



FEB 06, 2024

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Protocol Citation: Hui Xie 2024. Sinai SCENT TMC - Olink NPX manager for Olink Data Analysis.

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Protocol status: Working

We use this protocol and it's working

Created: Feb 02, 2024

Last Modified: Feb 06, 2024

PROTOCOL integer ID: 94624

Sinai SCENT TMC - Olink NPX manager for Olink Data Analysis

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ABSTRACT

To describe the procedures for using Fluidigm “Real-Time PCR Analysis” software and “Olink NPX manager” software to analyze Olink Assay data and generate final result reports. Fluidigm Real-Time PCR Analysis software generates the Ct value of the PCR reaction and Olink NPX manager software calculates the Normalized relative Protein Expression value (NPX) from the Ct value.

MATERIALS

- Computer
- Fluidigm Real-Time PCR Analysis
- Olink NPX manager

SAFETY WARNINGS



Not Applicable

Raw data Ct value analysis

- 1 Copy the Biomark File (entire Chip-run folder corresponding to the Olink assay) from the Biomark Computer to USB drive, the size (~1.3 GB). There should be always 2 sub-folders and 4 files in the Chip run folder.

 Cals	8/29/2017 5:33 PM	File folder
 Data	8/30/2017 11:21 AM	File folder
 ChipRun.bml	8/29/2017 5:09 PM	BML File 343 KB
 ChipRun.bml.orig	8/29/2017 5:09 PM	ORIG File 343 KB
 ChipRun.Processed.bin	8/29/2017 5:09 PM	BIN File 18,006 KB
 ChipRunLog.txt	8/29/2017 5:09 PM	Text Document 1,471 KB

- 2 Connect USB drive from above step with HIMC computer with installed Fluidigm Olink PCR analysis software
- 3 Start the Fluidigm "Real-Time PCR analysis software (Version 4.5.2)"
- 4 Open the Chip run under "Quick Tasks". This will redirect to a new window. Locate and double click the Chip run file (.bml-format) in the copied folder in Step 1.

Notes: Only one Chip run-file can be open at a time. Before opening a new run-file, the first chip run file must be closed.

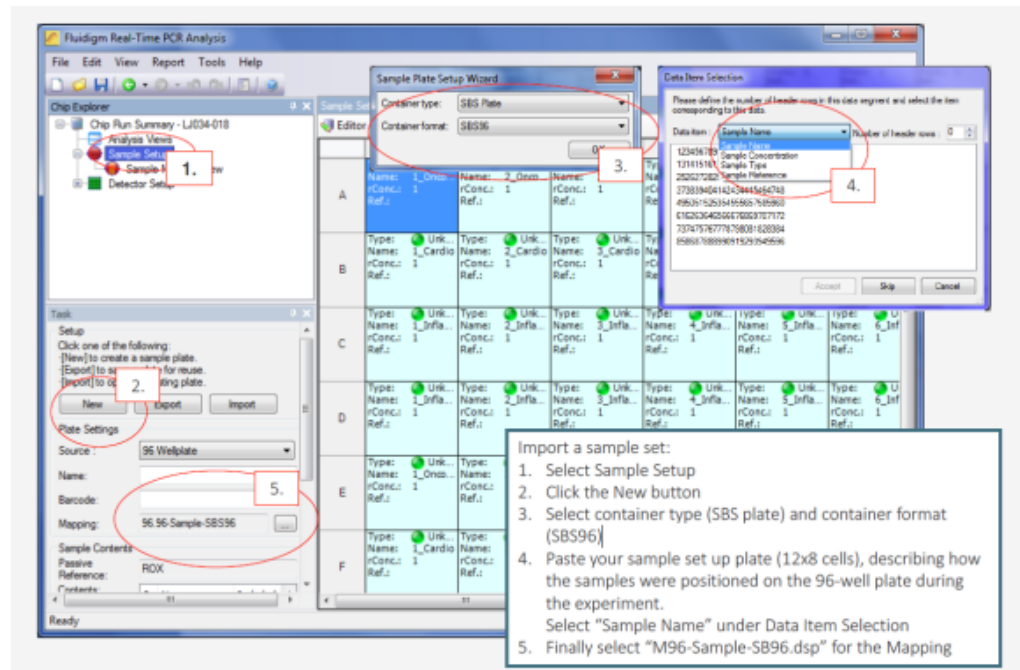


5 Import a sample set.

- 5.1 Prepare a 96-well-plate sample layout for the Olink Assay in excel according to HIMC format and save in HIMC shared drive or Google drive.
- 5.2 Double click Sample Setup in the Fluidigm Olink PCR analysis software. Select container type (SBS plate) and container format (e.g. SBS96) in the Sample Plate Set-up Wizard, press OK.
- 5.3 Copy the Olink assay sample layout from the plate layout excel file saved in Step 5.1, right click well A1 in the Sample Set-up "Editor" panel.
- 5.4 Select "Sample Name" under Data Item Selection.

- 5.5** Write Run name in the “Name” section under plate settings. Select mapping will open a new window. Different mapping files will be seen in that window.

- 5.6** Select “M96-Sample-SB96.dsp” for the Mapping.



6 Import a Detector setup:

- 6.1** Select Detector Setup and Click the New button.

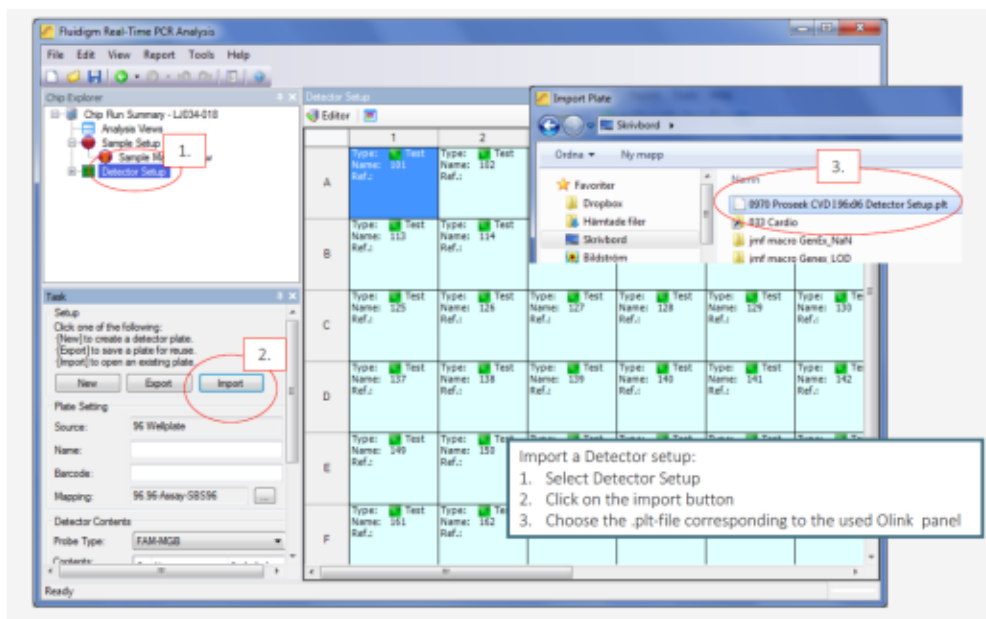
- 6.2** Click on the import button. Choose the .plt-file corresponding to the Olink panel kit used for the assay (i.e. for Inflammation panel use “Mapped Inflammation I 96x96 Detector Setup.plt”).

Notes: Check the Olink kit contents Paper insert for Lot # and the proper .plt file information.

