Microarray Toolkit

With the Partek Flow Microarray Toolkit, you can import and analyze microarray data with the same ease as any sequencing analysis pipeline. This document covers the following parts.

- Supported Platforms
- Data Import
- Conversion to Aligned Reads
- Visualizing Microarray Probes
- Downstream Tasks
- References
- Additional Assistance

Supported Platforms

Partek Flow supports popular gene expression microarray platforms including Affymetrix GeneChips (.CEL) and Illumina BeadChips (.idat). You can also import text output from Illumina Genome Studio in the form of tab-delimited text files with probe IDs and AVGSignal values. For the latter, each sample should correspond to one text file.

For custom-made Affymetrix .CEL and Illumina .idat or AVGSignal data generated from Genome Studio files, refer to the documentation on Importing Custom Microarrays for additional instructions regarding the probe files.

Data Import

To import microarray files into a project, go to the *Data* tab of the project and select the **Import Data** button. If the data is not already in an existing Partek Flow project, select the **Automatically create samples from files** button. You will be presented with three different ways to upload your data files:

- Directly from your Partek Flow server
- From My Computer
- From a URL

The most efficient way of importing array data would be to import them directly from your Partek Flow server. When you select this option, navigate to the folder containing your data files. Valid file types will be selectable for upload. Click on the files you would like to import into the project and select the **Create sample** button (Figure 1).

Flow Dueue Projects V		順 🛛 John A Smith 🗸
<u>Home</u> > <u>Microarray project</u> > <u>Data</u> > Create samples from files		
Select files from ()		
Current directory 🕖		
/home/flow/FlowData/Microarray_Project Goto	6 files selected	
★ Server Computer	✓ Name	Size
Tome flow		10.57 MB
 FlowData DNA-seq 	GSM1489329.CEL	10.57 MB
a download	GSM1489330.CEL	10.57 MB
▶ I∎ genomic_data ▶ I≣ library_files	GSM1489334.CEL	10.57 MB
Microarray_Project	GSM1489335.CEL	10.57 MB
	GSM1489336.CEL	10.60 MB
Don't see your folder? Refresh folder list Back Create sample	Valid file types are vcf, idat, bz2, gz, tar, csfasta, ra qual, zip, bcf, sra, sff, fastq, fasta, txt, bam, sam or	w, csfastq, bpm, probe_tab, CEL, ·bgx files
Copyright © 2016 Parte	Incorporated. All rights reserved.	

Figure 10. Importing microarray files into Partek Flow

During the upload process, Partek Flow will determine the specific platform of your dataset. If they are commonly used chips, such as Affymetrix HTA arrays, the gene and transcript annotations associated with the platform will automatically be downloaded in the *Library File Manager*.

Once the samples have been uploaded, the *Data tab* will list each sample along with the *Platform* associated with each sample. If Affymetrix .CEL files were uploaded, it will also include the date that the array was scanned (Figure 2). This information may be helpful in assessing possible batch effects.

ome	> Mic	croarray	proj	ect		\$ *
naly	yses	Data	Log	g Project settings		
				Attribut	es	
	San	nple nam	ie	Platform	Scan date	
1	GSM	1489328	*	RaGene-1_0-st-v1	04/28/2011	
2	GSM	1489329	\$	RaGene-1_0-st-v1	04/28/2011	
3	GSM	1489330	\$	RaGene-1_0-st-v1	04/28/2011	
4	GSM	1489334	<u>ې</u>	RaGene-1_0-st-v1	04/28/2011	
5	GSM	1489335	<u>ن</u>	RaGene-1_0-st-v1	04/28/2011	
6	GSM	1489336	÷	RaGene-1_0-st-v1	04/28/2011	
Sho	w dat	a files			Download	
Imp Add Proje	oort da d a sys ect out	ata E stem-wid	Edit a le attr ctory	ttributes Assign ibute column Ma ibute flow/Flow	sample attribut anage attributes Data/project_out	es from a file put/Project_Microarray project (16.62 TB free) 🥜

Figure 11. Imported microarrays will automatically include Platform and, in some cases, Scan date in the Attributes page

You can now add additional sample attributes to your microarray project. For more information on how to set up your project, please refer to the documentation on *Creating a project*.

