



Real-Time PCR System

FastGene® qFYR Real-Time PCR System

User Manual

Version 1.0



### Copyright & Disclaimer, NIPPON Genetics EUROPE GmbH

The copyright of this manual is owned by NIPPON Genetics EUROPE GmbH. No individual or organization shall, in any form, copy, edit, publish, or translate the contents of this manual (including but not limited to words, trademarks, logos, keystroke icons, lists, images, data, etc.) into other languages without the authorization or written permission of the company. In violation of the above statement, the Company will investigate its relevant legal liability.

NIPPON Genetics EUROPE GmbH strives to ensure that the contents of this manual are correct when printed, but the company still has the right to modify the contents of this manual at any time without prior notice. The contents of the manual shall be subject to the changed contents.

User manual version number:	V1.0
User manual revision time:	September 2022
Type of instrument applicable:	FastGene® qFYR Real-Time PCR System
Software Release Number:	V 1.0
Instrument production date:	See production label

### Introduction

Welcome to the FastGene<sup>®</sup> qFYR Real-Time PCR System. Please read this instruction carefully so that you can use this instrument correctly. After careful reading, please keep it for future inquiry when necessary.

### **Product information**

Description:	Real-Time PCR System
Product model:	FastGene <sup>®</sup> qFYR Realtime PCR System
Software Name:	FastGene®qFYR Analysis Studio
Software Release Number:	V1
Product Size:	350 x 520 x 370 mm (length x wide x height)
Net Weight:	25 Kg



### **Enterprise information**

#### Statement of intended use:

This product is used to run real-time fluorescence PCR experiments (Real-time PCR) and to analyze the experimental data. The instrument is operated in a laboratory with the corresponding reagents, to carry out rapid and accurate quantitative and qualitative detection of target nucleic acids sequences from human and animal samples (such as blood, oral swabs, nasopharyngeal swabs, body fluids, etc.) or other analytes, or to conduct melting curves, genotype analysis, etc.

#### Special statement:

Before installing and using this instrument for the first time, please read the instructions in this manual carefully. Please take into consideration the possible scope of maloperation or non-recommended usage and pay special attention to the possible consequences.

#### Intended Usage:

This product is for Research-Use-Only (RUO).

#### User requirements:

This product must be operated by trained professionals, who have studied this user manual.



### **Table of Contents**

Table of Contents		
Chapt	er 1 Security and considerations	10
1.1	Security Symbols and Markings	10
1.2	General Instrument Safety and Precautions	10
1.3	Personal Safety and Precautions	11
1.4	Electrical Safety and Precautions	11
1.5	Environmental Safety and Precautions	12
1.6	Biosafety and Precautions	13
1.7	Electromagnetic Compatibility (EMC) Standards	13
Chapt	er 2 Product Overview	15
2.1	Scope of application	15
2.1.1	Intended Use	15
2.1.2	Product Intended Use Environment	15
2.2	Application Areas	15
2.3	Basic Principles	16
2.4	Main Functions	16
2.5	Product Composition	16
2.6	Product Structure	17
2.7	Instrument Parameters and Characteristics	19
2.7.1	Physical Specifications	19
2.7.2	Instrument Technical Parameters	20
2.7.3	Software Features	21
2.7.4	Chemistry Reagent Features	21
2.7.5	Consumables Specifications	21
Chapt	er 3 Installation and Transportation of Instruments	22
3.1	Transport and Storage	22
3.2	Unboxing	22
3.3	Packing List	23
3.4	Installation Requirements	23

Instrument Working Environment Requirements	23	
Computer Configuration Requirements		
Equipment Installation		
Equipment Appearance	25	
Release Instrument Transport Lock	27	
First Installation and Run Protocol	28	
Instrument Software Installation	29	
FastGene® qFYR Analysis Studio Software Installation	29	
Pre-preparation for boot	32	
Instrument Self-inspection	32	
System Start-up	32	
Experimental Preparation	33	
r 4 Operation to Run the Experiment	34	
Software Start	34	
Create New Experiment	36	
New Experiment or Loading existing files	36	
Experiment Properties	37	
Name	37	
Plastic Type	37	
Experiment type	38	
Chemistry	39	
Experiment Method	40	
Reaction volume	40	
Thermal lid temperature	40	
Basic setting	40	
Plate Setup	42	
Setup	42	
Quick Setup	44	
Remove Wells from the Reaction	44	
Run an Experiment	45	
Insert a Plate or Reaction tubes	45	
Run the Experimental Program	45	
Save Experiment	47	
	Computer Configuration Requirements Equipment Installation Equipment Appearance Release Instrument Transport Lock First Installation and Run Protocol Instrument Software Installation FastGene® qFYR Analysis Studio Software Installation Pre-preparation for boot Instrument Self-inspection System Start-up Experimental Preparation r 4 Operation to Run the Experiment Software Start Create New Experiment New Experiment or Loading existing files Experiment Properties Name Plastic Type Experiment Method Reaction volume Thermal lid temperature Basic setting Plate Setup Setup Quick Setup Remove Wells from the Reaction Run an Experiment Insert a Plate or Reaction tubes Run the Experimental Program	



Chapte	Chapter 5 Operation of Different Experiment Types 4		
5.1	Absolute Quantification (Standard Curve) Experiment	48	
5.1.1	Setup of a Standard curve	49	
5.2	Experiment of Melting Curve	52	
5.2.1	Setup of Melt Curve Analysis	52	
5.3	Relative Quantification Experiment????Cq?	53	
5.3.1	Setup of comparative Cq (???Cq) Analysis	53	
5.4	Genotyping Experiments	54	
5.4.1	Set Up of a Genotyping analysis	54	
5.4.2	Set Up of a High-Resolution Melt Analysis (HRM)	55	
5.5	Presence / Absence Experiments	56	
5.5.1	Set Up of a Presence / Absence analysis	56	
Chapte	r 6 Result Analysis and Data Export	57	
6.1	Result Tab	57	
6.1.1	Buttons in Toolbar	57	
6.1.2	Other Toolbars	58	
6.2	Result Overview	59	
6.2.1	Amplification plot	59	
6.3.1	Plot Properties	59	
6.2.2	Plot Settings	60	
6.2.3	Threshold Settings	60	
6.2.4	Baseline Settings	61	
6.2.5	Target Settings	61	
6.2.6	Results as Reaction Board	62	
6.2.7	Results as List	63	
6.2.8	Raw Amplification Plot and Raw Melt Peak Plot	64	
6.2.9	Standard Curve	65	
6.2.10	Melt Curve	66	
6.2.11	Gene Expression Map	67	
6.2.12	Analysis Settings	68	

6.2.13	Genotyping Maps	69
6.2.14	Raw Melt Curves of the High-Resolution Melt analysis	70
6.2.15	Aligned Melt Curves of the High-Resolution Melt analysis	71
6.2.16	Difference plot of the High-Resolution Melt analysis	72
6.2.17	Presence/Absence Plot	74
6.2.18	QC Summary	75
6.3	Data Export	77
6.3.1	Toolbars in Export Tab	77
6.3.2	Data Export	77
6.3.3	Printed report	79
Chapte	r 7 Instruments Maintenance	81
7.1	Instrument Cleaning	81
7.1.1	Cleaning the Instrument	82
7.1.2	Clean Sample-tray and Reaction-holes	82
7.2	Instrument maintenance	83
7.2.1	Keep Air-Circulation	83
7.2.2	Keep Stable Electricity Supply	83
7.2.3	Keep Clean	83
7.2.4	Waste Disposal	83
7.2.5	Fuse Replacement	84
7.2.6	Overheating Protection	84
Chapte	r 8 Troubleshooting	85



Symbol	Heading	Description
	Caution	This symbol is used to indicate that non- compliance with instructions or procedures may lead to physical injury or even death or could cause damage to the instrument. Consult the Operator's Manual.
	Biohazard	This symbol is used to indicate that certain precautions must be taken when working with potentially infectious material.
<u></u>	Hot surface	Indicates the presence of a hot surface or other high- temperature hazard and to proceed with appropriate caution.
4	Electricity hazard	Indicates the presence of high voltage at the instrument.
Ð	Appliance class 1	Indicates that insulation of voltage-carrying parts (= basic insulation) and connection of touchable metal parts to the protective conductor (= grounding) protects against excessive touch voltages
CE	CE symbol	This symbol indicates European conformity.
SN	Serial number	Identification of the serial number of the instrument.
M	Date of Manufacturing	The date of manufacturing will appear next to this symbol.
***	Manufacturer	The name and the address of the manufacturer will appear next to this symbol.
X	Disposal	Do not dispose of this product as unsorted municipal waste. Follow local municipal waste ordinances for proper disposal provisions to reduce the environmental impact of waste electrical and electronic equipment (WEEE).

### Safety markings used on instruments

### 1.2 General Instrument Safety and Precautions

The operator shall not disassemble the instrument, replace components or debug the instrument without the authorization of our company. If one of the above is needed, it must previously be approved by our company, and completed by trained professionals, otherwise unexpected consequences will occur.

Please avoid collision, damage the instrument and store carefully.

The user should immediately unplug the instrument from the power outlet and contact the supplier or ask our professional maintenance staff if one of the following is to happen:

- The instrument is subjected to rain or flooding.
- Abnormal sound or smell in the instrument.
- Collision or damage to the enclosure.
- The function of the instrument changed.

#### It is forbidden to carry or move the instrument during the operation of the instrument.

The openings on this instrument are ventilated. In order to avoid excessive temperature, **do not block or cover these vents during the operation of the instrument**, or use dust shields and any other substances to cover the surface of the instrument.

If you need to install or transport this instrument, please contact our company to request professional personnel or professional guidance. Otherwise, we will not bear any responsibility for the damage caused by the instrument.

After the installation of the instrument, the transport lock should be properly retained in accordance with the requirements of the technical personnel.

During the operation of the instrument, do not forcibly open the sample draw, otherwise it will destroy the biological safety of the instrument, its electromagnetic radiation, and other protective measures.

Do not force consumables (8 tubes / test tubes /96 well plates, etc.) not matched with this instrument into the sample draw.

Failure to use this instrument in accordance with the manufacturer's prescribed method may damage the protection provided by the instrument.

### 1.3 Personal Safety and Precautions

The weight of the instrument is heavy. Moving or lifting the instrument requires appropriate tools, methods, or cooperation with others to complete. Please avoid moving the instrument alone without the cooperation of others. Improper lifting of this instrument may cause physical injury, pain and/or instrument damage.

Do not touch the power plug, power cord or switch with wet hands.

#### High surface temperature:

During the operation of the instrument, the heating module of the sample draw may become very hot! During the experiment and after the experiment, do not touch the heating block directly with your hand or any part of the body to avoid scalding. Please open the sample draw after the heating block temperature drops to standby temperature and take out the sample.

### 1.4 Electrical Safety and Precautions

#### Prohibition

This instrument has a voltage harmful to the human body! At any time, before disassembling the instrument shell, please cut off the instrument power supply. It is forbidden to replace components when power lines are connected.



The shell of this instrument should be properly grounded through the power ground wire. Any damage to the internal or external grounding path of the instrument may be dangerous.

**Note:** If leakage is found, please cut off the power immediately and stop using the instrument. **Note:** Please cut off the power supply before moving the instrument.

Use the power cord provided with this instrument. If the original power cord is broken, worn, or disconnected, the same power cord should be replaced as soon as possible.

In order to avoid the danger of electric shock, the power cord of this instrument must be reliably grounded. The power cord of this instrument is a standard plug power cord. Please insert the power plug into the appropriate three-wire grounding socket (the external network power supply connected by the socket must be reliably grounded) to ensure the safe use of the instrument.

The electrical supply in which the instrument is located must be connected with ground wire.

Please carefully check whether the power connection is firm. Do not hard pull the power cord. When the plug is plugged into the power supply, make sure the plug is fully plugged into the socket.

Please keep the power plug and the power cable away from high temperature objects such as heater. When using this instrument, please do not place any items on the power cord, do not put the power cord in the place where people often walk.

The fuse type of this instrument is T10AH250VP, placed in the backup box at the power outlet at the back of the instrument. The use of improper fuses may damage the instrument's line system and cause fire. Before turning on the instrument power switch, check and make sure the fuse is properly installed.

In order to prevent fire hazards, use this instrument only with the indicated fuse. When replacing the fuse, please cut off the power supply, pull out the power cord, pry open the fuse box with a word screwdriver, and replace the old fuse with the same new fuse.

### 1.5 Environmental Safety and Precautions

This instrument is only suitable for indoor use, indoor ventilation should be good, and free of corrosive gas.

EDM may cause explosion hazard and is prohibiting the use of this instrument in the presence or possible presence of flammable and explosive gases.

The working environment temperature of this instrument should be between  $10^{\circ}$ C~ $30^{\circ}$ C and the relative humidity should  $\leq 85\%$ .

The working environment pressure of this instrument should be normal atmospheric pressure (altitude less than 2000 meters).

### 1.6 Biosafety and Precautions

#### **Biological hazard**

The sample object of this instrument is nucleic acid. In practice, please regard it as a biological sample with potential biological hazard. When handling and operating samples, generally applicable safety precautions shall be taken and appropriate protective goggles, clothing and gloves shall be worn.

Users are requested to comply with all applicable local or national regulations to complete the disposal of waste samples and contaminated materials.