New version of .plt file may need to be downloaded before analysis based on kit lot#. All the assay .plt files from Olink are downloaded from Olink website and saved on:

J:\gtherapy\HumanImmuneMonitoringCenter\Olink_proteomic2016\NPX_analysisSoftware\mt Sinai

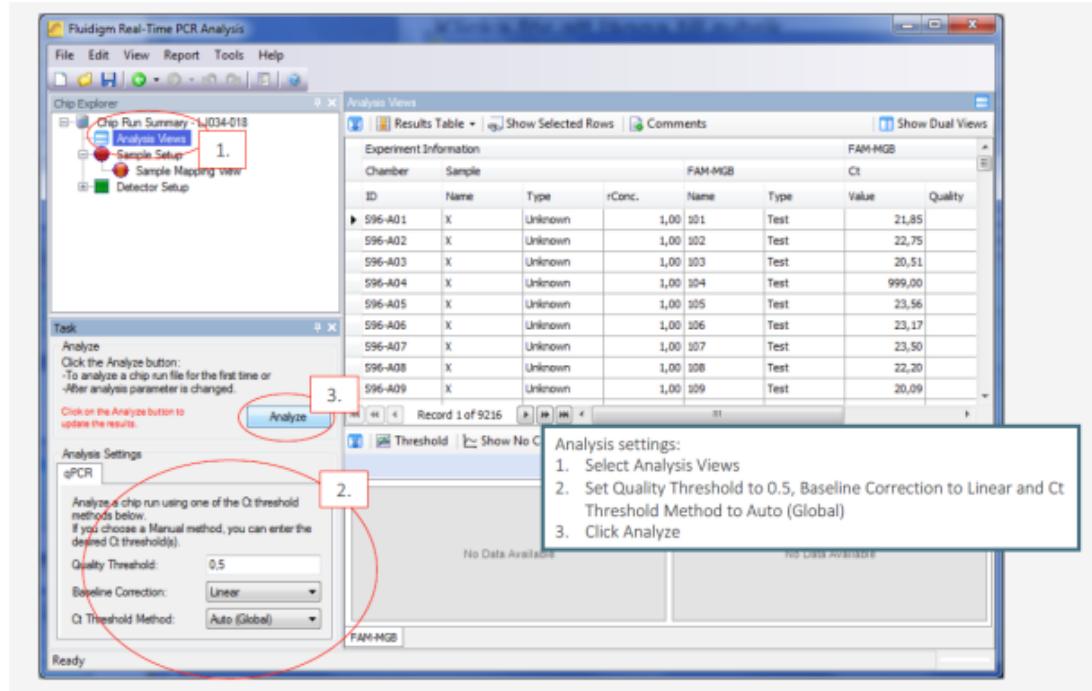
- 6.3** In the case of a project with different lot # kits, one will need to have the same reference samples in plates with biological samples for normalization purposes. However, if the “Data File No: XXXX” which comes in the paper with the kit is the same, then the data from the different kit lot numbers can be analyzed as a project.



7 Analysis settings:

7.1 Select Analysis Views/ Details Views.

7.2 Set Quality Threshold to “0.5”, Baseline Correction to “Linear” and Ct Threshold Method to “Auto (Global)”, (by changing the default settings) and Click Analyze button. These settings are common for all the Olink panels.

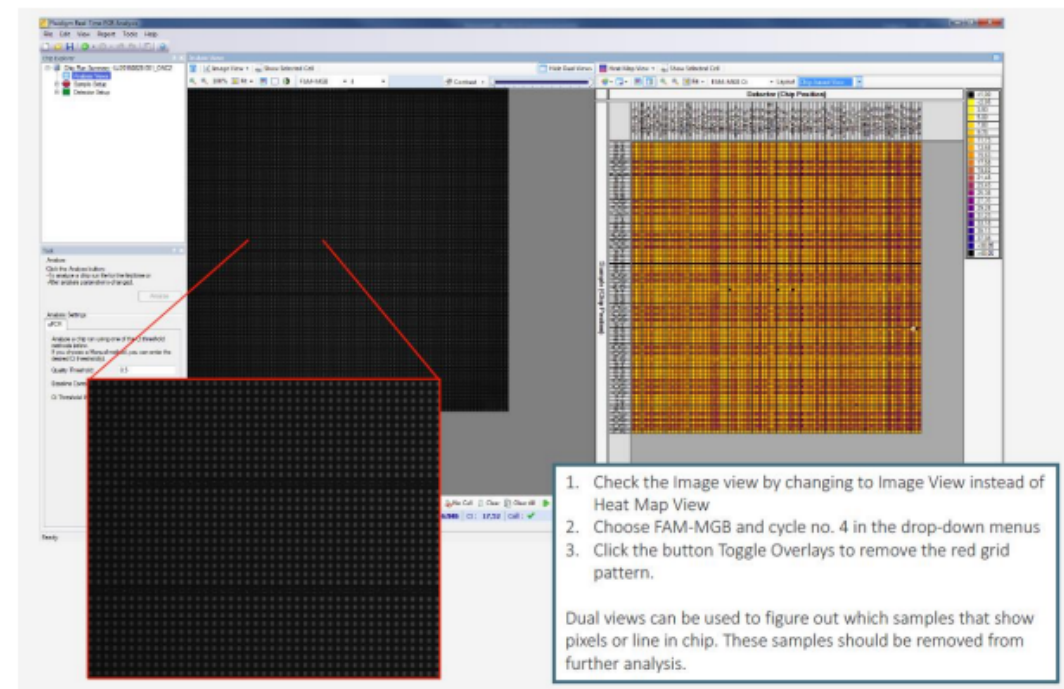
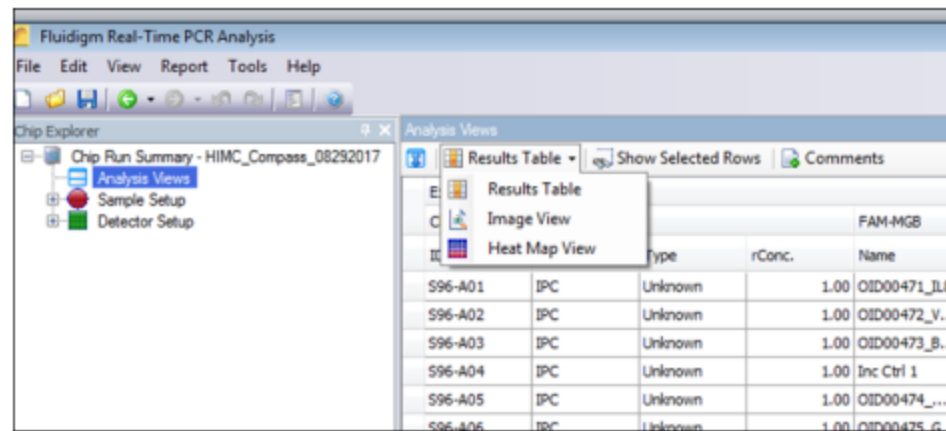


8 Data preview:

8.1 Select "Results Table and drop down. Select Heat Map View.

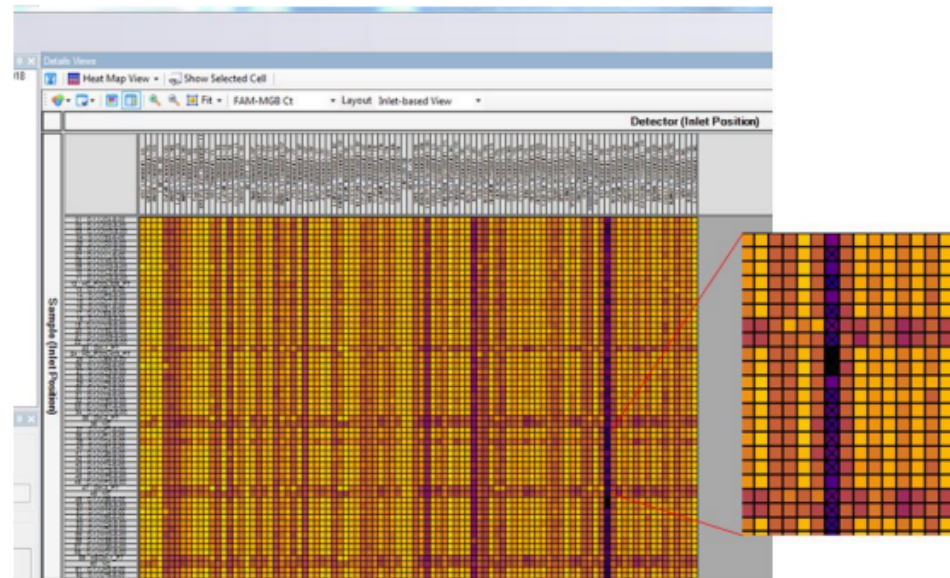
8.2 Review the Heat Map. The Negative Ctrl samples should give a darker “line” from left to right and IPC samples should give a brighter “line” from left to right (Bright color corresponds to more fluorescence, i.e. a lower Ct value and therefore a higher protein concentration). Check if any line of samples/assays has a Black square point that means the assay failed in that Chip well.

