BEIStack 02A Using Batch Effects Interface Assessments External Tod Casasent 2019-02-05-1520

1 Using Batch Effects Interface Assessments

This document focuses on explaining the components of the Batch Effects Interface (BEI) involved with creating a job, loading data, and running assessments. This document will not address statistical issues or "how to spot" batch effects.

The URL for your install should be provided to you, but will likely be something like:

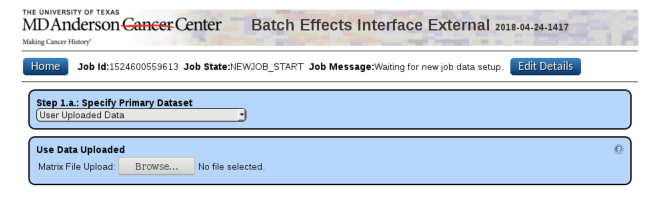
http://your-server.your-company.com:8181/BatchEffectsInterface/

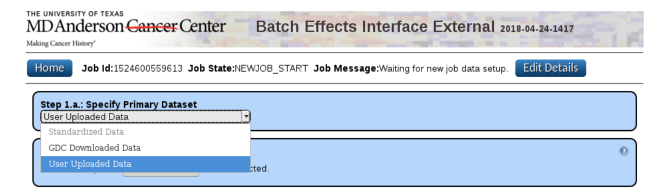
2 Starting a Job

When the Batch Effects Interface is first entered, the user is presented with a list of jobs (if any) and the option to Start New Job. (The external/public version of BEI does not support authentication and authorization.) The initial screen is shown below.



Upon selection "Start New Job", the user is presented with a new job page and the option to Update Job Info (covered later) and Step 1, selecting or uploading data, as shown below. The job id is shown in the Update Job Info section.





The next sections will follow each of these three options.

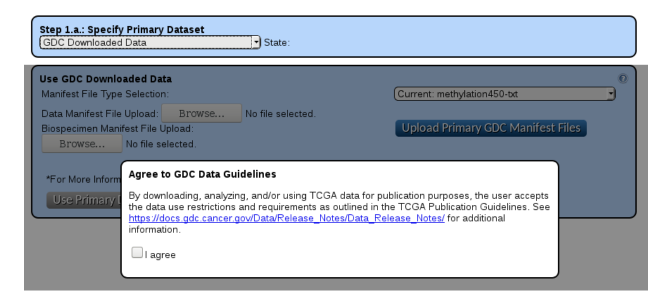
3 Downloading GDC Data

Upon selecting "Download Data from GDC", the user will be presented with the page below, asking them to agree to the TCGA and GDC release guidelines.

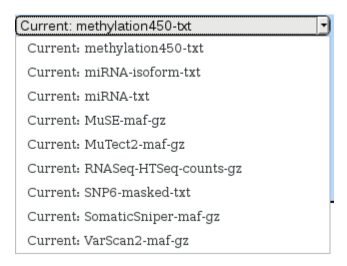
GDC Download Option:

Upload and use a manifest file from the GDC

Data Portal.



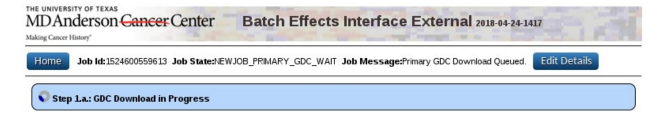
The user will need two manifest files from the GDC Data Portal https://portal.gdc.cancer.gov/. Data types supported for download and conversion are shown in the screenshot below. The manifest file corresponding to these data types is the "Data Manifest File".



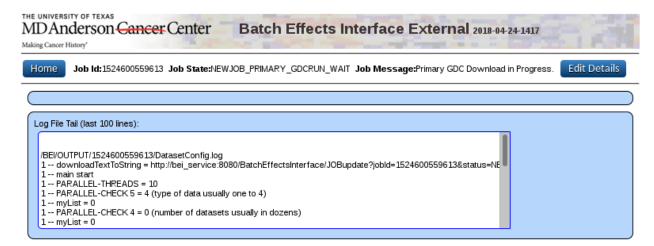
The second manifest file is the "Biospecimen Manifest File". This corresponds to the Data Category "Biospecimen" on the GDC site.

Currently, only batch information for the TCGA projects are supported.