This instrument shall be treated as waste with biological pollution hazard. Before reuse, recycling, or disposal, the instrument needs to be purified (including cleaning, disinfection and/or sterilization). This instrument shall be handled in accordance with relevant local or local regulations.

During any operation, if liquid samples overflow, appropriate disinfectants should be quickly used for disinfection, to avoid the spread of pollutants to laboratory personnel or contaminated instruments.

### 1.7 Electromagnetic Compatibility (EMC) Standards

This instrument should be installed and used according to the electromagnetic compatibility information in this chapter.

The electromagnetic compatibility of this instrument is classified into categories A and groups I.

- This instrument meets the emission and immunity requirements specified in the EN 61326-1:2013 and EN 61326-2-6:2013
- This instrument is designed and tested according to A kinds of instruments in the CISPR 11:2016
- Portable and mobile RF communication instruments may affect the use of this instrument. It is recommended to stay away from or close the portable and mobile RF communication instruments when using this instrument normally.
- 4. This instrument must be used with the power cord provided by our company, the length of the power cord is not more than 1.5 m

Except the accessories provided by our company, the use of other manufacturer accessories may lead to an increase in the emission of this instrument or a decrease in immunity.

- 1. Please evaluate the electromagnetic environment before this instrument is used.
- Do not use this instrument next to strong radiation sources (such as unshielded RF sources), otherwise it may interfere with the normal operation of the instrument.



**Note:** It is the responsibility of the user to ensure the EMC environment of the instrument so that the instrument can work properly.

### **Chapter 2 Product Overview**

### 2.1 Scope of application

#### 2.1.1 Intended Use

FastGene<sup>®</sup> qFYR Real-Time PCR Systems are used to run real-time fluorescence PCR experiments (Real-time PCR) and to analyze the experimental data. The instruments are operated in a laboratory, in conjunction with the corresponding reagents, for rapid and accurate quantitative and qualitative detection of target nucleic acids from human samples (e.g., blood, oral swabs, nasopharyngeal swabs, body fluids, etc.) or other analytes in the patient's body, or for melting curves, genotype analysis.

#### 2.1.2 Product Intended Use Environment

Institutions such as medical institutions, laboratories, and companies that meet the requirements of the product operating environment.

1. diagnostic laboratory use

2. General laboratory research use. Its users are professional and technical personnel of the hospital laboratory or general laboratory and need specialized training for PCR laboratory technology and instrument, software operation, and have skilled relevant operation skills.

### 2.2 Application Areas

- Basic scientific research
- Clinical detection of pathogens
- Genetic screening
- Cancer diagnosis
- Diagnosis and treatment of cardiovascular diseases
- The drug genome
- Public Health and Safety Surveillance
- Food Hygiene and Quarantine
- Customs Import and Export Quarantine
- Transgenic and Pathogen Detection in Animals and Plants
- and others

### 2.3 Basic Principles

PCR principle is similar to the natural replication process of DNA. Its specificity depends on oligonucleotide primers complementary to both ends of the target sequence. PCR consists of three basic reaction steps: denaturation-annealing-extension. Repeated cyclic denaturation-annealing-extending processes can amplify the target gene by millions of times in 1~2 hours, for a high detection sensitivity.

The principle of real-time fluorescence PCR is to add a fluorescent reporter (probe or dye) to the PCR reaction system. The target nucleic acid derived from DNA samples is qualitatively or quantitatively detected by fluorescence detection system and real-time monitoring of the fluorescence value. Real-time fluorescence PCR detection has higher specificity and sensitivity than end-point PCR detection.

### 2.4 Main Functions

FastGene® qFYR Real-Time PCR Systems are special instruments for real-time fluorescence detection. It integrates PCR amplification, fluorescence detection and data analysis. It can monitor the increase of fluorescence in each test tube in real time as the amplification increases. After the amplification, the software automatically processes the experimental data, carries on the quantitative / qualitative, melting curve or genotyping analysis to the sample, displays and prints the experimental results.

### 2.5 Product Composition

This product mainly consists of electronic control system, power module, temperature control system, detection system, shell components and software(V1).

### 2.6 Product Structure



### Diagram of the Structure of the Machine

Sample module
 Power indicator

9. Power switch

- Sample draw
  Status indicator
- 10. power cord socket

3. Inlet	4. Transport lock
7. In/Out button	8. Vent
11. USB interface	12. Fuse





Schematic diagram of temperature control module



Schematic diagram of optical path module



Schematic illustration of optical scanning patterns

## 2.7 Instrument Parameters and Characteristics

### 2.7.1 Physical Specifications

Instrument specifications	Size: 350 mm(L) × 520 mm (W) × 370 mm (H). Weight : 25 kg	
Packing specifications	Size: 710 mm(L) × 540 mm (W) × 580 mm (H). Weight : 36 kg	
Power supply specification	power supply voltage : AC 110-230 V Frequency : 50 Hz. Power : 750 w	
Communication specifications	USB ports	
Environmental conditions	Temperature : 10°C ~ 30°C. Humidity : ≤85% RH no condensation. Atmospheric pressure : 85.0 kPa ~ 106.0 kPa. Altitude : < 2000m	
Storage and transportation environment	Temperature : -20°C ~ 55°C, in the transport box. Humidity : ≤85% RH	
Running noise	Maximum noise when running instrument does not exceed 60dB (A)	



### 2.7.2 Instrument Technical Parameters

### Thermal parameters

Heat cycle	Peltier semiconductor	
Temperature control mode	Temperature control module	
Maximum ramp rate	3.6 °C/s	
Temperature accuracy	± 0.2 °C	
Temperature uniformity	± 0.2 °C	

Optical parameters		
Excitation source	Single color LED	
Detection device	High sensitivity MPPC (silicon photomultiplier tube)	
Fluorescent channel number	4 channels (2x FAM)	
	Channel 1	FAM/SYBR Green etc.
	Channel 2	VIC/JOE/HEX/TET etc.
	Channel 3	ROX/Texas Red etc.
Applicable to fluorescent	Channel 4	Empty
	Channel 5	Cy5 etc.
	Channel 6	FAM/SYBR Green etc.

Detection parameters		
Sample size	1-96 wells	
Sample volume	10-50 μl	
Repeatability of fluorescence intensity detection	CV ≤3%	
Detection of fluorescence intensity	CV ≤5%	
Sample repeatability	CV ≤ 3%	
Linear fluorescence intensity detection	A linear regression coefficient   r   ≥0.990 in 5 concentration gradients	
Sample linear correlation		

Sample linear correlation

A linear regression coefficient | ≥0.980 in 5 concentration gradients

#### 2.7.3 Software Features

**Software interface:** Wizard interface, intuitive sample plate layout and program settings, easy to use operation editing.

**Software functions:** absolute quantitation, relative quantitation, genotyping, HRM-analysis, and other analytical functions to meet the requirements of a variety of experiments.

Language Support: English.

**Data transmission:** The instrument can transmit the experimental data to the connected computer in real time during the experiment.

Overlong program settings: Each stage may include up to 100 steps and up to 10 cycles.

#### 2.7.4 Chemistry Reagent Features

PCR reagent: Open reagent platform, all quantitative, qualitative PCR reagents are applicable. Fluorescent dyes: Suitable for fluorescent dyes such as

> FAM/SYBR Green/ FastGene<sup>®</sup> IC Green VIC/HEX/TET/JOE ROX/Texas Red Cy5

#### 2.7.5 Consumables Specifications

0.1 ml clear/white PCR thin-walled single tube, flat-topped transparent optical tube cover.0.1 ml clear/white PCR thin-walled 8-well tubes, flat-top transparent optical tube cover.0.1 ml clear/white PCR thin-walled 96-well plate with no hemline or hemline.

**Note:** FastGene<sup>®</sup> qFYR Realtime PCR System is compatible with low 0.1 mL PCR flat top transparent tube cover reaction tube /8-well joint tube, no skirt or half skirt edge 96-well PCR reaction plate.

Do not use 0.2 mL PCR reaction tube and convex cover.

If the sample tubes are just a few, please use 8-well tubes for support on both sides, or use empty single tubes in the four corners to prevent uneven force on the plate slot.



## **Chapter 3 Installation and Transportation of Instruments**

### 3.1 Transport and Storage

Product transportation is carried out in accordance with the requirements of the product. Original packaging must be used in transportation to avoid damage to the instrument.

The instrument in the packaging and transport state, should be stored in the ambient temperature of -20°C~55°C. relative humidity less than 85%. and the air cannot contain corrosive agents.

The outer packaging of the product specifies the transport instructions allowed for this product, which are:

Packaged products are only allowed to be placed in the direction indicated by the arrow. Please hold the handling process lightly.

Maximum stack number is limited to four layers, do not exceed four layers stack.

#### Do not transport or store this product in rain or wet environment.



### 3.2 Unboxing

The outer packing of the product is cardboard box, filled with shock absorber foam. Please do not rush to open the equipment packaging after receiving the equipment. Carefully check the packaging's appearance for damage. Please check the content you received if parts are missing and damaged after unpacking. If the product is damaged during transportation, please do not use it, and contact NIPPON Genetics EUROPE or the distributor!

### 3.3 Packing List

	NAME	Unit
1	Real-Time PCR System FastGene® qFYR	1
2	Instrument power cable	1
3	Instrument data cable	1
4	Fuse	2
5	hexagon spanner	1
6	Packing List	1
7	USB Stick (including software and electronic user manual)	1

### 3.4 Installation Requirements

### 3.4.1 Instrument Working Environment Requirements

**Location:** The main unit must be placed on a stable, horizontal working table, avoiding direct sunlight, not near heating equipment. The instrument's environment must be at ambient temperature of 10~30°C, with a relative humidity less than 85%. For indoor use only.

**Surrounding environment:** Do not install the main unit next to instruments with strong electromagnetic interference or high induction factors, such as refrigerators, high-speed centrifuges, oscillators, etc.

The mainframe must be placed **at least 15 cm from the surrounding object or wall** for heat dissipation and ventilation of the instrument, and for the convenience of users to turn on and off the instrument power supply.

# Do not place the mainframe in a narrow space where it is difficult to operate and disconnect the power supply!

Power requirements: 110-220 VAC,50/60Hz, voltage fluctuations <10 V.

Power outlet: To prevent electrical shock, the instrument mainframe must be connected to a three-core grounding socket that meets the safety standards. The voltage is ~110-220 V (50-60 Hz) and the zero-ground voltage ≤5 V. If conditions permit, connect the instrument to a dedicated power outlet.

**Warning:** Incorrect grounding may cause electrical shock or instrument damage. Confirm that the input voltage meets the requirements of the instrument and that the fuse has been installed in accordance with the specifications.

### 3.4.2 Computer Configuration Requirements

The computer installed with the FastGene<sup>®</sup> qFYR Analysis Studio must have the following configuration:



Processor:	Intel or AMD dual-core CPU,2.8GHz
Memory:	4 GB
Hard disk:	500 GB
Network card:	10 M/100M adaptive network card (optional)
Display resolution:	1440×900 or higher
Operating system:	Windows 10 or above & Office Word/Excel 2007 or above.
Nate. The control co.	moutor of this instrument is not designed for online use. Connect

**Note:** The control computer of this instrument is not designed for online use. Connecting it to the network will have the risk of acquiring viruses and becoming the target of hacker attacks. NIPPON Genetics EUROPE is not responsible for any damage caused by an internet connection. It is not recommended to install other software in the control computer. There will be a potential risk of conflict with the instrument software module and may affect the reliability of the results. NIPPON Genetics EUROPE does not provide antivirus software. Therefore, if necessary, take preventive measures to ensure that there is no virus on the computer of this instrument.

### 3.5 Equipment Installation

The following contents are convenient for users to quickly complete the installation of the FastGene<sup>®</sup> qFYR Real-Time PCR System. Please be sure to operate according to the following contents. If you have any questions during the installation operation, please contact our support.

### 3.5.1 Equipment Appearance





### Status Lamp

There are power status light "POW" and system status light "STU" on the equipment's top part. The meaning of status lights is shown in the following:

Name	Status	Means			
POW	Blue light	Ready			
STU	Green light slowly flashing	Self-checking			
	Green light quickly flashing	Working			
	Green light on	In idle condition			
	Yellow	Caution, able to continue working			
	Red	Malfunction, stop working			

#### Equipment switch

The device has a power switch in the back. Turn on the instrument by turning the switch to the "I" position and switch off at the "O" position.

#### Accessing the Plate

The draw of the FastGene<sup>®</sup> qFYR Real-Time PCR System is controlled by the top button <u></u>. The plate holder draw will open as soon as you press the top button. The in-and-out operation needs to be carried out in the idle state. It will not work during a PCR and during the self-inspection. The system does not respond to the in-and-out request.



#### Data communication

The interface of the device's back panel is the USB, which is used for communication between the computer and the device.

#### Air inlet

The lower gap of the front panel of the equipment is a set of air intakes. Blocking must be prevented when running. Pay special attention to this gap for air inlet, **it is strictly prohibited to access this area with your hands**, as severe injuries can happen!

### 3.5.2 Release Instrument Transport Lock

In order to avoid the damage of important parts of the instrument during transportation, a transport lock is added. While the instrument is switched off, find the locking hole at the top of the instrument and remove the soft rubber cap. Use the M4 hexagonal wrench to the inner bolt, press down slightly to the bottom and rotate 90 degrees counterclockwise. This will loosen the wrench, and the transport lock will pop up due to the of an internal spring mechanism of the instrument (as shown below).