8.3 Check the Image view by changing to Image View instead of Heat Map View. Choose FAM-MGB and cycle no. 4 in the drop-down menus. Then Click the button Toggle Overlays (square) to remove the red grid pattern. Dual views can be used to figure out which samples that show pixels or line in chip. These samples should be removed from further analysis.



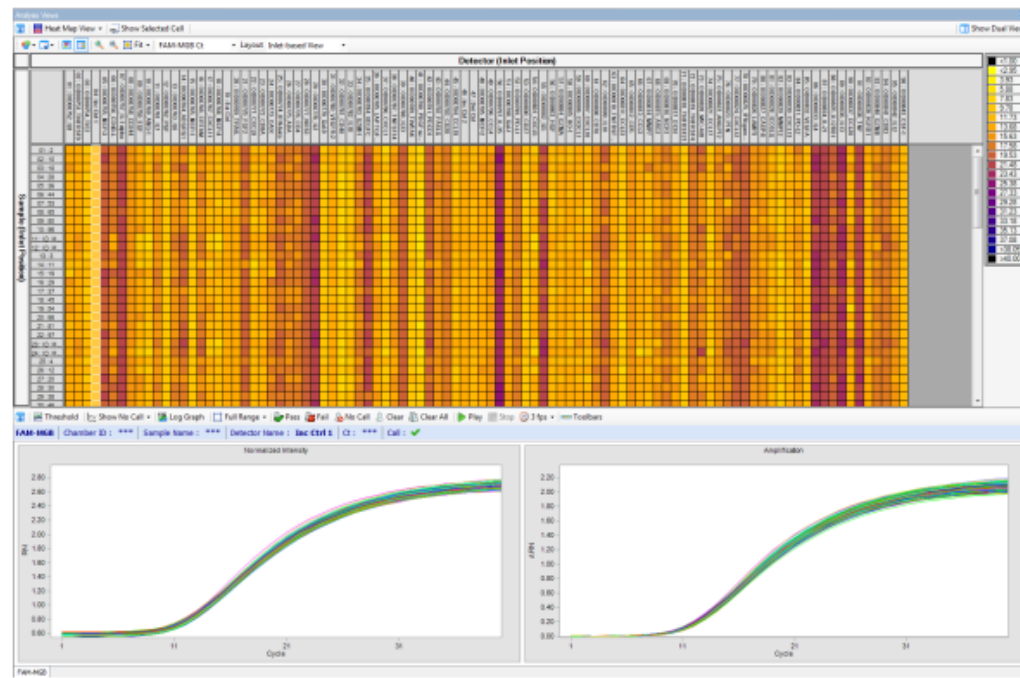
8.4

Check the Heat Map View by selecting the Heat map view tab on the top of detail view. If there is any analyte or sample has dark purple line with black crossed out square, then remove the data of this analyte/sample in the Olink NPX manager for further analysis. Follow the Step 28 to remove the data from NPX manager.

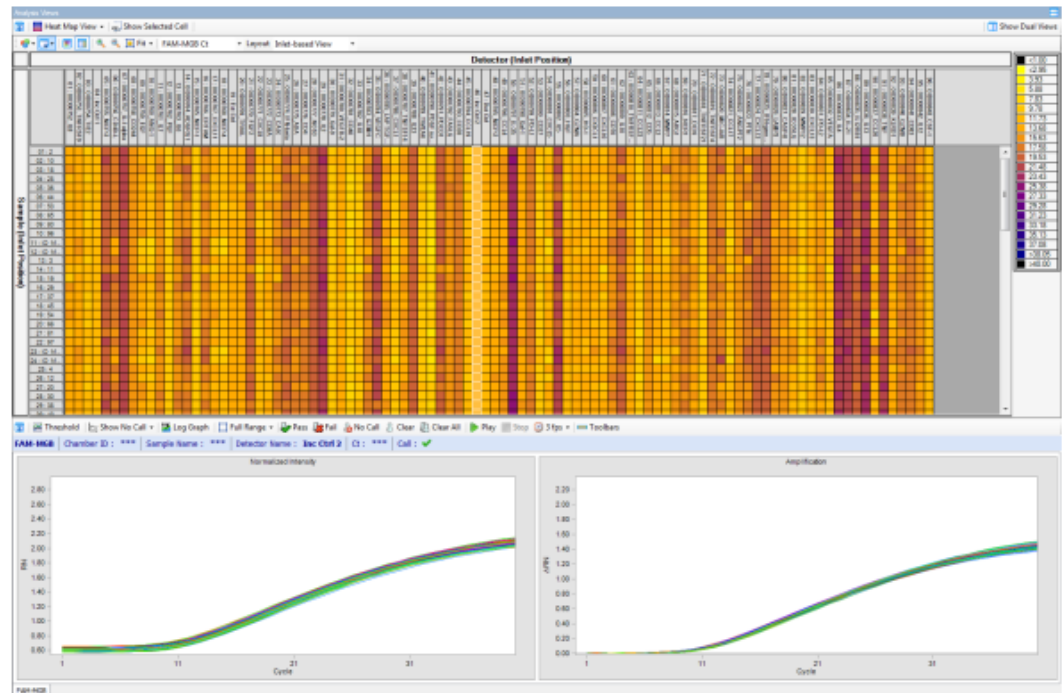


Note: The following figures are examples of different Ctrl/samples/analyte conditions and can be used for reference view only, while analyzing other plate data. These examples are taken from Olga_IO_10162017.

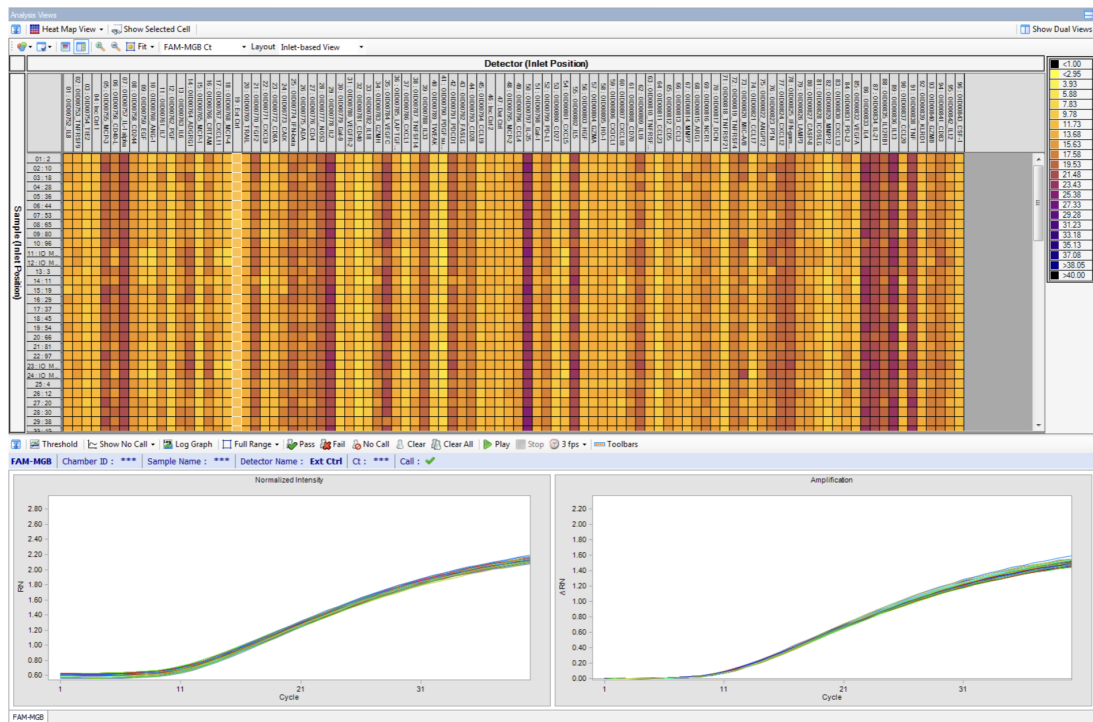
8.5 Select Incubation Ctrl-1 in the Detector on heat map and it will appear as the following figure with the amplification curves.