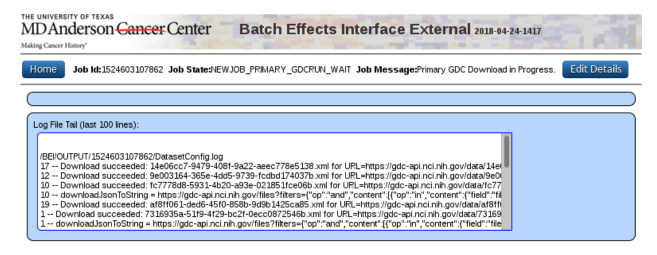
After uploading the Data Manifest File and Biospecimen Manifest File and pressing "Use Primary Downloaded GDC Data", the data will be downloaded and convert. The job will first go into GDC Data: Ready for download state. Depending on the setup, this could take a couple of minutes or more to process. The BEI Docker Stack is configured to run one download process at a time, due to memory constraints.



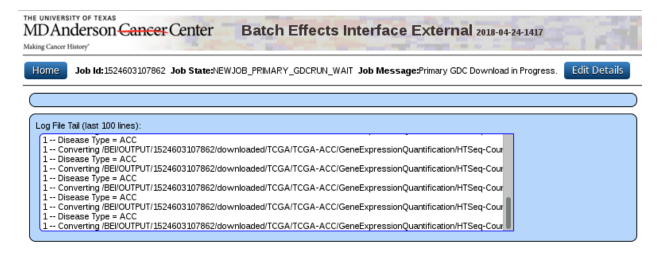
Once the download starts, the download log file will be tailed, as shown below.



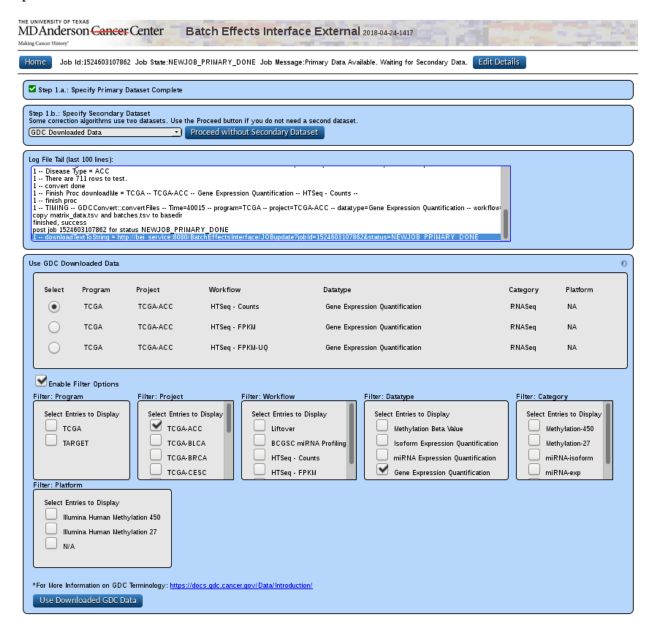
Here is another view of the log from further in the process.



Below, the downloaded files are being converted into a matrix format.

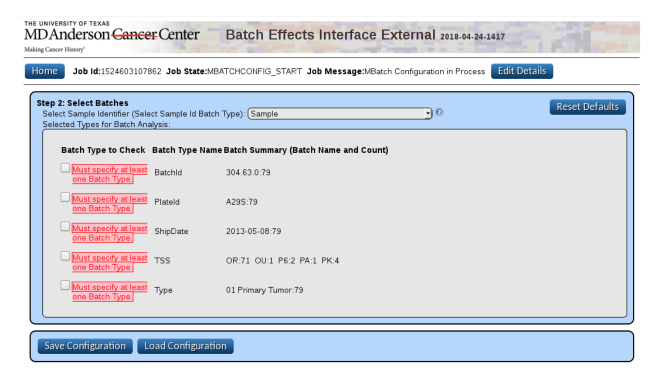


Once the download and convert are complete, the user will be shown the finished log, and the option to download or otherwise select a second dataset.

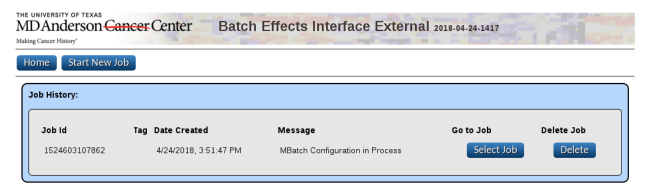


The secondary dataset is used for correction algorithms such as EBN-plus and RBN. For now, we will select the "Proceed without Secondary Dataset" button, to explore the basic assessment settings.

This will take the user Step 2: Select Batches for the MBatch Configuration, which is described later.



