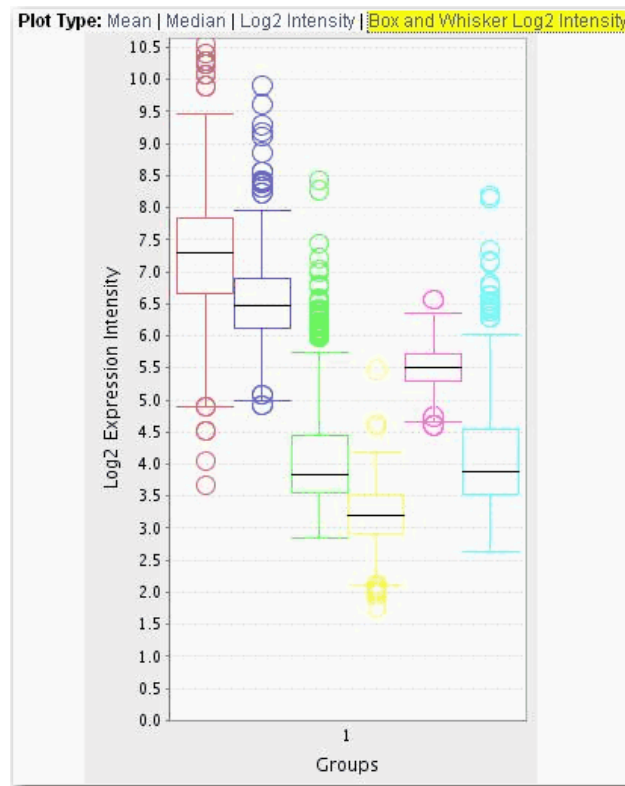


Figure 5.23 Box and whisker plot showing outliers

Boxplots can be useful to display differences between populations without making any assumptions of the underlying statistical distribution: they are non-parametric. The spacings between the different parts of the box help indicate the degree of dispersion (spread) and skewness in the data.

Analyzing Data with GenePattern

GenePattern is an application developed at the Broad Institute that enables researchers to access various methods to analyze genomic data. caIntegrator2 provides an express link to GenePattern where you can analyze data in any caIntegrator2 study.

Information is included in this section for connecting to GenePattern from caIntegrator2. Specifics for launching GenePattern tools from caIntegrator2 are included as well, but you may want to refer to additional GenePattern documentation, available at this website: http://www.broadinstitute.org/cancer/software/genepattern/tutorial/gp_concepts.html.

You have two options for using GenePattern from caIntegrator2:

- Option 1 – Use the web-interface of any available GenePattern instances.
 - a. To use the public instance from Broad, first register for an account at <http://genepattern.broadinstitute.org/gp/pages/login.jsf>.

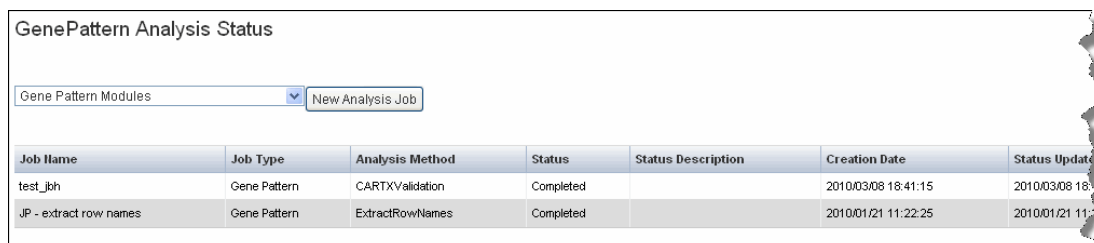
In calIntegrator2, enter the URL for connecting: <http://genepattern.broadinstitute.org/gp/services/Analysis>, then enter your user ID and password.

- Option 2 – Use GenePattern on the grid.

The GenePattern feature in calIntegrator2 currently supports three analyses on the grid: Comparative Marker Selection (CMS), Principal Component Analysis (PCA) and GISTIC-supported analysis.

Tip: If you are using the web interface to access GenePattern (option #1 listed above), then you can run other GenePattern tools in addition to CMS, PCA and GISTIC.

1. Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page.
2. Click **GenePattern Analysis** in the left sidebar of calIntegrator2. This opens the GenePattern Analysis Status page (*Figure 5.24*).



Job Name	Job Type	Analysis Method	Status	Status Description	Creation Date	Status Update
test_job	Gene Pattern	CARTXValidation	Completed		2010/03/08 18:41:15	2010/03/08 18:41:15
JP - extract row names	Gene Pattern	ExtractRowNames	Completed		2010/01/21 11:22:25	2010/01/21 11:22:25

Figure 5.24 GenePattern Analysis Status page

3. Select from the drop-down list the type of GenePattern analysis you want to run on the data.
 - **GenePattern Modules** – This option launches a session within GenePattern from which you can launch analyses. See [GenePattern Modules](#) on page 93.
 - **Comparative Marker Selection (Grid Service)**. This option enables you to run this GenePattern analysis on the grid. See [Comparative Marker Selection \(CMS\) Analysis](#) on page 95.
 - **Principal Component Analysis (Grid Service)**. This option enables you to run this GenePattern analysis on the grid. See [Principal Component Analysis \(PCA\)](#) on page 98.
 - **GISTIC (Grid Service)**. This option enables you to run this GenePattern analysis on the grid. See [GISTIC-Supported Analysis](#) on page 101.
4. Click the **New Analysis Job** button to open a corresponding page where you can configure the analysis parameters.

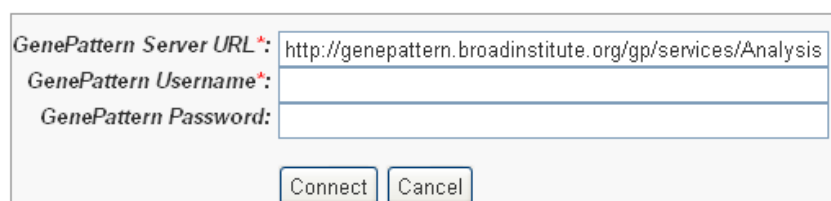
GenePattern Modules

Note: To launch the analyses described in this section, you must have a registered GenePattern account. For more information, see <http://genepattern.broadinstitute.org/gp/pages/login.jsf>.

To configure the link for accessing GenePattern from calIntegrator2, open the appropriate page as described in *Analyzing Data with GenePattern* on page 91.

1. Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page.
2. Click **GenePattern Analysis** in the left sidebar of calIntegrator2. This opens the GenePattern Analysis Status page.
3. Make sure **GenePattern Modules** is selected in the drop down list. Click **New Analysis Job**.
4. In the GenePattern Analysis dialog box (*Figure 5.25*), specify connection information, described *Table 5.1* and click **Connect**.

GenePattern Analysis



GenePattern Server URL*:

GenePattern Username*:

GenePattern Password:

Figure 5.25 Dialog box for configuring the link to GenePattern

Fields	Description
Server URL	Enter any GenePattern publicly available URL, such as http://genepattern.broadinstitute.org/gp/services/Analysis .
GenePattern Username	Enter your GenePattern user name.
GenePattern Password	Enter your GenePattern password.

Table 5.1 Fields for selecting GenePattern configurations

5. After logging in with the GenePattern profile, the dialog box expands to include fields for defining your GenePattern analysis..