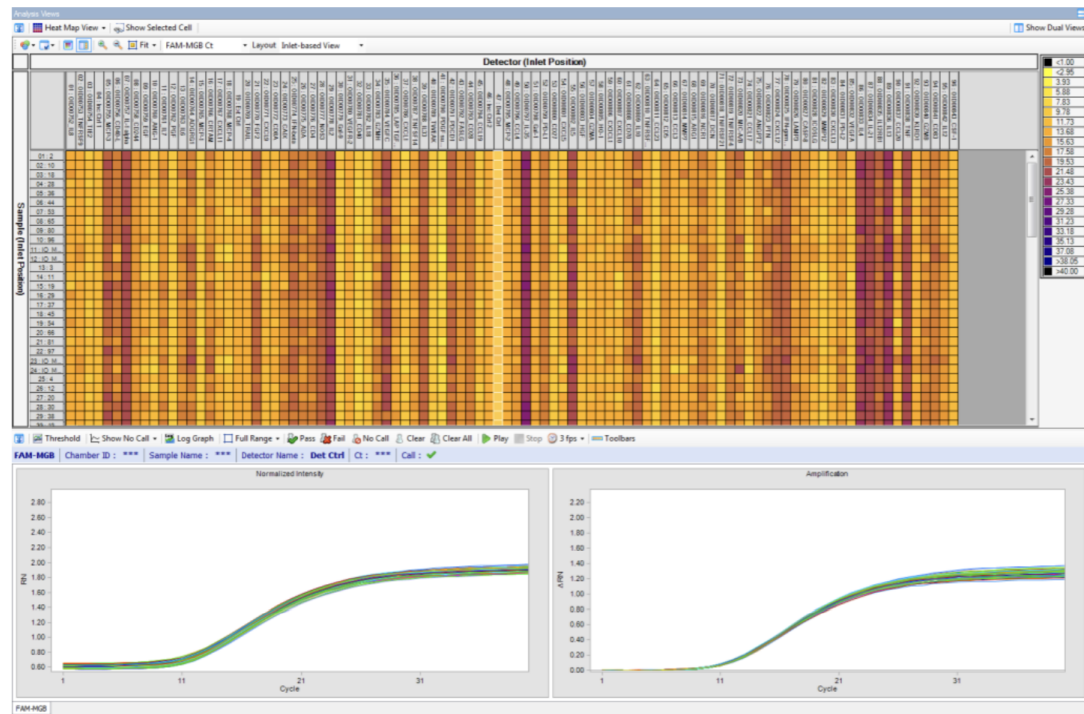
8.6 Select Incubation Ctrl-2 in the Detector on heat map and it will appear as the following figure with the amplification curves.



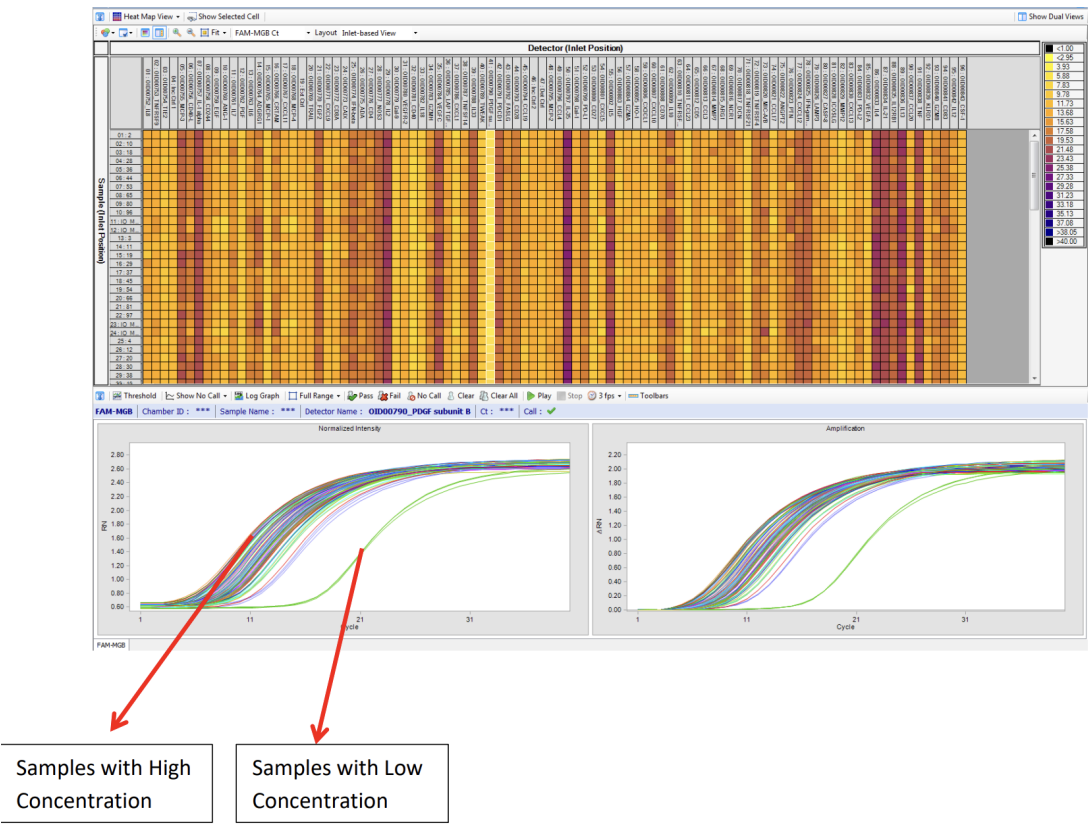
8.7 Select Extension Ctrl in the Detector on heat map and it will appear as the following figure with the amplification curves.



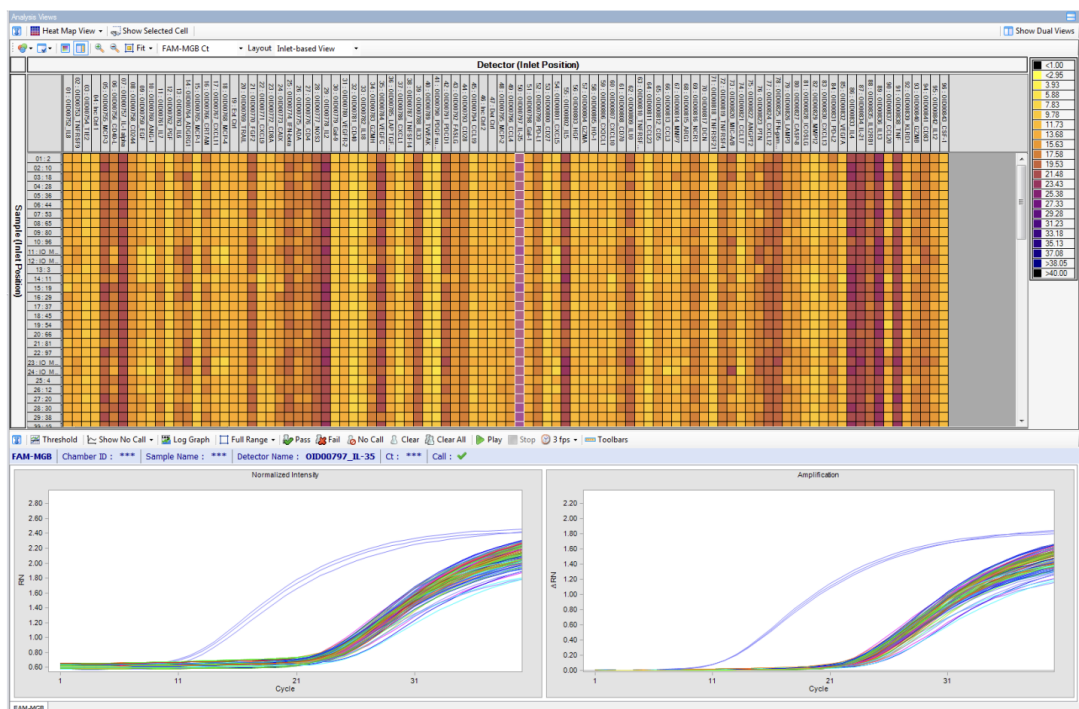
- 8.8** Select Detection Ctrl in the Detector on heat map and it will appear as the following figure with the amplification curves.