From here, hit the "Home" button, and you can see the job we just created which says "MBatch Configuration in Process".



4 Using Your Own Data

For this example, we will use data provided by the user. From the main job list, hit "Start New Job". The "User Uploaded Data" is selected by default for Step 1.a. Specify Primary Dataset.

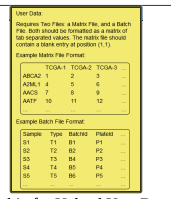


MBatch uses two different files (available from Standardized Data or to be provided by the user), for which the package provides code to read the files. The formats are described in the tooltip for User Data Uploaded section.

4.1 Standardized Data "Data Matrix" Format

The Standardized Data "Data Matrix" format is a tab delimited file. The first line of the file begins with a tab and contains sample identifiers. For Standardized Data, the sample identifiers are TCGA bar codes. Each subsequent row begins with a Feature Identifier and is followed by numeric data. Feature Identifiers are specific to the platform and explained later, but can be values such as Hugo Gene ids, probe ids, or microRNA identifiers.

This extract from the Data Matrix format shows four sample ids and five feature ids. Note that the first blank cell indicates the starting tab for the sample identifiers line. The features (left-most column) can be any set of unique strings. For proper processing, the rows and columns should be sorted.



Tooltip for Upload User Data

| | TCGA-OR-A5J2- | TCGA-OR-A5J3- | TCGA-OR-A5J6- | TCGA-OR-A5J7- |
|--------------------|----------------|----------------|----------------|----------------|
| | 01A-21-A39K-20 | 01A-21-A39K-20 | 01A-41-A39K-20 | 01A-21-A39K-20 |
| 14-3-3_beta-R-V | 0.211404 | -0.14778 | 0.220188 | -0.02738 |
| 14-3-3_epsilon-M-C | -0.03151 | -0.12861 | -0.0762 | -0.02275 |
| 14-3-3_zeta-R-V | -0.01203 | 0.032791 | -0.34541 | 0.136629 |
| 4E-BP1-R-V | 0.589134 | 0.365167 | 0.297887 | 7.34E-05 |

4.2 Standardized Data Batch File Format

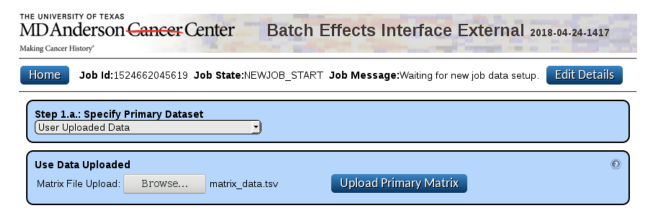
The Standardized Data Batch File format is also a tab delimited file. The first line of the file contains the sample id column id and batch type identifiers, none of which should contain spaces. The first entry should be the "Sample" column, which contains sample ids. For TCGA data (from the DCC and the GDC), the other batch type identifiers are Type, BatchId, PlateId, ShipDate, and TSS.

| Sample | Туре | BatchId | PlateId | ShipDate | TSS |
|-----------------------------|------|---------|---------|----------|--------------------|
| | | | | 5/7/201 | OR - University of |
| TCGA-OR-A5J2-01A-21-A39K-20 | 1 | 304 | A39K | 4 | Michigan |
| | | | | 5/7/201 | OR - University of |
| TCGA-OR-A5J3-01A-21-A39K-20 | 1 | 304 | A39K | 4 | Michigan |
| | | | | 5/7/201 | OR - University of |
| TCGA-OR-A5J6-01A-41-A39K-20 | 1 | 304 | A39K | 4 | Michigan |
| | | | | 5/7/201 | OR - University of |
| TCGA-OR-A5J7-01A-21-A39K-20 | 1 | 304 | A39K | 4 | Michigan |

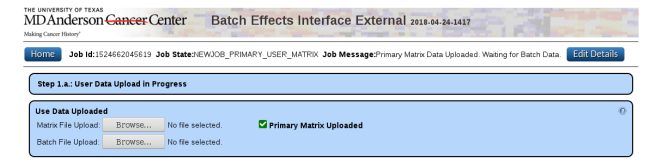
4.3 User File Uploads

The first part of Step 1.a. is to upload a matrix file. For this example, we downloaded the results of the GDC Download Example and extracted that file. (The results archive is described later.)

