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**Intro**

qbase+ is software developed by Biogazelle for analysis of quantitative PCR data. It is based on the geNorm (Vandesompele et al., Genome Biology, 2002) and qBase technology (Hellemans et al., Genome Biology, 2007) from Ghent University.

The software runs on a local computer with either Microsoft Windows, Apple OS X or Linux operating system and is compatible with virtually all qPCR instruments. The first version was released in March 2008, with frequent updates; the latest version is 3.1.

The software is intended for relative quantification (whereby different normalization strategies developed by the Biogazelle founders are available) and for gene copy number analysis. The geNorm module for determining the expression stability of candidate references and the optimal number of reference genes for accurate normalization is a much-improved version of the original algorithm. The implemented analysis wizard allows the straightforward step-by-step analysis of every qPCR experiment. The statistical wizard built into qbase+ makes it easy for any biologist to come to expert results and reliable conclusions.

I'm convinced using qbase+ will lead to more reliable results in a shorter period of time!

Jan Hellemans, co-founder and CEO
Installation and licensing

Installation

Different versions of qbase+ are available for installation. It is important to download the version matching your operating system (Windows, Mac or Linux) and bitness (32-bit or 64-bit). If compatible with your system, we advise to select the 64-bit version as it allows the use of larger amounts of memory to support analysis of very large experiments.

qbase+ requires JAVA to be installed on your system. Many systems already have JAVA installed. Not every version of JAVA would be suitable. It needs to be a 32-bit JAVA for 32-bit versions of qbase+ and a 64-bit JAVA for 64-bit versions of qbase+. qbase+ does require JAVA 1.8 or more recent. On Windows, a suitable version of JAVA will be co-installed as part of the installation procedure of qbase+.

To install qbase+ on Windows and Mac, double click on the downloaded installer and follow the instructions (Figure 1). The procedure for Linux is somewhat different. Instead of using an installer, the downloaded file needs to be unzipped and moved to an appropriate location. In addition, the execution bit needs to be set before qbase+ can be launched.

Figure 1: qbase+ setup wizard
First time users

We offer a trial version of qbase+ for evaluation purposes. For a period of two weeks, all premium functionalities will be available.

Every installation of qbase+ includes various carefully selected and representative data sets for:

- gene expression analysis
- microRNA profiling
- geNorm pilot reference gene stability analysis
- gene copy number analysis
- ChIP-qPCR

When starting qbase+ for the first time, the program will prompt you for the email address used for website registration (Figure 2). Based on these credentials, qbase+ will verify the availability of a valid license linked to your account. A working internet connection is required for this purpose.

Using a basic or premium license

No user interaction is required in qbase+ when updating or upgrading your license. Once the new license becomes available on qbase+’s license server (typically as the results of a new purchase) it is immediately being used by qbase+.

Advanced memory settings

The following section is only relevant for a minority of users performing high-throughput qPCR analysis (tens of thousands of datapoints per experiment). It’s also restricted to users with at least moderate IT skills and administrator rights on the computer running qbase+.

Despite recent optimizations in qbase+’s calculation engine, some very big experiments may consume more memory than what is made available to qbase+ by default. This will result in swapping to a hard disk drive, thereby significantly affecting the performance of qbase+. Larger experiments can still be properly
processed by changing the amount of memory that may be allocated to qbase+ in a so called ini-file.

For both Linux, Windows and Mac, we recommend to modify the 'qbase+.ini' file to take advantage of more RAM memory (limited to around 1.5 Gb in a 32-bit environment). This file can be found in the installation directory of qbase+, 'C:\Program Files\qbase+' or 'C:\Program Files (x86)\qbase+' on a Windows system for the 64 and 32 bit versions, respectively. To find the settings file on a Mac system, right click (or CTRL-click) on the qbase+ application, choose 'Show Package Contents' and Browse to Contents -> MacOS.

Open the file with a text editor, locate the line with "-Xmx1024m" and modify the number and unit depending on the amount of RAM available (e.g. "-Xmx3g" on a system with 4 Gb RAM, or "-Xmx7g" on a system with 8 Gb of RAM; general guideline: 1 Gb (g) less than the amount of RAM in Gb; make sure there is no space in between Xmx and the number). These settings need to be updated with every new installation of qbase+. On most systems, administrator rights are required to update these settings.
Definitions and concepts

General Definitions

**Target**
The generic term used for the DNA sequence to be quantified. It typically refers to a gene, but could also refer to a specific transcript or a genomic locus. Two target types are recognized: targets of interest and reference targets.

**Cq value**
Cq is the MIQE standard name for Ct (cycle threshold), Cp (crossing point), or other instrument specific quantification value name.

**Run**
Runs are collections of qPCR data coming from a single plate, array, rotor or chip (depending on the instrument being used).

**Experiment**
The set of data (stored in one or multiple runs), annotations and settings used for qPCR data analysis.

**Project**
Logical groups of related experiments.

**Inter-run calibration**
A calculation procedure to detect and remove inter-run variation.

**Sample**
In qbase+, a sample is defined as the nucleic acid dilution on which the qPCR measurement is performed. According to this definition, a serial dilution made from a single RT reaction or DNA sample would consist of different sample names.

**RDML**
A universal, open and XML based data format for the exchange of qPCR data, recommended by the MIQE guidelines.

**Reference target**
The targets that will be used to normalize the qPCR data. In the context of gene expression analysis, this target type used to be referred to as housekeeping gene.
**geNorm**

The algorithm developed by Biogazelle co-founder prof. Jo Vandesompele (Genome Biology, 2002) to determine the expression stability of selected (candidate) reference targets. It can be used in geNorm pilot studies to identify the optimal set of reference genes to be used in follow up studies, or later on to verify the expression stability of selected reference targets.

**Sample property**

Two types of sample properties are used in qbase+. There is a set of predefined sample properties such as sample name, sample type and quantity. Custom sample properties can be added to link (usually annotation) information to samples. For each sample, a sample property value can be entered. For example, the sample property 'gender' may have sample property values 'male' and 'female'.

**Normalization**

Normalization is the procedure by which technical sample specific variation (e.g. differences in total amount of cDNA) is corrected for. Normalization is typically performed by means of reference genes, but alternative approaches are available.

**Result scaling**

The results of a relative quantification analysis do not have a particular scale. Because of this they can be multiplied (or devided) by a common factor. This process will alter the actual values, but not the sample to sample relationships (i.e. the fold changes). It does not alter the results but may facilitate interpretation, for example by making all results relative to a control sample.

**Workspace**

The workspace is the central location in which qbase+ stores all its information: data, annotation and settings.
**Overview of qPCR workflow**

Where does qbase+ fit in the qPCR workflow?

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![Diagram of qPCR workflow](image)

Figure 3: Researchers spent a considerable amount of time in preparing samples, optimizing assays and setting up the actual experiments. Data analysis is far too often an underestimated step in the qPCR workflow. qbase+ is a reliable, easy, fast and flexible data analysis platform, outcompeting the time-consuming, error-prone and traditional way of doing calculations.
The analysis wizard in qbase+

qbase+ has an analysis wizard to guide you through the most essential steps of your data analysis. Starting from a new or existing experiment sample, target and run information can be added or updated.

Depending on the final aim of your analysis, the wizard guides the user through the relevant quality control parameters, amplification efficiency settings, normalisation methods and scaling parameters.

- Chapter Analysis wizard page 16

Steps in qbase+

1) start new experiment

A new experiment needs to be created every time you want to analyse a new data set. This is like making a new Excel file to process your qPCR data. Such an experiment will not only store your data, but also all its annotation, calculation parameters and quality control preferences.

- The concept of an experiment – chapter Experiments on page 24

2) import Cq values

qbase+ processes Cq values measured by a qPCR instrument and exported by that instrument’s software. Once such a table with Cq values for every well is exported, it can be imported in an experiment.

- The concept of a run – chapter Runs on page 24
- Importing run data – chapter Runs on page 26
- What data are supported? – chapter Can I analyze data from my qPCR instrument in qbase+? on page 85
3) add / review sample and target naming

Runs need to be properly annotated before experiments can be analyzed. Not only will your data be meaningless without annotation, it is simply essential for qbase+ to start any calculations.

- Appoint sample and target names – chapter Run annotation page 33
- Creating new samples – chapter Sample annotation page 35
- Creating new targets – chapter Target annotation page 38
- Apply/copy run layouts – chapter Run annotation page 34

4) add extra sample info

Adding extra sample information will allow better graphical representation of data (e.g. sample groups), statistical analysis of data, etc. There are three ways by which sample information can be provided to qbase+:

1. as part of the import of a run
2. by manually adding a sample
3. by importing a sample list.

- Creating new samples – chapter Sample annotation page 35
- Sample properties – chapter Sample annotation page 36
- Custom sample properties – chapter Sample annotation page 37

5) adjust calculation parameters

Four types of parameters can be defined allowing the most appropriate calculations for your experimental set-up:

1. the type of amplification efficiency correction
2. the most appropriate normalization method
3. the method to calculate average Cq values
4. the target scaling mode.

- Calculation parameters – chapter Calculation parameters and quality control settings page 41
- amplification efficiency – chapter PCR efficiency correction page 50
- How is PCR efficiency calculated and used for relative quantification? – chapter FAQ page 87
6) adjust quality control settings

Quality control is an important aspect of qbase+. By adjusting the quality control settings you will be able to estimate the quality of the data and/or define criteria for automatic exclusion of certain data points.

- Quality control settings – chapter Calculation parameters and quality control settings page 43
- How to flag bad technical replicates based on a standard deviation threshold? – chapter FAQ page 90
- How to exclude PCR replicates that do not meet quality control criteria? – chapter FAQ page 90

7) perform quality control

qbase+ contains several types of quality control that can be accessed after adjusting the appropriate settings, including technical replicates, positive and negative controls, stability of reference targets and sample quality control.

- Technical replicates – chapter Quality controls page 53
- Positive and negative controls – chapter Quality controls page 55
- Stability of reference targets – chapter Quality controls page 56
- Sample quality control – chapter Quality controls page 57

8) perform inter-run calibration

Inter-run calibration (IRC) is a calculation procedure to detect and remove (often underestimated) inter-run variation. These calculations are typically needed whenever samples need to be compared that are measured in different runs.

- When do I need IRC? – chapter FAQ page 89
- The inter-run calibration concept – chapter Inter-run calibration page 45 and chapter FAQ page 89
- Inter-run calibration in qbase+ – chapter Inter-run calibration page 47
9) inspect results

qbase+ offers several possibilities to visualise qPCR experiment results. Depending on your needs you can inspect commonly used bar charts, correlation plots and/or results tables. Dedicated analysis modules allow you to perform a geNorm analysis (including automated interpretation), statistical analysis (see further) or copy number analysis.

- Target bar charts – chapter Bar charts, correlation plots and results tables page 58
- How to interpret the target bar chart? – chapter FAQ page 88
- Multi-target bar chart – chapter Bar charts, correlation plots and results tables page 60
- Correlation plot – chapter Bar charts, correlation plots and results tables plot page 61
- Result table – chapter Bar charts, correlation plots and results tables page 62
- What is the meaning of CNRQ value in the result table? – chapter FAQ page 86
- Why does qbase+ ask to log transform the data when exporting the results table? – chapter FAQ page 90
- Special application: geNorm – chapter geNorm page 76
- What is the difference between Reference target stability and geNorm? – chapter FAQ page 91
- What is the difference between "M" and "geNorm M" in qbase+? How are the calculations different? – chapter FAQ page 92
- What to expect when performing a geNorm analysis in qbase+? – chapter FAQ page 92
- Special application: copy number analysis – chapter Copy number analysis page 82
- Export Datasets and Tables – chapter Exports page 73

10) perform statistical analysis

qbase+ has a built-in intuitive statistical wizard to perform commonly used statistical tests on the results that are calculated in a single experiment. The stat module is specifically tailored towards the typical needs of biologists performing qPCR analysis.

- Using the statistical wizard – chapter Statistics page 62
- Specify the goal of your analysis – chapter Statistics page 63
- Statistical result table and conditions of use – chapter Statistics page 67
- Statistical background – chapter Statistics page 70
- Does the statistical wizard perform a log transformation? – chapter FAQ page 91
Getting started

User interface

qbase+ has 2 main views: the **analysis wizard** and **Expert mode**. Switching between the two views is possible with the 'Launch wizard'- 'Close wizard' buttons. The analysis wizard allows for straightforward step-by-step analysis of typical qPCR experiments. Experienced users, or users with very specific research needs, might prefer the expert mode enabling direct access to all parameters.

Analysis wizard

The analysis wizard integrates the complete workflow of individual steps needed for the correct basic analysis of a qPCR experiment. Existing qbase+ experiments can be analyzed again. All individual settings are also accessible via the expert mode, and will be discussed in more detail throughout this manual.

Analysis wizard start page

There are three ways to start the analysis of qPCR data:

1. Start a new empty experiment and fill it with qPCR data (i.e. import run data)
2. Analyze an experiment that was previously created and that is stored within qbase+’s workspace
3. Analyze an experiment that was previously created and that is available as an experiment file that can be imported

Import run page

Runs are collections of qPCR data coming from a single plate, array, rotor or chip. The "import runs" button launches a wizard supporting the import of one or multiple run files in either the RDML format or an Excel, .txt or .csv file containing a table with Cq values. Detailed requirements for file formats are discussed in more detail in the section 'Import formats'.

Sample / target list

In qbase+, a sample is defined as the nucleic acid dilution on which the qPCR measurement is performed. Target is the generic term used for the DNA sequence to be quantified. It typically refers to a gene, but could also refer to a specific transcript or a genomic locus. Samples and targets have a unique name and a set of properties. These can be qbase+ specific properties (e.g. target type: target of interest or reference target) or custom properties (e.g. age, treatment, passage number, ...). Some information may already be derived from the run files. Additional properties can be set manually or imported by means of sample or target lists. More information on how to use these lists can be found in the section 'Samples and targets'.
Run annotation

Every datapoint (well) should be annotated with a sample and target name. This information may be derived from the run files if they were properly annotated in the qPCR instrument software. The run annotation can be reviewed, corrected and completed by manually editing run annotation. In cases where runs have the same layout (for either samples or targets) as a previously annotated run, that layout can simply be copied from the annotated run to the unannotated run. Detailed information can be found in the chapter 'Annotation'.

Aim

Different types of experiments require different settings and different analyses. By selecting the proper analysis type, qbase+ will only show the relevant settings. Experienced users, or users with very specific research needs (e.g. ChIP-qPCR) may exit the wizard and continue the analysis in the fully flexible Expert view. Again, more information on all different settings will be covered throughout the manual.

