## **Evaluation of DSP Normalization Methods – Tutorial**

Chi Wang and Daheng He Markey Cancer Center, University of Kentucky (November 2022) The App provide a convenient interface for users to normalize the data of NanoString GeoMx protein assay in multiple ways, and evaluate their performance in order to choose an optimal normalization method for downstream analyses. The initial input panel on the left-hand side of the App looks like this:

Evaluation of DSP Normalization Methods							
(	Tutorial and sample data can be found here )						
Step 1: upload raw DSP data (.csv only), with the 1st column being protein names:							
Browse	No file selected						
Step 2: selec	et negative control(s) (e.g., Ms lgG1 / Ms lgG2 / Rb lgG):						
Step 3: selec	et housekeeping control(s) (e.g., GAPDH / Histone H3 / S6):						
Step 4: uploa	ad your annotation table (.csv only):						
Browse	No file selected						
Step 5.a: sel	ect the 1st subtype for pair-wise comparison:						
Step 5.b: sel	ect the 2nd subtype for pair-wise comparison:						
Step 6: view	results in each panel for the sample:						
N/A	•						
<u></u> Downloa	ad normalized data (.xlsx)						

The users are expected to provide necessary info in this panel following the indicated steps 1 to 6 sequentially. At the last step, i.e., Step 6, upon making a

choice of sample of inquiry, results will be automatically generated in each panel on the right-hand side of the panel. Here are step-by-step graphical instructions on each input step:

### Step 1: upload raw DSP data:

This step is to upload the raw DSP expression data, which has only been ERCC-normalized, to remote server. As indicated in the input statement, currently the App can only process the raw expression data table in csv (comma-separated values) format. In preparing the csv file, please make sure that the first column is for protein IDs, and the rest of columns for protein expression values of ROIs. The basic structure of raw expression data input should look something like this in Excel:

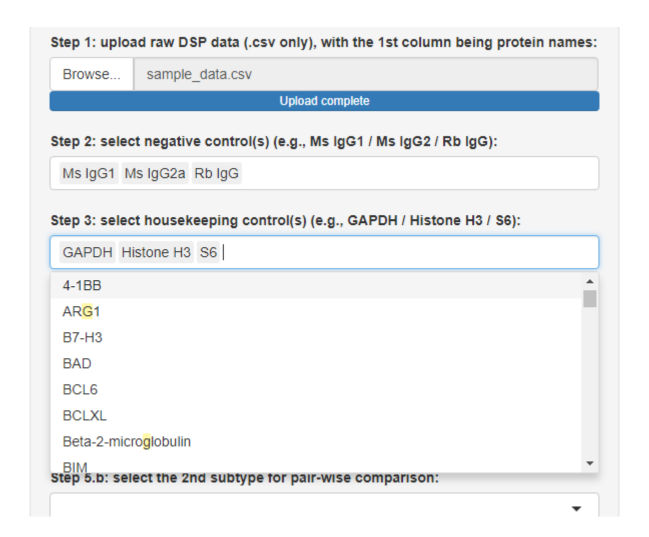
Protein_Name	ROI_001	ROI_002	ROI_003	ROI_004	ROI_005	ROI_006	ROI_007	ROI_008
Ms IgG1	22.6	18.58	28.51	17.25	12.52	16.47	22.59	1441.5
GAPDH	1223.9	1225.8	1692.9	747.7	786.8	1164.6	1300.4	7952.1
BRAF	2.05	6.12	2.95	1.38	3.54	1.35	4.22	3669.9
BIM	43.01	49.84	57.49	30.16	25.11	39.31	41.66	29.49
S6	2816.8	2718.9	3711.9	1857.4	1942.1	2922.3	3711.9	42.92
BCLXL	391.76	436.63	510.67	252.43	229.19	377.8	501.67	42.16
BCL6	46.58	56.34	82.24	33.64	26.59	52.18	45.48	42.5
ARG1	89.84	89.22	128.8	45.19	58.02	67.92	95.82	112.3
4-1BB	42.55	44.78	62.85	25.55	14.28	41.72	32.19	540.63

When the raw expression datafile in csv format is ready, just click on the "**Browse...**" button to initiate the local file path browsing and uploading process. You are only allowed to upload a single csv file at a time.