GenePattern Analysis

Figure 5.26 GenePattern module options

6. Enter information for the following fields. Fields with a red asterisk are required:

- a. **Job Name*** – Enter a unique name for the analysis
- b. **Analysis Method** – Select any method from the drop down list. Click Analysis Method Documentation for descriptions of the different analysis methods.
- c. **Data*** – All genomic data is selected by default. Select from the list any list that has been created for this study.
- d. **cls*** – Select any annotation field

The CLS file format defines phenotype (class or template) labels and associates each sample in the expression data with a label. It uses spaces or tabs to separate the fields. The CLS file format differs somewhat depending on whether you are defining categorical or continuous phenotypes:

- Categorical labels define discrete phenotypes; for example, normal vs tumor).
- Continuous phenotypes are used for time series experiments or to define the profile of a gene of interest (gene neighbors).

Note: Most GenePattern modules are intended for use with categorical phenotypes. Therefore, unless the module documentation explicitly states otherwise, a CLS file should define categorical labels.

- e. **prediction.results.file** – Enter the name of this file which is part of the output from a GenePattern module.
7. Click **Perform Analysis**. Based on the analysis method you select, you may be asked to add more information for the analysis. For more information, refer to

the GenePattern Help site: <http://genepattern.broadinstitute.org/gp/getTaskDocCatalog.jsp>

Once the analysis is launched, calIntegrator2 returns to the GenePattern Analysis Status page where you can monitor the status of your current study which is listed in the Analysis Method column as well as view information about other GP analyses that have been run on this study.

Job Name	Job Type	Analysis Method	Status	Status Description	Creation Date	Status Update Date	Action
test job	Gene Pattern	GctToPcl	Processing Locally		2010/03/10 17:29:38	2010/03/10 17:29:38	
jp - test web services	Gene Pattern	ExtractRowNames	Completed		2010/02/25 17:16:52	2010/02/25 17:21:15	Delete View Results 190676
jp - test	Gene Pattern - Grid	Comparative Marker Selection	Error Connecting		2010/02/25 17:10:45	2010/02/25 17:14:05	Delete Download Input
JP - test CMS	Gene Pattern - Grid	Comparative Marker Selection	Error Connecting		2010/02/16 13:09:31	2010/02/16 13:11:24	Delete Download Input
JP - test preprocess	Gene Pattern	PreprocessDataset	Completed		2010/02/16 13:03:44	2010/02/16 13:07:04	Delete View Results 187493

Figure 5.27 GenePattern Analysis Status page displays a list of GenePattern analysis performed on the current study

If you choose to access GenePattern in this way, you can continue to use GenePattern tools from within that application. See GenePattern user documentation for more information.

Tip: If you run these analyses within GenePattern itself, you may be able to view results in the GenePattern visualization module. Click **View Results** on the row where the results are listed. If you run them on the grid from calIntegrator2, your results will be available only in spreadsheet and XML format.

You can run GenePattern analyses for Comparative Marker Selection, Principal Component Analysis and GISTIC-based analysis on the grid if you choose.

Comparative Marker Selection (CMS) Analysis

The Comparative Marker Selection (CMS) module implements several methods to look for expression values that correlate with the differences between classes of samples. Given two classes of samples, CMS finds expression values that correlate with the difference between those two classes. If there are more than two classes, CMS can perform one-vs-all or all-pairs comparisons, depending on which option is chosen.

For more information, see the GenePattern website: http://www.broadinstitute.org/cgi-bin/cancer/software/genepattern/modules/gp_modules.cgi.

To perform a CMS analysis, follow these steps:

1. Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page. You must select a study saved as a subject annotation study, but which has genomic data.
2. Click **GenePattern Analysis** in the left sidebar of calIntegrator2. This opens the GenePattern Analysis Status page.

3. In the GenePattern Analysis Status page, select **Comparative Marker Selection (Grid Service)** from the drop down list and click **New Analysis Job**. This opens the Comparative Marker Selection Analysis page ([Figure 5.28](#)).

Comparative Marker Selection Analysis

Job Name*:

Preprocess Server*:

Comparative Server*:

Clinical Queries*: Must select two clinical queries, which are used to group the samples into two separate classifications to run against ComparativeMarkerSelection. The queries selected here have been previously saved by the user. Selected queries will result in the processing of only those samples which are mapped to patients in the saved query result.

All Available Queries:

Selected Queries:

Filter flag: ☐

Preprocessing Flag*:

Min Change*:

Min Delta*:

Threshold*:

Ceiling*:

Max Sigma Binning*:

Probability Threshold*:

Num Exclude*:

Log Base Two: ☐

Number Of Columns Above Threshold*:

Test Direction*:

Test Statistic*:

Min Std*:

Number Of Permutations*:

Complete: ☐

Balanced: ☐

Random Seed*:

Smooth Pvalues: ☐

Phenotype Test*:

Figure 5.28 Comparative Marker Selection analysis parameters

4. Select or define CMS analysis parameters, described in [Table 5.2](#). An asterisk indicates required fields. The default settings are valid; they should provide valid results.

CMS Parameter	Description
Job Name*	Assign a unique name to the analysis you are configuring.
Preprocess Server*	A server which hosts the grid-enabled data GenePattern PreProcess Dataset module. Select one from the list and caIntegrator2 will use the selected server for this portion of the processing.
Comparative Server*	A server which hosts the grid-enabled data GenePattern Comparative Marker Selection module. Select one from the list and caIntegrator2 will use the selected server for this portion of the processing.

Table 5.2 Comparative Marker Selection analysis options

CMS Parameter	Description
Annotation Queries and Lists*	All subject annotation queries and gene lists with appropriate data for the analysis are listed. Select and move two or more queries from the All Available Queries panel to the Selected Queries panel using the Add > and Remove < buttons. Note: The [SL] and [Q] prefixes to list names indicate “Subject Lists” or “Saved Queries”. A “G” in the prefix indicates the list is Global. For more information, see Creating a Gene List on page 58.
Filter Flag	Variation filter and thresholding flag
Preprocessing Flag*	Discretization and normalization flag
Min Change*	Minimum fold change for filter
Min Delta*	Minimum delta for filter
Threshold*	Value for threshold
Ceiling*	Value for ceiling
Max Sigma Binning*	Maximum sigma for binning
Probability Threshold*	Value for uniform probability threshold filter
Num Exclude*	Number of experiments to exclude (max & min) before applying variation filter
Log Base Two	Whether to take the log base two after thresholding; default setting is “Yes”.
Number of Columns Above Threshold*	Remove row if n columns are not \geq than the given threshold In other words, the module can remove rows in which the given number of columns does not contain a value greater or equal to a user defined threshold.
Test Direction*	The test to perform (up-regulated for class0; up-regulated for class1, two sided). By default, Comparative Marker Selection performs the two-sided test.
Test Statistic*	Select the statistic to use.
Min Std*	The minimum standard deviation if test statistic includes the min std option. Used only if test statistic includes the min std option.
Number of Permutations*	The number of permutations to perform. (Use 0 to calculate asymptotic P-values.) The number of permutations you specify depends on the number of hypotheses being tested and the significance level that you want to achieve (3). The greater the number of permutations, the more accurate the P-value. Complete – Perform all possible permutations. By default, complete is set to No and Number of Permutations determines the number of permutations performed. If you have a small number of samples, you might want to perform all possible permutations. Balanced – Perform balanced permutations
Random Seed*	The seed for the random number generator.

Table 5.2 Comparative Marker Selection analysis options

CMS Parameter	Description
Smooth P-values	Whether to smooth P-values by using the Laplace's Rule of Succession. By default, Smooth P-values is set to Yes , which means P-values are always less than 1.0 and greater than 0.0.
Phenotype Test*	Tests to perform when class membership has more than 2 classes: one versus-all, all pairs. Note: The P-values obtained from the one-versus-all comparison are not fully corrected for multiple hypothesis testing.