Technical quality control

A well executed qPCR experiment should have PCR replicates with very similar Cq replicates, no Cq values for negative controls and Cq values in a predefined range for positive controls. qbase+’s quality control evaluates the data in function of user defined thresholds for the maximum allowed variability on PCR replicates (maximum of 0.5 cycles by default), the allowed range of positive control Cq values and the negative control threshold. The latter is defined as the minimal difference in Cq value between the sample with the highest Cq value and the negative control with the lowest Cq value. The default value of 5 cycles corresponds to permitting up to 3% of background signal.
Amplification efficiencies

The basic formula for relative quantification (RQ=2^ddCq) assumes 100% amplification efficiency (E=2). More advanced methods correct for variable, assay specific amplification efficiencies. These efficiencies can be calculated from standard curves included in the experiment, or manually provided in case they were determined before.

Normalization method

Normalization is the procedure to correct for technical variations in total cDNA concentrations so as to distinguish samples with upregulated genes from samples that merely have the same expression level with a higher cDNA concentration. Several methods are available to correct for this type of variation. Normalization based on the measurement of one or more reference (historically referred to as housekeeping) genes is the most common option. For screening studies including a large number of genes, global mean normalization is a valuable alternative. Alternatively, for specific study types, users can provide custom normalization factors such as for example the cell count. When normalizing with multiple reference genes, the reference target stability values (M and CV) offer a measure for the suitability of the selected reference genes for proper normalization. Good reference genes have an M < 0.5 while M values up to 1 are acceptable for more difficult samples.

Scaling

By default qbase+ results are scaled to the average across all unknown samples per target. Relative quantities can be scaled arbitrarily, as long as the same scaling is applied to all samples. Rescaling will alter the relative quantity values, but not the relative fold changes in expression levels between samples. Rescaling is often used to facilitate interpretation of results, e.g. by setting the expression of a control sample or group of reference samples to 1 which makes all results relative to that sample or sample group.

Analysis

From this page you might visually inspect the results, export the results for downstream processing outside qbase+ or start a wizard to guide you through the statistical interpretation of the results.
Expert mode

The expert mode consists of three main windows (Figure 4): the project explorer (left window), the main window (upper right window) and the notification window (lower right window). The project explorer allows users to browse to the information they want to open in the main window, whereas the notification window contains tabs with information on alerts and program errors.

![Project Explorer](image)

**Figure 4:** Three main windows in the expert mode

Project explorer

The project explorer is a hierarchical data organizer that allows users to navigate through the experiments, settings and results. The elements in the project explorer tree can be opened by double-clicking, or by selecting 'open' from the context menu (this menu appears when clicking the right mouse button after placing the cursor on top of the item). This window can be minimized to get more screen space by clicking (-) in the upper right corner of the project explorer window, or by making use of the context menu ('Minimize') when right-clicking on the tab. The 'always on' situation can be restored by clicking (≠). An alternative way of working is to display the project explorer whenever it is needed, and to have it hidden otherwise. This can be achieved by clicking (≠); as soon as an item of the main window is selected, the project explorer disappears.

The width of the project explorer can be controlled by moving the mouse on the border between the project explorer and the main window (after which the pointing arrow becomes a horizontal arrow), followed by dragging the border.
Main window

When an item is opened from the project explorer, it will show up in the main window. Double-clicking a tab will maximize this window, and the situation can be restored by double-clicking the tab again. As an alternative, (esc) and (pageup) can be used, respectively.

Tabs can be dragged and dropped from one window to another window. Tabbed windows can also be placed next to (or underneath) each other, by click-dragging the tab until the ‘multiple tabs’ icon turns into a solid black arrow (Figure 5).

![Main window](image)

Figure 5: Displaying multiple windows in the main window

By default, qbase+ recalculates everything immediately (e.g. intermediate and final results, quality controls, and specific analyses) as soon as something changes in the data or data-analysis settings. Therefore, having multiple open windows next to each other is practical for evaluation of the effect modifying the settings.

For large datasets (>10,000 wells), it is recommended to disable automatic recalculations in order not to lose time upon recalculation of every minor change. The option to prevent automatic recalculations is accessible in the preferences window (in the main menu bar, go to: Windows > Preferences). Without automatic calculations, a recalculate button will appear in the qbase+ toolbar (recalculate), which needs to be clicked to execute the calculations.

Items can be closed by clicking on the close tab symbol (x), which appears when moving the mouse cursor on top of a tab, or by making use of the right-click context menu. The context menu also contains other features like ‘Close All’, ‘Close Others’ and ‘Duplicate window’. The latter can be used to duplicate a window in the main window.
window. Just like the project explorer, the main window can be minimized to get more screen space.

The header of each tab contains a path, describing what is being displayed and the position within the project explorer. This information may be essential to allow for proper identification of results, e.g. when bar charts for the same target in two different experiments are shown side by side.

**Notification window**

The notification window informs about the progress of calculation intensive steps and the occurrence of program warnings and errors. Several items (alerts, progress view, error log) can be added to this window (in the main menu bar, go to: Windows > Show View).

The alert window gives clues on potential issue with experiment design or data-analysis, e.g. no reference targets are appointed, targets are spread across runs (necessitating inter-run calibration), technical PCR replicates are spread across runs (not compatible with inter-run calibration), etc. By inspecting the Alert window, many problems can be quickly identified.

In the error log window, a small menu is accessible (top right) in which the user can set the information that has to appear in the log list (Figure 6). Log files can also be exported. A new window will appear in which the file name and destination folder has to be indicated. In case of problems, these error logs can help our support officers to identify the cause.

![Figure 6: Exporting error log](image)

**Menu and command bar**

Most options in the menu or command bar are quite self-explanatory, and are covered in various chapters where they are being used. Some options will be explained in some detail here.

The switch workspace option allows for the selection or creation of an alternative workspace. Briefly, it determines where all the qbase+ data is being stored. More information on workspaces is available in the Data management chapter.
Preferences

Calculations
By default, qbase+ will recalculate results with each change of data, annotation or settings. This is convenient for typically sized experiments, but may lead to slow performance in high throughput experiments. The size at which automatic recalculation becomes more a burden than a blessing depends on the experiment designs, calculation settings and computer performance. Any recent computer (less than 3 years) should be able to deal with experiments of up to ten 384-well plates without hick-ups or recalculation delays.

For larger experiments or older computers, it may be required to turn off automatic recalculation. When doing so, a 'recalculate' button is added to the command bar. It needs to be clicked whenever changes need to take effect. This includes opening of an experiment.

General
qbase+ allows the analysis of multiple experiments in parallel. This may be convenient for the comparison of results. However, it may also cause confusion to some users or slow down qbase+ when multiple large datasets are opened in parallel. Users can set the appropriate behavior when opening another experiment according to their needs and preferences.

Each window contains a descriptive path on the top, so you know to what experiment the window pertains. Because of this, the option is provided to include this path on chart prints.

Show sample in charts
Depending on their sample type, certain samples may be excluded from charts to avoid cluttered and absurdly scaled charts. By default, unknown samples (of interest) and positive controls will be shown. Standard (curve) samples and negative controls (all types) will not be shown. These defaults can be changed in this window, and overruled by making sample specific 'sample visibility' settings (Figure 7).

Figure 7: Preferences for sample visualization
Startup and shutdown

By default, qbase+ uses a single workspace. Advanced users that want to switch between workspaces may use the option to have qbase+ ask for the workspace to be used at startup.

The analysis wizard is launched when starting qbase+. Unchecking this option will launch qbase+ with the user interface of the three main windows.

Data management

qbase+ stores all its data in a global workspace, obviating the need to deal with storing and updating individual experiment files. Within this workspace, data and settings are hierarchically organized into projects, experiment and runs.

Workspace

Saving of data, annotations and settings typically occur in the background without any user intervention. All these data are stored in a global qbase+ workspace. Most users will store their data in a single workspace using the default location in the file system (see further). For advanced users with specific requirements it is possible to create alternative workspaces. Users may create or switch between workspaces by using the ‘Switch workspace’ option in the ‘File’ menu. If other workspaces have been used in the past, a short list of those workspaces will be shown. Otherwise, only the ‘Other’ option is available that allows browsing for a workspace. If a new empty folder is chosen, it will be treated as the location for a new workspace (Figure 8). Every new workspace will contain one project with a fixed set of demo experiments (ChIP-qPCR, Copy number analysis, Quick guide, Statistics on quick guide, geNorm pilot experiment, microRNA expression profiling).

![Workspace Launcher](image)

Figure 8: Creating a new empty workspace

By default the workspace is stored in \Users<username>\workspace on Windows or OS X. Its location can be retrieved by looking at the path shown when using the ‘Switch workspace’ option. This information can then be used to copy, move or backup the workspace. For this, simply treat the workspace as a regular folder.
It is possible to store the workspace on a network drive or in the cloud (e.g. on Dropbox). In view of the continuous data access that is required, this may not be a favorable approach when working with larger datasets. One should definitely refrain from sharing a workspace between different qbase+ users because the software has not been designed for parallel use. Simultaneous access to the same workspace may lead to inconsistent or even corrupted workspaces.

Projects

Projects are logical groups of related experiments. In a way, they have a role similar to that of folders in a file system. As such, a project has no direct implication on the way results are calculated in any of the experiments contained in the project.

To create a new project, click the command in the qbase+ toolbar and select 'create project'. Alternatively, use the 'new project' option in the context menu of any existing project. Freshly created projects are empty containers in which new experiments can be started or imported.

Experiments

Experiments are the set of data (stored in runs, see further), annotations and settings used for qPCR data analysis. Several experiments can go into the same project, but a single experiment should not contain data from unrelated qPCR measurements. Merging distinct experimental data into a single experiment will cause confusion at best, and may result in misinterpretations at worst. Because of this, it is important to put unrelated datasets in different experiments.

To create a new experiment, click the command in the qbase+ toolbar and select 'create experiment'. Alternatively, use the 'new experiment' option in the context menu of any of your projects. Newly created experiments are empty containers with default settings, but no data.

Runs

Runs are collections of qPCR data coming from a single plate, array, rotor or chip (depending on the instrument being used). An experiment may consist of a single run corresponding to a 96-well plate that is not completely used, or may comprise tens of runs of high-throughput qPCR instruments.

Runs cannot be created in qbase+, they can only be imported. Single color runs are a 1-on-1 representation of the plate as it was ran. Multiplex plates, on the other hand, are being split up in different runs (each run name with the probe color as a suffix) to allow for proper editing and visualization of the layout for the different colors.
**Data exchange**

qbase+ supports several ways to exchange data, at the level of a workspace, project or experiment.

It is possible to exchange entire workspaces. Although it encompasses the entire dataset, it is reserved for special situations. It can only be done by looking up the workspace location, and completing the exchange by copying its corresponding folder in the file system. This type of exchange is not part of qbase+.

Projects can be exchanged in the RDML data standard. This is ideally suited for publishing an entire dataset, or in general for the exchange of data with collaborators that don’t use qbase+. The drawback is that not all qbase+ settings can be saved in the RDML data format. Because of this, results may no longer be the same after having exported into and imported from an RDML file.

The best way to exchange data between qbase+ users is the export/import of experiment files. These files do not only contain the data, but also all qbase+ related annotations and settings. The files are in Extensible Markup Language (*.xml) format and can be easily compressed (zipped) to make them substantially smaller.

Both projects and experiments can be exported using the upward pointing arrow in the command bar. After having chosen the export type, one should select the project or experiment for export and provide a location and file name to which to save the data. Data export is also available as a context menu option for projects or experiments.

**Data import**

qbase+ can import five different types of data: projects, experiments, runs, samples and targets (Figure 9). The first two are mainly intended for the exchange of data, run import is used to get the primary data into qbase+, whereas the import of samples and targets is intended for purpose of data annotation. Any import can be initiated by clicking the downward pointing arrow in the command bar. The import procedure is completed by selecting an import type and a file to be imported.
Projects and experiments

qbase+ can import entire projects saved in the RDML format, or experiments in a qbase+ specific format (*.xml). See ‘Data management’ for more information on projects and experiments.

When importing projects or experiments, some restrictions apply to their names. Only the following characters are allowed: all alphanumerical characters (0-9, a-z, A-Z), space, _, -, $, #, :, ^, . and the Greek letter mu (µ). Illegal characters such as brackets and slashes should be removed or replaced.

Runs

For most users there is no need to modify the data file as qbase+ supports export files from the majority of the real-time PCR instruments and accompanying data analysis software. qbase+ cannot read the binary files that are used by the various instrument specific data collection softwares, nor can it analyze raw fluorescence data. Instead, a data file containing quantification cycle (Cq) values needs to be exported from the data collection software. Cq the MIQE standard name for Ct (cycle threshold), Cp (crossing point), and other instrument specific quantification values. For most instruments, these tabular export files (Microsoft Excel or delimited text) can be directly imported in qbase+. More information on the supported instruments and a description of the file formats can be found in a dedicated section of the qbase+ website.

If your instrument is not supported, you are advised to modify your file so it matches the generic simple, (former) qBase or RDML (real-time PCR data mark-up language) [Lefever et al., Nucleic Acids Res, 2009; http://www.rdml.org] file format. RDML is a universal, open and XML based data exchange format recommended to be used according to the MIQE guidelines [Bustin et al., Clin Chem, 2009].

Import file formats can be either a tab delimited text file (.txt), a comma or semicolon separated value file (.csv), a Microsoft Excel file (.xls or .xlsx). Please note that OpenOffice.org Calc (.ods) and proprietary or binary instrument files are not supported.

Runs are imported into experiments, and all information from one or multiple runs is stored in a qbase+ experiment. The import procedure does not alter the original data file. Whereas imported Cq values cannot be modified, other types of information (such as target and sample names) can be modified after import.
Open experiment
Open the project (ँ) and experiment (०) of interest in which the run(s) need(s) to be imported. If needed, create a new experiment as follows: right click on the project (ँ) in the qbase+ project explorer tree in which you want to start a new experiment and click new experiment.

Launch import wizard
Start the import wizard by clicking the import button (१) in the command bar or by using the right-click context menu. Choose Import run and click Next.

Select the experiment in which the run needs to be imported in the top part of the import wizard and browse for the file(s) to be imported (Figure 10). Multiple runs can be imported at once using the 'Shift' or 'Ctrl' key while selecting the data files.

![Image of import wizard](image)

Figure 10: Selection of runs to be imported in active experiment

Runs can only be imported into loaded (open) experiments. Experiments that are not open are not available for selection.

Users with a basic license can import and analyze no more than ten 96-well runs within one experiment.
Run file format

qbase+ will try to recognize the format of the selected import files. If only one format matches your file(s), it will be selected and the quick import option is enabled.

Figure 11: Auto detection of run file format

Quick import is disabled if the file is not recognized or if multiple formats match your run file. To support further development of run importers, qbase+ offers the option to upload your run file to the qbase+ support team (Figure 12). Your data will be treated confidentially, and will only be used to improve the importers of qbase+. It is advised to complete the extra information fields because it does support our effort to improve the qbase+ run importers. There is no need to upload the same, or identically formatted files, over again.