STU light status will change from slow flicker to green light on, unlocked successfully.



### 3.5.3 First Installation and Run Protocol

Insert the USB data cable and the power cable (as shown in figure A below) and connect data cable to the computer. After turning on the power supply, press the power switch at the back end of the instrument to turn on the instrument, and wait until the power indicator light (POW) is constantly on, and the system begins self-checking process. When the system self-test is completed, the instrument's normal state lamp (STU) is green (as shown in figure B). Press the plate remove button and remove the support plate in the sample platform (as shown in figure C). The installation steps are completed.



### 3.6 Instrument Software Installation

### 3.6.1 FastGene® qFYR Analysis Studio Software Installation

You need to install the FastGene<sup>®</sup> qFYR Analysis Studio software in your computer before using the FastGene<sup>®</sup> qFYR Real-Time PCR System. You can find the installation software in the provided USB-drive.

Please refer to the following steps to complete the installation process:

1. Double-click the Setup icon



- 2. Unzip file interface by clicking on "Extract" extract file
- 3. At the pop-up installation interface, click on next to continue



4. Select the destination folder of the installation





5. Click install to complete the program installation



6. You need to pre-install drivers in the system before using FastGene<sup>®</sup> qFYR Real-Time PCR System and the FastGene<sup>®</sup> qFYR Analysis Studio software. After installation of the FastGene<sup>®</sup> qFYR Analysis Studio software, please click on the install button in the window below:



7. Please click on install:



8. The installation will be confirmed:



9. Please close the window of the driver installer and press Next on the window below:

🚧 qFYR Analyzer Studio Setup		
and a	Device Drivers	
	Select the drive to be installe model	d according to the instrument
	FastGene qFYR series	Instal
AIX		
VA		
	< 8adi	Next > Caricel

10. The installation is complete by clicking Finish





### 3.7 Pre-preparation for boot

#### 3.7.1 Instrument Self-inspection

This instrument has the function of self-inspection. Before use, the user should run the instrument's self-inspection to ensure that the instrument can work normally during the experiment.

- 1. Turn the power switch on at the back of the instrument and turn on the instrument.
- The instrument automatically enters the self-inspection mode: The automatic inspection includes the instrument version, whether the electricity is in normal working condition, whether the power supply is normal or not, and the position of the initializing motor, etc.
- After the self-inspection, the top light will constantly show a green color. The instrument will be on standby until used.

**Note:** Make sure that instrument transport lock has been removed, the control computer and other external instruments have been correctly connected and that instrument's power supply has been connected correctly.

Note: In case of not passing the self-inspection, please turn the machine off and again on for selfinspection. If the instrument still cannot pass self-inspection, please contact the support.

#### Do not shake the instrument.

#### 3.7.2 System Start-up

- a) Start the computer
- b) Turn on the power switch of the backboard of the instrument, and wait until the power status light POW is blue and always on



c) Wait for the system to self-check. During the self-check, the status light STU is green flashing state. After the successful self-check, green light will shine constantly.

### 3.7.3 Experimental Preparation

#### Good laboratory practice for PCR and RT-PCR

When preparing PCR or RT-PCR amplification samples:

- Use only clean personal protection equipment and lab coat
- Whenever you suspect that the gloves are contaminated, it is recommended to change them
- ☑ Set up separate areas and special equipment and supplies for the following purposes:
  - Sample preparation and reaction settings
  - Product amplification and analysis
  - To avoid contamination, do not bring amplified products into the reaction setting area
- ☑ Carefully open and cap all sample tubes to avoid splashing or spraying samples
- Regularly clean the experimental table with proper disinfectants
- ☑ Use calibrated pipettes and anti-aerosol tips
- Prepare the reaction mixture according to the manufacturer's recommendations for the main mixture and the mixture to be tested
- Include excess volume in the calculation to provide excess volume of master mixes for losses during reagent transfer
- Dilute the sample and any standards with TE buffer or water
- ☑ Be careful when diluting samples and standards, errors or inaccuracies in the dilution process directly affect the accuracy of the data
- Place the diluent and assay mixture in the refrigerator away from light until they are ready for use. Overexposure may affect fluorescent probes or dyes
- Before use:
  - Thoroughly mix the master mix by rotating the bottle
  - Resuspend the assay mixture by vortexing and then centrifuge briefly.
  - Thaw any frozen samples on ice. After thawing, resuspend the sample by vortexing, and then centrifuge the tube briefly.

#### Guide to set reaction in plate or tube

- Make sure that the liquid in each tube is at the bottom of the tube without bubbles. Otherwise centrifuge the well plate again
- ☑ Ensure that the plate or tube is properly sealed
- Keep the reaction plate or reaction tube in a dark environment at 4° C until it is ready to be loaded into the instrument



- Keep the bottom of the plate clean. Liquid and other contaminants on the bottom of the plate will contaminate the sample and cause abnormally high background signals
- ☑ If necessary, use permanent markers or pens to mark on the side of the test tube and plate, do not use highlighters



### **Chapter 4 Operation to Run the Experiment**

This section will describe in detail how to set up and run experiments using FastGene<sup>®</sup> qFYR Analysis Studio software.

### 4.1 Software Start

1. Double-click the desktop software icon to launch the software



2. Interface of the software startup



#### 3. User login

After the software starts, the system Login dialog box will automatically pop up (see below). Select the user by changing the "User Name". If no user has been defined, you can proceed with "guest". Click on the "Log In" button to start the software.

Login		×
Enter your us	er name and passwo	rd to log in.
User Name:	guest	*
Password:		
	Log In	Exit Application

#### 4. Adding a user

Log in as Administrator. Select "admin" in the User Name and enter the Administrator password. The default password is <u>admin</u> (Please let the administrator of the instrument change the password as soon as possible).

Select "Users" in the pop-up window.



The next window will let you add, edit ("Update"), or delete users.

🕅 Users		_		×		
Add	Update	Delete				
User Name	ne Password		Ro	e		
guest			Normal			
admin	admin	admin		Administrator		
			_			
				ОК		

5. Startup interface will be shown:





### 4.2 Create New Experiment

The Startup interface is used to create a new experiment, run, or open previously saved templates to create a new experiment.

### 4.2.1 New Experiment or Loading existing files

	Eile	<u>E</u> dit	<u>A</u> nalysis	<u>T</u> ools	<u>H</u> elp		
	New Experiment			Experiment Setup	Ctrl+N		
Experime <u>n</u> t Setup	Open			Ctrl+0	From Template		

- 1. Click the "Experiment Setup" button at the bottom right of the startup interface or select "New Experiment" from the menu bar to start a new experiment setup.
- Click "From template" in the menu bar to create a new experiment by loading a previously saved experiment template.
- Click the "Open" button at the bottom right of the startup interface or select "Open" from the menu bar to open previously saved experiment or template files.



The file types are as follows:

- a) "\*.qpt" file is a template file for experiment operation and contains information about operating conditions and reaction plate settings.
- b) "\*.qps" file is a complete raw data file, including operating conditions, reaction plate settings and experimental results (see Chapter 5).
# 4.3 Experiment Properties

The "Experiment Properties" interface is used to define the relevant properties of the experiment according to the actual experiment. This includes the name and type of the experiment, as well as the selection of the reaction tube and PCR reagent.

of the loss had	tota fam ann				- 8 )
ý mana	CHI2HA, *				
Preparation	Experiment Properties				8 -
	Nam Industries Parkinge Lapshare type Denting Tempto type - ysteed	UR2 11 04,52055 Faclines of V3 hours Welse Notated Care And Annual - The Salary & Salaria - 20168	* * * *		
	Pails nandacture - spillow Inally retries - spillow	diela	×	Lies a standard for interesting sparsing sparsifier .	

#### 4.3.1 Name

The default setting for the experiment name is the current time and date, which can be changed as needed. Special characters are not supported.

#### 4.3.2 Plastic Type

The instrument is compatible with the common fluorescent quantitative reaction tube types, including 0.1 mL PCR single tube, 8-well strip, and 96-well plates. White tube or transparent tube (including frosted tubes) can be selected according to the reaction tube material. Reading of the wells will be adjusted according to the selected reaction tube type. Compared with transparent tubes, white tubes can reduce the loss of optical signal and crosstalk. Therefore, we recommend to use white opaque reaction tubes and plates to obtain better experimental data.





# 4.3.3 Experiment type

Select the experiment type according to the actual purpose of the experiment. The software will associate the subsequent experiment settings according to the selected experiment type. For the experiment type, please refer to the following table:

Type of experiment	Explanation
Standard Curve (Absolute quantification)	Absolute quantification based on a standard curve is a real-time quantitative PCR analysis method suitable for researchers who need to determine the actual concentration or copy number of target genes. For absolute quantification, a template solution with a known concentration needs to be serially diluted several times. The obtained data is used to generate a standard curve. The standard curve is plotted with each concentration and corresponding Cq value. The Cq value of the unknown sample is then compared with this standard curve to determine its quantity.
Relative Standard Curve or Comparative Cq (ΔΔCq) (Relative quantification)	Relative quantification is suitable for most gene expression studies. It can analyze the up-regulation or down-regulation of target gene expression levels in control samples and one or more experimental samples. With this technique, there is no need to accurately determine the copy number, but to focus on the fold change compared to the control. The $\Delta\Delta$ Cq method is a very common method of relative quantitative analysis. It compares the results of experimental samples including controls (such as untreated or wild-type samples) and normalization targets (such as housekeeping gene expression). Using this method, the Cq of the target gene in the same two samples can be compared based on the Cq of the normalization gene in the test sample and the control sample. The resulting $\Delta\Delta$ Cq value can be used to determine the fold difference in expression.
Melt curve or High Resolution Melt	The melt curve experiment is used to determine the melting temperature (Tm) of the PCR product detected by an intercalating dye, such as SYBR <sup>™</sup> Green dye. Gene sequences with different base compositions or lengths have different Tm, so different reaction products can be determined according to the Tm value, including non-specific product. During a high-resolution melt analysis, the change in fluorescence is detected at very small temperature increase steps, allowing the identification of SNPs and mutations. For the detection of unknown or new mutations, saturated DNA combined with dyes is usually used to genotyping by high resolution melting curve.
Genotyping	Real-time fluorescence quantitative PCR can provide different methods for genotyping analysis. For the detection of known mutations, the most commonly used basic method is the end point genotyping using enzyme cleavage hydrolysis probe, and the more advanced method is the fusion curve genotyping using hybrid probe. The method can be easily improved and applied to the detection of various other SNP.
Presence or Absence	Protocol using a probe-based assay to control the presence or the absence of a target.

## 4.3.4 Chemistry

The instrument is compatible with most of the fluorescent quantitative PCR reagent consumables in the market. You can choose the probe method (TaqMan™ reagent) or the dye method (SYBR™ Green reagent) to carry out subsequent experiments according to the specific experimental design. For the comparison of the probe method and the dye method, please refer to the following table:

Methods	Advantages	Disadvantage	Scope of application
Dye-based Method	Widely used Sensitive Convenience Cheap	Difficult-to- design primers Non-specific binding	Suitable for quantitative analysis of various target genes in scientific research, research of gene expression, research of transgenic animals
Probe-based Method	High specificity Good repeatability	Expensive Only suitable for specific targets	Pathogen detection, disease resistance gene research, Drug efficacy assessment, diagnosis of genetic diseases



# 4.4 Experiment Method

The "Method" interface is used to set the specific parameters of a quantitative PCR, including reaction volume, stage, step, temperature, time, and cycle number, etc. The above parameters can be carried out according to specific experimental needs and corresponding reaction reagent requirements.

(a teams)	2022-11-04	13							
	Experimen	4 Method							🖰 tee 🔹
	Vitera	20 pl Gauge 2010							
		and a second	Inge	POR	Steps		Mell Curve Stage		
			1000 1000	21417 16.00 0.00	1	(001) (001)	1	964%	
			/				1	/	
			ante		1 col	7	100	100 Mar	
	٢	t n n n n n n n n n n n n n n n n n n n	/		ener Banger	5.) -	es en es es en es	-	۵
		Sept.	Shipi	Stapt	Yopi	Magel	Step2	Step (Dissolution)	
					- 12				

#### 4.4.1 Reaction volume

The default volume is 20  $\mu$ L. Volumes starting from 1, up to 50  $\mu$ L are supported.



#### 4.4.2 Thermal lid temperature

Thermal lid temperature is fixed to 100°C and cannot be changed.



#### 4.4.3 Basic setting

The values of time, temperature, heating rate (melting curve mode) can be set by entering a value in the value area or clicking the arrows . Temperature can additionally be changed by clicking and dragging \_\_\_\_\_\_ in the temperature curve.

Note: The minimum value of the time setting is 00:01 minutes.



The temperature is selected

- 2. Light up the camera icon to select the data collection point
- 3. The camera is turned on by default to collect data after the extension is completed
- 4. If you need to add a reaction step in a certain stage of PCR, hover the mouse to the bottom left or right step area and click "+" to add a step before or after the current step. The "-" button removes the selected step.



5. If you need to add a reaction stage of the PCR, hover the mouse to the bottom left or right stage area and click "+" to add a stage before or after the current stage and select the type of stage (Hold, Cycling, Melt Curve, Infinite Hold) that should be added. The "-" button removes the selected stage.