Table 5.2 Comparative Marker Selection analysis options

- When you have completed the form, click **Perform Analysis**.

calIntegrator2 takes you to the JobStatus/Launch page where you will see the job and its status in the Status column of the list (*Figure 5.29*).

GenePattern Analysis Status (draft)

Gene Pattern Modules New Analysis Job

Job Name	Job Type	Status	Creation Date	Status Update Date
Well-diff vs adjuvant chemo	Comparative Marker Selection	Processing Locally	2009/08/14 11:48:35	2009/08/14 11:48:35
Filter out non-interesting genes	Gene Pattern	Completed - View 122444	2009/08/14 10:16:29	2009/08/14 10:19:47

Figure 5.29 The progress of a GenePattern analysis that has been launched displays in the status column of page

- When the job is complete, the system displays a completion date on the GenePattern Analysis status page. Click the **Download** link. This downloads zipped result files to your local work station. The number of files and their file type will vary according to the processing. The results format is compatible with GenePattern visualizers and can be uploaded within GenePattern.

Principal Component Analysis (PCA)

Principal Component Analysis is typically used to transform a collection of correlated variables into a smaller number of uncorrelated variables, or components. Those components are typically sorted so that the first one captures most of the underlying variability and each succeeding component captures as much of the remaining variability as possible.

You can configure GenePattern grid parameters for preprocessing the dataset in addition to PCA module parameters. For more information, see the GenePattern website: http://www.broadinstitute.org/cgi-bin/cancer/software/genepattern/modules/gp_modules.cgi.

To perform a PCA analysis, follow these steps:

- Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page. You must select a study with gene expression data.
- Click **GenePattern Analysis** in the left sidebar of calIntegrator2. This opens the GenePattern Analysis Status page.

3. In the GenePattern Analysis Status page, select **Principal Component Analysis (Grid Service)** from the drop down list and click **New Analysis Job**. This opens the Principal Component Analysis page ([Figure 5.30](#)).

Principal Component Analysis (draft)

This form submits a job which analyzes samples using the GenePattern Principal Component Analysis module.

Job Name - Please enter a job name.
Principal Component Analysis Server - Select a PCA grid service from the dropdown.
Clinical Queries - Select saved Clinical queries to specify which samples will be processed.
Enable Preprocess Dataset - (Optional) Check this to display and configure preprocessing parameters.

* Job Name:

* Principal Component Analysis Server: Default Broad service - <http://node255.broad.mit.edu:6060/wsrf/services/cagrid/PCA>

* Clinical Queries: Clinical Queries enable the user to specify which samples will be processed using PCA. The queries selected here have been previously saved by the user. Selected queries will result in the processing of only those samples which are mapped to subjects in the saved query result. If multiple queries are selected, all of the sample from each saved query are processed PLUS the results set will be classified according to those queries. (One class per selected query.)

All Available Queries

gender female
Never smoke

Add >
< Remove

Selected Queries

Enable Preprocess Dataset: ☐
(check to display preprocess parameters)

Perform Analysis

Figure 5.30 Principal Component Analysis parameters

4. Select or define PCA analysis parameters, described in [Table 5.3](#). You must enter a job name and select an annotation query, but you can accept the other default settings..

PCA Parameters	Description
Job Name*	Assign a unique name to the analysis you are configuring.
Principal Component Analysis Server*	A server which hosts the grid-enabled data GenePattern Principal Component Analysis module. Select one from the list and calIntegrator2 will use the selected server for this portion of the processing.
Annotation Queries*	All annotation queries display in this list. Select one or more of these queries to define which samples are analyzed using PCA. If you select more than one query, then the union of the samples returned by the multiple queries is analyzed.
Cluster By*	Selecting rows looks for principal components across all expression values, and selecting columns looks for principal components across all samples.

Table 5.3 PCA analysis options

5. If you want to preprocess the data set, click **Enable the Preprocess Dataset**. This opens an additional set of parameters ([Figure 5.31](#)), discussed in [Table 5.4](#). The preprocessing is executed prior to running the PCA.

Enable Preprocess Dataset: ☒
 (check to display preprocess parameters)

* **Preprocess Server:** Default Broad service - http://node255.broad.mit.edu:6060/wsrf/s

Filter flag ☐

* **Preprocessing Flag:** no-disc-or-norm ▼

* **Min Change:** 3.0

* **Min Delta:** 100.0

* **Threshold:** 20.0

* **Ceiling:** 2.1

* **Max Sigma Binning:** 1

* **Probability Threshold:** 1.0

* **Num Exclude:** 0

Log Base Two ☐

* **Number Of Columns Above Threshold:** 1

Figure 5.31 Parameters for pre-processing parameters for PCA

PCA Preprocessing Parameters	Description
Preprocess Server*	A server which hosts the grid-enabled data GenePattern PreProcess Dataset module. Select one from the list and calIntegrator2 will use the selected server for this portion of the processing.
Filter Flag	Variation filter and thresholding flag
Preprocessing Flag	Discretization and normalization flag
Min Change	Minimum fold change for filter
Min Delta	Minimum delta for filter
Threshold	Value for threshold
Ceiling	Value for ceiling
Max Sigma Binning	Maximum sigma for binning
Probability Threshold	Value for uniform probability threshold filter
Num Exclude	Number of experiments to exclude (max & min) before applying variation filter
Log Base Two	Whether to take the log base two after thresholding
Number of Columns Above Threshold	Remove row if n columns no \geq than the given threshold

Table 5.4 Parameters for preprocessing data sets for PCA

6. When you have completed the form, click **Perform Analysis**.

- When the job is complete, the system displays a completion date on the GenePattern Analysis status page. Click the **Download** link. This downloads zipped result files to your local work station. The number of files and their file type will vary according to the processing. The results format is compatible with GenePattern visualizers and can be uploaded within GenePattern.

GISTIC-Supported Analysis

Note: The GISTIC test option displays only if the study contains copy number or SNP data. For more information, see [Configuring Copy Number Data](#) on page 36.

The GISTIC Module is a GenePattern tool that identifies regions of the genome that are significantly amplified or deleted across a set of samples. For more information, see http://www.broadinstitute.org/cgi-bin/cancer/software/genepattern/modules/gp_modules.cgi.

To perform a GISTIC-supported analysis, follow these steps:

- Select the study whose data you want to analyze in the upper right portion of the calIntegrator2 page. You must select a study with copy number (either Affymetrix SNP or Agilent Copy Number) data.
- Click **GenePattern Analysis** in the left sidebar of calIntegrator2. This opens the GenePattern Analysis Status page.
- In the GenePattern Analysis Status page, select **GISTIC (Grid Service)** from the drop down list and click **New Analysis Job**. This opens the GISTIC Analysis page ([Figure 5.32](#)).

GISTIC Analysis

This form submits a job which analyzes samples using the GenePattern GISTIC module.

Job Name - Please enter a job name.

GenePattern Server URL / GISTIC Server - Select whether to use the GISTIC web service or grid service and provide or select the service address. If the web service is selected, authentication information is also required.

Clinical Queries - (Optional) Select a saved Clinical query to specify which samples will be processed.

Exclude Sample Control Set - (Optional) Select a Control Sample Set to be excluded from the Clinical Query.

* Job Name:

☒ Use GenePattern GISTIC Web Service

* GenePattern Server URL:

* GenePattern Username:

GenePattern Password:

* GISTIC Server:

For the Clinical query parameter below, choose either "All Samples" or a clinical query. If "All Samples" is selected, then all samples will be used. If a clinical query is selected, only those samples which map to the subjects in the clinical query results will be used. The clinical queries in this list have been previously saved by the user. Control samples can be excluded from this processing by selecting a control set name in the Exclude Sample Control Set dropdown.