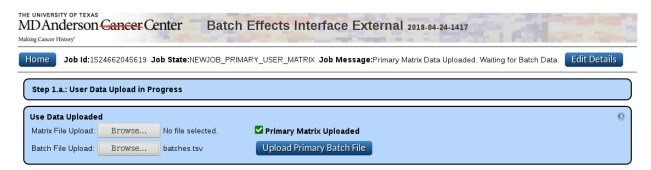
Pick Browse, select the matrix_data.tsv file (BEI will accept any file name provided and rename the file when uploaded) and click Upload Primary Matrix. The below screenshot shows the upload process.



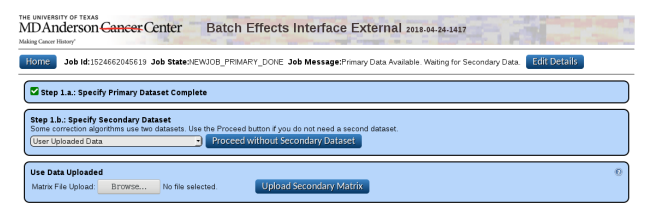
Then the user will be presented with the controls to upload a batch file (named batches.tsv in the downloaded archive).



Browse is used to select the file, after which an "Upload Primary Batch File" button allows the file to be uploaded.



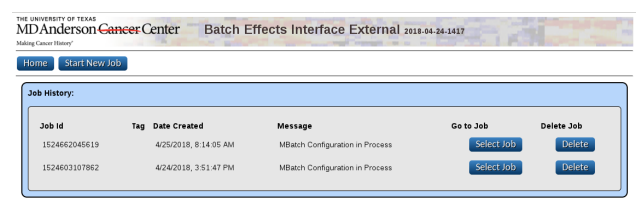
After the batch file is uploaded, the user is given the option of providing a second dataset.



We will select the "Proceed without Secondary Dataset" option. This will take the user Step 2: Select Batches for the MBatch Configuration, which is described later.

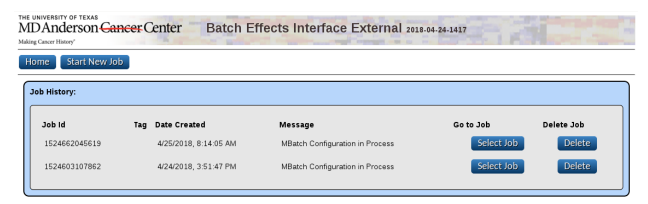


Then, we hit Home, to return to the job list, which now has two jobs.



5 Job Details

The current state of the job list is a little confusing. We have two jobs, in the same state, and with mostly similar Job Ids. Here, we will set the Job Details to make the jobs easier to tell apart. Let's start with the bottom job from 4.24, which is the one that used GDC Downloaded data. Click "Select Job" for that job.



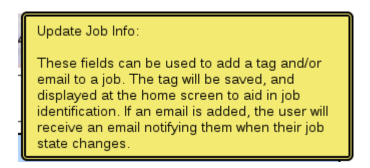
This takes us to the MBatch Configuration page.



But for now, just click the "Edit Details" button, which takes us to the Edit Details page.

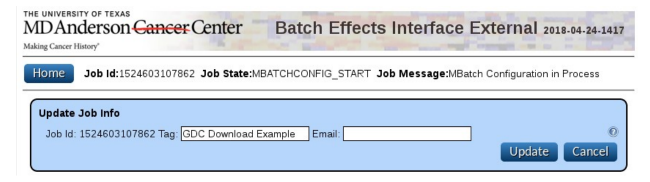


On this page, we hover over the tooltip question mark to see the help notice.

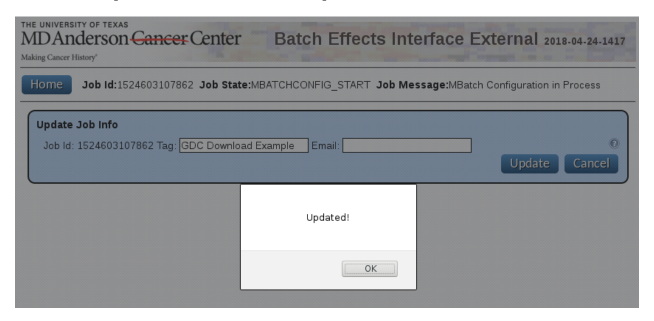


Since we are using the External Version, we have a Tag to describe the job, and an email field. Note, that emails may or may not be supported by your local installation.