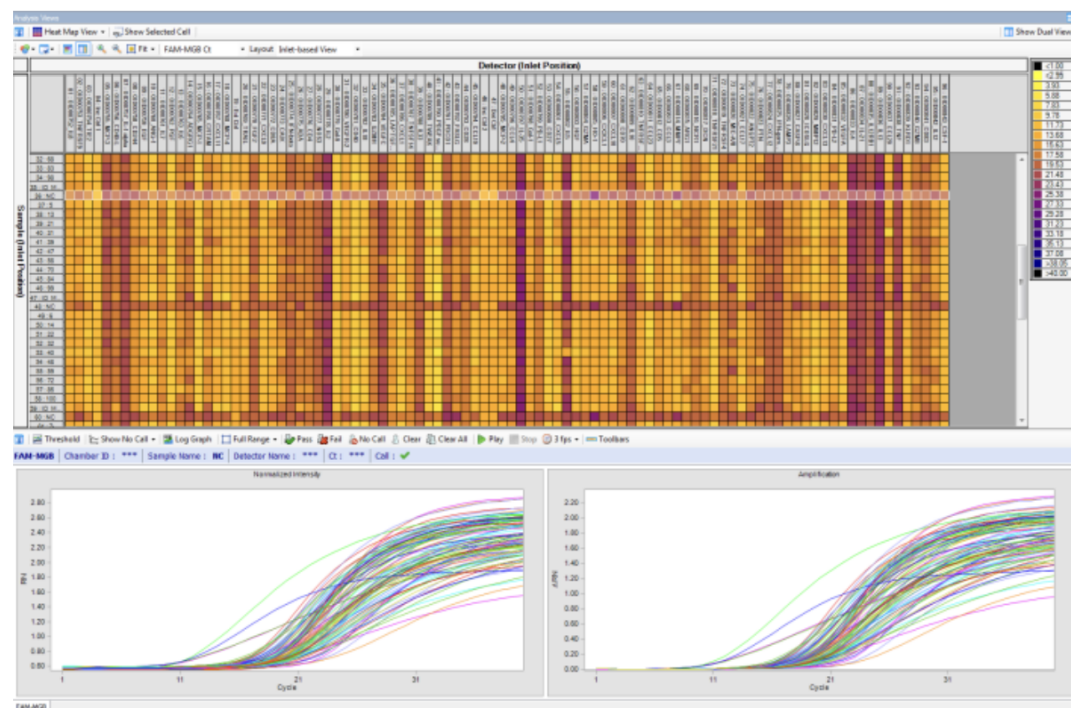
- 8.9** Select an assay with bright yellow color in whole column in the Detector on heat map and it will appear as the following figure with the amplification curves. This is the highly expressed/more concentration of protein.



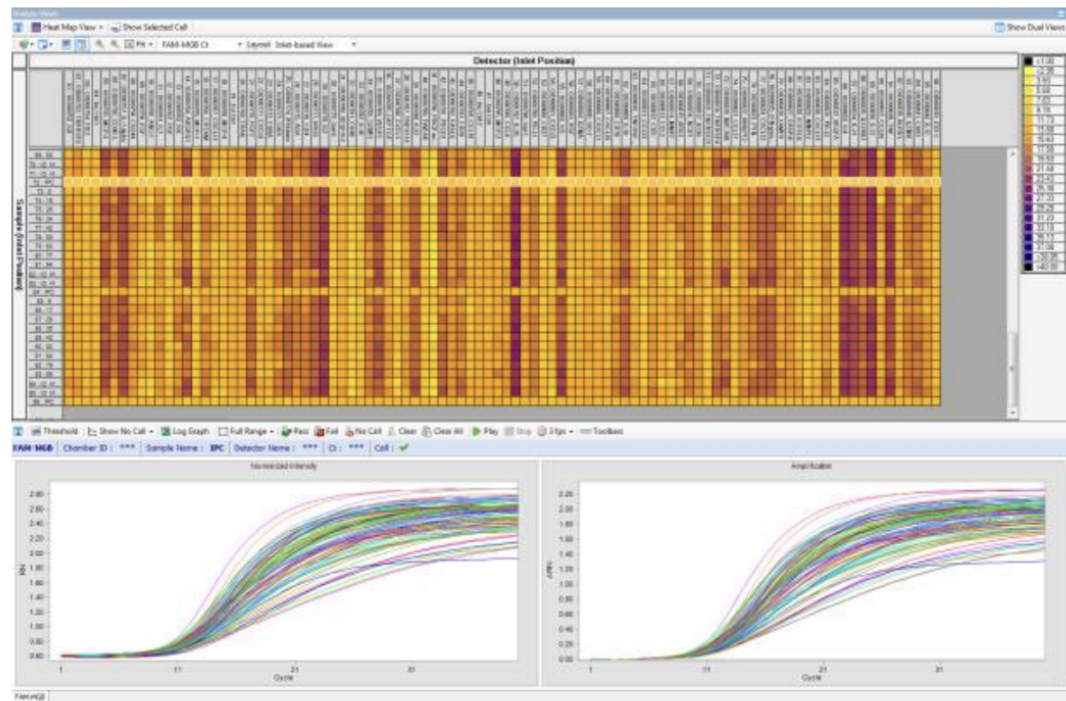
8.10 Select an assay with Dark purple color in whole column in the Detector on heat map and it will appear as the following figure with the amplification curves. This is the lowest expressed/low concentration of protein.



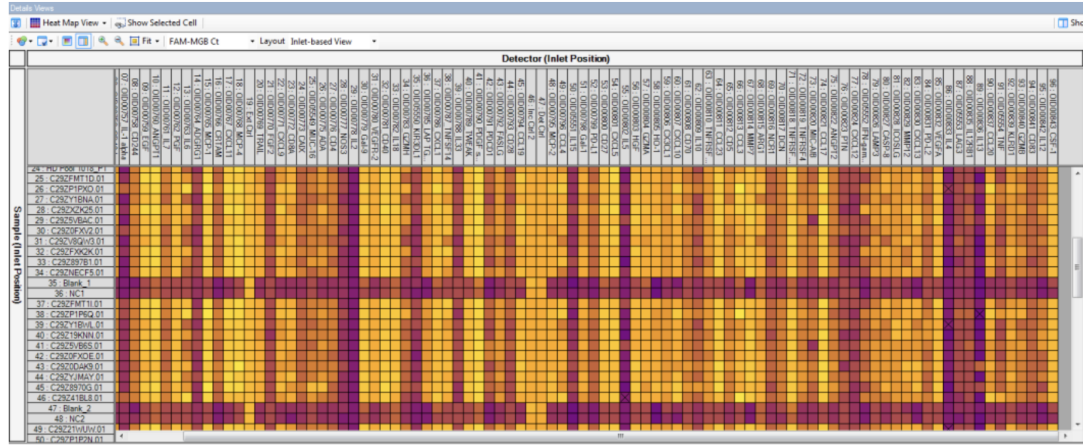
- 8.11** Select NC (Negative Ctrl) in Samples on heat map and it will appear as the following figure with the amplification curves. This should always be purple (dark line) color because it is negative Ctrl.



- 8.12** Select IPC (Positive Ctrl) in Samples on heat map and it will appear as the following figure with the amplification curves. This should always be yellow (bright line) color as it is positive ctrl.



The X marked wells/squares shown in below heat map picture in Analyte IL-4 indicates there is no Data.