### Step 2: select negative control(s) and Step 3: select housekeeping control(s):

The App will monitor the uploading process. Once the expression data has been uploaded successfully in step 1, the App will automatically fetch the protein names from your data and make them available for step 2 and step 3 in the form of dropdown menu, in which you may browse and click to make multiple choices to let the App know which proteins are the negative controls and housekeeping controls (typically there are 3 negatives controls and 3 housekeeping controls). To

speed up the protein names searching, you may simply type the first two or three letters of each wanted protein name in the text box, the App will narrow down the list of names that match your typing. Here is what step 2 and step 3 look like during operation:



### Step 4: upload your annotation table:

As an important component of DSP data set, you need to provide an additional table in csv format, which provides annotating info for each ROIs in the raw expression table, such as the area and nuclei count of each ROI, which ROIs belong to the comparison group "A" and which belong to the comparison group "B". The uploading of ROI annotation table is very similar to the uploading of raw

expression data -- just click on the "**Browse...**" button to initiate the local file path browsing and uploading process.

In our realistic practice of DSP data analysis, we have notices that in many DSP data multiple subjects were involved in DSP data collection, and the tissue slides from each subject were often prepared and measured at different times, under different lab conditions, or even by different machines, these technical variations together with the genuine biological difference among subjects often make the subject-related heterogeneity in DSP data quite sizeable, and may significantly affect the conclusions of downstream analyses. Such a heterogeneity must be properly identified and handled. Therefore, we are making it mandatory to provide the subject info in the annotation table, even if your data really just come from a single subject.

Each column in the annotation table corresponds to a specific kind of annotating info, such as ROI area, nuclei count, subgroup and subject each ROI belongs to. You need to arrange your annotation table in such a way that its rows, from top to bottom, are in exactly the same order as those columns (i.e., ROIs) from left to right in the raw expression data you uploaded in step 1. The basic structure of annotation table input should look something like this in Excel:

subject	area	count	group
A20_102_A40	91208.3	546	COVID19 Positive
A20_102_A40	99797.2	561	COVID19 Positive
A20_102_A40	118124.3	658	COVID19 Positive
A20_102_A40	58731.3	344	COVID19 Positive
A20_104_A31	75969.5	251	COVID19 Negative
A20_104_A31	67738.7	294	COVID19 Negative
A20_104_A31	71717.9	372	COVID19 Negative
A20_104_A31	55765.7	309	COVID19 Negative

As shown above, a complete annotation should contain 4 columns (the order of these columns in your annotation table do not matter), named exactly as "subject", "area", "count", "group", all are case-sensitive. Among the 4 columns, the "subject" and "group" columns are mandatory. In the case of single subject without a pre-specific subject ID, you still need to have a column named as

"subject", and just put some arbitrary short string, for example "S1", as the subject's ID in all of the row entries of the column. The "area" and "count" columns, on the other hand, are optional. If these two types of information were not recorded at all in the data collection steps, then you can simply ignore these columns, and just submit an annotation table with only the two mandatory columns "subject" and "group". In the absence of either or both of the two optional columns, the App will skip the evaluations that require the information. There is yet another situation, in which you must ignore the "area" column — sometimes the DSP data may be collected from the ROIs with identical areas, in which case the area info cannot provide any meaningful value in the evaluation of normalization methods, and more importantly, providing an "area" column with all identical values will make several internal statistical evaluations divergent, triggering unexpected error messages and failure of the App.