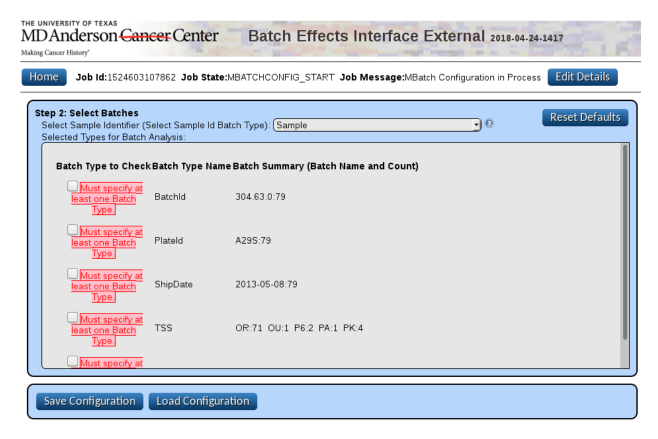
We will leave the email blank, but for the Tag put "GDC Download Example".



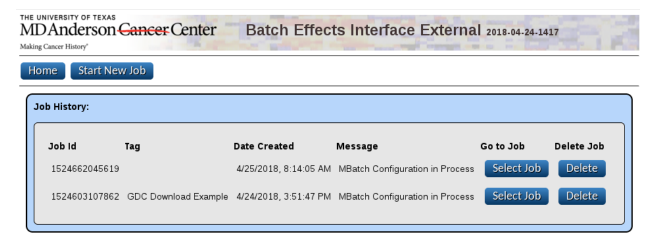
Then we hit "Update", which lets us know the update occurred.



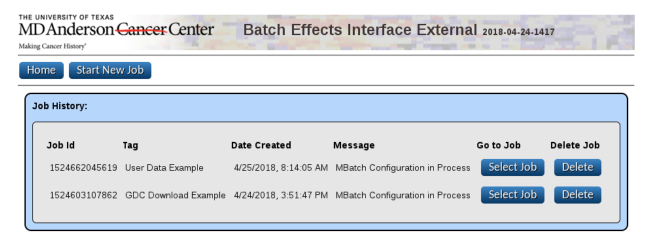
And returns us to the job page for that job, which right now is the MBatch Configuration page.



If we hit "Home", we get the job list, which shows the updated Tag for this job.

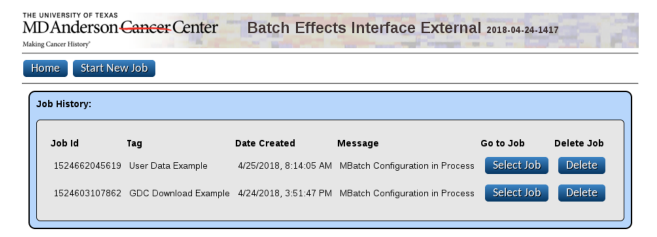


We can do the same process and add the "User Data Example" to the other job.



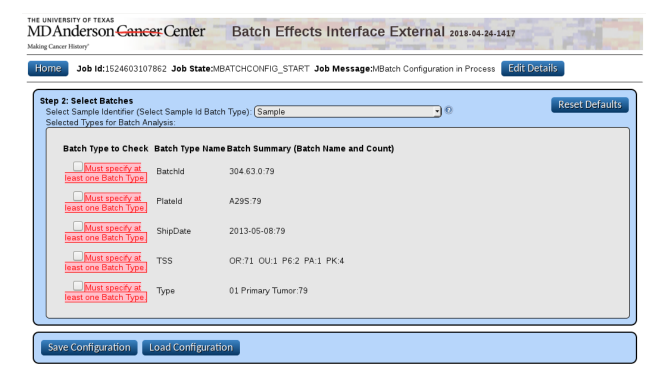
6 Configuring Assessments

This is based on the GDC Download data from above, but will be similar for each data type selected. From the Job List page, hit "Select Job" for the GDC Download Example job.

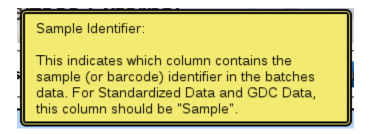


6.1 Step 2

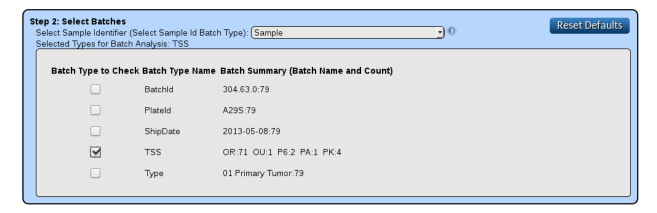
Below is the initial step in setting up assessments, Step 2: Select Batches, where the user selects the "Sample Identifier" from the batch file. For TCGA/GDC data, this is always "Sample". If the batch file does not contain a Sample column, the user will be prompted to select the Sample Identifier column from the Batch file.