Clinical query:

* Exclude Sample Control Set:

* Amplifications Threshold:

* Deletions Threshold:

* Join Segment Size:

* QV Thresh:

* Remove X:

cnv File:

Figure 5.32 GISTIC analysis criteria

4. Select or define GISTIC analysis parameters, as described in [Table 5.2](#). You must indicate a Job Name, but you can accept the other defaults settings, which are valid and should produce valid results.

GISTIC Parameters	Description
Job Name*	Assign a unique name to the analysis you are configuring.
GISTIC Server*	A server which hosts the grid-enabled data GISTIC-based analysis module. Select one from the list and calIntegrator2 will use the selected server for this portion of the processing.
Refgene File*	Enter or select the cytoband file to use in the analysis. Allowed values: {Human Hg18, Human Hg17, Human Hg16}. Default = Human Hg16.
Annotation Query	All annotation queries display in this list as well as an option to select all non-control samples. Select an annotation query if you wish to run GISTIC on a subset of the data and select all non-control samples if wish to include all samples.
Amplifications Threshold*	Threshold for copy number amplifications. Regions with a log2 ratio above this value are considered amplified. Default = 0.1.
Deletions Threshold*	Threshold for copy number deletions. Regions with a log2 ratio below the negative of this value are considered deletions. Default = 0.1.
Join Segment Size*	Smallest number of markers to allow in segments from the segmented data. Segments that contain fewer than this number of markers are joined to the neighboring segment that is closest in copy number. Default = 4.
QV Thresh[hold]*	Threshold for q-values. Regions with q-values below this number are considered significant. Default = 0.25.
Remove X*	Flag indicating whether to remove data from the X-chromosome before analysis. Allowed values = {1,0}. Default = 1(yes).

Table 5.5 GISTIC analysis parameters

GISTIC Parameters	Description
cnv File	<p>This selection is optional.</p> <p>Browse for the file. There are two options for the CNV file.</p> <p>Option #1 enables you to identify CNVs by marker name. Permissible file format is described as follows:</p> <p>A two column, tab-delimited file with an optional header row. The marker names given in this file must match the marker names given in the markers_file. The CNV identifiers are for user use and can be arbitrary. The column headers are:</p> <ol style="list-style-type: none"> 1. Marker Name 2. CNV Identifier <p>Option #2 enables you to identify CNVs by genomic location. Permissible file format is described as follows:</p> <p>A 6 column, tab-delimited file with an optional header row. The 'CNV Identifier', 'Narrow Region Start' and 'Narrow Region End' are for user use and can be arbitrary. The column headers are:</p> <ol style="list-style-type: none"> 1. CNV Identifier 2. Chromosome 3. Narrow Region Start 4. Narrow Region End 5. Wide Region Start 6. Wide Region End

Table 5.5 GISTIC analysis parameters

5. When you have completed the form, click **Perform Analysis**.
6. When the job is complete, the system displays a completion date on the GenePattern Analysis status page. Click the **Download** link. This downloads zipped result files to your local work station. The number of files and their file type will vary according to the processing. The results format is compatible with GenePattern visualizers and can be uploaded within GenePattern.

CHAPTER 6

ADMINISTERING USER ACCOUNTS

This chapter describes the process for creating and managing user accounts in calIntegrator2. It also discusses the processes for managing ownership and access to studies in calIntegrator2.

Note: The options for performing user management tasks are visible in calIntegrator2 on the left sidebar of the browser only if you have these Admin privileges.

Administering calIntegrator2 User Accounts Using UPT

Note: If you are interested in registering an account in calIntegrator2, see *Registering as a New calIntegrator2 User* on page 6.

In calIntegrator2, all tasks related to creating and managing user accounts can be performed only by a calIntegrator2 administrator using the CBIIT User Provisioning Tool (UPT) v. 4.2. The following sections discuss the use of the UPT for performing these tasks. For further information about UPT, see Chapter 3 of the CSM 4.2 Programmer's Guide located here: https://gforge.nci.nih.gov/docman/view.php/12/18945/caCORE_CSM_v42_ProgrammersGuide.pdf

The UPT is a separately installed application which serves as the user management interface for all National Cancer Institute CBIIT Life Sciences Distribution (LSD) applications, including calIntegrator2. The UPT application is the central point for all user management functionality within calIntegrator2. You can use UPT to add new users and to apply user group assignments to the calIntegrator2 database directly. The UPT groups can refer to predefined groups such as Study Manager or Study Investigator, which determine what roles the user has.

The following terms are used both in this chapter and in the UPT to define user-related roles:

- **User** – a person who is accessing calIntegrator2. The user has an associated account and user ID.

- **User Group** – a group of users, typically grouped by organization and role, for example, “Columbia University Study Managers”
- **Protection Group** – a group of studies given a secure status and typically grouped by organization, for example, “Columbia University Protected Studies”.

Steps for Creating User Access to caIntegrator2

The following steps summarize the process for establishing user access to caIntegrator2:

1. A potential user requests a user account in caIntegrator2. See [Registering as a New caIntegrator2 User](#) on page 6.
1. You, as a caIntegrator2 administrator, check if the **User** already exists in caIntegrator2. If not, create the new user. See [Creating a New caIntegrator2 User](#) on page 106.
2. Check if the requestor's **User Group** already exists in caIntegrator2. If not, create a new **User Group**. See [Creating a New User Group](#) on page 108.
3. Check if the **Protection Group** (e.g. “Columbia University Protected Studies”), containing the studies to which this user wants access currently exists. If not, create a new **Protection Group**. See [Creating a New Protection Group](#) on page 109.

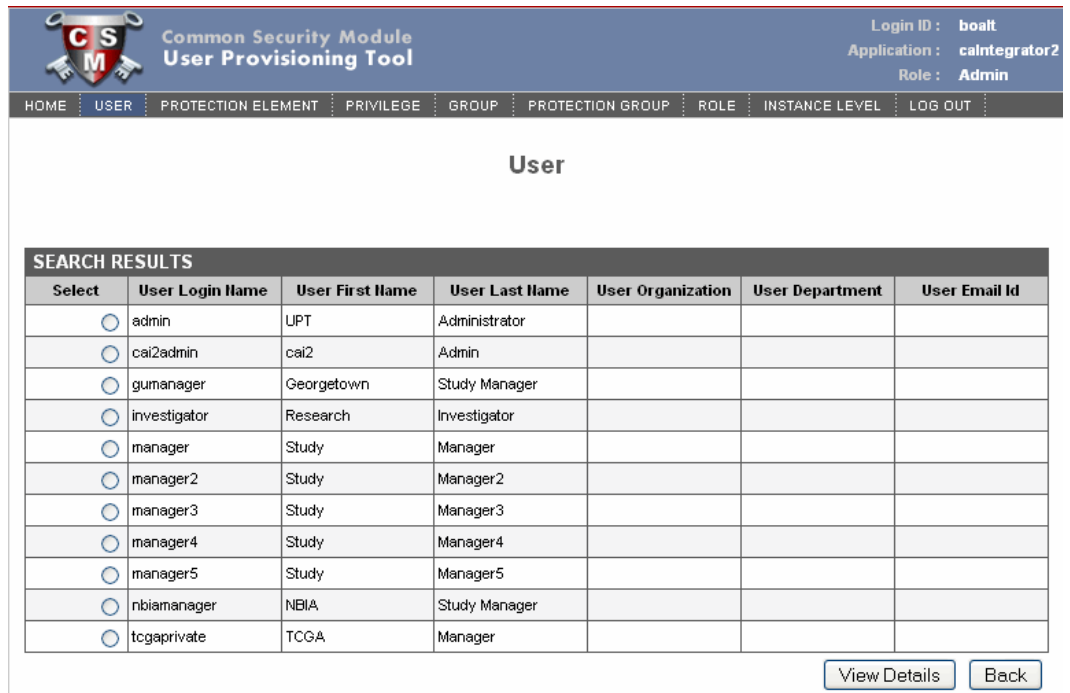
Note: If the Protection Group already exists, contact the Organizational Contact person to confirm that it is OK to give this person access to this Protection Group.
4. Give the requestor's **User Group** access to the **Protection Group**. See [Assigning a User Group to a Protection Group](#) on page 110.
5. Add the **User** to the **User Group**. See [Adding a User to a User Group](#) on page 113