Figure 12: Uploading unrecognized import file

If a file is not recognized automatically, it may still be imported by manually selecting the corresponding instrument in the last step of the run import wizard.
**Extract sample and target information**

If an annotated run (containing Cq values, sample and target names, and quantities in case of standard samples) is imported, a sample and target list is automatically generated that can be accessed in the project explorer (see chapter 'Run annotation'). For instruments that provide only a single information field per well, there exists a functionality to extract a sample name, target name, or both based on a given delimiter (Figure 13). This allows you to provide the sample and target well annotation already in your real-time PCR instrument software.

![Figure 13: Extract well information](image)

For example, well information “SampleN|GeneZ” can be imported as sample name “SampleN” and target name “GeneZ” based on the delimiter “|” in this case. Other available delimiters are: ‘-’, ‘,’ ‘.’, ‘_’, and ‘@’.

This functionality is only available for instruments that restrict export to a single information field per well: iCycler, MyiQ, iQ5, Opticon Monitor, Rotor gene, LightCycler, LightCycler 480, LightCycler 1536 and MX300xP.
Complete run import

Click ‘import’ to complete the import run wizard. You will see that the runs are added to your experiment and that they are automatically processed (unless you disabled automatic recalculation in the Preferences window, see higher).

An individual window can be opened for each imported run by double clicking the run in the project explorer tree. This allows editing of sample and target information for each well. In addition, the sample and target list in the project explorer tree is automatically updated with the sample and target names provided they are included in the import files.

More information on the annotation of runs with target and sample names, sample types and quantities for the standard samples can be found in chapter 'Run annotation'.

Samples and targets

qbase+ provides all the functionality for full annotation of wells (see ‘Run annotation’ chapter). Samples and targets are created as part of the import of an annotated run, or should be manually created in qbase+ (in case an unannotated run was imported). Since users typically already have lists of samples and targets, it may be more convenient to import those lists rather than to enter the information in qbase+. This is especially true if the information not only lists sample names, but also a series of sample property attributes (e.g. group to which a sample belongs).

The format of sample and target import files, as well as example files can be found on the qbase+ website. The import procedure for both is very similar. We here describe the procedure for the import of sample lists.

Start Import samples wizard

Right-click in the project explorer on samples (or any sample in the experiment project explorer) to start the import samples wizard. The import button ( ) in the toolbar will also lead to this wizard if selecting samples.

Select sample list

Select the experiment in which the sample list needs to be imported. By default, the experiment selected in the project explorer is indicated as the target experiment. Browse for your sample list file and click the next button.
Select properties to be copied

Select the properties to be copied from the sample list. Sample name and type (e.g. ‘unknown’, ‘standard’, ‘no template control’, etc.) are mandatory; other properties are optional (Figure 14). Click the finish button to proceed.

Figure 14: Selection of sample properties to be imported

Annotation

qbase+ support annotation at different levels: project, experiment, runs, samples and targets. Whereas annotation of projects and experiments is descriptive in nature, run annotation is important and impacts calculations.

Project and experiment annotation

Projects and experiments have two types of annotation: their name and description. Both can be modified by opening their properties windows: ‘Annotations’ or ‘Experiment properties’ for projects and experiments, respectively. When changing a project or experiment name, the text will turn blue if it is a valid name (some special characters are not allowed). A red background indicates that an invalid name has been entered. By pressing ‘Enter’ on the keyboard or clicking the save icon the name will be updated and the text will turn black again.

A shortcut for renaming projects or experiments is available in their respective context menus. Alternatively, hitting the ‘F2’ key while a project or experiment is selected allows changing its name as well.

Project and experiment descriptions are optional fields. These fields are often left blank because they seem superfluous at the moment. However, in the long run it may be beneficial to add a concise description to your projects or experiments. We recommend to add some background information (aims of the study, etc.) or other types of valuable information like references to lab notebooks. The description field
can hold multiple lines of information. Therefore the 'Enter' keyboard button cannot be used to save the information. Use the save icon to save changes to the description field (blue text will turn black).

Run annotation
Runs need to be properly annotated before experiments can be analyzed. Not only will your data be meaningless without annotation, it is simply essential for qbase+ to start any calculations.

Runs can be annotated before or after import in qbase+. If an annotated run (i.e containing a sample and target name for every well that has a Cq value) is imported, qbase+ will take over this annotation and generate a sample and a target list. All annotation can still be edited afterwards. In contrast, editing of Cq values is explicitly not supported by qbase+, although one can choose to exclude certain data points from calculations. Runs that were not annotated before import should be annotated in qbase+, more specifically in the run editor.

The qbase+ run editor does not support well annotation by typing in sample or target names. Sample and target names need to be selected from a dropdown list at the top to annotate wells. This approach avoids spelling mistakes that are often hard to spot and may undetectably lead to erroneous results.

Open run
Open a run by double-clicking its name in the project explorer, or use the open option in the run context menu. The Run window contains a properties tab at the bottom in which you can change the run name or alter the date on which the run was generated. It also contains information regarding the format of the run. Samples and targets can be appointed to wells in the matrix tab that depicts the run layout with an annotation bar on top (Figure 15).
Appoint sample and target names

Select one or multiple wells in the run layout for which you want to add a sample, and use the sample dropdown list in the upper part of the window to search for the corresponding sample name. Multiple wells can be selected by click-dragging, by using the 'Shift' or 'Ctrl' keyboard buttons, or by selecting an entire row or column by clicking on the row or column header, respectively. As soon as a sample is appointed to a well, its name and type appear in the well (unknown, positive control, standard, or 3 types of negative control). Note that the sample type can be modified in the run annotation bar on top of the run layout. In a similar way, one or multiple wells can be assigned to a certain target. To this end, the target dropdown list is used.

Although sample and target names cannot be typed directly into the wells, it is possible to annotated wells with a new sample or target name by using the '<new sample>' or '<new target>' option at the top of the dropdown boxes. This is basically a shortcut for creating a new sample or target in the project explorer, and then returning to the run editor to apply it to selected wells.
Quantities
For 'standard' (curve samples), an input quantity must be defined in the corresponding field in the run annotation bar. In contrast to sample and target information, quantities are not selected from a dropdown list but entered as a numerical value. Note that quantity values cannot be added to sample types other than 'standard'. Also, since quantities are a sample property, changing the quantity value for a standard sample in one well will alter the quantity value for that sample in all wells in which it occurs. A sample quantity can also be entered by double-clicking on the name of a 'standard' (curve) sample in the project explorer.

Exclusion
In the run layout window, you can select the wells that should to be excluded from analysis. This option does not erase the Cq value nor the sample/target annotation, meaning that excluded wells can later be re-included in the analysis if needed.

Note that empty wells and wells without Cq value are automatically excluded by qbase+ and cannot be included by the user.

Clear
The options clear sample and clear target can be used to remove sample and target names, respectively, from selected wells. It does not remove any other well information. The clear wells option removes both the sample and target annotation.

The 'undo' and 'redo' options, available in the command bar, can be used to restore accidentally removed annotation.

Apply/copy run layouts
To speed up run annotation, qbase+ enables copying the layout of samples, targets or both from one run to another. When for instance the same set of samples is measured in two different runs using the same layout (this is, the samples are measured in the same position in both runs), sample annotation of a run only needs to be performed once, saving you valuable time. This functionality can also be used across different (open) experiments.

Start the ‘Apply run layout wizard’ by clicking the corresponding option in the context menu of the run (or selection of multiple runs) for which you want to edit the run information. Then, select the run that serves as a template. Note that only runs can be selected from open (loaded) experiments. If needed, cancel the current procedure and right-click on closed (unloaded) experiment and select load experiment. Finally, select the properties that need to be copied (Figure 16). If annotation information is already present in the destination run, a warning will be displayed to notify you that information will be overwritten.
Sample annotation

There are three ways by which sample information can be provided to qbase+: as part of the import of a run, by importing a sample list or by manually adding a sample.

Creating new samples

There are four ways to create new samples:

1. by making use of the right-click context menu in the project explorer (right click on the samples node and select new sample)
2. by starting the new wizard by clicking on the command bar
3. by using the menu bar (File > New)
4. available in the run window only: select <new sample> from the sample drop down box

In case of option 2 and 3: choose sample and subsequently select the experiment to which the sample belongs. Click finish to complete the new wizard. Note that samples can only be added to loaded (open) experiments. Experiments that are not loaded are available for selection, but no sample can be added.
**Sample properties**

Upon creation of a new sample, the sample window opens (Figure 17). This window can also be accessed later by double-clicking the sample of interest. It allows for reviewing and editing of sample annotation, including the sample name and type (unknown sample, negative control, positive control, or standard), a custom normalization factor and options for visibility in charts.

![Sample window](image1)

Figure 17: Sample type and information

When editing the sample description, blue text indicates unsaved entries. Modifications can be saved by clicking the disc, or by closing the sample window. In the latter case, a Save resource window opens that allows to save the modifications. A red background indicates that illegal characters have been used (see higher for overview of accepted characters).

Quantity values are linked to samples of type 'standard'. Since one sample can only have one quantity, the different dilutions of a given template should be given different sample names (e.g. std1, std2, …), each with their quantity. If the import contains different quantities for a single standard (curve) sample, qbase+ will only retain the last value that is imported. You will need to manually edit sample names and quantities.
The custom normalization factor option allows the provision of a sample specific custom normalization factor. This option enables normalization based on a user provided value such as cell number count or mass/volume of the sample. Such normalization can only be performed if a custom normalization factor is entered (or imported) for all samples of interest.

The sample visibility in charts option allows indicating whether a specific sample or sample type has to be shown in charts. By default, negative controls and standard curve samples will not be shown. This results in more intuitive charts and gives more control on the content and look of your results.

Preferences can be modified in top menu (Window > Preferences). By default, only unknown samples and positive control samples are shown in the bar chart. Sample specific settings overrule the preferences.

Custom sample properties

In addition to the generic sample properties that have been described above, qbase+ also supports custom properties. These allow the annotation of samples with information that may vary between users and from one experiment to another. Examples include treatment, passage number, cell type, etc. Custom sample properties are always composed of a property and a property value, with properties being common across all samples and property values that may vary between samples. For example: treatment - control, treatment - low concentration, treatment - high concentration, or cell type - chondrocyte, cell type - osteoblast.

Custom sample properties are useful to:

- group results in the bar charts
- rescale results to a group (e.g. control samples)
- perform statistical analyses by comparing grouped samples (i.e. samples that have the same custom sample property)

Custom sample properties can be entered manually or imported as part of a sample list. To add or edit custom sample properties, open the sample properties window that is located in the annotations section. This window contains a list of samples that may or may not have custom samples attached to them (Figure 18). Use the buttons at the bottom of the window to add, remove or rename custom properties. On creation of a new custom property, an extra column is added to the table. Sample specific property values can be entered in this column.
Custom sample properties can also be reviewed in the properties tab of a sample window. This window does however not support editing of custom sample properties.

**Target annotation**

There are three ways by which target information can be provided to qbase+:

1. as part of the import of a run
2. by importing a target list
3. by manually adding a sample.

**Creating new targets**

There are four ways to create new targets:

1. by making use of the right-click context menu in the project explorer (right click on the targets node and select new target)
2. by starting the new wizard by clicking on the command bar
3. by using the menu bar (File > New)
4. select <New target> from the target drop down box (available in the run window only)
In case of option 2 and 3: choose **target** and subsequently select the experiment to which the target needs to be added (Figure 19). Click **finish** to complete the new wizard. Note that targets can only be added to loaded (open) experiments. Experiments that are not loaded are available for selection, but no targets can be added.

![Figure 19: Create new target in experiment ...](image)

**Target properties**

On creation of a new target, the **target** window opens. This window can also be accessed later by double clicking the target of interest. When creating a new target, the **target** window opens in the **properties** tab. For existing targets, the **target** window opens in the Bar chart. For reviewing and editing of target annotations one needs to activate the **properties** tab.

There are two types of target information that are essential to qbase+: the target name and the target type. The remaining information fields are there for annotation purposes only.

**Target type**

Two types of targets are used in qbase+, **targets of interest** and **reference targets**. Reference targets (also referred to as housekeeping genes, a name deprecated by the MIQE guidelines, Bustin et al., 2009) are used for normalization. The target type can be defined via the **target properties window** (see above), or by using the **set target type** option from the context menu of selected targets in the **project explorer** window. Multiple targets can be simultaneously appointed as **reference targets** or
as targets of interest by selecting all of them using the 'Shift' or the 'Ctrl' (Windows) or 'Command' (MacOSX) keyboard buttons.

Note that all targets labeled as reference targets are used in the multiple reference gene normalization procedure (Vandesompele et al., 2002). To exclude one or more reference targets, change their type into target of interest, which causes the software to treat them as if they were targets of interest.

Renaming and clean-up

Existing sample or target names can be modified (rename) by double clicking their name in the project explorer. A window opens where you can change the name (and other properties). Alternatively, a name can be modified using the menu bar (Edit > Rename), via the context menu or by using the 'F2' keyboard button (Figure 20).

Figure 20: Sample renaming

It is not possible to rename a sample or target to a name that is already in use. If a given sample (or target) exists with two different names (e.g. due to typing error), the replace option in the sample (target context menu should be used to replace the incorrect sample (target) name with the correct one (to be selected from a drop down list containing existing names).

Samples and targets can be deleted via their context menu (cursor needs to be on the sample or target that should be deleted) or by selecting and clicking the delete icon ✗ in the command bar. Multiple samples or targets can be simultaneously deleted by selecting all of them.

Some run import types provide default sample/target names for all wells if annotation has not been performed in the qPCR instrument software. This may lead to long lists of samples/targets that clutter the experiment and impede proper run annotation. qbase+ contains a clean-up function that removes all unused samples or targets. It is typically applied after having cleared all incorrect run annotations. The clean-up function can be found as an option in the context menu of samples and targets, but not in the context menus of individual samples or targets.
Calculation parameters and quality control settings

The calculation parameters and quality control settings windows can be opened by double clicking their respective icons under the settings node of your experiment in the project explorer tree. Similar settings can be found at the level of a project. These default experiment settings will not affect any existing experiment within that project, but will be used as the default values when creating new experiments.

Calculation parameters

The calculation parameters window (Figure 21) contains four types of parameters setting the type of amplification efficiency correction, the most appropriate normalization method, the method to calculate average Cq values or the target scaling mode.

![Figure 21: Calculation parameter window](image)

The first box in the calculation parameters window allows defining the PCR amplification efficiency to be used for calculations: one default amplification efficiency for all targets, a target specific amplification efficiency, or a target and run specific amplification efficiency. More information about the amplification efficiency correction types can be found in the chapter ‘PCR efficiency correction’.

The second box in the calculation parameters window allows defining the most appropriate normalization strategy. By default, the ‘reference target(s)’ normalization approach is selected, as this is the most commonly used strategy ([Vandesompele et al., 2002]).
In addition to normalization using one or multiple reference genes, qbase+ supports a range of different normalization procedures to suit the specific needs of different types of experiments.

**Reference targets**

This is qbase+'s default normalization procedure. Depending on the number of selected reference targets, relative quantities will be normalized by the relative quantity of a single reference target or by the geometric mean of the relative quantities for all reference targets.