6. The number of cycles can be changed by entering a value in the value field or clicking the arrows. The standard number of cycles is 40.



The ramp rate can be set in the drop-down list at the beginning of each step. Ramp rate is given at temperature change within the sample volume and can be set between 0.5 and 4 C/s.



 The temperature interval can be set in the drop-down list on the camera icon during a melt curve analysis in which the fluorescence signal is measured. The smaller the value, the more data points are collected. Standard value is 0.2 °C/detection step.





# 4.5 Plate Setup

The "Plate Setup" interface is used to set the properties of the reaction wells, including assigning samples, test targets, reaction well tasks and other properties to each reaction well. This will make the identification of the sample and the subsequent analysis possible.

the Let And	tos por por		- 10
() Hans	Sabila, *		
	Assign Targets and Samples	Q. Artiss - E	- tee -
	B *** * 11 # 0 # #	Serve Quick Serve	
		- Surgers (B) AM (\$ A	the v
		Lange Spec	
	•		
		- Tarpets (B) AM (Q, A)	- 10
		Anno Annos Annos Anno Annos	-
		D Trapit 1 EVM -	3 H
	Wate Contract I The Contract I Income I Contract I		
			-

Note: The plate setup can be performed before, during and after the experiment.

### 4.5.1 Setup

Define and assign well properties in the "Setup" window of the plate setup:

- 54	ntup	Quick Setup						
-	Sam	ples			🕀 Add	2	Action	w.
			Sample Name		Comments			
0		Sample 1						3
-	Targ	A LOCAL			⊕ A66			*
		Name rget 1		Reporter	Convention	Tank	Grapelity	
			SYBR					

1. Click Hadd, to add samples and target items

- 2. Click on the right  $\boxed{\times}$  to remove samples and targets items
- 3. In (well plate view), select the reaction well according to the corresponding well position of the actual reaction tube

gn Targets and Samples					d Artis	8	Law
19 Ves +		Setup	Quink Setup				
1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	M M	- 54	riphes		(E) AM	d) Alle	
				Saragen Ranker	Committe.		•
		0	Sample 1				. 3
		- Ta	peta		● A44	S. Action	
			Acre	Superior .	Contractor	-	
		•	Target 1	\$108		1	1

- Assign the sample (nucleic acid source) and the target item (gene of interest) to the selected well position and click the text box of each field to name the sample and target gene for actual test.
- 5. When you enter a new sample or target item in the "Setup" window, the software will automatically fill the Reporter with default values (FAM/SYBR) and quenching groups (NFQ-MGB) and assign tasks (unknown). Change the dye or probe used accordingly. After the setting is completed, check the corresponding samples and target to assign the set attributes to the corresponding well positions selected previously.
- 6. Assign tasks in the window of the "Plate Setup" tab:

- Targets	E.		E Ads	2	Action	٣
	Area	Reporter	Commands	Task	Questiy	
🔲 📕 Tarp	e 1	CVS				×
				•		

- 7. Select the reaction well in  $\blacksquare$  (well plate view).
- 8. In the inspection item, select the check box of the inspection item, and then select the task from the drop-down list.

Reaction type	Duty
Unknown (sample to be tested)	U
Negative control	N
Positive control	٩
Standard	S
Homozygous alleles 1/1	1
Homozygous alleles 2/2	<sup>2</sup> 2
Heterozygous alleles 1/2	12
Positive Control - IPC	Ι
Blocked IPC control	X

After the above settings are completed, assign the corresponding samples and targets items to the corresponding selected wells.

### 4.5.2 Quick Setup

"Quick Setup" is a function module for quick and easy setup of samples and targets. After selecting the well position to be set, you can click the sample text box and target text box to manually enter the name. If settings have been made in the "Setup", you can select the options defined in the "Setup" through the drop-down menu to assign attributes to the corresponding well positions.

Setup	Quick Setup		
Sample			
Target			
Well Come	nerts -	Wet Calment	
Setup	Quick Setup		
	Quick Setup	Providencial	0
Setup Sample Target	Quick Setup	Service	

### 4.5.3 Remove Wells from the Reaction

Select the wells to be removed, click the right mouse button, and select "Clear". For multiple wells, drag select these wells, then click the right mouse button to select "Clear".

1	2	3		5	6	7	8	9	10	11	12
		Sarryse 5 (1) Targer 1	Sampin 3. [0] Tanpat 1	Sampie 5 (2) Tanpat 1	Sample 3. [0] Target 1	Tampia 1 (II) Tampia 1	Karrysa 3 (12) Tarpet 3	Klempse 3 (U) Tanpet 1			
		Sample 1	Sampia 1 Ul Target I	Semple 1	Sergin 1 D Target 1	Jaroph 1 El Target 1	Seruria 1 (III) Terpir 1	Sergie 1	Copy		
						1			Paste Paste only sar Fill Column Fill Row	riples	
									Clear		
									Zoom In Zoom Out Fit Plate		
									Full Screen		
									Save As		
								_	Print Preview. Print		
lle: O tak	14	Negative (	Controlt 0								82 Em

# 4.6 Run an Experiment

### 4.6.1 Insert a Plate or Reaction tubes

- 1. Press the access button on the top of the instrument.
- 2. The instrument's heating block will be ejected.
- Load the reaction tube or plate onto the 96-well block. Note: When the drawer is closed, the heating lid will apply the appropriate pressure to securely place the tube band in the module.
- 4. Press the access button again to close the heating block.
- After the operation is completed, press the access button to eject the heating block, remove the reaction tube or plate from the heating block, and then close the door by pressing the access button.

Watch out! Risk of personal injury. During the operation of the instrument, the heating block can reach a temperature of  $100 \circ C$ . Allow it to cool down to room temperature before processing.

### 4.6.2 Run the Experimental Program

1. In the "Run" tab of the desktop software, click "Start Run". After clicking "Start Run", a save dialog box appears, select the data file storage location and name.

Love .	8 202		Course of the														
	Ran	Cont	nol														Latings 🗄 See
	F	3	Rate														
	15	View										::: 0	*				
								See							х	Multicomponent Plot	
							1	Sale in	E esperim					<u>()</u>	10		
				and a second	and the		10 12		A 200 H A 200 H A 200 H	22,4000004	fast 1014						
								-									
					in an	-		-									
	1			and the second	and the second				Pennet		15-25-0-0	100			500		
					-	-		-	File of Spe	1 het D	loaner 5	ingle New 71	1.0	۲	Canol		
					-	-	1		and do no	-							
				-	and in		-		-	-	and day		#bainbeg			Carle (Dely frame)	

After saving, the instrument starts to run the program, and the real-time heating lid temperature and reaction plate temperature will be displayed in the upper left corner of the running interface.



		an C		100												
						- 24		1967	24.57		1823	-18.83*C				
				Dam	ing .	States		1000	100	100.10		10000				
		-	View	.*										a s	1	
	h								COM 1			12000			un, Multicomponent Plot	
												-			12	
		11 T			-		-	-			1		-		18 18 18	
	4			4			1	1		10.44	and the owned	uting turn Unleg the s	-	Int.	*	
					-			E tant		ann an Rìomra			-	-	T-	
					1			-		an in	-				6 ia 14 18	
	ł			123	-		and the second	-	-	and a	10-34	-			10 10 10	
	•				and the second					1000		11				
	H							and the second		anise is an	-				tain (bestives)	
	Ľ			-						-						
-																
							Run	Cont	lor							

3. When the temperature of the heating lid reaches 100 ° C, the real-time heating lid's and reaction plate's temperature, and the cycle number of the reaction will be displayed in the upper left corner of the running interface.

Run Control		STOPRUN	Save 🗸
FastGene qFYR 1/40	Cover Temperature:100.04°C Block Temperature:90.36°C Block temp setting 60°C	Estimated Time Rema	ining: 124 min 1 sec
	Multicomponent Plot		
11.000			
10.000			
8.000			
8,000			
7,000			
§ 5.000			
§ 5.000			
4,000			
5.000			
2,000			
1.000			
	* Cycle (Data Points)		
SYDA			

**Note:** In the initial reaction cycles, the original data map is most of the background or baseline. When the amplification reaction enters the exponential growth phase, the curve will rise.

# 4.7 Save Experiment

Click the "File" option in the menu bar, choose "Save" or "Save as" to save the original experimental data. At each step of the experiment process, you can choose to save the experiment in the upper right corner of the interface. The extension of the experiment file saved before running is .qpt (experiment running template). The extension of the experiment file saved after the operation is completed, is .qps (original data file), which contains all the original operation data and analysis results.





# **Chapter 5 Operation of Different Experiment Types**

FastGene® qFYR Real-Time PCR System experiments are divided into seven steps:

- 1. Create an experiment
- 2. Set Experimental Properties
- 3. Set the operating conditions
- 4. Set the samples and targets
- 5. Start the experiment
- 6. View and analyze the results
- 7. Export the experimental results

Please see Chapter 4 for how to create, run, analyze, and review an experiment. The next chapter explains how to operate the different types of experiments within the software.

# 5.1 Absolute Quantification (Standard Curve) Experiment

Absolute quantification is a real-time fluorescence quantitative PCR analysis method suitable for researchers who need to determine the actual copy number of targets. To carry out absolute quantification, the target template solution with known concentration should be diluted several times continuously, amplified by real-time fluorescence quantitative PCR, and a standard curve is generated by using the obtained data. A standard curve is drawn with the concentration of each target and the corresponding Cq value. Cq values of unknown samples are then compared with this standard curve to determine their concentration or copy number.

The standard curve method is used for absolute quantification, and the template standard with known concentration or quantity is required, which means that the template should be accurately quantified before the curve is generated. Quantitative accuracy is directly related to the quality of the standard curve. The template used to generate the standard curve and the method used to quantify the template are the basis of the experiment. The accuracy of continuous dilution is very important, and the RT and PCR efficiency of target template and continuous dilution of actual samples are very important.

Although you may need uniform pure templates for initial copy number determination and standard curve generation, it is best to use target templates as similar as possible to experimental samples. Since steps such as nucleic acid extraction and reverse transcription can affect the dynamic range of the reaction, the same processing steps as the experimental samples should be adopted.

In the standard curve experiment:

- Software generates standard curves using data from standard dilution series.
- Software uses standard curves to calculate the number of unknown samples.

Type of response:

- Many samples can be analyzed in standard curve experiments, but each sample needs its own standard curve.
- The following table lists the types of sample settings involved in the standard curve experiment:

Type Setting	Example Description
Standard	contains a known number of target samples
Unknown	sample to be tested
No template control (NTC)	water or buffer replacement template, no target amplification should occur in NTC hole

The accuracy of quantitative experiments increases with the increase of repeated reactions (usually containing at least three repeats). For accurate detection, at least five dilution points in a wide range of standard quantities should be prepared. The range of the standard quantity can be adjusted according to the expected level of presence of the gene.

	The following is optional	
Template type - optional	OTHER	۲
Plastic manufacturer - optional	Enter plastic manufacturer here	
Sealing method - optional	OTHER	۲

#### 5.1.1 Setup of a Standard curve

1. Select experiment type "Standard Curve".

Experiment type	Standard Curve	~
Chemistry	Standard Curve	
	Relative Standard Curve	
	Comparative Cq (ΔΔCq)	
	Melt Curve	
Template type - optional	High Resolution Melt	
Plastic manufacturer - optional	Genotyping	
riasue manuracturer · optional	Presence/Absence	

- 2. Choose the used Plastic type and Chemistry.
- 3. Add name and optional information.
- After the Method setting is completed, enter the Plate section, or click Next in the lower right corner of the page to enter the well plate setting interface.
   Note: The plate setup can be set before, during, or after the reaction operation.



5. Right-click in any area of the plate view and select "Define and Set Up standards".

	A A Copy Sale Factor of purposed Relations	i		-	*	 4	4	B) Determined Technik Same Lange Technik Support: B' theoretic large for the conduct room. Tage [1] B Technik Standard point 1 or y strates (2) Researched 1 or y strates (2) Researched 1 or y strates (2) Researched 1 or y strates (2) Bearrandel 1 or y strates (2) Bearra
*	Ra Ren Chen Zaseri In Zaseri In Ka Rase Rad Reser Red Rasers, Red, Defen and Set Ur Steel	wa						I. Main 3.3 Stability: - Stability: - Stability:     Main 2.4 Stability: - Sta
•								
Wester Comment	ine Contract B		-				and long	Ann Des Over

6. Select the target, which should be used for generation of the standard curve.

Target 1 🔽			
Target 1	* Model:	Singleplex	v
Target 2	]	Singleplex	
Target 3		Multiplex	)
	Target 1 Target 2	Target 2 Target 3 Define the	Target 1 * Model: Singleplex Target 2 Singleplex Define the

**Note:** Singleplex means that there is only one gene to be tested in each reaction well, and Multiplex means that there are two or more genes to be tested in each well.

Set the number of points and replicates, dilution factor and starting quantity according to your experiment.