Creating a New caIntegrator2 User

To create a new User in caIntegrator2, follow these steps:

1. Login to UPT as a caIntegrator2 Admin.
2. First, search to see if the user already exists. Click the **User** menu option.
3. On the User page that opens, click **Select an Existing User**.

4. Use the form and search for the user. If you define no criteria, UPT returns a list of all caIntegrator2 users currently in the system (*Figure 6.1*).



The screenshot shows the UPT interface with a navigation bar at the top. The navigation bar includes a logo, the title "Common Security Module User Provisioning Tool", and user information: "Login ID : boalt", "Application : caIntegrator2", and "Role : Admin". Below the navigation bar is a menu with options: HOME, USER, PROTECTION ELEMENT, PRIVILEGE, GROUP, PROTECTION GROUP, ROLE, INSTANCE LEVEL, and LOG OUT. The "USER" option is selected.

The main content area is titled "User" and displays a table of search results. The table has the following columns: Select, User Login Name, User First Name, User Last Name, User Organization, User Department, and User Email Id. The table lists 12 users, each with a radio button in the "Select" column.

Select	User Login Name	User First Name	User Last Name	User Organization	User Department	User Email Id
<input type="radio"/>	admin	UPT	Administrator			
<input type="radio"/>	cai2admin	cai2	Admin			
<input type="radio"/>	gumanager	Georgetown	Study Manager			
<input type="radio"/>	investigator	Research	Investigator			
<input type="radio"/>	manager	Study	Manager			
<input type="radio"/>	manager2	Study	Manager2			
<input type="radio"/>	manager3	Study	Manager3			
<input type="radio"/>	manager4	Study	Manager4			
<input type="radio"/>	manager5	Study	Manager5			
<input type="radio"/>	nblamanager	NBLA	Study Manager			
<input type="radio"/>	tcgaprivate	TCGA	Manager			

At the bottom right of the table, there are two buttons: "View Details" and "Back".

Figure 6.1 A list of current caIntegrator2 users displays in UPT after a user search

5. If the user does not already exist (is not listed in the search results), then create a new user. To do so, select the **User** menu option again, then click **Create a New User**.

This opens the page for creating a new caIntegrator2 user (*Figure 6.2*).

Common Security Module User Provisioning Tool

Login ID : boalt
Application : caIntegrator2
Role : Admin

HOME USER PROTECTION ELEMENT PRIVILEGE GROUP PROTECTION GROUP ROLE INSTANCE LEVEL LOG OUT

Enter the details to add a new User. The **User Login Name** uniquely identifies the User and is a required field. The **User First Name** and **User Last Name** identifies the User. The **User Organization**, **User Department** and **User Title** provides his work details. The **User Phone Number** and **User Email Id** provides the contact details for the User. The **User Password** can be entered if the same schema is also going to be used for Authentication. The **User Start Date** and **User End Date** determine the period for which the User is a valid User.

* indicates a required field

ENTER THE NEW USER DETAILS	
*	User Login Name <input type="text"/>
*	User First Name <input type="text"/>
*	User Last Name <input type="text"/>
	User Organization <input type="text"/>
	User Department <input type="text"/>
	User Title <input type="text"/>
	User Phone Number <input type="text"/>
	User Password <input type="password"/>
	Confirm Password <input type="password"/>
	User Email Id <input type="text"/>
	User Start Date <input type="text"/> (MM/DD/YYYY)
	User End Date <input type="text"/> (MM/DD/YYYY)

Add Reset Back

Figure 6.2 UPT page for creating new user details

6. Enter details for the following required fields:

- **User Login Name**
- **User First Name**
- **User Last Name**
- **User Password**

Caution: If the requestor is an LDAP user, then the User Login Name must match the LDAP login ID AND the User Password field must be left blank. If the requestor is not an LDAP user, then provide a password.

- **User Organization**
- **User Department**

7. Click **Add** to confirm the new user.

Creating a New User Group

You can assign a user group to a protection group. The advantage of working with a user group is that you do not have to assign roles to each user individually. You can assign users to a user group to which you assign a role, and then assign that user group to the protection group, or you can assign a role collectively to a protection group after it is created.

To create a new User Group in calIntegrator2, follow these steps:

1. Login to UPT as calIntegrator2 Admin.
2. First search for an existing group that the user wishes to join. Click the **Group** menu option.
3. On the Group page that opens, click **Select an Existing Group**.
4. Use the form and search for the group. If you define no criteria, UPT returns a list of all calIntegrator2 groups currently in the system
5. If a user group does not already exist, then create a new user group. Click the **Group** menu option, then click **Create a new Group**.
6. On the form that opens (*Figure 6.3*), enter a unique **Group Name** and a description, if appropriate. Click **Add**.

Note: The recommended naming convention for a new User Group is *[insert organization name] Study [insert role]s* Example: "Columbia University Study Managers".

Common Security Module
User Provisioning Tool

Login ID : boalt
Application : calIntegrator2
Role : Admin

HOME USER PROTECTION ELEMENT PRIVILEGE **GROUP** PROTECTION GROUP ROLE INSTANCE LEVEL LOG OUT

Enter the details to add a new Group. The **Group Name** uniquely identifies the Group and is a required field. The **Group Description** is a brief summary about the Group.

* indicates a required field

ENTER THE NEW GROUP DETAILS

* **Group Name**

Group Description

Add Reset Back

Figure 6.3 UPT page for creating a new group

Creating a New Protection Group

If you prefer that a study or group of studies have limited access, you can assign a user to a particular protection group and assign roles which allow the users in the protection group study access. A protection group provides security or limited access for studies listed there.

To create a new Protection Group in calIntegrator2, follow these steps:

1. Login to UPT as calIntegrator2 Admin.
2. Click the **Protection Group** menu option.

- On the page that opens, click **Create a New Protection Group**. The page opens for defining PG Group details (*Figure 6.4*).

Common Security Module
User Provisioning Tool

Login ID : boalt
Application : calintegrator2
Role : Admin

HOME USER PROTECTION ELEMENT PRIVILEGE GROUP PROTECTION GROUP ROLE INSTANCE LEVEL LOG OUT

Enter the details to add a new Protection Group. The **Protection Group Name** uniquely identifies the Protection Group and is a required field. The **Protection Group Description** is a brief summary about the Protection Group. The **Protection Group Large Count Flag** is used to indicate if the Protection Group has a large number of associated Protection Elements.

* indicates a required field

ENTER THE NEW PROTECTION GROUP DETAILS

* **Protection Group Name**

Protection Group Description

Protection Group Large Count Flag ☐ Yes ☒ No

Add Reset Back

Figure 6.4 UPT page for creating a new protection group

- Enter a unique **Protection Group Name** and Description, if appropriate. Click **Add**.