Custom annotations

Partek Flow also supports custom-made Affymetrix and Illumina arrays. Once the data files have been uploaded, you will also need to supply a custom annotation file which contains information on the probe sequences associated with the array. Please refer to the document on Importing Custom Microarrays for additional information on how to create and upload these annotations.

Conversion to Aligned Reads

Once the microarray data has been uploaded, the *Microarray intensity* data node will appear in the *Analyses* tab. To convert this data to aligned reads, select the **Microarray intensity** data node and select the aligner you would like to use for conversion (Figure 3).

Flow Dueue V Projects V	🥊 John A Smith 🗸
<u>Home</u> > Microarray project	\$ ~
Analyses Data Log Project settings	
	× Microarray intensity
Microarray	
intensity	STAR
	GSNAP
	SHRiMP 2
	Bowtie
	Bowtie 2
	BWA
	TMAP
	TopHat
	TopHat 2
	Download data (63 MB)
Import a pipeline	Project disk space
Copyright © 2016 Partek Incorporated. All rights reserved.	

Figure 12. Converting Microarray intensity to Aligned reads

Selecting an aligner will bring up the *Microarray Conversion options* page (Figure 4). Under the *Select probe sequence file* section, make sure that the correct *Chip name* is specified. For non-custom arrays, these should be set automatically to the platform detected upon importing the array.

Select probe sequence file	
Chip name	MTA-1_0
	Change selection
Select coverage depth	
Nominal read coverage depth (in millions)	20.0
Advanced options	
Option set	Default Configure
Back Next	

Figure 13. The Microarray Conversion options page

If for some reason you wish to override that selection, then click the **Change selection** link and select the chip name in the drop-down menu or select **New chip...** if you would like to upload a different chip annotation (Figure 5).

Select probe sequence file			
Chip name	RaGene-1_0-st-v1 ▼ AG RaGene-1_0-st-v1 New chip		

Figure 14. Selecting a different probe sequence file

In the Select coverage depth section, you can set the nominal read coverage depth for your array. This is by default set at 20 million reads. However, you can adjust this number for better accuracy or if you are evaluating low-expressing genes. If you are comparing the microarray data with a corresponding NGS dataset, you can use this option to scale your data to a more comparable order of magnitude. Take note that setting this to a higher number will result in longer computation times overall.

Advanced options

Clicking the *Configure* link under the *Advanced options* section to view the advanced options that can be modified. These are generally transformations applied to microarray data (Figure 6). Note that both interrogating and control (if present) probes are used during fitting and adjustment.

Advanced options X			
Sequence options			
Consolidation of identical probe 🕧 sequences	mean v		
Specify annotation sequence format 🕧	none v		
Sequence correction transform			
Sequence correction transform 🕧			
GCCN sequence correction transform 🕧			
Signal space transformation (SST)			
Signal space transformation (SST) 🕧	✓		
Scale intensities floor 🏼 👔	1		
Scale intensities ceiling 🕧	1000000 🗘		
Scale intensities low 🏼 👔	20		
Scale intensities high 🏼 🕧	50000		
Scale intensities detect previous 🧃 transform	65535 🜩		
RMA background transform			
RMA background transform 🏼 🕧	✓		
Quantile normalization			
Quantile normalization 🕧			
Apply Save as new Cancel			

Figure 15. Advanced options for Microarray conversion

The Specify annotation sequence format drop-down menu allows you to specify the format of the probe sequences associated with your microarrays. Please consult with the manufacturer of your microarrays to determine the exact specification of their probe annotation files. The options available for sequence annotation include:

- None: Assumes that the sequences in the microarray annotation are the probe sequences printed on the array
- Complement: Assumes that the sequence is for the region being interrogated by the array
- Reverse complement: Assumes that the sequence is the reverse complement of the interrogated sequence. This is particularly useful if you have primer sequences as annotations

The Consolidation of identical probe sequences option is available to take into account probes that have identical sequences within the same array. This is particularly common in microRNA arrays, where identical mature miRNA sequence can exist for multiple organisms.