Define the standard curve	Standard Curve Preview
*# of Points: 5 (5 Recommended)	+ <
* # of Replicates: 3 (3 Recommended)	1E0 2E-1
* Starting Quantity. 10 (Enter the highest or lowest standard quantity for the standard curve.)	4E-2 8E-3 1.6E-3
* Serial Factor: 1:5 v (Select a value from 1:10 to 10×.)	
5 Points X 3 Replicates = Required Wells	]

The parameters of the standard curve can be set according to the specific experimental requirements and conditions:

- Standard concentration dilution gradient: recommended at least five different dilutions.
- Three replicates per dilution are recommended.
- Starting Quantity: Enter the maximum or minimum quantity value of the standard, excluding units (units containing copies, copies/µL or ng/µL) and the dilution factor.

#### 8. Select and arrange standard wells.

	inge wells for the s										
vrange stan	dards in: 🗌 Colum	ns 🔳 Rows									
Jse Wells:	Automatically Sele	ct Wells for Me	e 🔲 Let M	Me Select W	ells						
1	2 3	4	5	6	7	8	9	10	11	12	15 Required Wells / 15 Selected Wel
·											A1,A2,A3,B1,B2,B3,C1,C2,C3,D1,D2
											,D3,E1,E2,E3
		i —									

- The software can automatically select the reaction well by columns or rows or the operator can select their own reaction well according to the need and arrange the well position of the standard product.
- After the standard well position distribution is set, the software will arrange the standard well position according to the above setting, and display the corresponding well position arrangement and the number of wells.
- The software will automatically apply the above settings to the plate setup, and automatically define the above well position as the standard property (as shown below), so that the plate setup of the standard is completed.





# 5.2 Experiment of Melting Curve

### 5.2.1 Setup of Melt Curve Analysis

1. Select experiment type "Melt Curve".

Experiment type	Standard Curve						
Chemistry	Standard Curve						
	Relative Standard Curve						
	Comparative Cq (ΔΔCq)						
	Melt Curve						
Template type - optional	High Resolution Melt						
Plastic manufacturer - optional	Genotyping						
	Presence/Absence						

- 2. Choose the used Plastic type and Chemistry.
- 3. Add name and optional information.
- After the Method setting is completed, enter the Plate section, or click Next in the lower right corner of the page to enter the well plate setting interface.
   Note: The plate setup can be set before, during, or after the reaction operation.
- 5. Melting Mode:
  - a. For analysis of nucleic acids, select "Nucleic Acid" here
  - b. For analysis of protein thermostability, select "Thermal stability assay" here

Melt mode	Nucleic Acid	*
Chemistry	Nucleic Acid	
	Thermal stability assay	

# 5.3 Relative Quantification Experiment $(\triangle \triangle Cq)$

Relative quantification is suitable for most gene expression studies and can analyze the up- or down-regulation of target gene expression levels in calibrated (normal) samples and one or more experimental samples. Using this technology does not require accurate determination of copy number but focuses on multiple changes compared with calibration samples.  $\Delta\Delta$ Cq method is a very common method of relative quantitative analysis, which can be used to compare the results of experimental samples, including calibrators (such as untreated or wild-type samples) and standard products (such as housekeeping gene expression), which can be used. Using this method, the target gene in the same two samples can be compared according to the Cq of the standard (normal) gene in the test sample and the calibration sample Cq, and the  $\Delta\Delta$ Cq value obtained can be used to determine the multiple difference of expression.

Relative quantitative experiments are often used to:

\*Compare gene expression levels in different tissues.

\*Compare the expression level samples of genes in treated and untreated samples.

\*Compare the expression levels of genes of interest in different genetic backgrounds.

\*Analysis of the conditions of gene expression over time under specific treatment.

#### 5.3.1 Setup of comparative Cq (△△Cq) Analysis

1. Select experiment type "Comparative Cq  $(\Delta\Delta Cq)$ ".

Experiment type	Melt Curve	×.
Melt mode	Standard Curve	
	Relative Standard Curve	
Chemistry	Comparative Cq (ΔΔCq)	
	Melt Curve	
	High Resolution Melt	
Template type - optional	Genotyping	
remplace type optional	Presence/Absence	

- 2. Choose the used Plastic type and Chemistry.
- 3. Add name and optional information.
- After the Method setting is completed, enter the Plate section, or click Next in the lower right corner of the page to enter the well plate setting interface.
   Note: The plate setup can be set before, during, or after the reaction operation.



# 5.4 Genotyping Experiments

In the study of gene variation, in order to obtain a large number of characteristic markers, a large number of individuals must be genotyped. The reasons for the differences in individual genomes at different levels include single nucleotide polymorphisms (SNP) and DNA covalent modification. more than 90% of the gene sequence variants that cause individual differences belong to single nucleotide polymorphisms (SNP). SNPs mainly refer to DNA sequence polymorphisms caused by mutations at the nucleotide level of the genome frequency of at least one allele in the population is not less than 1%, including conversion of single base, transversion and insertion and deletion of single base: in addition, changes in DNA covalent bond modifications, such as DNA methylation.

Real-time fluorescence quantitative PCR can provide different methods for genotyping analysis. For the detection of known mutations, the most commonly used basic method is the end point genotyping using enzyme cleavage hydrolysis probe, and the more advanced method is the fusion curve genotyping using hybrid probe. For the detection of unknown or new mutations, saturated DNA combined with dyes is usually used to genotyping by high resolution melting curve. In this chapter we will introduce the method of allele analysis of known mutation sites using the TaqMan<sup>™</sup> probe endpoint method. The method can be easily improved and applied to the detection of various other SNP.

Common genotyping applications:

- \*Genetic identification of genetic diseases.
- \* Identification of tumor susceptibility genes and hotspot mutations.
- \*Identification of pathogen gene mutations and drug resistance.
- \*Animal and plant forensics, breeding research.

#### 5.4.1 Set Up of a Genotyping analysis

1. Select experiment type "Genotyping".

Experiment type	Genotyping	×
Chemistry	Standard Curve	
( ) ( ) ( ) ( ) ( ) ( ) ( ) ( ) ( ) ( )	Relative Standard Curve	
	Comparative Cq (ΔΔCq)	
	Melt Curve	
Template type - optional	High Resolution Melt	
Plastic manufacturer - optional	Genotyping	
	Presence/Absence	

2. Choose the used Plastic type and Chemistry.

Note: Genotyping experiments can only be performed using probe-based chemistry.

3. Add name and optional information.

After the Method setting is completed, enter the Plate section, or click Next in the lower right corner of the page to enter the well plate setting interface.
 Note: The plate setup can be set before, during, or after the reaction operation.

### 5.4.2 Set Up of a High-Resolution Melt Analysis (HRM)

1. Select experiment type "High-Resolution Melt".

Experiment type	Genotyping	
Chemistry	Standard Curve	
	Relative Standard Curve	
	Comparative Cq (ΔΔCq)	
	Melt Curve	
Template type - optional	High Resolution Melt	
Plastic manufacturer - optional	Genotyping	
	Presence/Absence	

- Choose the used Plastic type and Chemistry.
   Note: Genotyping experiments can only be performed using dye-based chemistry. Please use an HRM-specific saturating dye.
- 3. Add name and optional information.
- After the Method setting is completed, enter the Plate section, or click Next in the lower right corner of the page to enter the well plate setting interface.
   Note: The plate setup can be set before, during, or after the reaction operation.
- 5. For HRM experiments controls can be defined during the plate setup (optional).

Click Add to add controls and select wells to assign controls similar to assigning samples and targets

		H Add	3 Action	*
	Control Name		-1	
Control 1				×
	Control 1			Contra Name

# 5.5 Presence / Absence Experiments

Presence / Absence experiments are endpoint experiments that can be performed to determine whether a target is present or absent in a sample. This type of experiment can be used for the detection of pathogens.

Presence / Absence experiments can only be performed using a probe-based assay and data is collected at the end. The amount of accumulated target is quantified for each sample. For quantification, the normalized intensity of the reporter dye (Rn) is used and displayed.

### 5.5.1 Set Up of a Presence / Absence analysis

1. Select experiment type "Presence/Absence".

Experiment type	Genotyping 🗸
Chemistry	Standard Curve
í í	Relative Standard Curve
	Comparative Cq (ΔΔCq)
	Melt Curve
Template type - optional	High Resolution Melt
Plastic manufacturer - optional	Genotyping
	Presence/Absence

2. Choose the used Plastic type and Chemistry.

Note: Presence/Absence experiments can only be performed using probe-based chemistry.

- 3. Add name and optional information.
- After the Method setting is completed, enter the Plate section, or click Next in the lower right corner of the page to enter the well plate setting interface.
   Note: The plate setup can be set before, during, or after the reaction operation.
- 5. For Presence / Absence experiments four types of tasks can be applied to the plate setup: Unknown-IPC (well contains unknown sample and IPC (internal positive control)), control (well contains a positive control and IPC), NC (negative control)-IPC (well contains no-template control and IPC) and NC-blocked IPC (well contains no-template control and IPC) and IPC blocking agent).

# **Chapter 6 Result Analysis and Data Export**

This chapter explains the data overview, analysis, and export.

# 6.1 Result Tab

# 6.1.1 Buttons in Toolbar

Button	Name	Function
<b>K</b> 3	zoom in	Zoom in for plate overview and curves
3 K	zoom out	Zoom out for plate overview and curves
3	Return	return to last step
$\blacksquare$	Plate view	Plate overview for results
	List	Experiment results by list
Ð	Send to printer	Print current window image file
	Сору	Copy current window as image file
	Save as file	Save current window as file (.jpg, .svg, .png, .emf, .pdf)
ഷ്ട്	Plot Properties	Plot properties set up
	Hide plot result	Hide plot result
	Display plot result	Display plot result
	Show plot settings	Show plot settings



# 6.1.2 Other Toolbars

Toolbar	Name	Function
Analyze	Analyze	Click to analysis result after changing plate or plot settings
کی Action 🖌	Action	Click to select sample or target
Save v	Save	Save current results as .qps file
View V	View	View current results by plate or list format
Group by V	Group by	Click to show results by different groups
< >	Expand/Shrink	Expand/Shrink plate or image view

# 6.2 Result Overview

FastGene® qFYR Analysis Studio will analyze experiment results by default settings after an experiment is finished, and show amplification plot in the Result tab.

Attention: After omitting wells or changing any settings, please click Analyze to obtain new results.

### 6.2.1 Amplification plot

1. Enter the Amplification plot tab to view the amplification curve.

Raw Melt Plot	Melt Peak Plot	QC Summary	Gene Expression
Amplificati	on Plot	Raw Ampli	ification Plot

 Overview amplification curve and evaluate curve shape. A typical amplification curve includes four different parts:





### 6.3.1 Plot Properties

Select <sup>off</sup>, open the Plot Properties settings. In the "General"," X Axis"," Y Axis" tab, you can adjust the title, name, font, color, range and much more, as shown below. Select "Save" if any adjustment is made. The adjusted curve will be shown after saving.





### 6.2.2 Plot Settings

1. Select 🗟 to enter the Plotting Settings and to adjust plot type, graph type, threshold etc.

Plot Type	ARn vs Cycle	Graph Type	Linear	*	Plot Color	Target	۲
	ARn vs Cycle						
Target:	Rn vs Cycle	april .					
Thresh	Cq vs Well	9971	EA.	(o-1)	indire.		
080	w: Threshold						
I Hid	e unselected curv	es					
Ena Ena	ble baseline supp	ression					

 Use the drop-down menu to choose the Plot Type. The default setting is "Rn vs cycle", but "△Rn vs cycle", "Cq vs well" are available as well.

Plot Type	Description	Purpose
Rn vs cycle	Rn value is the normalized fluorescence	Recognize and checking unnormal
	value.	amplification well.
Cq vs well	Cq value is the cycle number which	Recognize the unnormal amplification
Cq vs well	fluorescence signal achieved threshold.	well.

3. Graph Type can be changed from "Linear" to "Log" by using the drop-down list. The default setting is "Linear" graph.

Plot Type ARn vs Cycle 💌 Graph T	ype Linear	Plot Color Target	. 4
	Linear		
Target: Lock AMP V	Log		
Threshold: Auto 1.339971	Aoto	Bandere	
5how: Threshold -			
Hide unselected curves			
Enable baseline suppression			

Note: Plot settings can also be adjusted using the tool bar above the graph.

Target: AMP ▼ Plot Type ΔRn vs Cycle ▼ Graph Type Linear ▼ Plot Color Target ▼

#### 6.2.3 Threshold Settings

The software will calculate the threshold value by an automatic model. However, it can still be adjusted by unchecking "Auto" and changing the value. Additionally, the threshold line in the graph can be dragged. The new Cq value will be automatically calculated and displayed after the threshold value is adjusted. The threshold line will not be displayed if "Show Threshold" is not selected.



### 6.2.4 Baseline Settings

The software automatically sets the baseline in the first few cycles of the reaction, usually withing 3 to 15 cycles. If you need to set the baseline manually, remove the automatic threshold check, you can drag and drop the baseline (green triangle) directly on the curve and change it within the 1<sup>st</sup> to 15<sup>th</sup> cycle. In case of an early Cq value, it is suggested that the baseline should be set in the straight area of the curve, and when the baseline setting range is too wide, fluorescence can be detected in the 10<sup>th</sup> cycle, which may cause a curve drop and Cq delay.