Note: The recommended naming convention is *[insert organization name here] Protected Studies*. Example: "Columbia University Protected Studies".

Assigning a User Group to a Protection Group

To give a User Group access to a Protection Group (a group of protected studies), follow these steps:

- Login to UPT as calIntegrator2 Admin.
- Find the user group that you want to assign. Click the **Group** menu option and click **Select an Existing Group**. In the page that opens, click **Search**. If you define no criteria, UPT returns a list of all calIntegrator2 groups currently in the system (*Figure 6.5*).

Group

SEARCH RESULTS		
Select	Group Name	Group Description
<input type="radio"/>	Study Managers Group 3	Study Managers who can create/modify any Group 3 studies.
<input type="radio"/>	Study Managers Group 4	Study Managers who can create/modify any Group 4 studies.
<input type="radio"/>	Study Managers Group 5	Study Managers who can create/modify any Group 5 studies.
<input type="radio"/>	NCI Study Investigators	Study investigators for the NCI studies.
<input type="radio"/>	NCI Study Managers	Study Managers who can create/modify any NCI studies.
<input type="radio"/>	Platform Manager Group	The platform manager group.
<input type="radio"/>	TCGA Study Managers	Study Managers who can create/modify any TCGA studies.

View Details Back

Figure 6.5 UPT page showing Group search results

3. Select the radio button next to the group name you want to assign to the Protection Group. Click **View Details**. This opens the Group Details page (Figure 6.6).

Common Security Module
User Provisioning Tool

Login ID : boalt
Application : calintegrator2
Role : Admin

HOME USER PROTECTION ELEMENT PRIVILEGE GROUP PROTECTION GROUP ROLE INSTANCE LEVEL LOG OUT

Update the details of the displayed Group. The **Group Name** uniquely identifies the Group and is a required field. The **Group Description** is a brief summary about the Group. The **Update Date** indicates the date when this Group's Details were last updated.

GROUP DETAILS	
* Group Name	NCI Study Managers
Group Description	Study Managers who can create/modify any NCI studies.
Group Update Date	09/24/2009 (MM/DD/YYYY)

Update Delete Back

Associated Users Associated PE & Privileges Associated PG & Roles Assign PG & Roles

Figure 6.6 UPT page showing details for a selected group

4. Below the group details, click **Associated PG & Roles**. The page that opens displays any PG to which the user group is already assigned (Figure 6.7).

Group, Protection Group and Roles

SELECTED GROUP

Group Name
NCI Study Managers

Select the **Protection Group** association which to be removed for the selected **Group** or whose **Roles** Association needs to be updated.

SEARCH RESULTS		
Select	Associated Protection Group Name	Associated Role Name
<input type="radio"/>	NCI Protected Studies	STUDY_MANAGER_ROLE

Remove PG & Roles Associated Roles Back

Figure 6.7 UPT page that shows any PGs to which the select user group is assigned

5. Below the group name, examine if the Protection Group of your choice is already listed there. If so, this means your user group is already assigned to the protection group of choice, and you can skip the remainder of the steps in this section. If the Protection Group is not listed there, then click **Back**.

- Back on the User Group details page, click **Assign PG & Roles**. This opens the Group, Protection Group and Roles Association page ([Figure 6.8](#)).

Group, Protection Group and Roles Association

SELECTED GROUP	
Group Name	NCI Study Managers

Select a single **Protection Group** to associate with the selected **Group**.

AVAILABLE PROTECTION GROUPS
Platforms
Protected Studies for Group 3
Protected Studies for Group 4
Protected Studies for Group 5
TCGA Protected Studies

ASSIGNED PROTECTION GROUP

Select **Roles** which are to be associated with the selected **Group**.

AVAILABLE ROLES
PLATFORM_MANAGER_ROLE
STUDY_INVESTIGATOR_ROLE
STUDY_MANAGER_ROLE

ASSIGNED ROLES

Figure 6.8 UPT page for assigning user group to a protection group and selected roles

- From the list of Available Protection Groups, highlight your PG of choice and click **Assign**.

Now you can assign a role to the user. The caIntegrator2 Roles are defined in [Table 6.1](#):

Role Name	Role Definition
STUDY_MANAGER_ROLE	Assigning this role allows the user to modify existing studies, create new studies, and deploy existing studies.

Table 6.1 Names and definitions for caIntegrator2 roles

Role Name	Role Definition
STUDY_INVESTIGATOR_ROLE	Assigning this role allows the user to search the study, save queries about the study and perform analyses.
PLATFORM_MANAGER_ROLE	Assigning this role allows the user to create and delete array platforms for the entire calIntegrator2 installation. Caution: Array platforms are shared by all users and studies in the calIntegrator2 installation. A user with this role can affect the platforms that are used by all users and studies in the calIntegrator2 installation.

Table 6.1 Names and definitions for calIntegrator2 roles

8. If this user group is a group of study managers, then select STUDY_MANAGER_ROLE. If this user group is a group of study investigators, then select STUDY_INVESTIGATOR_ROLE. Click **Assign**.
9. Click **Update Association** at the bottom of the page. This completes the assigning of the user group to the protection group you chose.

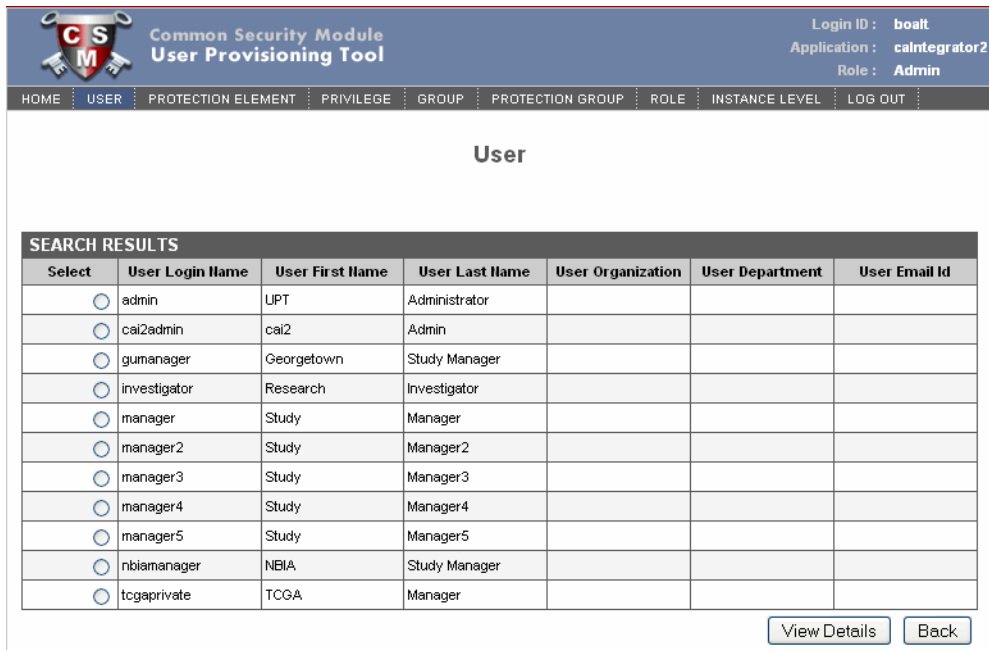
Note: If a **User** has the STUDY_MANAGER_ROLE role for more than one **Protection Group**, then any study that the **User** creates will be assign to each of those **Protection Groups**.