**Global mean - premium license only**

The global mean normalization procedure was initially developed for the normalization of extended miRNA screening experiments where it was shown to be a superior alternative to the commonly used small nuclear and small nucleolar RNAs as reference targets (Mestdagh et al., Genome Biology, 2009). It is however useful for any experiment in which a sufficiently large set of unbiased genes is quantified. It is based on the same principles commonly used for microarray normalization. The results for a given sample will be normalized by the geometric mean of the relative quantities of all the targets that are expressed in that sample. The global mean normalization method in qbase+ is an improved version of the Mestdagh et al. method by giving equal weight to each target [D’haene et al., Methods in Molecular Biology, 20112].

**Global mean on common targets - premium license only**

Similar to the 'Global mean' normalization procedure with the exception that the normalization factor will only be based on the targets that are expressed in all samples.

**Custom value**

This option enables normalization based on a user provided value such as cell number. In this method, a custom normalization factor should be provided for every sample, or be imported as a sample property.

**None**

This option is included to enable qPCR data-analysis for which normalization may not be appropriate, e.g. absolute quantification or single cell analysis.

The third box in the calculation parameters window allows defining the method by which average Cq values are calculated. In addition to the (default) arithmetic mean on replicated Cq value measurements, qbase+ also supports the calculation of median Cq values. The median Cq value is a more robust measure than the arithmetic mean when confronted with outliers and having at least 3 PCR replicates. Median results (without outlier removal) are a good alternative to arithmetic averages calculated on datasets in which outliers have been removed.
The fourth box in the calculation parameters window allows setting the scaling of the normalized relative quantities (NRQ values) and the corresponding target bar charts. Please note, that target scaling will not affect the result of the analysis. By default, the results are scaled to the average across all unknown samples per target, which means that the average across all unknown samples is set to one. Similarly, when selecting the option scale to maximum, scale to minimum, or scale to sample the maximum, minimum, or a particular sample are set to 1. scale to group is only possible after having defined sample groups in the Sample properties window. scale to positive control is useful for copy number analysis and allows you to indicate the copy number in your calibrator samples (see chapter 'Copy number analysis').

Quality control settings

Quality control is an important aspect of qbase+. The program contains several types of quality controls (see dedicated manual chapters on quality control for more information). The parameters for these quality controls can be defined in the quality control setting window (Project Explorer > Settings > Quality control settings) (Figure 22).

Figure 22: Quality control settings window
The first box in the quality control settings window contains parameters that do not affect calculations, but merely set the thresholds beyond which data are flagged for low quality. The quality control parameters include a threshold for maximum replicate variability, a range of acceptable Cq values for positive controls, a delta-Cq value for the interpretation of negative controls, and two thresholds for assessment of reference target stability.

In the second box in the quality control settings window contains criteria for automatic exclusion of certain data points. These settings do impact the final experiment results. The 'Difference to negative control sample < …' is used to automatically exclude data points that could be significantly impacted by the signal found in the negative control. The 'Well with too high Cq > …' and 'Well with too low Cq > …' are used to automatically exclude data points in a Cq range with inaccurate results. Similarly, the 'Target-sample combination with average Cq value > …' can be used to automatically exclude replicates with an average Cq in the range with inaccurate results. The latter two options are particularly relevant when analyzing whole genome miRNA expression profiles with the new global mean normalization approach.

Data points that have been automatically excluded are grayed out in the replicate quality control. These data points, like those that have been manually excluded, will not be used for calculations. In contrast to manually excluded data points they cannot be re-included in the replicate quality control screen or the run editor, i.e. they are strictly linked to the auto exclusion settings.

Specific for qPCR-based copy number analysis (premium license only) is the definition of the thresholds for the lower boundary for normal copy and the upper boundary for normal copy in the third box. These thresholds are used for conditional bar coloring for easy detection of deletions and amplifications and are by default set to 1.414 (geometric mean of 1 and 2 copies) and 2.449 (geometric mean of 2 and 3). These default settings are recommended for a diploid organism (like human, mouse and rat), and may need to be adjusted for polyploid organisms.
Inter-run calibration

Inter-run calibration is a calculation procedure to detect and remove (often underestimated) inter-run variation. Whenever samples need to be compared that are measured in different runs, one should be cautious of this potential bias. Importantly, inter-run calibration is needed for each gene separately. Detailed information is available in the original qBase paper (Hellemans et al., Genome Biology, 2007).

The basic principle is that the experimenter measures one or (preferentially) more identical samples in different (to be calibrated) runs, in addition to the other samples that are spread across the runs. The results for these identical samples (so-called inter-run calibrator or IRC samples) can then be used to quantify and correct inter-run variation. By measuring the difference in Cq or normalized relative quantity between the IRCs in the different runs, it is possible to calculate a correction or calibration factor to remove the run-to-run difference, and proceed as if all samples were measured in the same run.

The inter-run calibration concept

In a relative quantification study, the experimenter is usually interested in comparing the expression level of a particular gene between different samples. Reliable estimates for relative expression levels can only be obtained by minimizing and correcting technical variations between samples and measurements. It is well recognized that variations in the target nucleic acids input amount between samples need to be corrected by means of normalization, typically using one or multiple reference genes. Run-to-run variation within a series of measurements for a given gene is a second type of technical variation that needs to be avoided, minimized or corrected for.

We advise to follow the sample maximization method (Figure 23), which dictates that all samples (or as many as possible) for a given gene should be analyzed in the same run. The sample maximization strategy does not suffer from technical (run-to-run) variation between samples, and therefore does not require inter-run calibration to be performed. Since reference genes, independently of the other assays, quantify the relative concentration of nucleic acids between samples there is no need to repeat the measurement of the reference genes in every run.
If the sample maximization approach cannot be followed because of experimental design (e.g. prospective studies where not all samples are available at the start of the study), or in studies where the number of samples exceeds the number of available wells in a given run, technical run-to-run variation will arise when comparing samples between runs. This variation will need to be corrected for by means of inter-run calibration (Hellemans et al., Genome Biology, 2007).

If not all samples are analyzed in the same run, the experimenter needs to analyze inter-run calibrators (IRCs), which are identical samples that are measured in different runs. It is advisable to use multiple IRCs because it will give more precise results. In addition, a failed calibrator does not ruin the entire experiment if two or more are available. We recommend 3 IRC samples (analogous to the use of 3 reference genes for normalization).
Different types of input material can be used to create IRC samples: real samples for which sufficient material is available to measure them in different runs; pools of samples; synthetic templates (e.g. external oligonucleotide standards, see Vermeulen et al., Nucleic Acids Research, 2009); etc. Serial dilutions (standard curve) often make good IRC samples (make sure to give each dilution a different sample name in qbase+).

Importantly, inter-run calibration should be performed on a gene per gene basis. It is not sufficient to quantify the inter-run variation for a single assay and to extrapolate this result to other assays. The experimenter should perform inter-run calibration for all genes/assays.

**Inter-run calibration in qbase+**

qbase+ performs inter-run calibration after data normalization. This does not only result in fewer calculations, but provides more flexibility as a new batch of cDNA can be synthesized from the inter-run calibrator RNA samples while still remaining a valid inter-run calibrator. It does however put some limitations on the design of the study.

When designing a qPCR experiment and determining how measurements will be spread across runs, different approaches can be chosen: 1) all samples for a given gene within a single run, 2) all genes for a given sample within a single run, 3) both samples and genes are spread across runs. The first approach does not require inter-run calibration, the second approach is 100% compatible with the inter-run calibration that is being performed by qbase+. Inter-run calibration for the third approach is only valid when specific requirements are met: the same set of inter-run calibrators is used in every run (and for every gene) and samples are grouped into runs in the same way across the entire experiment.

qbase+ recognizes PCR replicates based on identical sample and target names. For qbase+ not to interpret IRC data points as PCR replicates, the IRCs should have a different sample name in each run. Assuming sampleA is used as inter-run calibrator, it could be named sampleA_1 in the first run, sampleA_2 in the 2nd run, etc. The same procedure should be used for all genes within your experiment (to make sure normalization is performed correctly).

Some researchers deliberately split PCR replicates over 2 runs. Although this approach is supported by qbase+, it is not compatible with inter-run calibration. If inter-run variation is a concern that you want to correct for by means of inter-run calibration, you should not introduce this type of variation by spreading replicates across run.
If samples are analyzed across multiple runs for a given target and if no IRC samples have been appointed, the user will be notified by an alert message in the alerts window (Figure 24) and by the appearance of the message 'IRC Missing' in the calibration factors window.

Figure 24: Alert for potential inter run variation

Open the interrun calibration window
Open the interrun calibration window from the project explorer (intermediate results > interrun calibration) by double clicking it, or by using the open option from its context menu. Two tabs are available in this window: interrun calibrators and calibration factors. The calibration factors tab is shown by default (Figure 25). Because no IRC information is available, all calibration factors (CF) are set to 1.

Figure 25: Defining inter run calibrators
**Appoint IRC samples**

Create the first IRC by clicking in the *interrun calibrators* window. Upon clicking, a new IRC (default name: IRC 1) will appear in the left part of the *interrun calibration* window. The right part of the window provides an overview of the samples that have been linked to the selected inter-run calibrator (top), as well as a list with remaining samples from which additional ones can be linked to the selected IRC (bottom).

Select the first sample and click (or double click on the sample). The sample will appear in the *interrun calibrator details* section. Other samples can be appointed to this IRC in the same way. Samples can be removed by clicking . Additional IRC samples are added by repeating this procedure.

**Evaluate the calibration factors**

Once all IRC samples are defined, calibration factors for each target can be found in the *calibration factors* tab. When selecting a target from the Targets list, its run-specific calibration factors are shown, as well as the corresponding propagated error. qbase+ reports the calibrated normalized relative quantities (CNRQ values) in real-time.

Quality control on inter-run calibration is only possible when more than one inter-run calibrator has been included in the experiment. All inter-run calibrators should measure the technical variation between the runs to the same extent (Figure 26). If one inter-run calibrator returns a substantially different interpretation of the technical variation, the validity of this inter-run calibrator should be interpreted with care. The *replicate quality control* window may help revealing technical problems with such an inter-run calibrator. Exclusion of an inter-run calibrator is perfectly valid as long as it is excluded from all runs for that particular gene, i.e. the same set of inter-run calibrators should be used in all runs.

![Figure 26: Interpretation of inter run calibration factors](image-url)
Quality control and results

PCR efficiency correction

qbase+ employs a universal and flexible quantification model (Hellemans et al., Genome Biology, 2007). Depending on the selected option in the amplification efficiency box of the calculation parameters window, this model will either equal the delta-delta-Ct method described by Livak and Schmittgen or the efficiency corrected model described by Pfaffl. The former assumes 100% amplification efficiency for all targets, whereas the latter supports correction for target specific amplification efficiencies.

There are several ways to approach PCR efficiencies. One could assume that all assays perform perfectly. This is the most straightforward, but not the recommended approach. We advise to assess the amplification efficiency for all targets. The gold standard for efficiency correction is the use of standard curves. This approach is supported by qbase+. Alternatively, any of the various algorithms that have been described to estimate the PCR efficiency based on a single amplification curve might be used. Unfortunately, many of these methods do not provide a precise or an accurate estimate of the efficiency. One should therefore be cautious in using these single reaction PCR efficiency values; it is generally recommended to calculate the mean efficiency of all reaction wells in which the same target is amplified (more information in Karlen et al., BMC Bioinformatics, 2007 – Jan Ruyter). Since qbase+ does not read raw fluorescence values, this latter approach can not be supported.

qbase+ uses the base of the exponential function (E value) as the amplification efficiency value for relative quantification (see formula 5 in Hellemans et al., Genome Biology, 2007). This base is the efficiency value + 1, e.g. an E value of 1.95 for 95% efficiency (efficiency value of 0.95).

Use one default amplification efficiency

When using the same amplification efficiency for all targets, either as an assumption or because this has been proven before, one should select the 'One default amplification efficiency' option in the calculation parameters window. The actual value for E (efficiency + 1) can also be entered in this window. If known, the error on the estimated efficiency could also be provided. Default values for E and SE(E) are 2 and 0, respectively. These values correspond to what is assumed in the delta-delta-Cq quantification model as originally described by Livak and Schmittgen (Methods, 2001).
Use target specific amplification efficiencies

The gold standard method for PCR efficiency estimation is a serial dilution of nucleic acid template that is as similar to the samples under investigation as possible (e.g. a mixture of cDNA from a representative set of your samples). We have observed that the use of small synthetic templates (e.g. comprised of the first 30 and the last 30 bases of the amplicon sequence) is a cheap and easy alternative that results in equivalent efficiency estimates.

qbase+ offers the possibility to make use of target specific amplification efficiencies by selecting the ‘target specific amplification efficiencies’ option in the calculation parameters window. This option may be used for experiments in which serial dilutions or standard curves are included, or whenever these efficiency values have been determined before.

The E value can be calculated from the slope of a serial dilution as follows: $E = 10^\left(-\frac{1}{\text{slope}}\right)$ (with an E of 2 being perfect, indicating 100% efficiency). In our own experiments, we aim for E values in the range of 1.90 – 2.10 (PCR efficiency between 90 and 110%) with standard errors typically below 0.01 (1%). The Y-intercept is only meaningful when actual concentrations or template copy counts are used to define the quantities for standard samples. It may be used to estimate the Cq value that is expected for qPCR reactions with just a single molecule of template. Cq values beyond this Y-intercept value may indicate primer dimer formation (for SYBR reactions) or inhibited amplification. The Y-intercept may also provide valuable information about the Cq range in which a considerable degree of sampling noise may be expected; this is down to 3 or 4 cycles below the Y-intercept. It may also yield additional insights in the performance of the assay. High Y-intercepts typically coincide with low amplification efficiencies. If these occur for assays with good efficiencies (90%–110%) they may indicate problems to generate the first PCR copies, e.g. due to strong secondary structures in or around the primer binding sites.

The amplification efficiency window can be opened by double clicking ‘amplification efficiency’ in the project explorer (Figure 27). Depending on the option chosen in the calculation parameters window either the computed or user-defined is greyed out (disabled). With ‘Target specific amplification efficiencies’ set to user-defined the user has the opportunity to manually enter an efficiency value for each target by providing a ‘user-defined’ E-value.
By unticking the box in front of an outlier well, certain dilution points can be excluded from the calculation of the efficiency value. By hovering over a data point in the standard curve, the identity of an outlier reaction is easily determined.

**Make amplification efficiencies run specific**

If a target is measured in different runs, and a serial dilution for that target is present in each run, qbase+ offers the possibility to use a run-specific target amplification efficiency. Hence, each Cq value will be converted to a relative quantity (RQ) using the efficiency measured in that particular run.

**Quality controls**

Quality control is an important feature in qbase+. The program contains several types of quality control that can be accessed by double-clicking the quality control icon in the project explorer:

- quality control on technical replicates
- quality control on positive and negative controls
- quality control on the stability of reference targets
- quality control on different sample specific characteristics

The parameters for quality control can be defined in the quality control setting window (Project Explorer > Settings > Quality control settings).