By default, this is set to take the *mean* of the intensities the probes and utilizes that to calculate the read count for the probe sequence. Clicking on the drop-down menu allows you to either turn off this feature (*none*) or use the *median* instead (Figure 6).

Import options		
Nominal read coverage depth (in millions)	0	20.0
Consolidation of identical probe sequences	i	mean 🔻
Advanced options		mone mean
Option set	De	fault ▼ Configure

Figure 16. Changing the way identical probes are consolidated

Clicking the *Sequence correction transform* checkbox corrects intensity values for sequence-specific effects. It is turned off by default and we only advise that you select this option if there are wide differences in the GC content of your probes.

Both the GCCN sequence correction transform and Scale intensity transforms were developed by Affymetrix [1]. GCCN scales the intensities with respect to the difference in probe affinity associated with Guanine and Cytosine (GC) content. On the other hand, the Scale intensity transform was developed improve inter-platform comparisons between microarray and RNA-seq data. Specifically, the algorithm stretches the intensity distribution to a common range with a power law mapping that decompresses fold change ratios. This ultimately simplifies fold change comparisons between different technologies.

The *RMA background transform* check box applies robust multi-array average (RMA) normalization to your microarray samples [2]. This is a well-accepted method of background correction for microarray intensities.

The Quantile normalization check box performs quantile normalization on all microarray intensities. This is important in making sure that signals from any two arrays are comparable.

Any combination of these advanced options can be saved as its own option set. Once the option set has been created, it will be available in the *Option set* drop-down menu for use in future projects.

After you have finished configuring all microarray conversion options. Click the **Next** button to configure the parameters for the aligner you have chosen to use. Please refer to the documentation on *Alignment* for more information about configuring aligners.

Visualizing Microarray Probes

To visualize microarray probes in Partek Flow's *Chromosome viewer*, click the **Select tracks** button at the top left corner of the viewer. Select the **Probe** intensities check box to display microarray probes (Figure 8).



Figure 17. Displaying microarray probes in Chromosome viewer

This will display microarray probes aligned to the reference genome (Figure 9). By default, the probes are colored by their intensities with darker shades signifying higher probe intensities.

 Selection details 		78K SNORA17 U6 U1 Sir2a5
Sample GSM1489336	Ensembl Transcripts release 83	
Position 5:167173745-167173770 Browse on UCSC	->>>>>>>>>>>>>>>>>>	
Read length 25	->>>>>	$\rightarrow \rightarrow $
		Sic2a5
Strand Forward	Probe intensities: STAR - 2.4.1d (ethanol)	178
Average intensity	Probe intensities: STAR - 2.4.1d (saline)	
17.14	- 60	120
Sequence	Reference sequence 🛛 A 🗖 C 🗖 G 📕 T 🔲 N	Tend blidden ferene is to size
GG BLAST this sequence	167173366 167173575	таск повен (zoom in to view) 1671†3784 1671†3993 167174203

Figure 18. Viewing probe intensities

Clicking a specific probe will display additional information about that probe including mapping position and sequence length. If the probe intensities are consolidated for specific categorical attributes, the average probe intensity is also displayed. For more details on how to use the *Chromosome viewer*, please consult its documentation.

Downstream Tasks

The Convert to aligned reads task generates an Aligned reads data node. This then allows you to perform most tasks that can be performed on that data node type in Partek Flow. These include Quantify to transcriptome, Normalize counts and Differential gene expression (GSA) analysis. Please consult the documentation on a specific task for specific instructions on their use.

References

- 1. Affymetrix Whitepaper on Microarray normalization using Signal Space Transformation with probe Guanine Cytosine Count Correction. Accessed February 17, 2016
- 2. Irrizarry RA, Hobbs B, Collin F et al. Exploration, normalization, and summaries of high density oligonucleotide array probe level data. *Biostat.* 2003; 4:249-264.

Additional Assistance

If you need additional assistance, please visit our support page to submit a help ticket or find phone numbers for regional support.