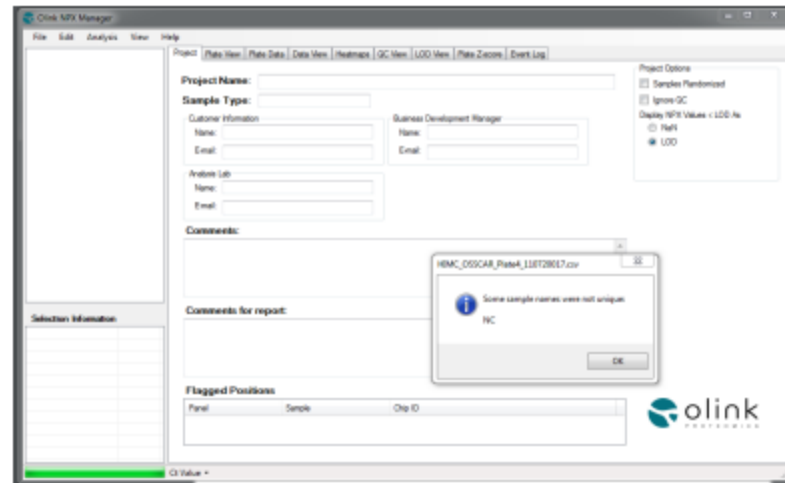


Data Export

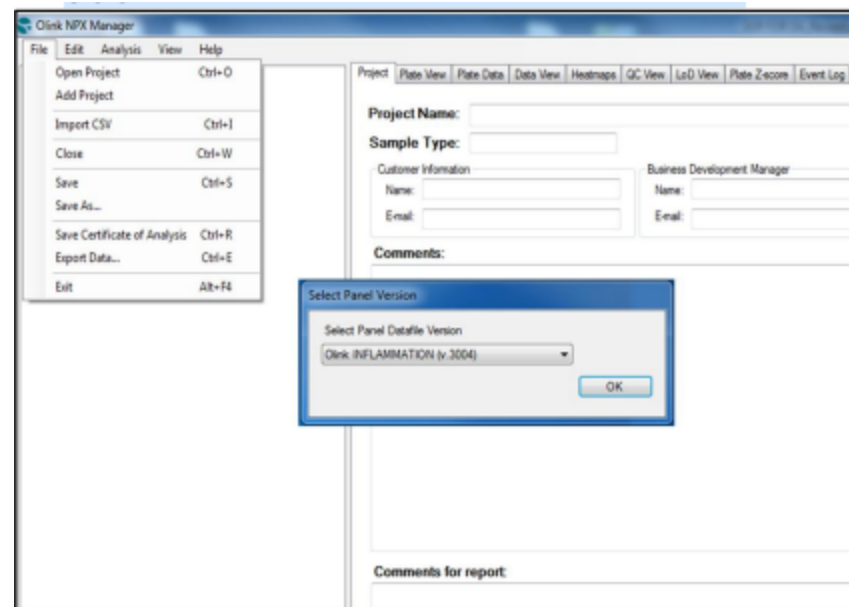
- Select File>Export. Make sure to choose Heat Map Results (*.csv) for File format from "Save as Type" drop down. Name the excel file according to HIMC format "HIMC_PI_Date.Ct.csv"

Press OK.

Note: Multiple files within the same project can be selected for import at the same time.



13.2 Next screen is for the “Select Panel Version”

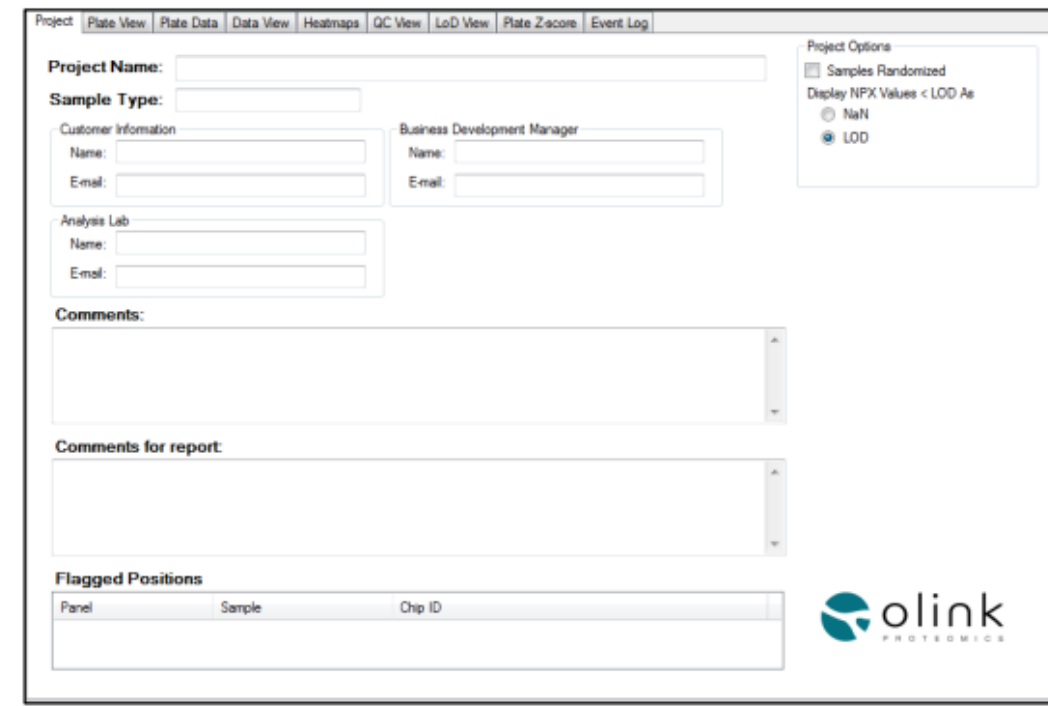


13.3 Each imported run is automatically annotated with the correct Olink Panel. Click OK

14 Enter project information:

14.1 Go to the “Project”. Project Name will be displayed in the data export file and the Certificate of Analysis (CoA). Information about Sample type, customer, business development manager and analysis lab will be displayed in the CoA. Comments will be saved in the Olink NPX Manager Project (oaf) file. Comments for report and Flagged Positions will be displayed in the Certificate of Analysis (CoA).

14.2 Fill out required Project information.



The screenshot shows the 'Project' tab of the Olink NPX Manager software. It includes fields for 'Project Name' and 'Sample Type'. Under 'Customer Information', there are fields for 'Name' and 'Email'. Similarly, under 'Business Development Manager' and 'Analysis Lab', there are fields for 'Name' and 'Email'. There are two large textareas: 'Comments' and 'Comments for report'. A 'Flagged Positions' table is partially visible. On the right, there is a 'Project Options' section with a radio button for 'Display NPX Values < LOD As LOD' (which is selected). The olink logo is at the bottom right.

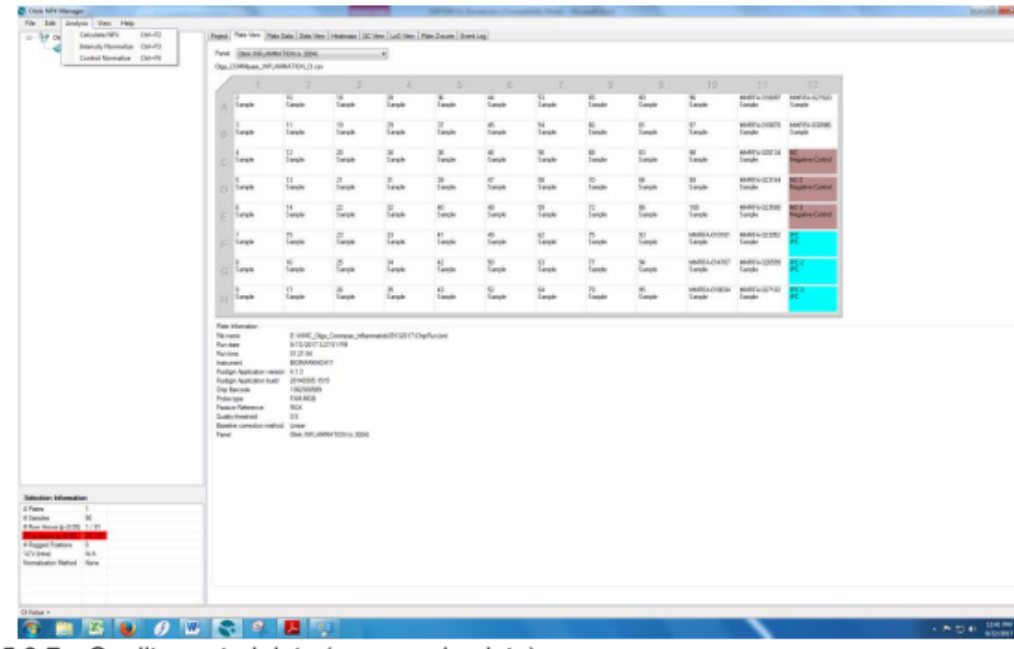
15 Go to “Plate View” Verify sample and data annotation.

