# **Release Notes**

To install or upgrade to the latest version of Partek Flow software, follow the directions outlined in the Installation Guide.

### 11.0.24.0529

- · Allowed to add multiple colors for numeric variable coloring on scatter plot
- In normalization task, allowed to use numeric attribute to divide or subtract from data
- · Added Leiden algorithm in graph-based clustering task
- Added support to import Seurat 5 object
- · Added support to import Seurat object saved as .qs format
- Added filter feature option in SVD task
- Added function to allow to specify multiple urls to import fastq files
- Improved filter based on GSEA enrichment score dialog
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:24.0529.62

### 11.0.24.0414

- Added CellphoneDB task to analyze cell-cell communication process
- Added display transcripts location ability on 10X Genomics Xenium data visualization
- Improved trim adaptor task to be more memory efficient
- Improved file browser when specify a location and click go button, it will scroll to the location on the left panel
- Moved all tasks in Peak analysis section to Region analysis section
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:24.0414.55

# 11.0.24.0325

- · Added more options for variant filter by fields
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:24.0325.52

#### 11.0.24.0317

- · Added hierarchical cluster analysis task to similarity matrix data node
- Added statistical analysis task to cell descriptive stats
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:24.0317.49

# 11.0.24.0311

- Improved convert Seurat to matrix dialog to allow to configure meta data
- Added more options in filter annotated peaks dialog
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:24.0311.45

# 11.0.24.0226

- · Improved differential analysis report filter option on log2 ratio to allow user to specify a range to filter
- Improved Normalize to housekeeping genes dialog to display both gene ID and gene name
- Added data repository management on Settings page
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:24.0301.39

# 11.0.24.0204

- Added long reads aligner minimap2 and pbmm2
- Simplified the Data Viewer configuration dialogs
- Added Scree plot and component loadings table to the PCA report by default
- · Added Poisson regression and Negative binomial regression as options in the Differential analysis task
- · Minor bug fixes

Latest docker image: registry.partek.com/rtw:24.0204.33

#### 11.0.24.0102

- · Automatically generate spatial report on analysis tab after spatial data import
- Added alt-splicing task option on Salmon's transcript count data node
- Added STARsolo task for 10x Genomics scRNA-seg data
- Updated FreeBayes to v1.3.6
- PCA task default feature filter method is changed to variance from vst
- · Filter peaks task is moved to Filtering section from Peak analysis section on the menu
- Make the enable/disable user account as separate privilege from add user account for admin
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:24.0102.29

# 11.0.23.1204

- Enabled MACS peak detection report visualizations in the Data viewer
- Improved the algorithm for the normalize to housekeeping genes task
- · Added more gene annotation options in SNV task report
- Fixed the Venn diagram issue on some of the browsers
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:23.1204.23

# 11.0.23.1105

- Added search function in Help menu to directly search the content in documentation page
- Added option to choose a number of features to use in PCA computation
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:23.1105.16

### 11.0.23.1023

- · Improved differential analysis report layout
- Add impute missing value task
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:23.1023.14

# 11.0.23.1009

- Improved pre-alignment QC report
- · Fixed differential analysis report table download issue
- · Minor bug fixes

Latest docker image: registry.partek.com/rtw:23.1009.11

# 11.0.23.0918

- The user interface now boasts a modern appearance
- Enhanced the import wizard for greater intuitiveness and user-friendliness
- Added support for SomaLogic ADAT bulk protein data
- Improved the file browser for fast access to recent server uploads
- Bolstered file protection to prevent accidental deletions when used by other projects
- Added support for the 10x Genomics Xenium platform
- Introduced support for the Nanostring CosMx platform
- Added a new task for Correlation analysis between different assays
- Improved scatterplot selection with a handy painting mode
- Added the ability to display a high resolution Visium image with a new Spatial imaging report task for quick viewing
- Added a manual alignment option for tissue image visualization
- Streamlined the import process for multiple files per sample, e.g. import all 3 sparse matrix (feature-barcode-matrix) files per sample for all samples at one time
- Added more available information on the Annotated region report
- Improved the Seurat object conversion to be more intuitive
- Added a more detailed report to the Adapter trimming task
- Creating pseudobulk data by pooling single cells has been made easier with increased functionality
- Optimized the differential analysis dialogs for improved usability
- Changed the default normalization method for bulk RNA-Seq data to Median ratio (DESeq2)
- Improved the Detect fusions and Trim bases tasks to be more intuitive

- Allowed the Single-cell QA/QC task to be performed on individual samples
- Minor bug fixes

Latest docker image: registry.partek.com/rtw:23.0918.1

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# Additional Assistance

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