More information about post-PCR quality control can be found in the paper ‘The Importance of Quality Control During qPCR Data Analysis’ (D’haene and Hellemans, International Drug Discovery, 2010).
**Technical replicates**

qbase+ automatically deals with technical replicates or repeated measurements, which are recognized as different PCR wells with an identical sample and target name. Importantly, we highly recommend measuring all PCR replicates in the same run. If PCR replicates are measured in different runs, an alert will be displayed in the alert window (‘PCR replicates (wells with identical sample and target name) are spread across runs. This assumes that you have controlled for inter-run variation.’).

The Cq values of all (non-empty) wells of a replicate group are averaged at the very beginning of the calculation process. Outliers can be excluded so they do not take part in the calculations.

**Inspecting replicate quality control**

The replicates window (Project explorer > Quality control) contains an overview of all replicate groups (defined by wells with the same target and sample name) (Figure 28). The summary on top of the list contains the pass rate of the replicates and informs the user about the overall replicate quality of the experiment.

![Replicates window](image)

Figure 28: Table of replicates failing to meet the set quality criteria

qbase+ can selectively display the failed replicates, i.e. replicates that do not meet the quality threshold as defined in the quality control settings. This is done by checking the ‘failing’ option in the well selection box. Other visualization options include ‘all’ and ‘excluded’. The first option may be disabled for very large experiments. The last option can be used to reinclude data points that have been incorrectly excluded, or to inspect the wells that have been excluded by means of auto exclusion.

In larger experiments it may be interesting to inspect the distribution of replicate variability. A delta-Cq frequency histogram can be inspected in normal or in cumulative mode by clicking on the corresponding tabs (frequency or cumulative) at the bottom of the replicates window.
Excluding bad replicates

Failing replicates can be excluded from further calculations by removing the tick in front of the well (second column in the table). The number of failed replicates is determined by the quality of the data in combination with the quality control settings. qbase+ flags bad replicates based on a user-defined maximum allowed difference in Cq value. This value can be defined in the quality control settings window in the replicate variability box. By default, the replicate variability threshold is set to 0.5, which means that the difference in Cq value between the replicate with the highest Cq value and the replicate with the lowest Cq value (\( \Delta Cq \)) must be smaller than 0.5 cycles.

If a more stringent quality control is used (which is equal to a lower threshold value), more replicates will fail the quality control: the list of failed replicates will be longer and the summary will report a lower pass rate.

Individual PCR replicates should only be excluded if there is a good, objective reason to do so (e.g., abnormal melt curve, no sample added). When in doubt, keep all replicates, as the higher replicate variability will simply result in a larger propagated error on the final result. With 3 or more PCR replicates one could look for statistical outliers among the PCR replicates. Alternatively, one could simply opt to use the median Cq instead of the arithmetic mean of Cq values in the calculation parameters window. This approach results in a more robust measure for the average Cq without the need for meticulous inspection and correction of all replicates failing the quality control.

PCR replicates (repeated measurements of the same sample in the same run) are useful for a number of reasons:

- it allows for quality control on the technical reproducibility of the qPCR data
- provides better accuracy
- allows the generation of results when individual qPCR reactions failed

Strict guidelines on the optimal number of PCR replicates cannot be given since it depends on the purpose of the study (e.g., diagnostics versus research), the quality of the PCR assay, the precision of the qPCR instrument, the Cq determination method, and the pipetting skills of the operator. Ultimately, it all comes down to determine how much confidence is needed for a given data point.

Important to note is that the biological variability is often much larger than the technical variability. Therefore, it is acceptable to omit PCR replicates when the sample size is sufficiently large or when a screening experiment is run (as is often the case for miRNA expression profiling studies).
By placing the quality control settings window next to the replicates window, you can see the effect of changing the replicate variability threshold in real-time. More information on the flexible use of windows in qbase+ is available in the 'User interface' chapter.

**Positive and negative controls**

The second type of quality control is evaluation of the positive and negative sample controls. The +/- controls window (Project explorer > Quality Control) shows a summary of the QC results and a list of the samples that can be filtered for those that fail the quality control (Figure 29). qbase+ flags suspicious no-template control (NTC) samples based on a user defined threshold that can be found in the quality control settings window. The default value is flagging of non-negative NTC results that are less than 5 cycles away from the sample of interest with the highest Cq value. For positive controls, a range of acceptable Cq values can be set in the quality control settings window. The default range includes any meaningful Cq value, i.e. Cq between 5 and 35.

![Figure 29: List of data for all negative controls in the +/- controls window](image)

In order to get a result for the positive and negative controls, positive and negative control samples should be appropriately labeled in the sample list. If not, no results are shown in the +/- controls window. More information on the annotation of samples can be found in the 'Annotation' chapter.

An amplification signal in the NTC sample indicates a potential contamination issue or formation of primer dimers. Such problems can be ignored as long as the difference in Cq value between the NTC and the sample with the highest Cq value is sufficiently large. For example, a Cq value difference of 5 corresponds to a fold difference of about 32, indicating that approximately 3% of the signal in your samples may be caused by these unwanted signals (well below the technical error on PCR replicates). Smaller differences between the NTC and the unknown samples should be avoided.
Stability of reference targets

When using the multiple reference target normalization approach (Project explorer > Settings > Calculation parameters), reference target stability is the third type of quality control. The user is able to choose a minimal acceptable reference target stability by defining a threshold value for two indicators of expression stability of the used reference genes: the geNorm expression stability value of the reference gene (M) and the coefficient of variation of the normalized reference gene relative quantities (CV) [Vandesompele et al., Genome Biology, 2002; Hellemans et al., Genome Biology, 2007]. Both parameters are automatically calculated by qbase+ during the analysis. More information on finding the best set of reference genes in a given experimental condition can be found in the 'geNorm' chapter.

In order to get a result for the reference target stability, at least two reference targets have to be appointed. No quality control results will be shown in the reference target stability window (Figure 30 – top) if no reference targets or just a single one have been appointed. The algorithm used to evaluate the reference target stability does require all samples to have results for all their reference targets. In case of missing data, no stability measures can be calculated and NaN is shown in the reference target stability window. Either deleting those sample(s) or making the reference target with missing data a target of interest.

The threshold value for M and CV can be set in the quality control settings window (Figure 30 – bottom). By default, these thresholds are set to 0.5 for the M value and 0.2 for the CV value. M and CV values passing the QC are highlighted in green in the reference target stability window while failing ones are highlighted in red.

Figure 30: Quality control on reference target stability
Sample quality control

The sample quality control window covers three ways to evaluate whether certain samples fall outside the expected range in terms of their normalization factor, the fraction of detected targets or the average Cq value. The tabs at the bottom of this window can be used to switch between the different options to quality control samples.

Normalization factors

Inspection of normalization factors allows you to inspect possible experimental problems. Using approximately equal amounts of equal quality input material and stably expressed reference genes, the normalization factor values should be similar for all samples. Evaluation of normalization factors may reveal samples for which less than the expected quantity of DNA was detected by the reference target assays. Possible explanations include:

- a known shortage of template DNA
- an unexpectedly low cDNA quantity resulting from RNA degradation or failed reverse transcription
- the presence of inhibitors leading to impaired PCR amplification
- differential expression of reference targets in this sample resulting in an unreliable normalization

A variation of 2- to 3-fold is generally acceptable (this is the experimental variation that you want to remove in the normalization process). Any higher variation should be treated with care (Figure 31).

Figure 31: Quality control on sample normalization with a potential issue in sample 13
The results of this quality control window are identical to those in the normalization factors window as part of the intermediate results in the project explorer. In addition to the graphical representation used for sample quality control, that window can also visualize results in tabular format. Use the tabs at the bottom to switch graphical and tabular format. This table can be exported by using the export normalization factors option in the normalization factor context menu, or by using the export wizard launched by clicking the upwards pointing arrow in the command bar.

**Detected targets**

Assessment of the fraction of targets being detected per sample is typically applied in screening experiments comparing the expression levels of larger numbers of genes (including miRNAs). The results of this analysis may be used as an alternative to the normalization factor analysis in experiments that have been normalized with the global mean approach.

**Mean Cq**

Inspection of mean Cq values per sample is a third alternative for analysis of the consequences of sample quantities. It may also be used to reveal challenges with lowly expressed genes, having Cq values close to the detection limit.

**Bar charts, correlation plots and results tables**

qbase+ supports several ways to investigate qPCR experiment results. Most will use target bar charts for the comparison of expression levels for a given gene between different samples. In addition to this, qbase+ also supports the comparison of expression profiles between different genes, either in a multi-target bar chart or in a correlation plot. All results are also available in tabular format, most relevant for further downstream processing.

Without altering the interpretation of the results, the actual values can still be changed by selecting a different scaling option. By default, results are scaled to the average for all the samples of a given target using the mathematics described in the qBase paper. It may be more convenient to apply rescaling to a given sample or sample group, often a reference sample or a group of normal controls.

**Target bar charts**

qbase+ has two ways to investigate qPCR results for a given target in bar charts. Similar charts are obtained by double clicking a target in the project explorer or by opening the target bar chart window located in the analysis section of the project explorer (Figure 32). The charts and the options are identical between both approaches. They do however differ in the fact that the first approach is specific for the selected target, whereas the latter approach allows for easy switching between different targets. One approach may be more convenient for the interpretation of
just a single target or for the side-by-side comparison of two targets. The other approach is better suited for the serial analysis of one target after the other.

Figure 32: Target bar chart

The **target bar chart** plots the relative quantities of one sample compared to the others. The scale of these graphs is meaningless – the ratios between samples are of importance. The error bars on the bars represent the standard errors of the relative quantities. The size of these error bars has been calculated by propagating all relevant errors: replicate variability, uncertainty on the estimated amplification efficiency and the error on the normalization factors. The mathematics behind these error calculations are described in the qBase paper. Because of the multiple sources of errors it is possible that the results for a sample with just one replicate do have error bars.

The bottom of Target bar chart window contains a few options to change the way results are visualized. The Y-axis can be plot in linear or logarithmic scale. In view of the nature of relative expression levels (expressed as fold changes) it makes perfect sense to plot the result in log scale. On the other hand, many users may be more familiar with the linear scale.
The ordering and grouping options affect the way samples are plot on the X-axis. Samples can be ordered alphanumerically or according to their expression levels (small to big or big to small). The option to group samples is available in experiments in which custom sample properties have been entered. Samples can be ordered according to two properties. The second property is nested within the first. By default grouping only affects the order of samples, but not the results. By selecting the ‘plot group averages’ instead of ‘plot individual samples’ the default behavior is changed to plot the average result across all samples within a group. In case of two properties, the average is calculated over all samples with the same value for both properties.

**Multi-target bar chart**

The multi-target bar chart window supports the visualization of the relative expression levels between samples for multiple targets (Figure 33). It can be used to compare expression profiles between targets, but is explicitly not designed to compare expression levels between targets. Because of the way relative quantities are being calculated, all relationships between targets are lost. Although alternative mathematical models may suggest results can be compared between targets, it is hard to assure the reliability of these comparisons because of the significant impact that can be caused by subtle differences in PCR reaction kinetics.

![Multi-target bar chart](image)

Figure 33: Multi-target bar chart
In contrast to the single-target bar charts described above, results are only visualized after making some settings. The setup tab of the multi-target bar chart is used to select the targets to be visualized and compared in the chart tab. The setup tab contains two lists, one with available targets on the right and the other with selected targets on the left. Targets can be selected by double clicking them in the available targets list, or by selecting one or multiple targets followed by a click on the leftwards pointing triangle. Targets can be deselected for comparison using an analogous procedure. The upwards and downwards pointing triangles at the left of the selected targets list can be used to change the order of the targets.

In addition to the Y-axis scale and the X-axis ordering options that are in common with the target bar chart described above, the Multi-target bar chart does support zooming in and out on the X-axis.

**Correlation plot**

Open the correlation plot window by clicking on the corresponding entry in the analysis node of the project explorer (Figure 34). Select the targets to be compared at the bottom of the window.

![Figure 34: Correlation plot](image-url)
Correlation coefficients (Pearson or Spearman) are only calculated on the samples that are shown in the correlation plot (see Sample visibility settings). When the plot is shown in logarithmic scale, the correlation coefficients are calculated based on the 10log transformed results. As Spearman rank correlation is based on ranked results, the correlation coefficients do not change in function of the axis scale.

We recommend Spearman rank correlation for a small sample size (< 24 data-points) or for clearly non-normally distributed data. Pearson's correlation analysis requires normally distributed data; as such, it is sensitive to outliers, especially when the sample size is small. It is generally recommended to do statistics on log-transformed gene expression levels.

Result table

Open the result table window by clicking on the corresponding entry in the analysis node of the project explorer. The table shows the calibrated normalized relative quantities (CNRQ values) for every sample-target combination, together with their associated uncertainty expressed as standard errors. Sample-target combinations without any value are not included in the experiment. 'NaN' is shown for combinations that are included in the experiment, but for which no results could be calculated – typically because of missing data.

Statistics

qbase+ has a built-in intuitive statistical wizard to perform commonly used statistical tests on the results that are calculated in a single experiment, i.e. quality controlled and rescaled (inter-run calibrated) normalized relative quantities (CNRQ values). The stat module is specifically tailored towards the typical needs of biologists performing qPCR analysis. This philosophy is reflected in a wizard based approach that minimizes the use of statistical lingo, asks only minimal input from the user, and provides the required information to guide the user to appropriate statistical test and accompanying settings.

Importantly, the stat module is much more than a p-value generator; it also calculates relevant values such as the mean value per sample subgroup or the fold change (ratio) between 2 groups, always with accompanying 95% confidence interval and graphics.

Using the statistical wizard

Prepare your data before launching the stat wizard by properly annotating your experiment (including custom sample properties), performing basic relative quantification analysis according to the appropriate settings and by thoroughly evaluating the different built in quality controls. Start the wizard by clicking Stat wizard (Σ) in the project explorer tree on the left, under Analysis () > Statistics (Σ:). The wizard guides users to results in 4 easy steps.
Specify the goal of your analysis

The first step in the wizard enables specification of the goal of your analysis (Figure 35). The selected goal guides the user to the statistical test that will be applied at the end of the wizard.

![Statistics wizard - step 1](image)

Figure 35: Statistics wizard - step 1

The “Mean comparison” goal leads to:
- Unpaired t-test
- Paired t-test
- Mann-Whitney test
- Wilcoxon signed rank test
- One-way ANOVA

The “Target correlation” goal leads to:
- Pearson correlation
- Spearman correlation

The “Survival analysis” goal leads to:
- Cox proportional hazards
Define sample groups

Here, the user either defines the sample groups that need to be compared, or restricts the analysis to a subset of the samples. Such sample groups are based on samples with identical sample property values (Figure 36). Consult the 'Annotation' chapter for more information on custom sample properties.

