Release Notes

To install or upgrade to the latest version of Partek[®] Genomics Suite™ software, follow the directions outlined in the Installation Guide.

PGS7.21.1119

Bug fixes:

- Support MacOS 12
- Minor bug fixes

PGS7.20.0831

Bug fixes:

· Minor bug fixes

PGS7.19.1125

Bug fixes:

- Improved memory efficiency
- Minor bug fixes

PGS7.18.0723

General Improvement:

Microarray methylation workflow to make it more intuitive

Bug fixes:

· Minor bug fixes

PGS7.18.0518

General Improvement:

- Implemented DMRCate algorithm for differential methylated regions detection
- Added false discovery rate report in html format
- Add options to delete rows contains missing value only during text import

Bug fixes:

Minor bug fixes

PGS7.18.0402

General Improvement:

- Sped up multidimensional scaling function
- Added chi-square analysis for categorical data
- Improved Fisher exact test on integer columns with two distinct values
- Improved handling beta value as 0 or 1 during M value transformation, instead of output "?", we set the value as 1e-6 or 9e-6 by default

Bug fixes:

- Fixed GEO downloader issue
- · Fixed scatterplot set cutoff line and regression line button issue
- Fixed a dot plot close issue
- Bug fixes on OSX

PGS7.18.0130

General Improvement:

- Improved the methylation workflow add region methylation analysis workflow
- · Added correlation analysis across two spreadsheets when they have the same set of samples

Bug fixes:

· Bug fixes on OSX

PGS7.17.1222

General Improvement:

- Improved the ANOVA contrast dialog; added the option to output difference between the two groups instead of fold change
- Added new options on the Illumina methylation array importer to allow filtering of probes based on detection p-value and XY chromosomes
- Improved Illumina methylation array analysis workflow to compute differential methylation sites based on M value and report the difference of M and beta values

Bug fixes:

- Fix Tk event crashes on OSX
- Other minor bug fixes

PGS7.17.1103

General Improvement:

- · Sped up genome view with large annotation files
- Added option to specify output file name on task dialogs
- Added option to save merged spreadsheet to a new spreadsheet

Bug fixes:

- · Fixed a hierarchical clustering viewer bug on Mac version
- Other minor bug fixes

PGS7.17.1018

General Improvement:

- · Partek spreadsheet can only be saved as binary format
- Partek spreadsheet can be exported as text format. The name of the exported spreadsheet will not be changed
- "Import Text File" will not change the input text file, the imported file will be saved as user-defined name in binary format
- When there are unsaved spreadsheets, "Save Project..." and "Export Zipped Project..." will give a warning message to save each unsaved spreadsheet
- When cloning a spreadsheet, the newly generated spreadsheet name is unique by default
- Re-ordered the File menu to improve usability
- · Added a function to impute missing values with random numbers from a defined distribution
- Improved the "Save Image As..." dialog on Mac and Linux
- Added support for save image as JPEG on Mac and Linux
- Added file type selector for open and save file dialogs on Mac
- · Violin Plot displays the properties tab by default
- Gene Expression workflow "Import from Affymetrix CEL Files" and File menu "Import Affymetrix .CEL Files" produce the same output spreadsheet for gene expression arrays
- Improved graphics for visualizations on Mac

Bug fixes:

- Fixed a bug on "Split Column" dialog on Mac
- Fixed a bug in "Import Affymetrix .ARR File..."
- · Other minor bug fixes

PGS7.17.0918

General Improvement:

- Most of the computations are multi-threaded, which drastically increases the speed
- Viewers generated from spreadsheets are new tabs within the same window frame by default; viewer windows can be docked back to the main window frame
- · Workflow panel is visible in any invoked viewers, order of the workflow steps is more intuitive
- Updated with modern icons and graphics color palette
- Mixed model ANOVA uses REML by default
- Sample ID factor is not included in the model of Alt-splicing ANOVA, GO ANOVA, pathway ANOVA
- Apply button are removed from most of the dialogs
- · Improved interface is cleaner and more user friendly
- New functional normalization method in the Illumina methylation array importer (450K and 850K)
- Improved SWAN normalization in the Illumina methylation array importer
- · Allows import of different versions of the Illumina methylation arrays when using the same set of library files
- Improved text file importer by adding wizard for specifying the properties of the spreadsheet

- Removed repeated measures menu option, it is recommended to use ANOVA doing repeated measures analysis
 Simplified menus and dialogs, removed obsolete functions, improved wording
 Automatically downloads annotation files, even from sites with HTTPS or redirects (e.g. Affymetrix and GEO)
 Logged data is detected and appropriate defaults are automatically set

Older release notes are available on our website at Partek® Genomics Suite™ Software Update Archive.