The tooltip text for Select Sample Identifier is shown below.



Here, we need to select the Batch Type to check for batch effects. TCGA ACC Gene Expression data does not have particularly interested batches. In this case, we will pick TSS, since it has 5 different batches, while all the others have one.



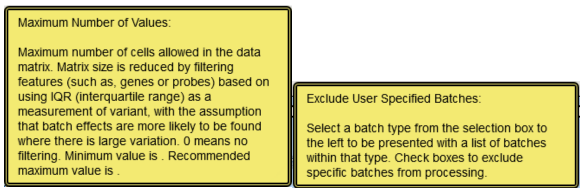
Once at least one Batch Type is selected, the rest of the MBatch Configuration steps are shown.

6.2 Step 3

In Step 3: Select Filtering Options, the user is asked how many data points to use and if there are batch to exclude.

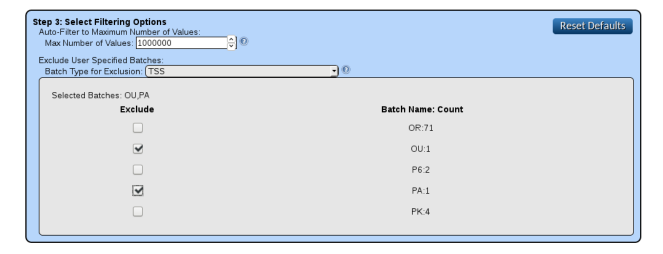


The Max Number of Values is the number of rows (features like genes) times the number of columns (samples). Features are dropped based on variance, since lower variance features are less likely to contain batch effects.



Tooltip Text for Step 3

Below, we reduced the Max Number of Values and selected to exclude two TSS batches, which have only one sample each. The user is not limited to selecting only the TSS batch type—any one batch type may be used for excluding data. The use-case for this is that a researcher has discovered a problem with a particular plate, so you elect to exclude it from processing.



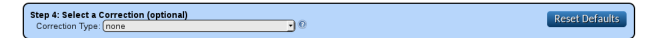
6.3 Step 4

In Step 4, a Correction is optionally selected. In this document, we will accept the default of "none" and only run assessments.

Correction Method:

Select a correction method and related settings for the correction. Note that not all correction methods are appropriate for all data and most will drop genes with NAs in their data. Some correction algorithms are based on correcting one set of data against a second data set.

Tooltip Text for Step 4



6.4 Step 5

Step 5 concerns values that affect PCA-Plus and DSC calculations. Each option has explanatory tooltip text, covered in detail below. For more details on PCA-Plus and the DSC value, please see http://bioinformatics.mdanderson.org/main/TCGABatchEffects:Overview#The DSC metric

sa

| Step 5: Select PCA-Plus/DSC Ar | guments | Reset Defaults |
|--------------------------------|----------|----------------|
| Number of DSC Permutations: | 2000 🖶 💿 | Neset Belautes |
| Number of DSC Threads (1-5): | 5 🔛 💿 | |
| Minimum DSC Batch Size: | 5 🔛 💿 | |
| DSC Random Number Seed: | 314 | |
| Maximum Number of Features: | 5000 🔄 ◎ | |

Tooltips for Step 5 are shown below.

| The Number of Dispersion Separability Criterion (DSC) Permutations refers to the number of permutations used to calculate the p-value associated with the DSC. In general, for TCGA/GDC-like data, we recommend no | Number of DSC Permutations: Permutations performed to calculate the DSC and associated p-values. Minimum of 500, recommended no less than 2000 for valid |
|---|---|
| less than 2000 permutations. | results. |
| The number of DSC threads is related to the size of your data and the memory and cores available on your machine. For most TCGA/GDC data sets, 5-10 threads and 16-32GB of memory should be sufficient. | Number of DSC Threads: Number of threads (usually equal to the desired number of cores) used to perform DSC calculations. Generally, use the default value. |
| The minimum batch size to consider for inclusion in DSC processing. Small outlier batches can greatly influence the outcome. We generally recommend 5 for TCGA/GDC-like data. | Minimum DSC Batch Size: Enter a minimum batch size acceptable as a usable batch (depends on correction algorithm). |
| The seed used for performing DSC permutations. This is needed in order to reproduce results from other researchers. | DSC Random Number Seed: Random number seed used in permutations and necessary for reproducible results. |
| The Maximum Number of Features (such as genes or probes) to use for DSC calculations. Reduction in size is based on IQR (interquartile range) as a measurement of variance. (Generally, batch effects are found in areas of high variance.) | Maximum Number of Features (DSC): Maximum number of features (such as, genes or probes) for DSC computations. Size is reduced based on using IQR (interquartile range) as a measurement of variant, with the assumption that batch effects are more likely to be found where there is large variation. Minimum recommended value is 1000. Maximum recommended value depends on your setup. |

