

Tutorial: Microarray-based expression analysis part I: Getting started

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Tutorial: Microarray-based expression analysis part I: Getting started

This tutorial is the first part of a series of tutorials about expression analysis. Expression analysis often requires advanced skills in statistics, but this tutorial is intended to show a straight-forward example of how to identify and interpret the differentially expressed genes in samples from two different tissues. If you are familiar with the statistical concepts and issues within expression analysis, you may find this tutorial too simplistic, but we have favored a simple and quick introduction over an exhaustive and more "correct" explanation.

The data comes from a study of gene expression in tissues from cardiac left ventricle and diaphragm muscle of rats [van Lunteren et al., 2008]. During this series of tutorials, you will see how to import and set up the data in an experiment with two groups (part I), to perform quality checks on the data (part II), to perform statistics and clustering to identify and visualize differentially expressed genes (part II), and finally to use annotations to categorize and interpret patterns among the differentially expressed genes in a biological context (part IV).

Importing array data

First, import the data set which can be downloaded from the Gene Expression Omnibus (GEO) database at NCBI: http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc= GSE6943&targ=gsm&form=text&view=data. After download, click Import () in the Tool bar and select the file. You will now have 12 arrays in your Navigation Area as shown in figure 1.



Figure 1: 12 microarrays have been imported.

Grouping the samples

The next step is to tell the CLC Main Workbench how the 12 samples are related.

This is done be setting up an **Experiment** (**III**). An **Experiment** is the central data type when analyzing expression data in the *CLC Main Workbench*. It includes a set of samples and information about how the samples are related (which groups they belong to). The **Experiment** is also used to accumulate calculations like t-tests and clustering.

First step is to set up the experiment:

Toolbox | Expression Analysis (🙀) | Set Up Experiment (🎼)



Select the 12 arrays that you have imported (see figure 2).

Figure 2: Select the 12 microarrays that have been imported.

Note that we use "samples" as the general term for both microarray-based expression values and sequencing-based expression values. Clicking **Next** shows the dialog in figure **3**.

 Select at least two samples 	Define experiment type
2. Define experiment type	Experiment Two-group comparison Unpaired Paired Multi-group comparison Unpaired Paired
	RVM-Seq expression values

Figure 3: Defining the number of groups.

Here you define the number of groups in the experiment. Since we compare heart tissue with diaphragm tissue, we use a two-group comparison. Leave it as **Unpaired**. Clicking **Next** shows the dialog in figure 4.

Name the first group Heart and the second group **Diaphragm** and click **Next** (see figure 5).

Here you see a list of all the samples you chose in figure 2. Now select the first 6 samples (by clicking in the group column of the first sample and while holding down the mouse button you drag and select the other five samples), right-click and select **Heart**. Select the last 6 samples, right-click and select **Diaphragm**. In this way you define which group each sample belongs to.

Click **Finish** and the experiment will be created. Note that the information from samples located in the **Navigation Area** is copied into the experiment, so they now exist independently of each other.

Set Up Experiment	×
 Select at least two samples and (optionally) an annotation file 	Assign group names
2. Define experiment type	
3. Assign group names	
	Assign names to groups Group 1 [Heart Group 2 Diaphragm]
?	← Previous → Next √ Finish X Cancel

Figure 4: Naming the groups.

🚾 Set Up Experiment							
1. Select at least two samples and (optionally)	Assign groups to the selected samples Please right-click onto each cell and assign a group						
an annotation rile	Sample 🔬	Group					
Define experiment type	GSM160089	<right-click assign="" group="" to=""></right-click>					
Assign group names	GSM160090	<right-click assign="" group="" to=""></right-click>					
4. Assign groups to the	G5M160091	<right-click assign="" group="" to=""></right-click>					
selected samples	GSM160092	<right-click accion_group="" to=""></right-click>					
	GSM160093	<ric dianhrann="" roup=""></ric>					
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	G5M160096						
	G5M160097	<right-click assign="" group="" to=""></right-click>					
	GSM160098	<right-click assign="" group="" to=""></right-click>					
	GSM160099	<right-click assign="" group="" to=""></right-click>					
	GSM160100	<right-click assign="" group="" to=""></right-click>					
? 9	Frevious	Next Finish X Cancel					

Figure 5: Assigning the samples to groups.

The experiment table

Once it is created, the experiment will be opened in a table as shown in figure 6.

The table includes the expression values for each sample and in addition a few extra values have been calculated such as the range, the IQR (Interquartile Range), fold change and difference values and the present counts for the whole experiment and the individual groups (note that absent/present calls are not available on all kinds of data).

Save the experiment and you are ready to proceed to the expression analysis tutorial part II.

🏢 * Heart vs. Dia 🕄	
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Rows: 15.923 Filter:										% ~ ₩ ?	
			Experiment							✓ Column width	
Feature ID 🔬	Total prese Range (orig I	[Exponitione			GSM160089		GSM160090			Manual 🚩
		IQR (origin	Difference (Fold Chang	Expression	Presence call	Expression	Presence call	Expre	Experiment level	
1367452_at	12	862,00	470,50	464,82	1,19	2.532,90	P	2.518,60	P	~	Analysis level
1367453_at	12	1.231,00	430,80	428,37	1,13	3.464,20	Р	3.197,40	Р		
1367454_at	12	536,50	349,10	153,85	1,10	1.620,80	Р	1.870,50	Р		Annotation level
1367455_at	12	2.196,20	1.352,90	-772,35	-1,17	5.512,50	Р	4.103,90	Р		 Group level
1367456_at	12	2.095,50	1.264,20	-1.205,65	-1,27	6.090,80	Р	5.352,20	Р		Heart
1367457_at	12	508,20	319,00	-64,73	-1,07	1.093,90	P	1.134,30	Р		Minearc
.367458_at	12	268,30	148,90	112,30	1,38	347,80	Р	223,90	Р		🗹 Diaphragm
.367459_at	12	3.993,80	2.434,30	2.557,35	1,36	7.665,80	Р	7.415,90	Р		Group columns
.367460_at	12	1.182,80	557,00	-484,73	-1,17	3.155,70	Р	2.946,90	Р		Means
1367461_at	12	485,70	280,20	184,35	1,29	507,00	Р	610,30	Р		
1367462_at	12	1.032,50	309,70	268,23	1,08	3.207,50	Р	3.371,30	Р		Present count
1367463_at	12	1.621,60	510,00	701,97	1,21	3.510,30	Р	3.050,30	Р		Select All
.367464_at	12	317,70	202,00	-111,67	-1,13	797,70	Р	1.038,90	Р		Decelect All
1367465_at	12	699,00	196,80	257,50	1,22	1.103,10	Р	1.281,80	Р		Deselect All
1367466_at	12	265,50	122,50	-54,67	-1,04	1.385,10	Р	1.321,30	Р		Sample level
1367467_at	12	1.780,60	453,40	414,95	1,12	3.561,60	Р	3.838,40	Р		
1367468_at	12	572,60	480,60	439,67	1,73	656,30	P	658,20	Р		
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5										>	
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Figure 6: The experiment table.

Bibliography

[van Lunteren et al., 2008] van Lunteren, E., Spiegler, S., and Moyer, M. (2008). Contrast between cardiac left ventricle and diaphragm muscle in expression of genes involved in carbohydrate and lipid metabolism. *Respir Physiol Neurobiol*, 161(1):41–53.