Target: DiLock AMP +	Target: D Lock AMP
Threshold: Auto 0.33947 E Auto Quedine	Threshold: Auto 0.33947 Auto Baseline
Show: Threshold -	Show: Threshold -
Hide unselected curves	Hide unselected curves
Enable baseline suppression	Erable buseline suppression
Amplification Flor	Amplification For 10 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

**Note:** Please do not forget to click Analyze after a baseline setting is changed.

### 6.2.5 Target Settings

You can select different targets, or all targets to be shown:

Plot Type ΔRn vs Cycle 💌 Graph Type Linear 💌 Plot Color Target	×	
Target Diock Al	Target: All V Plot Type ΔRn vs Cycle V Graph Type Linear V Plot Color Target	۷
Threshold:	All Amplification Plot	
Acto	2.6 Actg	
B2m	2.4 B2m	



### 6.2.6 Results as Reaction Board

To view the results in plate format, click  $\blacksquare$ .



In the plate view, the information of each well (incl. target, task, dye, sample) will be showed while the mouse is moved to and paused on each well. E.g., information of well B4 as in figure above.

. 4	2	3.	14	.8	4				44	86.	田田;	5 # + 7 # E 4 G 5
												Target: AMP ¥ Plet Type ДКл vs Cycle ¥ Graph Type Linear ¥ Plut Color Target ¥
												Amplification Plot
			and I	144	And Carolan	1444	144	1040				116
			10.11.00		Circle H	19.9.8		19-11-00				67 10
				_	-	1						
			and a	and a	14	313	213	Land Carl				
			4	140	14		14	140				10 10
			1.14		Carried.	043531	SUDN.	Self.R.				
			_	-	-	_	-					
			-14	14	18. An 1935	10.00	114	1988				\$ 12 10
												*
			140	104	1			140				
			10.040	14.000	11 Con 1940	- 834 IIII 204		14-11-0				
			<u> </u>	-	_	-	-	-				
			1000	100	- 5.00 Ca-5180	04E	L'ini					
			1	250	11.00	04.010	anata)	De Print.				10 er 82094"
												14
												**
												A AND D AND A CHARACTER ON AND A STALLED AND A STALLED AND

If a different well is selected, the amplification plot will be shown accordingly. E.g., the amplification curve of well E6 is shown after selecting well E6 in the plate format, as in the figure above.

Click buttons of Zoom in 🛐, Zoom out 🧚, Return to original size 🔍, to change the plate layout.

## 6.2.7 Results as List

To display the results as list, click

All the experiment information is listed, including well, sample, target, task, Cq, Cq mean, Cq SD etc.

E	View	¥	Group by	¥						Œ	
	Well	Omit	Flag	Sample Na.	Target Na	Task	Oyes	Cq	Cq Mean	Cq SD	Quantity Qua
18	86		0	0.01	AMP	STAND	SYBR	17.457	17.462	0.023	0.010
19	87	0	0	1:1.2	AMP	UNKNO	SYBR	17.449	17.433	0.041	0.000
20	88		0	1:1.2	AMP	UNKNO	SYBR	17.463	17.433	0.041	0.000
21	89		0	1:1.2	AMP	UNKNO,	SYBR	17.385	17.433	0.041	0.000
28	C4		0	1:2	AMP	STAND	SYBR	16.865	16.861	0.013	0.005
29	cs	0	٥	1:2	AMP	STAND	SYBR	16.846	16.861	0.013	0.005
30	C6		0	1:2	AMP	STAND	SYBR	16.870	16.861	0.013	0.005
31	07		0	1:1.3	AMP	UNKNO	SYBR	17.326	17.327	0.004	0.000
32	CB		0	1:1.3	AMP	UNKNO	SYBR	17.332	17.327	0.004	0.000
33	<b>C9</b>		٥	1:1.3	AMP	UNKNO	SYBR	17.323	17.327	0.004	0.000
40	D4		4	1:4	AMP	STAND	SYBR	15.847	15.887	0.037	0.002
41	D5	0	0	1:4	AMP	STAND	SYBR	15.892	15.887	0.037	0.002
42	D6		0	1:4	AMP	STAND	SYBR	15.921	15.887	0.037	0.002
43	D7	0	0	1:1.4	AMP	UNKNO	SYBR	17.318	17.312	0.017	0.000
44	DB		0	1:1.4	AMP	UNKNO	SYBR	17.325	17.312	0.017	0.000
45	D9		0	1:1.4	AMP	UNKNO	SYBR	17.294	17.312	0.017	0.000
52	E4		0	1:8	AMP	STAND	SYBR	15.100	15.102	0.019	0.001

The list can be grouped by the following:



Please note that sections 6.3.1 to 6.2.7 can be applied to all subsequent plot types.



### 6.2.8 Raw Amplification Plot and Raw Melt Peak Plot

Raw plots present the real time fluorescence signal change. The cycle number is shown on the X-Axis while the fluorescence signal value is plotted on the Y-Axis.



By entering the corresponding tabs multicomponent plots for the amplification stage or for the melt curve step will be shown.



# 6.2.9 Standard Curve

Standard curves can be used to verify the amplification efficiency of comparative Cq experiments or to perform an absolute quantitative PCR experiment.

1. Click Standard Curve



2. View standard



**Note**: The amplification efficiency (Eff%), slope, Y-Intercept, R2 and Error will be shown below the graph. A slope of -3.2~-3.5, a R<sup>2</sup>>0.99 and an Efficiency of 90 to 110% are indicators of a reliable experiment.

3. The default mode shows red dots as standard samples, while blue and green are unknown samples.



2.

### 6.2.10 Melt Curve

The melt curve shows the reduction in fluorescence caused by the denaturation of the DNA double helix during the final slow heating step (melt curve analysis).

1. Enter Melt curve plot in the menu.



**Note:** Melt curve plot is applied to check the specificity of PCR reaction and products: Two or more peaks are indicators of an unspecific amplification or of multiple amplicons.

3. Change the plot configurations



**Note:** The "Derivative reporter" plot displays the first derivative of fluorescence signal change according to the temperature. The  $T_m$  (Temperature of melt, the temperature of 50% of dsDNA unwinding to ssDNA) could be used to detect unspecific amplification and primer dimer.



The "Normalized reporter plot" displays the fluorescence signal value decreasing as temperature rises.

# 6.2.11 Gene Expression Map

Gene expression maps can be used to view the results of relative quantitation calculations and the gene expression profile.

1. Click Gene Expression

Amplificati	on Plot	Raw Amplification Plot			
Raw Melt Plot	Melt Peak Plot	QC Summary	Gene Expression		

2. All sample information is shown in the selected table

#	Omit	Sample	Target*1	Cq Mean	∆Cq Mean	∆Cq SE	ΔΔCq	RQ	RQ Min	RQ Max
1		Leber 1	mACTB	20.207						
2		Leber 2	mACTB	20.08						
3		Miz 1	mACTB	18.899						
4		Milz 2	mACTB	19.233						
5		Niere 1	mACTB	19.644						
6		Niere 2	mACTB	21.688						
7		Leber 1	mEef2	20.032	-0.175	0.111	0	1	0.808	1.237
8		Leber 2	mEef2	19.648	-0.432	0.064	-0.257	1.195	1.058	1.351
9		Miz 1	mEef2	18.845	-0.055	0.18	0.12	0.92	0.65	1.301
10		Milz 2	mEef2	19.218	-0.016	0.08	0.159	0.895	0.767	1.045
11		Niere 1	mEef2	18.822	-0.822	0.198	-0.647	1.566	1.07	2.291
12		Niere 2	mEef2	21.872	0.184	0.159	0.359	0.78	0.574	1.059
13		Leber 1	mGAPDH	26.103	5.896	0.489	0	1	0.39	2.564
14		Leber 2	mGAPDH	25.506	5.425	0.082	-0.47	1.386	1.183	1.623
15		Miz 1	mGAPDH	24.255	5.355	0.19	-0.54	1.454	1.008	2.098
16		Milz 2	mGAPDH	25.606	6.373	0.353	0.477	0.718	0.364	1.416
17		Niere 1	mGAPDH	25.525	5.881	0.257	-0.014	1.01	0.616	1.657
18		Niere 2	mGAPDH	31.116	9.428	0.208	3.533	0.086	0.058	0.129

3. Check the gene expression map:



**Note:** There are two plot types available: RQ vs Target (relative quantitation (RQ) values are grouped by target), and RQ vs Sample (relative quantitation (RQ) values are grouped by sample).



# 6.2.12 Analysis Settings

General Analysis settings can be changed by entering the Analysis settings in the menu toolbar:



Enter the tab Relative Quantification Settings to edit settings used for relative quantification of data, such as References, Efficiency of used primers, Outlier Rejection and RQ Min/Max calculations.

nalysis Settings fo	or 2022-08-11_G	iewebe_gDN	IA-murin				
Cq Settings	Relative Q	uantificati	ion Settings	Advanced Settings	Melt Curve Settings	CrosstalkSetting	
Analysis Type Select the type of	analysis to perfe	orm.					
Multiplex	Singleplex						
Select Reference Select the sample	e(s) to use as the ref	ference(s) for	r this experime	nt.			
Reference Sa	mple: Lebe	er 1 ¥					
Select Reference Select the target I necessary.		erence targe	t for this exper	ment. To use multiple endogenous	controls, select Multiple Contr	ols from the drop-down list, then select the c	ontrois as
Reference tar	get(s): mAl	ств	*				
Efficiency					Tarpet	Efficiency (%)	
2000000000	STON AS A	10100-111	e processo de	ACTB	Larget	100.0	
Enter percent	age values be	tween 1 a	COLUMN TO STATE	Eef2 GAPDH		100.0 100.0	
	plicates with ACo			o the value entered below. These	analysis settings apply only to mu	Itplex reactions.	
CReject Rep	viicates with s	pecified Δ	Cq				
∆Cq ≤ 1.0							
RQ Min/Max Ca Select the algorith		rmine RQ Min	and Max value	es (error bars).			
Confidence I	Level: 95.0	× % C	] Standard D	evations: 1 ¥			
						Cinci	Apply

### 6.2.13 Genotyping Maps

2.

The allelic discrimination plot clusters the normalized reporter fluorescence (Rn) of the allelespecific probes used in the SNP assay for the three possible genotypes (Allele 1 homozygous, Allele 2 homozygous, and Allele 1/2 heterozygous) as well as for negative controls (undetermined)

1. Click the allele identification map.





3. The allele identification map, the colors of different samples and the allele types represented by each color will be displayed.



2.

### 6.2.14 Raw Melt Curves of the High-Resolution Melt analysis

The melt profile of a high-Resolution Melt analysis (HRM) reflects the mix of amplicons present in the sample. Variations in sample characteristics like GC content, length, sequence, and heterozygosity are defining the melt curve characteristics of each amplicon.

1. Click the Raw Melt curves tab.

.

..........

...........



3. The software automatically defines the pre- and post-melt regions by pairs of vertical bars before and after the active melt region. If you need to set the regions manually, you can drag and drop the vertical bars of each region directly on the curve. The area between the pair of bars to the left of the active melt region is used to designate 100% fluorescence, where all amplicons are double-stranded. Respectively, the area between the bars to the right of the active melt region is used to determine 0% fluorescence, where all amplicons are single-stranded.

\*\*\*

Temperature("C)

------

01 NI 95

**Note:** Please do not forget to click Analyze after melt-region setting is changed.

### 6.2.15 Aligned Melt Curves of the High-Resolution Melt analysis

During HRM two types of melt curves can be observed. On the one hand, melt curves that differ from each other in melting temperature  $(T_m)$  of the amplicon, but are similar in curve shape. Usually, this kind of curve is generated by homozygous variant samples. On the other hand, melt curves that differ in shape from homozygote melt curves usually indicate heterozygous samples due to the presence of base-pairing mismatches (heteroduplexes) present in the PCR product mix.

1. Click the Aligned Melt curves tab



**Note**: A scaled view of the data is displayed in the aligned plot. Sequence variants with differences in their melt curve (e.g., homozygous vs. heterozygous variants) can be easily discriminated.



### 6.2.16 Difference plot of the High-Resolution Melt analysis

The difference plot is a tool to easily visualize small differences in melt curves of HRM analysis. Curves displayed here are generated by the differences between samples and a single reference.

1. Click the Difference Plot tab.



2. Check the difference plot.



References can be changed by using the drop-down menu and can be either pre-defined controls or single wells.

Reference	blue	
	blue	
rence Pl	brown	
	B6	
	B7	
	B8	
	C5	
	C6	
	C7	

4. The default number of variants in the software pre-set is three. If the actual number of variants is less, any slight difference among the replicates may lead the replicates of one
sample to be divided into different variation groups, even if they have similar curve characteristics. Default settings can be changed by opening the analysis settings in the menu toolbar.

//// qFYR Analysis Studio v1.0										
Eile	<u>E</u> dit	<u>A</u> nalysis	Tools	<u>H</u> elp						
🔂 Ho	me	Analys	sis Setting	s						
		Analyz	ze							

In the tab "HRM Settings" uncheck "Automatically determine the number of variant groups" and enter the desired number. Confirm by clicking Apply.

q Settings	Flag Se	ettings	Advanced	Settings	HRM Settings	Crosstalk Settings
Select an As	say		_	_		HRM Settings for SNPrs12913832, SYBR
Assay	Pre Start	Pre Stop	Post Start	Post Stop	# of Variant G	Number of Variant Groups
P15129138	AUTO	AUTO	AUTO	AUTO	3	Automatically set the pre-melt and post-melt regions.
						Pre-melt region (*C) Start Auto Stop Auto
						Post-melt region (*C) Start Auto Stop Auto
						##HRMAnalysisSettingsPanel.NumberOfVariantGroups
						Automatically determine the number of variant groups.
						Number of variant groups
						Remove all manual variant calls upon reanalysis.
	Ωps	n Assay	Spire	o Assa		

Settings can be saved as default for future experiments using this SNP assay by clicking





#### 6.2.17 Presence/Absence Plot

The Presence/Absence plot display the fluorescence intensity measured in each well.