15.1 Verify the plate layout. The layout can be changed by right-clicking on a selected well, or a selection of wells, and changing the well type to any of the available selections: Sample, Control, Negative Control, IPC and Not Used.

15.2 Select all 3 NC wells (NC, NC-2, NC-3) and mark them as Negative Control. Repeat Steps 15.1 and 15.2 for all imported plates.

16 Calculate NPX

- 16.1** Highlight an Olink Panel in the left pane tree node. Click on “menu Analysis” > “Calculate NPX”. Randomized studies with more than one sample plate will be intensity normalized if we select the “Samples Randomized” check box under the project tab before normalization or by clicking on Analysis” > Intensity Normalize v.2. Other study set ups will be IPC normalized by default. The “Intensity Normalization” will calculate the median of all the plates and individual plate NPX values will be normalized based on the median value of the all plates.
- 16.2** Default view is Ct Value. To view NPX values, select NPX Value in the status bar in the top left corner of the main window and go to Data view tap.



17 Quality control data (per sample plate)

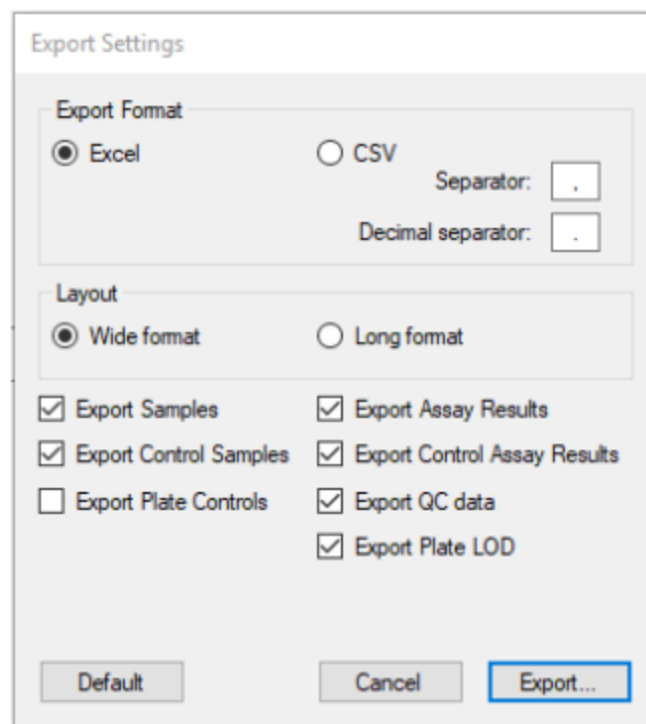
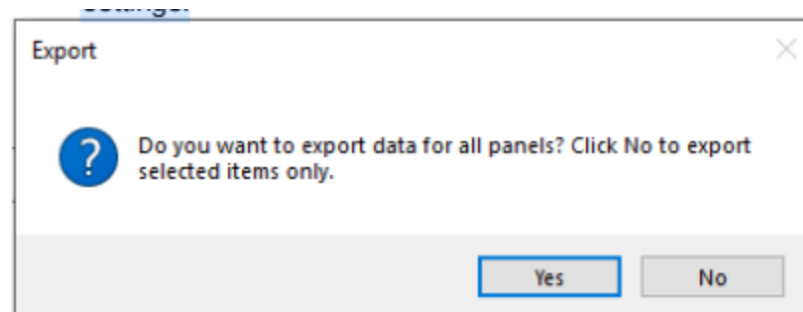
- 17.1** Select a plate in the left pane tree node and go to QC View

- 17.2** Check acceptance criteria Run QC:
- Number of flagged samples less than 1/6th of the total number of samples (16 samples in a full plate)
 - Standard deviation < 0.2 NPX for each of Incubation control 1, Incubation control 2 and Detection control
- 17.3** Acceptance criteria Sample QC
- Incubation Control 2 deviations from median of plate: < +/- 0.3 NPX
 - Detection Control deviation from median of plate: < +/- 0.3
 - NPX Samples that deviate more than +/- 0.3 NPX will be marked as flagged
- 18** Evaluate results in selection information (based on NPX values)
- 18.1** Plate ANOVA (if more than one sample plate per panel) < 10
- 18.2** Row/Column ANOVA: < 30 OK, 40-60 Investigate further
- 18.3** If more than 5 LODs highlighted yellow under LoD view tab, deviating from the values observed (+/- 2.5 NPX: which is Olink recommended value), check the following steps:
- If One of the negative Control samples is flagged in the NPX software, this control has to be marked "not used sample" before calculating the NPX
 - Check the Heat map view to show only negative controls to see if any one negative ctrl is different from other two that it has many dark "BLUE" values (= low Ct = high NPX which affects the LOD value) by going through the "assays".
 - In such a case remove that negative ctrl by marking it "Not Used" and reanalyze the NPX data.
- 19** Export data

19.1 Select NPX Value in the status bar in the topleft corner of the main window

19.2 Under Project Options in the Project tab, select whether NPX values below LOD should be exported as NaN or replaced with the value for LOD or Actual Value (below LOD). Select menu File > Export Data.

19.3 Click "Yes" if exporting data for all plates together and "No" for exporting data on selected plate only. Select desired export settings.



- 19.4** Enter filename and export format (Excel files *.xlsx or CSV-files*.csv) in the Save-dialog and click the Save button.
- 20** Export certificate of analysis: Select menu File > Report > Save Certificate of Analysis.

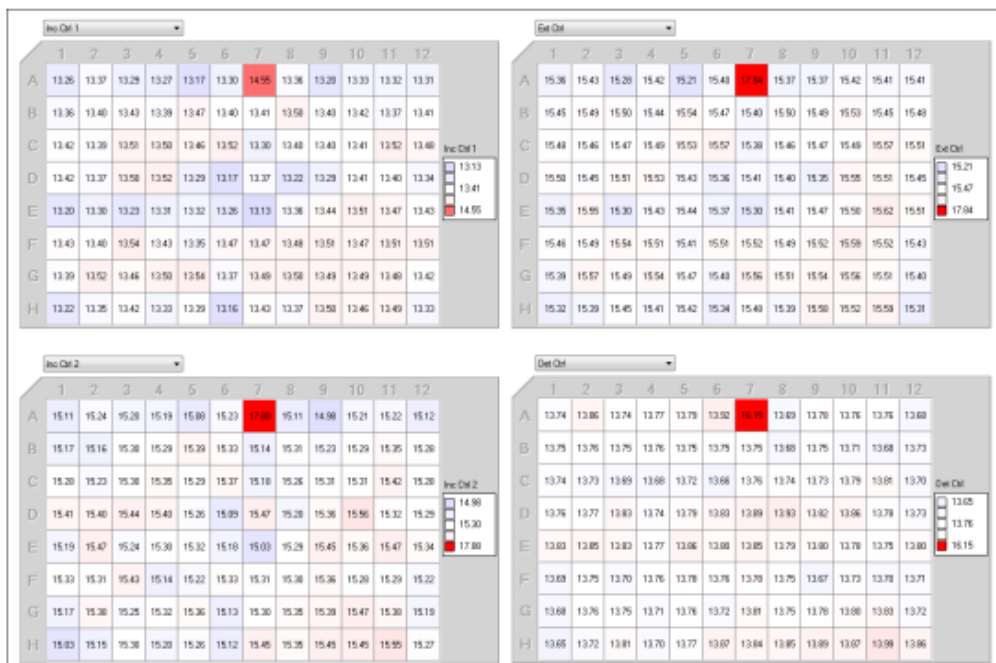
Analysis for Multi-plates Project Per Olink Panel

- 21** For randomized projects with more than one plate per Olink panel, follow Steps 11 through 15.2 for all plates one after another.
- 22** Imported plates are grouped by Olink panel and displayed in the left pane tree view. Plates can be reorganized in the tree view by drag-and-drop.
- 23** Then check the Samples Randomized box under Project Options to analyze data using Intensity Normalization. Intensity Normalization can also be applied by Clicking intensity Normalize v.2 under Analysis tab. Follow the Steps from 16 and further analysis.
- 24** Click the + sign next to an Olink Panel name to expand the tree node and reveal all imported runs for the Olink panel. Highlight the first run and go to the Plate View tab Verify the data file version
- 25** For using the NPX software do the multi-plate analysis, check the plate Z-score for each analyte. If one/more sample spots are far from the 0- line, then check the Ct value, NPX value and also check the Chip image in RT-PCR analysis software if there is any Chip line has issues- meaning dark line mentioned in Step 8.3. The sample/s should be removed from analysis.