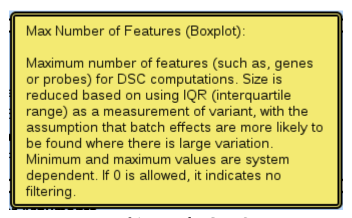
Tooltip Text for Step 5

Here we have reduced the number of permutations and the number of features, to make our run finish sooner.



6.5 Step 6

In Step 6, the number of features (such as, genes or probes) to be used with the boxplot assessment algorithm are designated. In general, we limit this to 5000 features, as more features takes excessive amounts of memory and processing time for TCGA/GDC-like data.

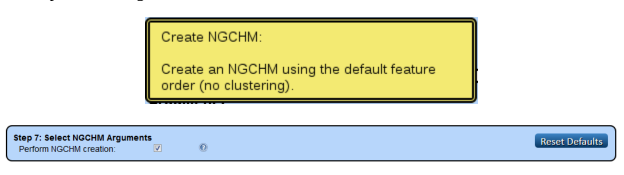


Tooltip Text for Step 6



6.6 Step 7

Step 7 is selecting the optional creation of an NGCHM (Next Generation Clustered Heatmap). Heatmap is built using default values.



6.7 Configuration Options and Do MBatch Run

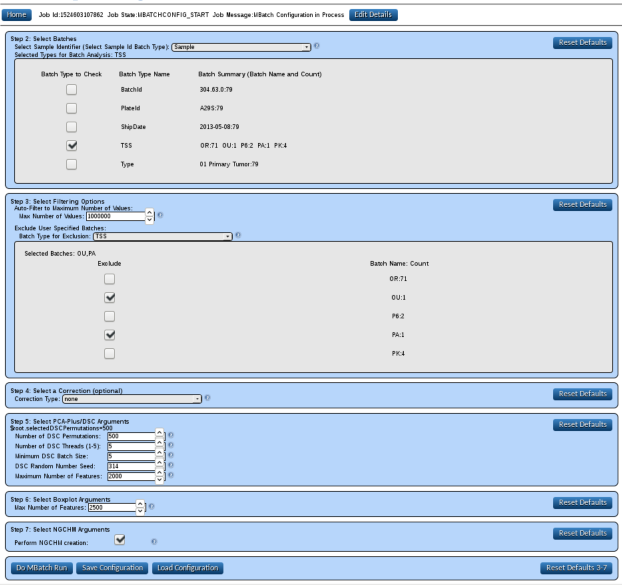
Two different sets of buttons appear at the bottom of the page. Until Step 3 is complete, the user has only the options of Save Configuration and Load Configuration. This allows the user to save the current MBatch configuration, if they need to come back to it later (for example, after getting a list of samples to exclude). Load Configuration loads the last saved configuration.



Once Step 3 is complete, the user can also press the Do MBatch Run button, to start the run.



6.8 Complete Page View



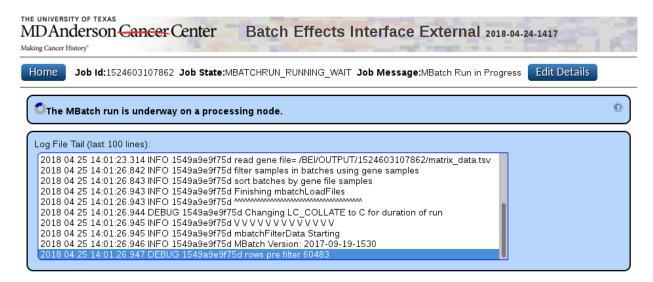
6.9 Do MBatch Run

After finishing setting up the assessments, the "Do MBatch Run" button is pressed. The page will update and go into the "MBatch Run Queued" state, seen at the bottom of the screenshot below. The page will automatically update as needed.