1. Enter the Presence/Absence plot tab.



2. Check the Presence/Absence results.



3. There are 4 types of plot views that can be chosen in the Plot settings: "All calls" – all calls are displayed, "Presence" – only samples where the target is present are displayed, "Absence" – only samples where the target is absent are displayed, and "Unconfirmed", only unconfirmed samples are displayed. Additionally, you can choose to display fluorescence intensity of the IPC target ("Show IPC") as well as fluorescence intensity of the IPC target in Negative Control-IPC wells and Negative Control-Blocked IPC wells ("Show Controls").



- 4. The software is calculating two types of thresholds for the Presence/Absence plot that are used for the call:
  - The "IPC Threshold" is calculated from Negative Control- Blocked IPC reactions.
  - The "Target Threshold" is calculated from Negative Control- IPC reactions.

- Targets are called regarding their fluorescence intensity:
  - o Call is present if fluorescence intensity is higher than the target threshold,
  - Call is absent if fluorescence intensity is lower than the target threshold, and fluorescence intensity of the IPC is higher than the IPC threshold,
  - Call is unconfirmed if fluorescence intensity is lower than the target threshold, and fluorescence intensity of the IPC is lower than the IPC threshold.

### 6.2.18 QC Summary

The QC Summary screen displays a list of defined experiment flags, including the flag frequency and location of flagged wells for the experiment.

1. Enter the QC Summary tab.

Amplificati	on Plot	Raw Ampli	fication Plot
Raw Melt Plot	Melt Peak Plot	QC Summary	Gene Expression

2. Check the quality of results.

Flag:	Description	Frequency	Wells
AMPNC	Amplification in negative control	0	
DRNMIN	Define acceptable delta Rn based on Cq range	0	
OFFSCALE	Fluorescence is offscale	0	
HIGHSD	High standard deviation in replicate group	0	
CQCONF	Low Cq confidence	0	
NOAMP	No amplification	0	
NOISE	Noise higher than others in plate	0	
SPIKE	Noise spikes	0	
NOSIGNAL	No signal in well		
OUTLIERRG	Outlier in replicate group	0	
EXPFAIL	Exponential algorithm failed	0	
BLFAIL	Baseline algorithm failed	0	
THOLDFAIL	Thresholding algorithm failed	0	
CQFAIL	Cq algorithm failed	0	
мтр	Multiple Tm peaks	10	C1, C3, D2, E2, E9, F2, F10, G1,

**Note:** For some experiment types QC settings can be changed by entering the analysis settings in the menu toolbar:

M qFYR Analysis Studio v1.0										
<u>F</u> ile	<u>E</u> dit	<u>A</u> nalysis	<u>H</u> elp							
🔂 Но	me	Analys	sis Setting	s						
D		Analy	ze							



F140	Description	Use	Attribute	Con	-	Value	Reject We
OFFSCALE	Fluorescence is offscale			1			0
NOSIGNAL	No signal in well	0	Threshold		*	100000.0	0
PCFAIL	Positive control failed						0
SMCLUSTER	Small number of samples in cluster		Number of data points in the cluster		*	2.0	0
AMPNC	Amplification in negative control		C4	4		35.0	
DRNMIN	Define acceptable delta Rn based on Cq range		Cq			35.0	
DRNMIN	Define acceptable delta Rn based on Cq range		ΔRn	4		6.2	0
CQCONF	Low Cq confidence		Cq Confidence		*	0.8	0
NOAMP	No amplification		Amplification algorithm result	4	*	0.1	0
NOISE	Noise higher than others in plate		Relative noise			4,0	0
SPIKE	Noise spikes		Spike algorithm result.	*	*	1.0	0
EXPFAIL	Exponential algorithm falled						0
BLFAIL	Baseline algorithm failed						0
THOLDFAIL	Thresholding algorithm failed						0
CQFAIL	Cig algorithm failed						0

## 6.3 Data Export

#### 6.3.1 Toolbars in Export Tab

Toolbars	Name	Function
Export	Export	Export data
🖾 Save 🗸	Save	Save as .qps file

### 6.3.2 Data Export

 File name: The default file name is the name defined in "Plot properties" or could be changed to customized name. English characters are supported, but not special characters.



 Export content: Include sample setup, Amplification data, Multicomponent data, Results, Melt Curve Result. The contents can be selected by box checking.

	Sample Setup	Amplification Data
	Multicomponent Data	Results
Content	Melt Curve Result	
	Melt Curve Stage: Stage 3 💌	
	Customize	

- The default option is "Unify above content into one file", which will be included as different sheets in one excel file. To export single files, choose "Split above content items into individual files".
- 4. File type: Select QPCR and .xsl, .txt or .csv.



- File Location: The default file location is desktop, which can be changed to other file location by click Browse...
- File Export: The experiment result will be saved and opened in Excel automatically after clicking Export.



7. The export is shown in Excel as below.

ACC 1011	0.1		1000			10.00101		1.1	1.0	45.1	M	10.000	0	1.00		18.00		T	- M-1	v	110011	1.0		2.2		1.48	AC
risto			<b>Due lated</b>					-									_										1.00
Course			2023.04.07	85 17 25	ALC OT																						
almaid Command																											
annual File fileme			( in starting	10000	Color State	and a Direct of	10.	Said street							_												
annual Plants			A PROPERTY.	Acres 1	Added Incom	Concession in	a casar y	pan na p		10.00	4444	Contractor		w11/4													
and the last	S		2022-345-14	1000	Charles and	-																					
arrest flat liter			2022-08-16	10.21.82	PRICEST																						
	1 Tarres				PM CEST																						
ermint Type			Shankard G	ALC: NO.																							
mant Sena No.	riber .		AlMAZIA																								
mant Type			FastCare 1	PY8.48																							
ampendicite			100.0																								
to manufacturer			PG-182214																								
real Stage Sieg																											
wail Shape-Shap																											
Acatum Cycle N	Martine .		64																								
a Vilume			C4. 19.8																								
p method			Athene !																								
Bernard and Dec			Contraction of the local division of the loc																								
an rection			18																								
where like Are			filment .																								
Cucle share C	added to bee	1000 C	orapit	1.0																							
el Cycle sittere C	Cil Pendhymia-	a petterned		462																							
plate type			\$CTA																								
Tante																											
Well Post	Orel	Danish Nan			Superior.	104 3	Di Mean	Ce 50	Quality	Guantity Is Q	ivertity S.Y.	Arban; 84775	ispectal (1	ingen			Cq Threehol		Easeline (		(Any Statu	Centents	Cq Carl	Fiscality	MILLIO	DIMMI	THE.
F.A.L	FALSON		artisty	N10	OVER	Undersonal	18,706									FALBOH		WWW.			incarehosi.				.94	N	
	PALSON		angly 1	MRC .	5V84	Underson	15,754									FALSON	1.012	WHIT			benchus.				.14	84	
2.62	FALSON	and a	angly 1	MPC .	Division .	(Indeferred	18,708									PALSON	1.012	Volume 1	- 3		incruit-m.					84	
	FALSON.	and a	artigity 1	NDC .	01100	(D-Selected)	18.706									FALSON	1.017	VUNUES.			Sectors have				N	84	
4.84				NPC .	0100	Underson	15,705									FALSON	1.012	WARE .		- 44	Board-m					84	
4.84					PTR.	Children of	18,206									ALSO!	1.012	Volume			Belanders.				8		
5.A6	FALSON	engly.														FALSON	1012	Distantion of the	-		- Buistylant						
5.45	FALSON	artisty .	engly 1	NIC	STORE											ALCON.	1.812	Martin .			beard as				2	2	
5.45 5.48 7.47	FALSON FALSON FALSON	engly .	ettphy (	MIC	SYSR.	(Address)	15,706																				
5.45 9.48 7.43	FALSON FALSON FALSON FALSON	erigity erigity		NTC .	2158	Understa	11,706												- 1								
1222	FALSON FALSON FALSON FALSON	erigity erigity erigity erigity	111	NIC NC	SYSR SYSR SYSR	Undergrowthe Undergrowthe	11.706								-	PACIENT	1.812	Volenii .			<b>Beardulan</b>				2	N.	
5.45 9.78 7.47 9.40 9.40 10.410	FALSON FALSON FALSON FALSON FALSON	erigity erigity erigity erigity erigity	1111	ATC ATC ATC	21/28 21/28 21/28	Underson Underson	11.706								-	ALSO!	1812	Voluel Voluel		- 1	Nonclass Nonclass				8-	N. N.	
5.45 5.40 7.47 8.40 10.40 10.40 11.415	FALSON FALSON FALSON FALSON FALSON FALSON		*****	NTC NTC NTC NTC	2756 2756 2756 2756	Undersonne Undersonne Undersonne	11.706 11.706 11.708 11.708									ALSOH FALSOH	1012	10448 10449 10449	-	-	Northan Northan Realthan				1	N N N	
5.46 5.80 7.47 8.40 10.40 11.417 12.412	FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON	******	*****		2198 2198 2198 2198 2198	Underson Underson Underson Underson	11.706 11.706 11.708 11.706									ALSON ALSON FALSON	1012				Roardian Roardian Reardian				1		
5.45 5.80 7.47 8.48 8.49 10.40	FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON	******	******		2788 2758 2758 2758 2758 2758 2758	Underson Und	11.706 11.706 11.706 11.706 11.706									ALSON ALSON ALSON ALSON	1.012				Inconduces Inconduces Inconduces Inconduces				and a		
5.46 5.88 7.47 8.48 5.49 10.419 11.419 12.419 13.819 14.802	FASDI FASDI FASDI FASDI FASDI FASDI FASDI FASDI FASDI	111111111111111111111111111111111111111	*****			Undersonne Undersonne Undersonne Undersonne Undersonne 15,708 11,512	11.706 11.706 11.706 11.706 11.706 11.706 11.706	1.236	121.000	121.918		19.865	1.360	2.790	101.948	ALSON ALSON ALSON ALSON ALSON	100				Inconduces Inconduces Inconduces Inconduces Inconduces						
5.46 5.48 7.47 8.48 8.48 8.48 10.419 11.417 12.412 17.81 14.82 16.82	FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	2013313333	NEC NEC NEC NEC NEC NEC NEC NEC NEC NEC	2758 2758 2758 2758 2758 2758 2758 2758	Underson Und	11.766 11.766 11.766 11.766 11.766 11.766 11.766 11.817 11.817	0,236	Q1.30 01.30	121.365		18.866	0.966	2,760	101.548	ALSON ALSON ALSON ALSON ALSON ALSON	1.012				Repetion Repetion Repution Repution Repution Repution						
5.46 5.48 7.47 8.40 9.40 9.40 9.41 10.41 10.41 10.41 10.01 14.00 16.00 1	FASDI FASDI FASDI FASDI FASDI FASDI FASDI FASDI FASDI	angly angly	2002333333333	NEC NEC NEC NEC NEC NEC NEC NEC NEC NEC	2758 5758 5758 5758 5758 5758 5758 2758 2	Undersonne Undersonne Undersonne Undersonne Undersonne 15,708 11,512	11.756 11.756 11.756 11.756 11.756 11.756 11.756 11.857 11.857 11.857	8.230 9.230 8.230				9.85	0.961	276	101.548 101.548	A.SOI A.SOI A.SOI A.SOI A.SOI A.SOI A.SOI A.SOI	1.812 1.812 1.812 1.812 1.812 1.812 1.817 1.887	10048 10048 10048 10048 10048 10048 10048			Inconduces Inconduces Inconduces Inconduces Inconduces						
5.46 5.48 7.47 8.40 9.40 9.40 9.41 10.41 10.41 10.41 10.01 14.00 16.00 1	FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON	engly engly engly engly engly engly engly Uling Uling Uling Uling	22002333333333	NEC NEC NEC NEC NEC NEC NEC NEC NEC NEC	2758 5758 5758 5758 5758 5758 5758 2758 2	Underson Und	11.756 11.756 11.756 11.756 11.756 11.756 11.756 11.857 11.857 11.857	8,238	125,000	120.000		10.055	0.966	2740	101.548 101.548	A.SOI A.SOI A.SOI A.SOI A.SOI A.SOI A.SOI A.SOI	1.812 1.812 1.812 1.812 1.812 1.812 1.817 1.887	10048 10048 10048 10048 10048 10048 10048			Representation Representation Representation Representation Representation Representation						
5.46 5.40 7.47 8.40 10.4	FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON	engly engly engly engly engly engly engly Uling Uling Uling Uling	22222333333333333333333333333333333333	NEC NEC NEC NEC NEC NEC NEC NEC NEC NEC	2788 5758 5758 5758 5758 5758 5758 2758 2	Underson Und	11.766 11.766 11.766 11.766 11.766 11.766 11.766 11.817 11.817	0,236	125,000	120.008		18.866	0.961	2,760	101.548	ALSON ALSON ALSON ALSON ALSON ALSON ALSON ALSON	1.812 1.812 1.812 1.812 1.812 1.812 1.817 1.887				Repetion Repetion Repution Repution Repution Repution						
5.46 5.40 7.47 8.40 8.40 9.40 9.40 9.40 9.40 10.40 10.40 10.00 10.	FA30H FA30H FA30H FA30H FA30H FA30H FA30H FA30H FA30H FA30H FA30H FA30H FA30H FA30H	engdy engdy engdy engdy engdy engdy engdy USing USing USing USing	22222233333333333333333333333333333333	NEC NEC NEC NEC NEC STANDA STANDA STANDA	2756 2756 2756 2756 2756 2756 2756 2756	Underson Und	11.766 11.756 11.756 11.756 11.756 11.756 11.957 11.957 11.957 11.957 11.957	0,230 0,230 0,230 0,230	10 10 10 10 10 10 10 10 10 10 10 10 10 1	120.000 120.000 120.000 120.000		19,866 19,855 19,855 19,855	0.961 0.962 0.962	2740 2740 2740 2740	101.548 101.548 101.548 101.548 101.548	ALSON TALSON TALSON TALSON TALSON TALSON TALSON TALSON	1.02 1.02 1.02 1.02 1.02 1.02 1.02 1.07 1.07 1.07	20149 20149 20149 20149 20149 20149 20149 20149 20149 20149			Nordan Rotal Rotalan Rotal Rotalan Rotalan Rotalan Rotalan Rotalan Rotalan Rotalan Rotalan Rot						11111
5.46 5.40 7.47 8.40 10.4	FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON FALSON	engely engely engely engely engely engely engely USINg USINg USINg USINg USINg	22222222222222222222222222222222222222		5756 5756 5756 5756 5756 5756 5756 5756	Underson Und	11.756 11.756 11.756 11.756 11.756 11.756 14.857 14.857 14.857 14.857	0,230 0,230 0,230	10 10 10 10 10 10 10 10 10 10 10 10 10 1	125.000 125.000 125.000 125.000 125.000		10,055 10,055 10,055	0.960	274	101.548 101.548 101.548	ALSON TALSON TALSON TALSON TALSON TALSON TALSON TALSON	1.02 1.02 1.02 1.02 1.02 1.02 1.02 1.02	10048 10048 10048 10048 10048 10048 10048 10048			Representation Representation Representation Representation Representation Representation Representation Representation Representation						