# Step 5.a: select the 1st subtype for pair-wise comparison and Step 5.b: select the 2nd subtype for pair-wise comparison:

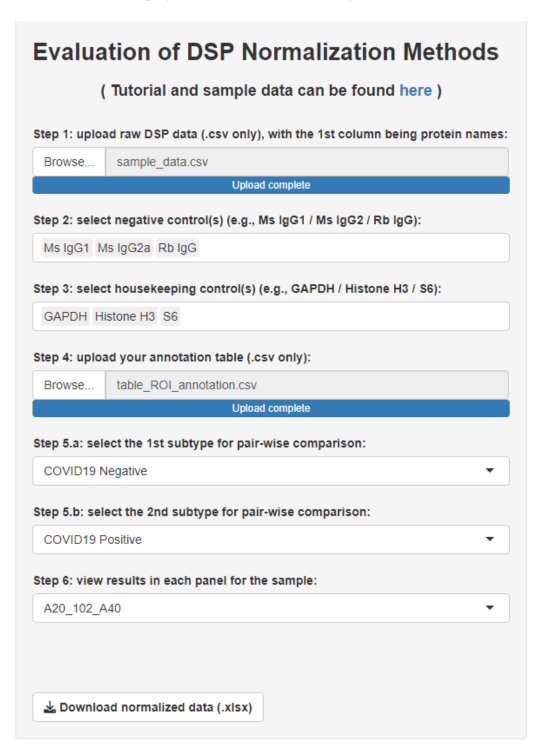
After uploading the ROI annotation table in step 4, the App will automatically scan the mandatory column "group" to summarize the unique values in it, and make them available for step 5.a and step 5.b in the form of dropdown menu, in which you may browse and click sequentially to let the App know between which two biologically distinct subgroups you would like to compare. In each of step 5.a and 5.b, you are only allowed to make a single choice of subgroup -- currently the App only supports two-group comparison.

### **Step 6: view results in each panel for the sample:**

After loading the ROI annotation table in step 4, the App will also scan the mandatory column "subject", summarizing its unique values in a dropdown menu, and by default choosing the first subject ID alphabetically as the input of subject for evaluation. At this stage, all of the required inputs are already fed to the App, therefore the evaluation processes in the background will be initiated automatically. You can start to browse each output panel on the right-hand side of the webpage to view its detailed output. You can choose a different subject from the subject menu, and all output panels will respond to the change of subject

simultaneously. Please note that in some panels, computing and generating figures may take a while, so please allow a few seconds for the updating of figures.

Upon providing all the necessary inputs to the App correctly, the input panel on the left-hand side of the webpage should look something like this:



#### Download normalized data:

Upon providing all the necessary inputs to the App from step 1 – 6, the App will generate the normalized DSP expression matrices based on the build-in normalization methods. There are in total 17 build-in normalization methods, but depending on your input, if the area and/or nuclei count info is absent, certain normalization methods requiring such info will not be available. You can simply click on the button "**Download normalized data (.xlsx)**" to download the normalized data in xlsx format containing multiple sheets, among which the first sheet is the table giving detailed descriptions of the normalization methods, those methods that are not available due to the absence of area or nuclei count info are marked with "(not available)", the rest of sheets are for the normalized expressions by each available normalization method. You can choose the data, which is normalized by the most optimal normalization method as decided by those evaluation metrics in the output panels, for your downstream analysis for more reliable conclusions.

### **Acknowledgement:**

In searching for relevant DSP datasets to test our models, the research groups of Desai N et al [1] and Delorey TM et al [2] kindly shared their DSP data and provided help with explaining their DSP data via private communications. We deeply appreciate their kind help.

### **Reference:**

- [1] Desai N, Neyaz A, Szabolcs A, Shih AR et al. Temporal and spatial heterogeneity of host response to SARS-CoV-2 pulmonary infection. Nat Commun 2020 Dec 9;11(1):6319. PMID: 33298930
- [2] Delorey TM, Ziegler CGK, Heimberg G, Normand R et al. COVID-19 tissue atlases reveal SARS-CoV-2 pathology and cellular targets. Nature 2021 Jul;595(7865):107-113. PMID: 33915569