Adding a User to a User Group

To add a user to an existing user group, follow these steps:

1. Login to UPT as calIntegrator2 Admin.
2. Find the user that you want to assign to a user group. Click the **User** menu option, then click **Select an Existing User**.

- Enter the name of the user you are looking for and click **Search**. If you define no criteria, UPT returns a list of all calIntegrator2 users currently in the system (Figure 6.10).



The screenshot shows the 'Common Security Module User Provisioning Tool' interface. The top navigation bar includes links: HOME, USER, PROTECTION ELEMENT, PRIVILEGE, GROUP, PROTECTION GROUP, ROLE, INSTANCE LEVEL, and LOG OUT. The user's session information is displayed as: Login ID: boalt, Application: calIntegrator2, Role: Admin.

The main section is titled 'User' and displays 'SEARCH RESULTS' in a table. The table has the following columns: Select, User Login Name, User First Name, User Last Name, User Organization, User Department, and User Email Id.

Select	User Login Name	User First Name	User Last Name	User Organization	User Department	User Email Id
<input type="radio"/>	admin	UPT	Administrator			
<input type="radio"/>	cal2admin	cal2	Admin			
<input type="radio"/>	gumanager	Georgetown	Study Manager			
<input type="radio"/>	investigator	Research	Investigator			
<input type="radio"/>	manager	Study	Manager			
<input type="radio"/>	manager2	Study	Manager2			
<input type="radio"/>	manager3	Study	Manager3			
<input type="radio"/>	manager4	Study	Manager4			
<input type="radio"/>	manager5	Study	Manager5			
<input type="radio"/>	nbiamanager	NBIA	Study Manager			
<input type="radio"/>	tcgaprivate	TCGA	Manager			

At the bottom right of the table are two buttons: 'View Details' and 'Back'.

Figure 6.9 UPT page showing a list of calIntegrator2 users

- Select the radio button next to the name and click **View Details** (Figure 6.10).



The screenshot shows the 'Common Security Module User Provisioning Tool' interface for editing a user. The top navigation bar and session information are the same as in Figure 6.9.

The main section contains instructions: 'Update the details of the displayed User. The **User Login Name** uniquely identifies the User and is a required field. The **User First Name** and **User Last Name** identifies the User. The **User Organization**, **User Department** and **User Title** provides his work details. The **User Phone Number** and **User Email Id** provides the contact details for the User. The **User Password** can be entered if the same schema is also going to be used for Authentication. The **User Start Date** and **User End Date** determine the period for which the User is a valid User. The **Update Date** indicates the date when this User's Details were last updated.'

The 'USER DETAILS' section contains the following fields:

*	User Login Name	manager
*	User First Name	Study
*	User Last Name	Manager
	User Organization	
	User Department	
	User Title	
	User Phone Number	
	User Password	••••••
	Confirm Password	••••••
	User Email Id	
	User Start Date	(MM/DD/YYYY)
	User End Date	(MM/DD/YYYY)
	User Update Date	09/24/2009 (MM/DD/YYYY)

At the bottom right are three buttons: 'Update', 'Delete', and 'Back'. Below these are four tabs: 'Associated Groups', 'Associated PE & Privileges', 'Associated PG & Roles', and 'Assign PG & Roles'.

Figure 6.10 UPT page showing details for a selected user

- Click the **Associated Groups** button at the bottom of the page. This opens the page where you can assign a user to a group (*Figure 6.11*).

User and Groups Association

SELECTED USER	
User Login Name	manager

Assign or Deassign multiple **Groups** for the selected **User**. To remove the complete association Deassign all the **Groups**.

AVAILABLE GROUPS
Study Managers Group 3
Study Managers Group 4
Study Managers Group 5
NCI Study Investigators
TCGA Study Managers

ASSIGNED GROUPS
Platform Manager Group
NCI Study Managers

Figure 6.11 UPT page for assigning a user to user groups

- Select the group(s) that you want the user to be in and click **Assign**.
- At the bottom of the page click **Update Association**. This completes the assigning of the user to the user group. Now the user will have access to any studies to which the user group has been given access.

Note: You can add a user to more than one user group. For example, a user could be assigned to “Columbia University Study Managers” as well as to “Columbia University Study Investigators”.

Changing a User Password

To change a password for a User, follow these steps:

- Confirm if the User is an LDAP user or not. If the User is an LDAP user, then this person must change their password using the NCI password change utility. Skip the rest of these steps.
- If the User is not an LDAP user, then continue with the rest of these steps.
- Login to UPT as calIntegrator2 Admin.
- Find the User that you want to change. Click the **User** menu option, then **Select an Existing User**.
- Enter the name of the user you are looking for and click **Search**. If you define not criteria, UPT returns a list of all calIntegrator2 users.
- Select the radio button next to the name and click **View Details**

7. Replace the **User Password** and **Confirm Password** fields with the new password (Figure 6.12).



Common Security Module
User Provisioning Tool

Login ID : boalt
Application : calIntegrator2
Role : Admin

HOME USER PROTECTION ELEMENT PRIVILEGE GROUP PROTECTION GROUP ROLE INSTANCE LEVEL LOG OUT

Update the details of the displayed User. The **User Login Name** uniquely identifies the User and is a required field. The **User First Name** and **User Last Name** identifies the User. The **User Organization**, **User Department** and **User Title** provides his work details. The **User Phone Number** and **User Email Id** provides the contact details for the User. The **User Password** can be entered if the same schema is also going to be used for Authentication. The **User Start Date** and **User End Date** determine the period for which the User is a valid User. The **Update Date** indicates the date when this User's Details were last updated.

USER DETAILS	
* User Login Name	manager
* User First Name	Study
* User Last Name	Manager
User Organization	
User Department	
User Title	
User Phone Number	
User Password	••••••
Confirm Password	••••••
User Email Id	
User Start Date	(MM/DD/YYYY)
User End Date	(MM/DD/YYYY)
User Update Date	09/24/2009 (MM/DD/YYYY)

Update Delete Back

Associated Groups Associated PE & Privileges Associated PG & Roles Assign PG & Roles

Figure 6.12 UPT page where you can edit user details, such as a password

8. At the bottom of the page click **Update**.

DATA IMPORT CONFIGURATIONS

This appendix describes configurations for importing data into a study.

Topics in this appendix include the following:

- [Subject Annotation Data Configuration](#) on this page
- [Delimited-Text Annotation Import](#) on this page
- [Annotation Field Configuration](#) on page 118
- [Sample Data Configuration](#) on page 118
- [Genomic Data Configuration](#) on page 119
- [Supplemental Files Configuration](#) on page 119
- [Imaging Data Configuration](#) on page 121

Subject Annotation Data Configuration

The following subject annotation data configuration information is collected:

- subject annotation Data Source (delimited text)
- Protocol Id (of study to import)
- For delimited text, see [Delimited-Text Annotation Import](#). For subject annotation files, one field must be identified as the subject identifier.
- See [Annotation Field Configuration](#) for details on specification of visibility and browse configuration.

Delimited-Text Annotation Import

Delimited-text annotation files must be in standard comma-separated value format. The file must include a header line that specifies the name for each field. Each row of data

must contain the same number of values as the header row. The file must include a column that will be designated as the identifier (e.g. subject identifier, sample identifier, etc.) for each row. Optionally a file may include a single column that will be designated as a time-point indicator. Each row must contain a unique combination of identifier and time-point indicator of a unique identifier if no time-point is included. An example of the content of a file including a time-point is shown below.

```
"patientId", "timepoint", "bloodPressure", "weight"  
"1234", "T1", "120/80", "180"  
"1234", "T2", "125/80", "190"  
"5678", "T1", "120/85", "200"
```

After upload of the file, the Study Manager must indicate for each field:

- Field type (identifier, timepoint indicator, text, integer, float or Boolean)
- After specification of these types, the file will be validated to ensure that the values are valid for the types selected and that the file conforms to the requirements given above.