Figure 36: Statistics wizard - step 2

Select targets

Statistical analysis can be performed on a single target (gene), on multiple targets, or on all targets measured in the experiment (excluding reference targets). If more than one target is being analyzed, qbase+ automatically suggests performing a multiple testing correction of the calculated p-values.

Make last settings

Technical questions with statistical terminology have been reduced to a minimum while maintaining statistical validity by recognizing the nature of the data that is being analyzed. For example, relative quantities that do not follow a log-normal distribution should be log-transformed to make them suitable for parametric analyses. Without user intervention, qbase+ will automatically logarithm transform the results prior to doing stats.
For a paired analysis (e.g. Paired t-test or Wilcoxon signed rank test), two sample properties are required: a grouping property (similar to unpaired statistical analyses) and a pairing property (unique to a paired analysis). The example below describes the required sample properties for a hypothetical experiment in which gene expression levels were measured in blood from 6 mice before and after treatment with a given compound. The first column contains the sample names, the second column contains the grouping property 'treatment' with values 'before' and 'after', the third column contains the pairing property 'mouse ID' with values denoting the ID of the mice that will be used to identify the samples that belong together, i.e. are paired and come from the same mouse. A pairing value can be a number or a letter combination, and must be identical and unique for the pair.
By progressing through the 4 steps in the wizard, the number of appropriate statistical tests (shown on the right side) will decrease until only one test remains (Figure 37) that will subsequently be used. At each step of the wizard, relevant information and alerts are displayed at the bottom of the wizard.

### Results

Each analysis is saved in the **statistics** (*Σ*) section of the **project explorer** tree with a unique name containing the statistical analysis performed followed by a serial number. These results can be deleted, renamed and exported (CSV, XLS or XLSX file type). Upon opening (double clicking) a stat result (*Σ*) in the **project explorer**, results are recalculated instantaneously. Hence, if data have changed, results will reflect that change (e.g. more samples were added to the experiment, or calculation settings were modified). Dramatic changes (e.g. removal of a target for which results were previously calculated) can result in a conflict, whereby an alert will be shown that results cannot be recalculated. In this situation, the user needs to restart the wizard and complete a new analysis.

A stat result window contains 3 tabs at the bottom: the table contains the calculated p-values and associated values, the chart tab provides a graphical representation of the results, and the settings tab summarizes the input provided and options selected using the stat wizard. Double clicking on a target or target pair in the table tab may bring you to the corresponding graph, depending on the statistical test. Each graph also has a dropdown list, for quick browsing through all results.

---

<table>
<thead>
<tr>
<th>sample name</th>
<th>treatment</th>
<th>mouse ID</th>
</tr>
</thead>
<tbody>
<tr>
<td>sample-01</td>
<td>before</td>
<td>mouse-A</td>
</tr>
<tr>
<td>sample-02</td>
<td>after</td>
<td>mouse-A</td>
</tr>
<tr>
<td>sample-03</td>
<td>before</td>
<td>mouse-B</td>
</tr>
<tr>
<td>sample-04</td>
<td>after</td>
<td>mouse-B</td>
</tr>
<tr>
<td>sample-05</td>
<td>before</td>
<td>mouse-C</td>
</tr>
<tr>
<td>sample-06</td>
<td>after</td>
<td>mouse-C</td>
</tr>
<tr>
<td>sample-07</td>
<td>before</td>
<td>mouse-D</td>
</tr>
<tr>
<td>sample-08</td>
<td>after</td>
<td>mouse-D</td>
</tr>
<tr>
<td>sample-09</td>
<td>before</td>
<td>mouse-E</td>
</tr>
<tr>
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<td>after</td>
<td>mouse-E</td>
</tr>
<tr>
<td>sample-11</td>
<td>before</td>
<td>mouse-F</td>
</tr>
<tr>
<td>sample-12</td>
<td>after</td>
<td>mouse-F</td>
</tr>
</tbody>
</table>

*Table 1: Pairing properties*
## Statistical result table and conditions of use

### Unpaired t-test, Mann-Whitney

<table>
<thead>
<tr>
<th>Column header</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target</td>
<td>Name of the target &lt;br&gt;The small icon in front of the target name can be clicked to collapse / expand the details for that target.</td>
</tr>
<tr>
<td>p</td>
<td>p-value, multiple testing corrected (when selected)</td>
</tr>
<tr>
<td>Property</td>
<td>Sample property used to define the subgroups</td>
</tr>
<tr>
<td>Comparison</td>
<td>2 values from property used to define the 2 subgroups &lt;br&gt;The direction of the comparison (A/B or B/A) can be altered by changing the Target scaling options in the Calculation parameters window. The selected group will be used as denominator.</td>
</tr>
<tr>
<td>Ratio</td>
<td>Fold change between nominator and denominator subgroup</td>
</tr>
<tr>
<td>95% CI low</td>
<td>lower value of the 95% confidence interval of the ratio</td>
</tr>
<tr>
<td>95% CI high</td>
<td>upper value of the 95% confidence interval of the ratio</td>
</tr>
<tr>
<td>Value</td>
<td>sample property value used to define one of the subgroups</td>
</tr>
<tr>
<td>Mean</td>
<td>mean value of the sample subgroup (value depends on 'Target scaling' option)</td>
</tr>
<tr>
<td>95% CI low</td>
<td>lower value of the 95% confidence interval of the mean value</td>
</tr>
<tr>
<td>95% CI high</td>
<td>upper value of the 95% confidence interval of the mean value</td>
</tr>
<tr>
<td>Datapoints</td>
<td>number of datapoints per subgroup</td>
</tr>
</tbody>
</table>

Table 2: Results for unpaired t-test and Mann-whitney test

**Note 1:** no equal variances are assumed in the unpaired t-test as Satterthwaite approximation is used

**Note 2:** non-symmetrical CIs are obtained because statistical analysis are performed on log transformed data
Paired t-test, Wilcoxon signed rank

<table>
<thead>
<tr>
<th>Column header</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target</td>
<td>Name of the target</td>
</tr>
<tr>
<td>p</td>
<td>p-value, multiple testing corrected (when selected)</td>
</tr>
<tr>
<td>Property</td>
<td>Sample property used to define the subgroups</td>
</tr>
<tr>
<td>Comparison</td>
<td>2 values from property used to define the 2 subgroups</td>
</tr>
</tbody>
</table>

The direction of the comparison (A/B or B/A) can be altered by changing the Target scaling options in the Calculation parameters window. The selected group will be used as denominator.

<table>
<thead>
<tr>
<th>Column header</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ratio</td>
<td>Fold change between nominator and denominator subgroup</td>
</tr>
<tr>
<td>95% CI low</td>
<td>lower value of the 95% confidence interval of the ratio</td>
</tr>
<tr>
<td>95% CI high</td>
<td>upper value of the 95% confidence interval of the ratio</td>
</tr>
<tr>
<td>Pairs</td>
<td>number of data pairs</td>
</tr>
</tbody>
</table>

Table 3: Results for paired t-test and Wilcoxon signed rank test

ANOVA

<table>
<thead>
<tr>
<th>Column header</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target</td>
<td>Name of the target</td>
</tr>
<tr>
<td>The small icon in front of the target name can be clicked to collapse / expand the details for that target.</td>
<td></td>
</tr>
<tr>
<td>p</td>
<td>Two-sided p-value, multiple testing corrected (when selected)</td>
</tr>
<tr>
<td>r2</td>
<td>Fraction of the overall variance (of all the data, pooling all the groups) attributable to differences among the subgroup means</td>
</tr>
<tr>
<td>Property</td>
<td>Sample property used to define the subgroups</td>
</tr>
<tr>
<td>Comparison</td>
<td>All combinations of 2 values from the grouping property</td>
</tr>
<tr>
<td>Ratio</td>
<td>Fold change between nominator and denominator subgroup</td>
</tr>
<tr>
<td>95% CI low</td>
<td>lower value of the 95% confidence interval of the ratio</td>
</tr>
<tr>
<td>95% CI high</td>
<td>upper value of the 95% confidence interval of the ratio</td>
</tr>
<tr>
<td>Significant</td>
<td>indication if 2 subgroups from 'comparison' are statistically significantly different (p&lt;0.05)</td>
</tr>
<tr>
<td>Value</td>
<td>sample property value used to define one of the subgroups</td>
</tr>
<tr>
<td>Mean</td>
<td>mean value of the sample subgroup (value depends on 'Target scaling' option)</td>
</tr>
<tr>
<td>95% CI low</td>
<td>lower value of the uncorrected 95% confidence interval of the mean value</td>
</tr>
<tr>
<td>95% CI high</td>
<td>upper value of the uncorrected 95% confidence interval of the mean value</td>
</tr>
<tr>
<td>Datapoints</td>
<td>number of datapoints per subgroup</td>
</tr>
</tbody>
</table>

Table 4: Results from ANOVA test

Note: there is no need for equally sized sample subgroups as the Tukey-Kramer post-test is used
**Spearman correlation, Pearson correlation**

<table>
<thead>
<tr>
<th>Column header</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target X</td>
<td>Name of the target in the X-axis</td>
</tr>
<tr>
<td>Target Y</td>
<td>Name of the target in the Y-axis</td>
</tr>
<tr>
<td>r</td>
<td>correlation coefficient</td>
</tr>
<tr>
<td>p</td>
<td>p-value</td>
</tr>
</tbody>
</table>

Table 5: Results for correlation analysis

Note: switching target between X and Y axis has no effect on p and r value.

**Cox proportionzal hazards**

<table>
<thead>
<tr>
<th>Column header</th>
<th>Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target</td>
<td>Name of the target</td>
</tr>
<tr>
<td>p</td>
<td>p-value</td>
</tr>
<tr>
<td>HR</td>
<td>hazard ratio, increase (or decrease if HR &lt; 1) in risk per log10 unit (equals 10-fold difference) increase in CNRQ value</td>
</tr>
<tr>
<td>95% CI low</td>
<td>lower value of the 95% confidence interval of the HR</td>
</tr>
<tr>
<td>95% CI low</td>
<td>upper value of the 95% confidence interval of the HR</td>
</tr>
<tr>
<td>Datapoints</td>
<td>number of datapoints used in the Cox model</td>
</tr>
</tbody>
</table>

Table 6: Results for Cox proportional hazards analysis
Statistical background

Calculations on logarithm transformed rescaled result table

As gene expression levels are typically log-normally distributed, all statistical calculations are done on the log10 transformed CNRQ values (Hellemans and Vandesompele, 2011). For easy interpretation of the statistical results, values are re-transformed to linear scale by taking the anti-logarithm. The statistical results are based on the rescaled results, hence, the mean values per subgroup (as well as their respective 95% confidence intervals) will change in function of the target scaling option selected in the project explorer tree (Settings > Calculation parameters > Target scaling).

Of note, rescaling has no effect on the p-values and fold changes (ratio).

One-sided and two-sided p-values

If you know the direction of the observed effect (e.g. gene expression levels are decreased after siRNA treatment) prior to generating the data and performing the statistical test, you can use a one-sided p-value. In all other cases, a two-side p-value is recommended. Note that if you selected a one-sided p-value and if the direction is opposite to what you expected, then you need to adjust the p-value by calculating 1-p (e.g. you expect a gene to be down regulated and therefore asked a one-sided p-value; upon analysis, you observe a p-value of 0.075 but the gene appears to be up regulated. Therefore, the true p-value is 0.925 (1-0.075)).

Multiple testing correction

It is highly recommended to correct the calculated p-values when testing multiple hypotheses at the same time (e.g. when performing the same test for multiple targets), in order to control the false positive rate. In qbase+, the false discovery rate (FDR) multiple comparison method is implemented (Benjamini and Hochberg, 1995). This method guarantees that only 5% of all tests that yield a p-value smaller than 0.05 are false positives.

While the FDR method is also applicable for correcting multiple ANOVA p-value results, an additional multiple comparison post-test is implemented in each individual ANOVA analysis when comparing the different subgroups. qbase+ uses the Tukey-Kramer method to correct for the pairwise group comparisons in an ANOVA test. Group comparisons that are statistically different in an ANOVA test (corrected p-value < 0.05) are marked in the stat result table.

Multiple testing correction does not affect the size of the confidence intervals (CI); the confidence intervals are in other words 'uncorrected' (this is also true for the ANOVA CI). Consequently, results that appear significant based on their confidence intervals (e.g. CI on ratio not overlapping 1) may have p-values larger than 0.05.
Approximate and Exact p-values for non-parametric test
qbase+ calculates exact p-values for non-parametric tests when the sample size is small and returns approximate p-values for large sample sizes (Spearman rank correlation: > 6 pairs; Wilcoxon signed rank: > 50 pairs; Mann-Whitney: sum of the 2 groups > 50 datapoints with each group having > 10 datapoints).

Are the data coming from a log-normal distribution?
The sample size (N) and population distribution (log-normal or not) are both important to select the proper statistical test. The following table helps with this selection. In short, when the sample size is sufficiently large (typically two dozen or 24 datapoints), the choice of parametric or non-parametric test is irrelevant. A parametric test can be safely used by relying on the central limit theorem that states that the sample means follow a normal distribution when the sample size is sufficiently large; a non-parametric test is sufficiently powerful when the sample size is large.

For smaller sample sizes (N<24), a parametric test can be safely used if the population distribution is log-normal (i.e. normally distributed when log-transformed).

For sample sizes smaller than 24, but larger than a technical threshold (T value, see further), a non-parametric test is always appropriate. When the underlying population distribution is not known, qbase+ will apply a non-parametric test as such a test is somewhat more conservative, resulting in slightly larger p-values. However, when p<0.05, you can be more sure that there is a real difference.

For sample sizes smaller than the technical threshold T, there is only one valid option, i.e. a parametric test in combination with the assumption of log-normality. Non-parametric tests for sample sizes smaller than T will always result in a p-value > 0.05. For a Wilcoxon signed rank test, there must be at least 6 pairs (T=6). For a Mann-Whitney test, the sum of the sample sizes of the 2 groups must be at least 8 (T1 + T2 = 8).
Are the data coming from a log-normal distribution?