Troubleshooting

- 26** EXAMPLE OF data set without any flagged samples by NPX manager but needed to be corrected manually

- When 1 or more samples have Ct value VERY different (darker red or darker blue color) from the rest of the wells on the plate but that sample is not flagged by NPX manager
- The sample is not flagged because the NPX-values for Incubation control 2 and Detection control were within accepted limits.
- All four controls are affected for that sample and when using the extension control to get the NPX value, the NPX values for Incubation control 2 and Detection control happen to be within accepted limits.
- When the extension control is affected in the same way as the other controls, meaning that all four controls have lower or higher Ct values than the rest of the plate, this difference can disappear in the normalization step.
- In this case, the sample should be marked as “failed” in the plate view tab and the data for that sample will disappear and will show “No Data” in data view tab. For example in DEP Culture samples inflammation Olink panel, see below:



The screenshot displays four tables representing different control groups for a DEP Culture sample. The top row contains Inc Ctrl 1 and Ext Ctrl. The bottom row contains Inc Ctrl 2 and Det Ctrl. Each table has columns labeled 1 through 12 and rows labeled A through H. The data is represented as numerical values. In Inc Ctrl 1, row H, column 7 is significantly darker red. In Ext Ctrl, row E, column 7 is significantly darker red. In Inc Ctrl 2, row H, column 7 is significantly darker red. In Det Ctrl, row H, column 7 is significantly darker red. These darker colors indicate extreme Ct values compared to the rest of the plate.

- 26.1** If sample/s is flagged in a row/column and all internal controls and analytes have high/low Ct values compared to that of the whole plate, it could be an error while adding detection mixture in those wells.
- 26.2** If same samples of the study or healthy donor plasma/serum samples were on all plates they will be used as control for analysis, Olink uses Plasma pool of 20 HD samples. The intra assay %CV for control sample is recommended <10% and that of inter assay is <15%.

NPX manager uses highest LoD value out of all plates LoD values.

- 27** Check if the chosen (highest) LoD affects more than 50% of the samples for that analyte/protein detected in all the plates meaning this high LoD value is used for more than 50% samples on all plate.
- 28** In this case, mark that assay failed for that Highest LoD value analyte on that one specific plate.
1. by selecting that plate on the left pane tree view, go to Data View
 2. Select analyte by RIGHT click on the analyte and mark as failed.
 3. A comment box will appear where associated comment can be made and all values for that analyte will be shown as "No Data".
 4. Now select Olink panel (all plates) on left pane tree view to recalculate.
 5. Confirm % of protein detected for the analyte in question with LOD.
- | Project | Plate View | Plate Data | Data View | Heatmaps | QC View | LOD View | Plate Scores | Event Log |
|---------|------------|------------|-----------|-----------|---------|----------|--------------|-----------|
| CX3CL1 | CXCL10 | CD70 | IL10 | TNFRSF12A | CD23 | CD5 | CD3 | |
| 13.91 | 11.13 | 16.96 | 13.71 | No Data | 8.91 | 15.69 | 12.20 | |
| 15.12 | 15.39 | 16.54 | 21.53 | No Data | 10.29 | 15.70 | 13.10 | |
| 12.94 | 13.37 | 17.89 | 21.44 | No Data | 9.88 | 16.10 | 12.96 | |
| 15.04 | 14.41 | 16.34 | 20.33 | No Data | 9.79 | 15.87 | 12.21 | |
| 13.99 | 13.03 | 14.95 | 19.59 | No Data | 9.72 | 16.39 | 13.68 | |
| 13.69 | 15.07 | 15.45 | 20.47 | No Data | 10.25 | 16.94 | 14.00 | |
| 13.85 | 12.56 | 15.90 | 18.86 | No Data | 9.19 | 15.65 | 13.64 | |
| 14.31 | 13.13 | 16.36 | 20.44 | No Data | 10.44 | 16.20 | 14.10 | |
| 19.58 | 21.46 | 19.22 | 21.80 | No Data | 19.09 | 20.12 | 18.30 | |
| 13.63 | 11.88 | 16.02 | 19.91 | No Data | 10.75 | 15.99 | 13.46 | |
| 12.31 | 13.52 | 15.14 | 19.92 | No Data | 9.91 | 15.48 | 12.79 | |
| 14.16 | 14.01 | 16.44 | 19.58 | No Data | 9.39 | 15.59 | 13.75 | |
| 13.78 | 11.14 | 14.35 | 18.93 | No Data | 9.61 | 15.93 | 13.51 | |
| 19.93 | 21.47 | 19.45 | 22.03 | No Data | 18.90 | 20.07 | 18.31 | |
| 14.91 | 14.48 | 17.03 | 21.18 | No Data | 10.32 | 15.01 | 13.33 | |
| 12.34 | 12.96 | 14.71 | 19.27 | No Data | 10.05 | 15.79 | 12.99 | |
| 11.77 | 12.07 | 15.88 | 19.42 | No Data | 9.49 | 16.37 | 12.49 | |
| 10.96 | 12.74 | 12.36 | 18.96 | No Data | 9.86 | 16.59 | 13.20 | |
| 19.75 | 21.56 | 19.51 | 21.79 | No Data | 18.88 | 20.11 | 18.52 | |
| 12.57 | 13.40 | 16.93 | 19.73 | No Data | 9.49 | 15.98 | 12.51 | |
| 12.42 | 12.66 | 14.87 | 19.19 | No Data | 9.70 | 15.58 | 12.30 | |
| 11.95 | 12.70 | 16.91 | 21.16 | No Data | 9.76 | 16.68 | 13.07 | |
| 14.00 | 12.78 | 15.72 | 19.60 | No Data | 9.48 | 15.10 | 9.45 | |
| 11.26 | 12.89 | 12.25 | 13.34 | No Data | 11.94 | 13.60 | 12.71 | |
| 14.30 | 13.45 | 15.07 | 19.83 | No Data | 10.40 | 16.76 | 13.62 | |
| 10.95 | 11.12 | 15.17 | 19.20 | No Data | 9.09 | 16.34 | 12.21 | |
| 14.18 | 13.67 | 16.65 | 18.55 | No Data | 10.08 | 15.60 | 12.97 | |
| 11.43 | 13.08 | 12.42 | 13.50 | No Data | 12.12 | 13.83 | 12.95 | |
| 16.70 | 21.07 | 16.95 | 21.49 | No Data | 19.23 | 20.13 | 8.11 | |
| 11.16 | 12.88 | 12.10 | 13.26 | No Data | 11.87 | 13.79 | 12.58 | |
- HOWEVER: optionally,
- Note: When we have too very different LODs between the two plates for some analyses and, therefore, lose the data points of one plate (with low LOD) due to the other plate, which has the higher LODs for that specific analytes, one can remove the "NEGATIVE CONTROL". AT THE Data View- Display only "Neg Ctrl", right-click the NC, NC2, NC3 for the specific analyte (depends on the variation- remove all or the only one w HIGH LOD) and select the "Mark as Failed". This will remove the high- LOD for that analyte.