#### 6.3.3 Printed report

Next to data export, a report of the complete data for an experiment, including graphs, can be saved, and printed.

- 1. In the result section, select the wells that should be reported
- 2. Access the main menu toolbar, enter the "File" drop down and click "Print Report"



3. A pop-up menu will open to select the data and graphs to be reported

- Fost Report		. *
Experiment Summary Gene Expression Summary (By Technical Repilcate) Gene Expression Rot for Technical Replicate(By Target)	Information about the experiment, including experiment name, experiment spipe. The name, user name, run information, and comments. A table of experiment name information in production, including sample, Cq Imaeni, Cq Intel and, dCq Imaeni, ACq Intel addo of ACG Intel ren. ADGS, IRCA INFO Intel and INO Imael. Minister experiment of marks. The samples a compared to the information sample.	
Orientation: @Hersental Orientation: Scale	Blaner Oth Oling2 Oling18	
Gene Expression Plot for Technical Replicate(By Sample	Relative quantity of target in the samples as compared to the reference sample.	
Orientation: # Horanna () Switcal Scale	# Liner Ola Olag 2 Olag 18	
AmpBfication Plot (Rn vs. Cycle)	detend autient occh number. Data calected during the cycling or amplification stage. Displays normalizet reporter (Nr) plotted apirist cycle romber. A balle of experiment results for each well, including sample, target, task, JAIn, Co, JACs.	
Plate Layout	An illustration of the wells in the reaction plate. Displays the contents assigned to each sell.	
QC Summary Mett Curve (Derivative Reporter)	A table of flags applied to wellk in the experiment, including flag description, frequency of occurrence, and a flat of flagstate wells. Data collected during the melt curve stage. Displays derivative reporter (-Rn*) plotted against temperature (*Cl.	
Mot Curve (Normalized Reporter)	Data collected during the melt curve stage. Displays normalized reporter (Rr) plotted against temperature ( *Q.	
Select All		
	Card Part Parton	nt Report

- Report can either be directly send to a connected printer by clicking Print Report or previewed and saved by clicking Print Preview.
- 5. "Print Preview" will open a new pop up and document can be reviewed and saved or printed. File format for saving is \*.pdf and file can be password-secured.



M Print Preview				×
	1 D D 🗋 🔍 🔍 5	i0% v		
			oFTR Analysis Skallo v1.0	
	Experiment	Results Report		
	2022-09-28_	_uniformity_AMP		
	Experiment Summary			
	Fits Name.2022.09.21, u Beperinst Type:Comparison Ran Started:09.2022 (T Ran Started:09.2022) Namber of Web sith Results:01 Benoment Type:FitsConf.00 Diggstopic and the start Results:01 Diggstopic and the start	Cq (ΔΔCq) 8:11:08 CEST 1:34:53 CEST /R-X6 006		
	Use	1/12	Press 1140-3022 12 0354 057	

Page 1 of 12

## **Chapter 7 Instruments Maintenance**

This instrument does not need a lot of maintenance in case of routine use. If this instrument is used for a long time, it is necessary to clean and maintain the instrument regularly to ensure normal instrument operation. Before cleaning the instrument, please read this chapter carefully. A correct maintenance and cleaning help to prolong the instrument's service life.

If you need to replace any instrument parts during the maintenance, please contact us for professional technical confirmation, and only replace the parts provided by our original factory or our official license agency.

## 7.1 Instrument Cleaning

Do not clean or maintain the instrument during electrification or operation.

Do not pour water or other solutions into the sample bin, sample module or any instrument component. When the instrument is electrified, the flow of liquid may lead to electric shock!

70% medical disinfectant alcohol is recommended for instrument cleaning and disinfectant. However, ethanol is a volatile liquid. Exposure can irritate the eyes, skin, and respiratory tract, and can lead to central nervous system dysfunction and liver damage. When cleaning with ethanol, please wear suitable protective goggles, clothing, and gloves.

**Note**: If the instrument user has any questions about the compatibility of disinfectants or cleaning solution with the materials contained in the equipment parts, please consult the manufacturer or official license agency.

During operation, the sample module may have high temperature. Before cleaning or maintaining, wait for the temperature to drop to room temperature to avoid scalding.

#### **Biological Hazards**

Please treat and operate all samples as potential biological hazards. If liquid samples overflow or splatter, please immediately take appropriate disinfectants for disinfection, to avoid contamination or damage to laboratory personnel.

Do not deal with and operate any potential biological hazards without any safety precautions.



#### 7.1.1 Cleaning the Instrument

Turn off the instrument and unplug the power cord.

Use a damp soft cloth to clean the instrument shell, use a mild cleaner to clean if necessary.

**Note:** Do not spray the cleaner directly on the instrument, which may cause the failure of the electronic instrument.

Please avoid using strong acid and alkali, more than 75% concentrated alcohol, organic solvents or strong detergent and other cleaning disinfectants to clean the instrument without our company's permission, it will destroy the surface coating.

#### 7.1.2 Clean Sample-tray and Reaction-holes

Please close the instrument and open the sample bin.

Use damp soft cloth to clean the sample tray, use mild detergent to clean it if necessary.

Please clean the reaction hole regularly, to ensure that the consumable test tube is in full contact with the reaction hole and the heat conduction is perfect.

Use cotton swabs dipped in mild detergent to clean the reaction hole.

Please close the heat cover after sample tray and reaction hole are completely dry. The reaction module with detergent on the heating surface may damage the instrument.

Do not spray the cleaner directly on the instrument, which may cause the failure of the electronic instrument.

## 7.2 Instrument maintenance

#### 7.2.1 Keep Air-Circulation

The instrument must be positioned with a possibility of free air circulation to ensure accuracy of the target temperature. Please check the placement area regularly to ensure a free air circulation, and no items that can affect the air flow.

#### 7.2.2 Keep Stable Electricity Supply

A stable power supply is required for the normal operation, please check it regularly to ensure that the supply voltage is consistent with the voltage requirement ( $\pm 10\%$  deviation is allowed). Ensure that the power outlet rated load is not less than the instrument requirement.

#### 7.2.3 Keep Clean

Sample tray, reaction modules and optical components contaminations may interfere with thermal cycling and data collection. Please refer to the following instructions to avoid contamination of this instrument:

Make sure the outside of the plates or tubes is clean before adding them to the instrument.

Please clean the reaction module and inner cover regularly, to prevent accumulation of dirt, biological hazards, or fluorescent solvents.

Please confirm the seal of the tubes or plates before running the reaction.

Do not put open reaction consumables or damaged lids into the sample tray. The reagent may leak out and cover the hot-lid surface, sample tray and reaction hole.

Do not run PCR reactions containing volatile reagents, which may lead to contamination in the hot-lid, sample tray and reaction hole.

When this instrument is not used for a long time, please unplug and cover with soft cloth or dustproof bag to prevent dust from entering.

#### 7.2.4 Waste Disposal

After each experiment, there is a large number of amplified products in the test tube, which should be treated as soon as possible according to relevant regulations, to avoid contamination of laboratories and instruments.

Do not open the test tube cap after removing from the reaction module! The high concentration of nucleic acid might pollute the laboratory.



#### 7.2.5 Fuse Replacement

When the circuit fails or is abnormal, current increase or rising of the current may damage important components. If the fuse is installed correctly, the fuse will break when the current rises to a certain height and heat, to protect the instrument or the user.



FastGene<sup>®</sup> qFYR Real-Time PCR System are equipped with two 10 A. When the fuse is damaged, the user can replace the fuse as follows:

- Cut off the mainframe power supply and unplug the power cord
- Open the fuse box lid with a screwdriver
- Check the inner fuse tube. If the middle is disconnected (two or one of them), it means that the fuse is broken
- Replace the new fuse tube (fuse type Φ5×20 mm -T10AH250VP)
- Install the replaced fuse seat under the socket.
- Note: Power must be turned off and removed before replacing fuse.



#### 7.2.6 Overheating Protection

The instrument heating system is equipped with overheating protection. When the heating system fails and the temperature value exceeds the upper limit of the allowable range, the protection will automatically disconnect, and cannot be restored. At this time, the heating system cannot continue to rise and cool normally. After the heating system broke down, user should stop using the instrument, contact the manufacturer for maintenance in time.

# **Chapter 8 Troubleshooting**

a) **Observation:** Switch on the instrument, but no reaction, and power light off.

Possible cause: Inconsistent connection with power cable

Recommended action: Switch off the power on the rear of instrument, re-plug power cable.





Inconsistent plug

correct plug

b) Observation: Shows "No Available Instrument" after click "Start Run".

Possible cause: Inconsistent connection between instrument and computer.

**Recommended action:** Unplug and plug the USB data cable again, re-start qFYR Analyzer Studio.







Inconsistent data cable

correct connection



c) Observation: There is no fluorescence signal appearing on "Multicomponent plot", while the sound of scanning can be heard.

**Possible cause:** The "sample" and "target item" columns on the "well-plate setup" interface are not filled in. The fluorescence signal is displayed before filled in both. If the user does not fill in "sample" or "target item ", the instrument will collect and store the fluorescence signal of all channels in 96 wells in the experimental file, but not display them on the experimental interface. It can be displayed after the user completes the" well plate setting "information.

 $\label{eq:commended} \textbf{action:} \ \mbox{Return to Plate Tab, complete the "Sample" and "Target" information.$ 

d) Observation: Error notice of "Not enough memory, please close some experiment".

Possible cause: Too many experiment files are opened.

**Recommended action:** Closed some inactive experiments, the error notice will be closed. It is recommended No more than six experiment files are opened simultaneously.



e) Observation: The ramp rate slowed down obviously.
Possible cause: The outlet of airflow was blocked.
Recommended action: Clean the tamper on inlet and/or outlet of airflow.



f) Observation: While scanning empty block, background signal of some wells is much higher than average.

Possible cause: There are contamination in these wells.

**Recommended action:** Clean the contamination, please refer to Section 5.3 Monthly maintenance.



**Observation:** Reaction reagent been evaporated
**Possible cause:** The tubes or seal on plate are not closed correctly.
**Recommended action:** Select better air-tight consumables and restart experiment.

Note: If the user cannot judge and troubleshoot the above, contact our company directly.

If the following circumstances occur, the power supply should be cut off immediately, contact the supplier or our company, we will assign qualified maintenance personnel to handle:

- Liquid flows inside the instrument.
- Instruments be rained or watered.
- The instrument has any abnormal sound or smell.
- Damage to the shell caused by instrument drop down.
- Instrument function changes obviously.





### NIPPON Genetics EUROPE GmbH

Production Permit Number:

Product Registration Certificate No./ Product Technical Requirements No.:

Production Enterprise: NIPPON Genetics EUROPE GmbH

Address: Mariaweilerstraße 28-30 | 52349 | Düren |Germany

Tel: +49 2421 554960

Web: www.nippongenetics.eu

NIPPON Genetics EUROPE GmbH

⋈→ +49 2421 554960
⋈→ info@nippongenetics.de
↔→ +49 2421 55496 11
↔ www.nippongenetics.eu