<table>
<thead>
<tr>
<th>Sample size (N)</th>
<th>Yes</th>
<th>No</th>
<th>Don’t know</th>
</tr>
</thead>
<tbody>
<tr>
<td>N ≥ 24</td>
<td>Parametric</td>
<td>Non-parametric</td>
<td>Non-parametric</td>
</tr>
<tr>
<td></td>
<td>appropriate</td>
<td>appropriate</td>
<td>appropriate</td>
</tr>
<tr>
<td>T ≤ N &lt; 24</td>
<td>Parametric</td>
<td>Non-parametric</td>
<td>Non-parametric</td>
</tr>
<tr>
<td></td>
<td>Appropriate if assumption of log-normal data is correct</td>
<td>Appropriate</td>
<td>Best choice when unsure about the log-normality of the data, and with a sample size too small to rely on the central limit theorem but sufficiently large for non-parametric testing</td>
</tr>
<tr>
<td>N &lt; T</td>
<td>Parametric</td>
<td>Non-parametric</td>
<td>Parametric</td>
</tr>
<tr>
<td></td>
<td>Appropriate if assumption of log-normal data is correct</td>
<td>Correct test, but can never yield a significant result</td>
<td>Situation to be avoided. Sample size too small to either rely on the central limit theorem or to yield significant result when using a non-parametric testing</td>
</tr>
</tbody>
</table>

Table 7: Selecting between parametric and non-parametric tests

Technical vs. biological error

The variation within each set of technical replicates is related to the fidelity with which the replicates were created, and is irrelevant to the hypothesis being tested (is there a difference between the groups?). Statistics should only be calculated for independent biological replicates, and never for the PCR replicates (Cumming et al., 2007). As such, the technical error (as shown in the Target bar chart and in the Result table) is not used for the statistical calculations. Technical errors are typically much smaller than the biological variability.

In qbase+ PCR replicates are automatically recognized as wells having identical sample and target names. Biological replicates are defined based on sample properties: all samples that have the same value for a sample property are considered biological replicates.

Confidence interval for non-parametric tests

Confidence intervals on mean value and ratio have to be interpreted with care when doing a non-parametric test. When the sample size is small (< 24 datapoints), the calculated mean or ratio and confidence intervals might be inaccurate or too optimistic, respectively.
Exports

Export Datasets and Tables
qbase+ has an extensive data export functionality. This chapter explains how raw and analyzed data can exported out of qbase+. The softwares also allows easy data-exchange between qbase+ users (.xml format) and is able to generate project files in the RDML (Lefever et al., Nucleic Acids Res, 2009) file format. RDML is a universal, open and XML based data exchange format recommended to be used according to the MIQE guidelines (Bustin et al., Clin Chem, 2009).

The software is only able to export data from opened experiments. The export wizard can be launched by clicking the export button ( ) in the command bar. The first step in this wizard is the selection of the dataset that is to be exported. Alternatively, the context menu of several elements in the project explorer (samples, targets, replicates, reference target stability, amplification efficiencies, normalization factors and results) can be used to export data.

Export Amplification Efficiencies
This export will generate a table (in .csv, .xls or .xlsx format) with for each target a series of efficiency related information:

- computed and user defined E value, with E = efficiency +1 (with an E of 2 being perfect, indicating 100% efficiency)
- computed and user defined standard error on E
- the coefficient of determination (r²) for the regression of the standard curve
- the slope of the standard curve
- the Y-intercept of the standard curve (only relevant for known standard curves with known quantities)

Export Experiment
qbase+ offers the possibility to export an entire experiment (.xml format) for easy data-exchange between qbase+ users. Such an experiment can be easily imported using the import function ( ) and by selecting the option 'Import Experiment'. Exchanging such .xml data files will not result in any loss of data, annotation or settings.

Export Normalization Factors
This export will generate a table (in .csv, .xls or .xlsx format) with for each sample its normalization factor. In case of reference gene normalization, the relative quantities for the selected reference genes will be included in the export as well.
Export Project
qbase+ is able to generate project files in the RDML ([Lefever et al., Nucleic Acids Res, 2009] file format. RDML is a universal, open and XML based data exchange format recommended to be used according to the MIQE guidelines ([Bustin et al., Clin Chem, 2009]). Please note, that some information cannot be saved in this file format (e.g. inter-run calibration).

Export Raw Data Table (RQ)
This export will generate a table (in .csv, .xls or .xlsx format) with for each sample its relative quantity for all targets. Empty cells indicate that the sample-target combination has not been measured, whereas 'NaN' indicates that the RQ value could not be calculated because of missing data (excluded data points or missing efficiency value). This export has the option to include, or leave out, the standard errors on the calculated RQ values.

Export Reference Target Stabilities
This export will generate a table (in .csv, .xls or .xlsx format) with reference target stability measures as shown in the corresponding window in qbase+. 'NaN' indicates that stability measures could not be calculated because of missing data.

Export Replicates (Cq)
This export will generate a table (in .csv, .xls or .xlsx format) with replicate Cq values as shown in the corresponding window in qbase+. The tickboxes used in qbase+ are replaced by a Y/N in the 'included' column.

Export Results Table (CNRQ)
This export will generate a table (in .csv, .xls or .xlsx format) with calibrated normalized relative quantities (CNRQ values). This dataset represents the final results calculated by qbase+ on which data interpretation can be performed. During export users have the options to

- include errors
- log transform the result
- include sample properties

The error bars that can be included represent the standard error on the CNRQ values. Log transformation can be applied during export to prepare the data for statistical analysis using parametric tests (see ‘Statistics’ chapter for more information on the need for log-transformation). Sample and target properties can be included in the export to facilitate downstream processing of the data. For sample properties, both the default and the custom sample properties are included at the end of the data table. For target properties, the target name and its type (reference target or target of interest) are included at the bottom of the data table.
Export Samples
This export will generate a table (in .csv, .xls or .xlsx format) with for each sample information about their type, a description, quantity, custom normalization factor, positive control quantity and a variable number of custom sample properties.

Export Targets
This export will generate a table (in .csv, .xls or .xlsx format) with for each target information about its type: 'REF' for reference targets and 'TOI' for targets of interest.

Export Statistics Results
This export will generate a table (in .csv, .xls or .xlsx format) with statistics results for the selected test in a format similar to what is shown in the corresponding window in qbase+. More information on the interpretation of this table is available in the 'Statistics' chapter.

Export Figures
qbase+ charts can be copied for pasting in other applications, or be exported in any of the following formats: .png, .jpg, .pdf or .svg. Both the copy and the export functions are available as options in the context menu of selected graphs.
Special applications

geNorm

geNorm analysis enables the selection of the optimal set of reference genes from a series of tested candidate reference genes (Vandesompele et al., Genome Biology, 2002). The qbase+ implementation of geNorm provides five great benefits compared to its predecessor in Microsoft Excel:

1. Calculation of relative quantities and geNorm analysis are combined in a single program to speed up analysis. Previously, many manual pre-calculations were needed.

2. Experiments with missing data are automatically processed in a way that has the lowest impact on the overall analysis through intelligent retention of as many data points as possible.

3. Whereas geNorm Excel could not distinguish the two most stably expressed genes, qbase+’s geNorm now allows ranking of candidate reference genes up to the single most stable gene.

4. Interpretation of a geNorm result is not always straightforward and requires a certain level of expertise. To accommodate this, qbase+ now includes an expert interpretation with recommendations on the number and nature of genes to be used for optimal normalization as well as information on the suitability of the selected genes.

- The geNorm implementation in qbase+ is at least 20 times faster than that in Excel.

If your final experiment contains many samples, it is highly recommended to perform a geNorm pilot experiment. Such a pilot experiment involves the analysis of a set of candidate reference genes (preferentially more than 8, coming from different pathways and belonging to different functional classes) on a representative set of samples (typically 10 independent samples). Use the sample maximization approach (Hellemans et al., Genome Biology, 2007) to avoid unwanted technical variation that may interfere with proper geNorm analysis.

It is important that the samples are representative for the final experiment; if one works with treated and untreated samples, or with different biological subgroups, an equal number of samples from these subgroups should be studied.
Import run data in a new empty experiment

In qbase+ qPCR run data are organized in experiments. To start data analysis, a new empty experiment needs to be created. Right click on the project (/bg) in the qbase+ project explorer tree in which you want to start a new experiment and click new experiment. If needed, create a new project first in a similar way (New > Project). An experiment name and description can be provided in the experiment properties window.

qbase+ experiments contain data from one or more runs. Run data can be imported by clicking the downward pointing arrow (bg) in the toolbar followed by the selection of the import Run option.

Selection of normalization strategy and reference genes

qbase+ supports a range of different normalization procedures to suit the specific needs of different types of experiments (premium license only). For a geNorm pilot experiment it is required to apply the multiple reference target normalization approach.

Open the settings node in the project explorer, double click on calculation parameters, and select the option reference target(s) (Figure 38).

Figure 38: Normalization strategy
Mark all candidate reference genes to be evaluated by geNorm as reference targets: select all genes to be tested from the **targets of interest** node and then use the **set target type** option from the context menu to change the target type into **reference target** (Figure 39).

![Figure 39: Set reference genes](image-url)
**Interpret geNorm results**

Open the **analysis node** in the **project explorer** and start an analysis by double clicking **geNorm**.

The **geNorm** window consists of three tabbed views (bottom): **geNorm M**, **geNorm V** and **Interpretation**.

The first view, **geNorm M** (Figure 40), shows a ranking of candidate genes according to their stability (expressed in geNorm M values) from most unstable genes at the left (high M value) to the best reference genes at the right (low M value). Note that the qbase+ implementation of geNorm allows ranking of candidate reference genes up to the single most stable gene, whereas its Excel predecessor could not make a distinction between the 2 most stably expressed candidate reference genes.

![Figure 40: geNorm M](image-url)
The second view, **geNorm V** (Figure 41), shows a bar chart that helps determining the optimal number of reference genes to be used in subsequent analyses. A $V_{n/n+1}$ value is shown for every comparison between two consecutive numbers ($n$ and $n+1$) of candidate reference genes. As a general guideline it is stated that the benefit of using an extra ($n+1$)th reference gene is limited as soon as the $V_{n/n+1}$ value drops below the 0.15 threshold, indicated with a horizontal green line.

**Figure 41: geNorm V**
The third view, interpretation (Figure 42), contains a summary and interpretation of the geNorm results. The first section contains an evaluation of the input data with respect to the numbers of samples and candidate reference genes included, possible missing data and appropriate experimental design. While the old geNorm algorithm only worked if there were no missing data, qbase+ will automatically exclude samples or targets from the analysis until the dataset is complete, using an intelligent approach that minimizes the effect of data exclusion.

The second section will summarize the automatic interpretation of the geNorm results, telling you exactly which reference genes to use in follow up studies without having to fully understand the geNorm M and geNorm V graphs. In addition, the stability of the selected reference genes will be compared against empirically determined reference values for acceptable geNorm M values ([Hellemans et al., Genome Biology, 2007]).
Copy number analysis

Import run data in a new empty experiment

In qbase+ qPCR run data are organized in experiments. To start data analysis, a new empty experiment needs to be created. Right click on the project (**) in the qbase+ project explorer tree in which you want to start a new experiment and click **new experiment**. If needed, create a new project first in a similar way (**New > Project**). An experiment name and description can be provided in the **experiment properties** window.

qbase+ experiments contain data from one or more runs. Run data can be imported by clicking the downward pointing arrow (⌄) in the toolbar followed by the selection of the **import Run** option.

Define calculation parameters

The parameters for the calculations can be defined in the **calculation parameters** window (**Project Explorer > Settings > Calculation parameters**). This is the place to select the appropriate normalization strategy and target scaling option.

The multiple reference target normalization strategy is used by default. This normalization approach is perfectly suited for copy number analysis. Although not as important as for gene expression analysis, normalization with multiple assays is also beneficial for copy number analysis since it improves the accuracy of the results.

The default option for target scaling is **scale to average**. For copy number analysis you should select the **scale to positive control** option (Figure 43).

Figure 43: Scale to positive control
Define control samples

At least two types of control samples should be included in every qPCR-based copy number analysis [D’haene et al., 2010]. As for all PCR based assays, a no template control should be included to detect the presence of contaminating DNA. Specific for qPCR-based copy number analysis, is the inclusion of reference samples (positive controls) with a known copy number. These control samples are used as a reference point (or calibrator) for determination of the true copy numbers. The inclusion of multiple reference samples will result in more accurate results (analogous to using multiple reference targets for more accurate normalization).

Positive and negative control samples should be appropriately labeled in the sample list. Open the sample properties window for the control samples and indicate the sample type (negative control or positive control). The positive control option allows you to indicate a sample specific copy number (e.g. 2 for a normal control or 1 for a deletion control) (Figure 44).

Figure 44: Quantity for positive controls

Please note that reference/calibrator samples with varying copy numbers can be used to provide greater flexibility in reference sample choice and to allow known deletions or duplications to be used both as a reference and as a point for quality control on the ability of the assay to accurately call deletions or duplications. In this example, we included three positive control samples with a known copy number: 2 normal controls (N1 and N2) and a deletion control (Positive).
Select reference genes

qbase+ supports the use of one single or multiple reference genes for normalization when choosing the reference target(s) normalization strategy. You only need to indicate which targets should be used for normalization by selecting them, followed by a right click on any of these targets and set target type to reference target. This procedure can be done for multiple targets simultaneously by holding the CTRL key (Windows) or Command key (Mac) while selecting the targets. As soon as reference targets (ZNF15 and GPR80 in this example) are defined as reference targets, the results for your target of interest will be available.

Quality control

The parameters for quality control can be defined in the quality control setting window (Project Explorer > Settings > Quality control settings ). These parameters do not affect the result of the analysis, but define the required precision and accuracy of the analysis.

More information on the generic quality controls in qbase+ is available in Quality controls. Specific for qPCR-based copy number analysis is the definition of the thresholds for the lower boundary for normal copy and the upper boundary for normal copy in the copy number analysis box (Figure 22). These thresholds are used for conditional bar coloring for easy detection of deletions and amplifications (see below) and are by default set at 1.414 (geometric mean of 1 and 2 copies) and 2.449 (geometric mean of 2 and 3). The default settings are recommended for a diploid organism (like human, mouse and rat).

Copy number analysis results

To visualize the copy numbers on a per sample basis, open the copy number window (Project Explorer > Analysis > Copy number). The identified copy numbers are visualized on a per sample basis and conditional bar coloring is applied for easy detection of deletions and amplifications (Figure 45: Copy number analysis). The Y-axis indicates the copy number. Use the Show sample drop down menu at the bottom to evaluate the results of the different samples.

The target bar charts also show the copy numbers, be it without conditional bar coloring in function of the copy number. These plots may be relevant if a given locus needs to be compared between different samples.
**Support**

**FAQ**

**What are the hard and software requirements for running qbase+?**

qbase+ is suited for high-throughput real-time PCR data-analysis on standard computers. We recommend at least 2 Gb RAM memory and a recent processor.

qbase+ is compatible with Microsoft Windows 7 and higher. qbase+ is also compatible with Linux and MAC OS X 10.8.3 and higher.