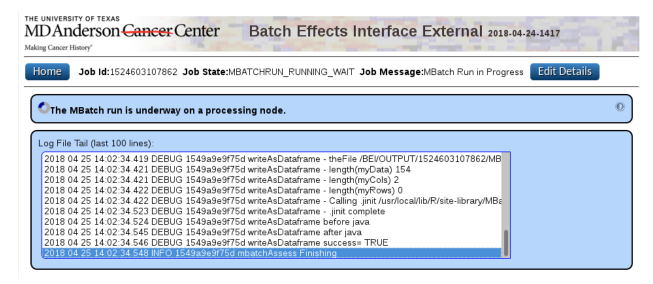


6.10 During the MBatch Run

The MBatch run starts, which could take up to 2 minutes in an unused system or until after other users in the queue are finished (which depends on your system load). After starting, the last 100 lines from the log file are displayed. Below shows the start of the process, with the two data files from the job being loaded. (Updating log files to have more user friendly output is also on the future list of enhancements.)



The assessment portion of MBatch finishes with the line marked below saying "mbatchAssess Finishing". The NGCHM creation follows this. The NGCHM creation log is not currently shown during processing, but is on the list to be added.



6.11 Finished Job

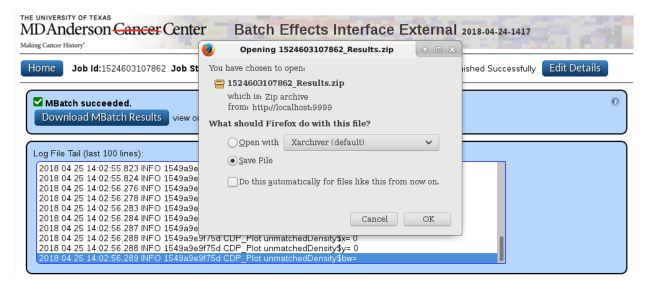
After a job has finished, the user is presented with a "Download MBatch Results" button that offers the ability to download the complete job directory, with data used, downloaded, configuration files, and results. (The Download option is described in more detail later.)

If successful, the user will also be presented with a link to the Batch Effects Viewer website, to view the results.

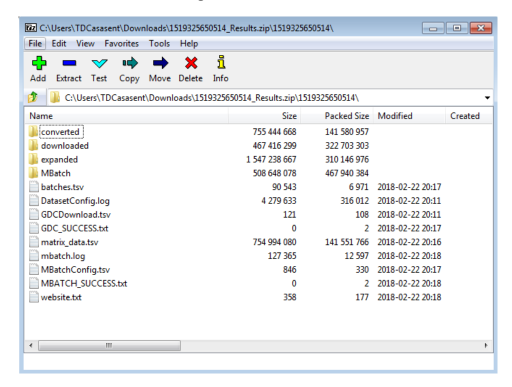


7 Download MBatch Results

If the user presses the Download MBatch Results button, the job directory will be compressed and sent to the browser. (The exact download process will vary by browser. This screenshot is Firefox setup to ask the user to Open or Save.)



Within the ZIP archive are the following directories and files.



The directories can mostly be ignored by users. The MBatch directory has results that are used for display by the Batch Effects Viewer. To save space, the converted, downloaded, and expanded directories can be deleted if desired. The downloaded directory contains the files downloaded from the GDC. The expanded directory contains the uncompressed version of the GDC files. The converted directory has the converted data, batch file, covariates file with information like demographics, and annotations file with information like the value added before log normalization, if performed.

| Filename | Contents |
|--------------------|---|
| batches.tsv | Batch file for data. See user data documentation for file format. |
| DatasetConfig.log | Log from downloading GDC data. This is not particularly user- |
| | friendly at present. |
| GDCDownload.tsv | Contains a description of the data downloaded from the GDC. |
| GDC_SUCCESS.txt | An empty "flag" file indicating successful completion of the |
| | GDC Download process. |
| matrix_data.tsv | Data file for data. See user data documentation for file format. |
| mbatch.log | Log from MBatch run. This is not particularly user-friendly at |
| | present. |
| MBatchConfig.tsv | Configuration file for MBatch assessments and corrections. |
| MBATCH_SUCCESS.txt | An empty "flag" file indicating successful completion of the |
| | MBatch run. |
| website.txt | Timestamp and links to the original job and output. |

8 View MBatch Results

For details on the Batch Effects Viewer website, please see http://bioinformatics.mdanderson.org/main/TCGABatchEffects:Overview

This document will be updated when a new version of the Batch Effects Viewer, currently in development, becomes available.

9 Job List

Pressing the Home button takes the user to the main page, with a list of jobs.

Clicking on the Select Job button takes the user to the results page, which also has the Update Job Info. Jobs can also be deleted. The external version of Batch Effects Interface has not security on viewing, running, or deleting jobs.