Annotation Field Configuration

For each annotation field (regardless of the source), the Study Manager must specify the following information:

- Annotation semantics: each annotation field (whether associated with a subject, image series, image or sample) must either:
 - be associated with an existing annotation definition known to the system,
 - be associated to an existing CDE in caDSR or
 - have sufficient semantic metadata recorded so that the field may be submitted for registration as a CDE in caDSR.
- Field authorization: Each field must be either declared publicly visible or restricted to a list of groups. The default will be the visibility settings given at the study level. For more information, see [Adding Subject Annotation Data](#) on page 20.
- Whether the field is to be included in the results list for a given entity type (i.e. Subject, Sample, Image Series or Array Data) when browsing data.
- Whether the field is to be included in simple single-input searches when browsing data.

Sample Data Configuration

Sample data may be uploaded from either caArray 2 or from delimited-text import. Samples imported from caArray 2 may have annotation updated by use of the delimited-text import functionality if sample annotation is required. Import from caArray 2 requires specification of the following information:

- caArray server hostname
- caArray server JNDI port

- caArray username
- caArray password
- Either the experiment identifier (to import all samples in the experiment) or a file containing a comma-separated list of samples in the format “experiment identifier”, “sample name”.
- Mapping of samples to subjects. This may be specified by a comma-separated list in the format “subject identifier”, “sample identifier” or by a regular-expression based mapping formula.

When samples are imported via delimited-text import, the time-point is associated to the sample itself. This means that each sample may be associated with only one time-point (i.e. multiple time-points for the same sample are invalid).

Genomic Data Configuration

All genomic data (i.e. array data) is imported from caArray 2. First the Study Manager must specify sufficient information to map study samples to caArray 2 samples. If all samples were imported directly from caArray 2 as described in Special Requirement: Sample Data Configuration, no further information is required for this step. If samples were imported via delimited-text, the Study Manager must specify

- caArray server hostname
- caArray server JNDI port
- caArray username
- caArray password
- A mapping of calIntegrator2 sample identifiers to caArray 2 samples, specified as a comma-separated list in the format “calIntegrator2 sample identifier”, “caArray 2 experiment identifier”, “caArray 2 sample name”.

The system will enable the Study Manager to navigate easily to the selected caArray 2 instance.

Next, the system will indicate the available platforms and array data types available for the study samples. The Study Manager will indicate which platforms and data types to import and for each platform/data type combination will indicate:

- Whether to import the data
- The visibility of the data; either public or restricted to a set of groups. Low-level genotyping data (raw data and normalized) will always have restricted visibility.

See also [Supplemental Files Configuration](#).

Supplemental Files Configuration

This section describes the format that must be used when creating supplemental files for use by calIntegrator2. The supplemental files described here are to be added to an experiment in caArray prior to configuring a study in calIntegrator2.

The file itself is a tab-delimited text file. The file extension can be anything, though users typically use .txt. The name of each supplemental file must be unique within a caArray experiment.

Inside the file, each row in the file contains the data from one reporter. Each column in the file must have a unique header name, that is, you cannot give two different columns the same column name.

There are two supported formats for these files: Single Sample Format and Multiple Sample Format.

Single Sample Format

- Minimum of two required columns
- One column must contain the reporter/probe name
- One column must contain the value be reported by the reporter
- The file can have additional columns, though other than reporter/probe name and value mentioned above, the rest will be ignored
- One single sample file for each sample in the experiment

An example of single sample format file is shown in [Figure A.2](#)

ProbeID	signal:Log2
A_14_P112718	0.01
A_16_P15000916	0.5166
A_16_P15001074	0.4965
A_16_P00000012	0.1553
A_16_P00000014	-0.5684
A_16_P00000017	NA
A_16_P00000021	-0.6415
A_16_P00000023	-0.6041
A_16_P00000027	-0.374
A_16_P00000033	-0.493
A_16_P35001586	-0.465
A_16_P15001533	0.1939
A_16_P00000060	-0.0576
A_16_P15001594	-0.0674
A_16_P00000082	0.1754
A_16_P00000090	0.0878
A_16_P00000099	-0.4532
A_16_P15001666	-0.1321
A_16_P00000112	0.8277
A_16_P00000114	0.1214
A_16_P00000127	0.8765
A_16_P00000136	0.1701
A_16_P00000140	-0.3096

Figure A.1 Example of single sample format file

Multiple Sample Format

- One column must contain the reporter/probe name.
- Each additional columns are the reporter values such that there is one column per sample.
- One multiple sample file for the whole experiment.

Note: Currently the multiple sample format is slower to load than the single sample format for platforms other than Agilent Copy Number. Future releases should show improvements in this performance.

An example of multiple sample format file is shown in [Figure A.2](#).

Hybridization Ref	Chr	Pos	TCGA-09-0364-01A-02D-0357-04 signal:Log2	TCGA-13-0723-01A-02D-0357-04 signal:Log2	TCGA-13-0757-01A-01D-0357-04 signal:Log2
A_14_P112718	1	554268		0.01	0.2111
A_16_P15000916	1	554287		0.5166	1.2929
A_16_P15001074	1	639581		0.4965	0.4769
A_16_P00000012	1	736483		0.1553	0.3047
A_16_P00000014	1	742533		-0.5684	0.3057
A_16_P00000017	1	746956	NA		0.3308
A_16_P00000021	1	757922		-0.6415	-0.2729
A_16_P00000023	1	769590		-0.6041	0.4084
A_16_P00000027	1	784458		-0.374	-0.2919
A_16_P00000033	1	792413		-0.493	-0.3545
A_16_P35001586	1	800905		-0.465	0.0274
A_16_P15001533	1	823964		0.1939	0.4682
A_16_P00000060	1	836543		-0.0576	0.3648
A_16_P15001594	1	842726		-0.0674	0.4352
A_16_P00000082	1	847646		0.1754	0.4735
A_16_P00000090	1	853295		0.0878	0.3196
A_16_P00000099	1	857406		-0.4532	-0.2435
A_16_P15001666	1	862519		-0.1321	0.0441
A_16_P00000112	1	865691		0.8277	0.9984
A_16_P00000114	1	868794		0.1214	0.1444
A_16_P00000127	1	875165		0.8765	1.0351

Figure A.2 Example of a multiple sample format file

The following software programs create the supplemental data format used by caArray:

- Affymetrix Expression Console – This software produces supplemental files. In Expression Console, use the “Export Result” function to create these files. Note that when you use an algorithm other than MAS5 to normalize the data (for example using RMA or Plier), Expression Console automatically creates a [...summary.txt] file that contains extra lines on top of the derived data results. The extra lines all start with a “#” to signify that it is a remark. These lines are ignored by calIntegrator2 parsing.
- Agilent GeneSpring GX – This software can export a results table in .txt format.

Imaging Data Configuration

The following imaging data configuration information is collected:

- NBIA grid server hostname (defaults to NCICB instance)
- NBIA grid server port (defaults to NCICB instance port)
- Protocol Id
- Mapping of NBIA Patients to subjects imported from subject annotation data source. This may be specified by a comma-separated list in the format “subject identifier”, “NBIA patient identifier” or by a regular-expression based mapping formula.
- Which annotation fields to import from NBIA.
- The system will enable the Study Manager to navigate easily to the selected caArray 2 instance.

Additional annotation for either images or image series may be imported using the delimited-text import functionality.

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