**Can I analyze data from my qPCR instrument in qbase+?**

In principle, qbase+ can read data export files (.txt, .csv, .xls or .xlsx) from any real-time PCR instrument software, as long as the user organizes the data into a format currently accepted by qbase+. It does not handle raw fluorescent data or instrument specific binary files. qbase+ directly reads export files containing Cq values from the following instruments:

<table>
<thead>
<tr>
<th>Brand</th>
<th>Instrument</th>
</tr>
</thead>
<tbody>
<tr>
<td>Agilent Technologies</td>
<td>MX3000P, MX3005P</td>
</tr>
<tr>
<td>Applied Biosystems</td>
<td>5700, 7000, 7300, 7500, 7900 SD2.2, 7900 SDS2.3, 7900 SDS2.4, StepOne, StepOnePlus, ViiA7</td>
</tr>
<tr>
<td>Bio-Rad</td>
<td>iCycler, MyIQ, iQ5, Opticon, Opticon 2,MiniOpticon, Chromo4, CFX96, CFX384</td>
</tr>
<tr>
<td>Eppendorf</td>
<td>Mastercycler ep realplex</td>
</tr>
<tr>
<td>Fluidigm</td>
<td>BioMark</td>
</tr>
<tr>
<td>Illumina</td>
<td>Eco System</td>
</tr>
<tr>
<td>Roche</td>
<td>LightCycler 1.5, LightCycler 2.0, LightCycler 480, LightCycler 1536</td>
</tr>
<tr>
<td>Wafergen</td>
<td>SmartCycler</td>
</tr>
</tbody>
</table>

In addition, three general and instrument independent formats are available:

- Simple
- qBase
- RDML
What are the basic calculations in qbase+?

qbase+ uses a generalized model of the delta-delta-Ct approach, thereby supporting the use of gene specific amplification efficiencies and normalization with multiple reference genes. All formulas of this model are detailed in Hellemans et al., Genome Biology, 2007.

1. Conversion of Cq values into relative quantities (RQs).
   1.1. Calculation of the average Cq value for PCR replicates.
   1.2. The average Cq value is transformed into the RQ using the average Cq across all samples for that given gene as a reference point, and a calculated or user defined amplification efficiency.

2. Conversion of RQ values into normalized relative quantities (NRQs).
   2.1. Calculation of the normalization factor (NF) for each sample is typically based on the RQs of the reference genes for that sample. Alternative normalization methods are available.
   2.2. The NRQ is calculated by dividing the RQ by the sample specific NF.

3. Conversion of NRQ values into CNRQs (calibrated normalized relative quantities).
   Only needed when expression levels for a given gene are compared between samples measured in different runs. Inter-run calibration is based on comparison of results for samples that have been measured for the same gene in different plates, i.e. inter-run calibrators (IRCs).
   3.1. Calculation of the calibration factor (CF) for a given gene in a run based on the RQs of the inter-run calibrators (IRCs) in that run.
   3.2. The CNRQ is calculated by dividing the NRQ by the CF.

4. Final CNRQ or NRQ results can then be rescaled according to various user preferences.
   This only changes the scale of the data, but not the fold changes between the samples. By default, expression levels are scaled to the average across all samples. Alternatively, expression levels can be rescaled to the sample with the lowest or highest expression, to a specific sample (e.g. untreated control), to the average of a certain group (e.g. all control samples), etc.

What is the meaning of CNRQ value in the result table?

CNRQ = Calibrated Normalized Relative Quantity

If one does not perform inter-run calibration, then CNRQ equals NRQ (Normalized Relative Quantity). This value represents the relative quantity (gene expression level, copy number, ...) between different samples for a given target (gene). It does not have an absolute scale, and is only meaningful in comparison to values obtained for other samples measured for that gene in the same experiment.
**Where is the calculate button?**

By default, qbase+ will automatically recalculate on every change (selection of reference genes, exclusion of replicates, changed settings, …). This approach offers the best user experience for typical experiments, but may not be the best solution for very large experiments that require too much computing time. In these cases, it is best to disable the “Recalculate automatically” setting in the preferences window. After having disabled this setting, a new icon is added to the command bar. Now, results are only recalculated when clicking that button.

**How do I save the changes to my experiments?**

There is no need to save an experiment because qbase+ will save all the new data, annotations, and other changes automatically in the background.

One exception to this approach is an update of text fields. These need to be confirmed (ENTER or click on save icon) before the changes are applied and saved. Unconfirmed changes can be recognised by the blue font (versus a black font for saved text).

**How is PCR efficiency calculated and used for relative quantification?**

The gold standard method for PCR efficiency estimation is a serial dilution of representative template (e.g. a mixture of RNA or cDNA from your samples). The PCR efficiency can be calculated from the slope of a serial dilution as follows: \( \text{PCR efficiency} = 10^{\frac{-1}{\text{slope}}} - 1 \). The formula to go from Cq values to relative quantities is \((E+1)^\Delta \text{Cq}\) (hence \(2^{\Delta \text{Cq}}\) for an assay with 100% PCR efficiency).

In qbase+, users should provide the base of the exponential function as amplification efficiency value for relative quantification if they want to correct for target specific efficiency. The base number \((E)\) is the efficiency value + 1, e.g. 1.95 for 95% efficiency (E value of 1.95).

Next to the gold standard method for PCR efficiency, there are a few algorithms (from the large number out there) that provide an estimate of the PCR efficiency based on a single amplification curve. Importantly, the calculated results should be precise and accurate (and many algorithms fail in this respect). Hence, various papers (see references below) point at the danger of using sample specific PCR efficiencies based on a single amplification curve (or even replicate measurements). The authors rather propose to average the sample specific efficiencies to obtain a target (gene) specific efficiency. This is also what we recommend to our users. LinRegPCR is a program that calculates Cq & efficiency values for fluorescent amplification curves & exports data in RDML file format that can be directly imported in qbase+.

References: Nordgård et al., Anal Biochem, 2006; Goll et al., BMC Bioinformatics, 2006; Karlen et al., BMC Bioinformatics, 2007.
How to interpret the target bar chart?

The target bar chart represents the Normalized Relative Quantities (NRQ) or the Calibrated Normalized Relative Quantities (CNRQ) values for all samples for a specific target. By default qbase+ expresses the results as fold changes relative to the average expression (NRQ) across all samples for that gene. If you want to make your results relative to a control or control group, you should set the target scaling correspondingly (in the Calculation Parameters window). This will make the relative expression for your control (group) 1, and the results for your other samples are then expressed as the fold change compared to your control (group).

How does qbase+ deal with replicates?

Technical (PCR) replicates

qbase+ automatically deals with technical replicates or repeated measurements. PCR wells with an identical sample and target name are recognized as technical replicates. At the very beginning of the calculation workflow, the software averages the Cq values of all (non-empty) wells of a replicate group. Outliers can be removed before calculations.

Biological replicates

It’s also possible to define biological replicates. Custom sample properties defining biological groups can be added as extra information to the samples (Sample properties window). They enable statistical analysis and grouping of results in bar charts.

Inter-run calibrators (IRCs)

Inter-run calibration is a calculation procedure to detect and remove inter-run variation. The experimenter measures one or more identical samples for the same target in different runs. These identical samples are called inter-run calibrators (IRCs) and they are used to detect and correct inter-run variation.

To avoid interpretation of inter-run calibrators as technical replicates, they should have a different sample name in the different runs (e.g. IRC1_a, IRC1_b, ...). Users should then indicate that a number of sample names actually refer to the same biological sample that is used as an inter-run calibrator (e.g. both IRC2_a and IRC2_b refer to the second inter-run calibrator, sample IRC2). This procedure is known as setting the inter-run calibrators.
What is inter-run calibration and how does it work?
Inter-run calibration is a calculation procedure to detect and remove inter-run variation. The experimenter measures one or more identical samples in different runs. These identical samples are called inter-run calibrators (IRCs) and they are used to detect and correct inter-run variation.

Detailed information is available in Hellemans et al., Genome Biology, 2007.

qbase+ is the only software that
- allows inter-run calibration using more than one IRCs making it more accurate
- performs inter-run calibration after normalization allowing the experimenter to re-synthesize cDNA from the inter-run calibrator RNA samples
- propagates the error introduced during the inter-run calibration procedure

To avoid interpretation of inter-run calibrators as technical replicates, they should have a different sample name in the different runs (e.g. IRC1_a, IRC1_b, ...). Users should then indicate that a number of sample names actually refer to the same biological sample that is used as an inter-run calibrator (e.g. both IRC2_a and IRC2_b refer to the second inter-run calibrator, sample IRC2). This procedure is known as setting the inter-run calibrators.

When do I need IRC?
Two different experimental set-ups can be followed in a qPCR relative quantification experiment.

Sample maximization method:
- As many samples as possible are analyzed in the same run. This means that different targets should be analyzed in different runs if not enough free wells are available to analyze them all in the same run.
- Preferred method because the experimenter is usually interested in comparing the expression level of a particular gene between different samples. The sample maximization method does not suffer from (often underestimated) technical (run-to-run) variation between the samples.

Gene maximization set-up:
- Analyzes multiple targets in the same run, and spreads samples across runs if required.
- Often used in commercial kits or in prospective studies.
Whatever set-up is used, inter-run calibration is required to correct for possible run-to-run variation whenever a target is screened for different samples across multiple runs.

**How to flag bad technical replicates based on a standard deviation threshold?**

qbase+ flags bad replicates based on a user defined maximum allowed difference in Cq values (defined in the Experiment quality control settings).

There is a relation between the standard deviation and difference in Cq (independent of the actual Cq values). In fact, the standard deviation increases 0.1 units per 0.14 cycle difference between duplicated reactions; similarly, the standard deviation increases 0.07 units per 0.1 cycle difference between duplicated reactions. Hence, if you want to flag bad duplicates that differ by more than 0.2 standard deviations, you need to use a 0.28 cycle difference. For triplicates, a standard deviation threshold of 0.1 means that the highest and lowest Cq value can only differ by maximum 0.1 Cq values from the middle point.

**How to exclude PCR replicates that do not meet quality control criteria?**

qbase+ flags bad replicates based on a user defined maximum allowed difference in Cq values (defined in the Experiment quality control settings), but does not automatically exclude outliers. This means that the experimenter needs to decide what to do with the failing replicates (in Replicates window).

In addition to the (default) arithmetic mean on replicated Cq value measurements, qbase+ also supports the calculation of median Cq values (option in the Calculation parameters window). The median Cq value represents a more robust measure than the arithmetic mean when confronted with outliers and having at least 3 PCR replicates. Median results (without outlier removal) are a good alternative to arithmetic averages with outlier removal.

**Why does qbase+ ask to log transform the data when exporting the results table?**

It is good practice to log transform the final gene expression results (i.e. the normalized relative quantities), in order to make the data distribution more symmetrical (as gene expression data is often log normally distributed). Together with the Central Limit Theorem, this allows the use of parametric statistical tests and calculations that rely on a distribution that resembles a normal distribution (e.g. classic t-test, confidence intervals, Analysis of Variance).

Note that log transformations will not have an adverse effect on non-parametric statistical tests.
Does the statistical wizard perform a log transformation?
Yes. As gene expression levels (NRQs) are typically log-normally distributed, all statistical calculations are done on the log transformed CNRQ values.

For easy interpretation of the statistical results, values are re-transformed to linear scale by taking the anti-logarithm. The statistical results are based on the rescaled results, hence, the mean values per subgroup (as well as their respective 95% confidence intervals) will change in function of the target scaling option selected in the Project explorer tree (Settings > Calculation parameters > Target scaling). Of note, rescaling has no effect on the p-values and fold changes (ratio).

Where does qbase+ save my data?
All experimental data stored in qbase+ are collected in a workspace. The location of the workspace on your computer can be verified in qbase+ > File > Switch workspace. The workspace is part of your file system and can be included in your backup procedures like any other file or folder.

To create a new workspace you can copy an existing workspace and give it a new name. You can easily switch between workspaces (qbase+ > File > Switch workspace). We do not recommend sharing workspaces on network drives since concurrent use of a single workspace is not supported, and may result in corrupted data.

When launching qbase+ for the first time after an upgrade from a previous version, the workspace (collection of all experimental data stored in qbase+) will be converted to a new data structure. A zipped backup copy will be generated in your workspace that allows for a rollback to a previous version of qbase+.

What is the difference between Reference target stability and geNorm?
geNorm was developed to support selection of the best set of reference genes from a list of measured candidate reference genes. It is typically applied to pilot studies that are only meant to select reference genes.

The reference target stability quality control can be applied to verify whether the selected reference genes allow for reliable normalization. This quality control can be applied to all experiments in which at least two reference genes are included.

In both cases, the calculation of the M-value is based on the same principle, described in (Vandesompele et al., Genome Biology, 2002). The geNorm algorithm uses the M-value calculation in an iterative process to zoom in onto the most stably expressed candidate reference genes.
What is the difference between "M" and "geNorm M" in qbase+? How are the calculations different?

The M-value calculations are identical and described in Vandesompele et al., Genome Biology, 2002. The M-value in the reference target stability window is determined using all reference targets and hence reflects the relative stability of all appointed reference genes in that particular experiment. This is typically the place where you investigate if the previously validated reference genes in your experiment are (still) good (on average 2-4 reference genes should ideally be used for final normalization in a given experiment).

The geNorm M value (geNorm analysis) is slightly different as it denotes the average M value of all remaining reference genes upon stepwise exclusion of the most unstable reference gene (highest M value). For more details, see Figure 2 in http://www.ncbi.nlm.nih.gov/pmc/articles/PMC126239/. The geNorm module is typically used for assessing the stability of a large set (ideally 8 or more) of candidate reference genes. This is what we call a geNorm pilot experiment.

What to expect when performing a geNorm analysis in qbase+?

geNorm is a popular algorithm to determine the most stable reference (housekeeping) genes from a set of tested candidate reference genes in a sample panel. This is what we call a geNorm pilot experiment. In brief, geNorm calculates the gene expression stability measure M for a reference gene as the average pairwise variation V for that gene with all other tested reference genes. Stepwise exclusion of the gene with the highest M value allows ranking of the tested genes according to their expression stability.

The underlying principles and calculations are described in Vandesompele et al. Genome Biology, 2002. Accurate normalization of real-time quantitative RT-PCR data by geometric averaging of multiple internal reference genes

The results in qbase+:

- A first chart (genorm M) indicates the average expression stability value of remaining reference genes at each step during stepwise exclusion of the least stable reference gene. Starting from the least stable gene at the left, the genes are ranked. (In the reference target stability window you see the values based on the calculation including all ref genes).
- A second chart (genorm V) indicates the pairwise variation V between two sequential normalization factors containing an increasing number of genes. A large variation means that the added gene has a significant effect and should preferably be included for calculation of a reliable normalization factor. We propose 0.15 as a cut-off value; below 0.15 the inclusion of an additional reference gene is not required. For example, if the V3/4 value is 0.22, then the normalization factor should preferably contain at least the 4
best reference genes. Subsequently, if the V4/5 value is 0.14, then there’s no real need to include a 5th gene in the normalization factor. Note: Please bear in mind that the proposed 0.15 value must not be taken as a too strict cut-off. The second graph is only intended to be guidance for determination of the optimal number of reference genes. Sometimes, the observed trend (of changing V values when using additional genes) can be equally informative. Anyway, 'just' using the 3 best reference genes (and ignoring this second graph) is in most cases a valid normalization strategy, and results in much more accurate and reliable normalization compared to the use of only one single reference gene.”

- Expert report with recommendations on the number and identity of genes to be used for optimal normalization as well as information on the relative suitability of the selected genes.