



# **RTPCR - 3096i** Real Time PCR System





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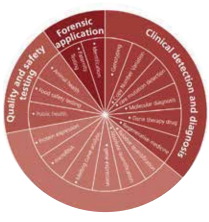
## Analytical Technologies Limited

An ISO 9001 Certified Company

www.analyticalgroup.net

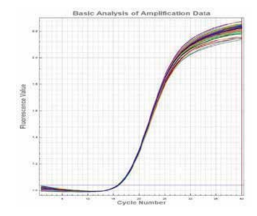


ATL RTPCR-3096i is based on global vision of product design concepts and manufacturing processes. It creatively combines Fresnel lens optical signal acquisition technology, time-resolved signal separation technology and unique temperature control technology. And it reaches international advanced level in sensitivity, multi-color crosstalk, temperature uniformity and accuracy. It supports the application of all common QPCR detection modes.



## Up to 6 fluorescence detection channels allowing multiplex PCR Simultaneous detection of 5 target genes in 96 samples

Simultaneous scanning of the six-channel shows that the standard deviation of the Ct value of the FAM channel is <0.07. No fluorescence signal in other channels.



#### **>>** Technical Innovation 1

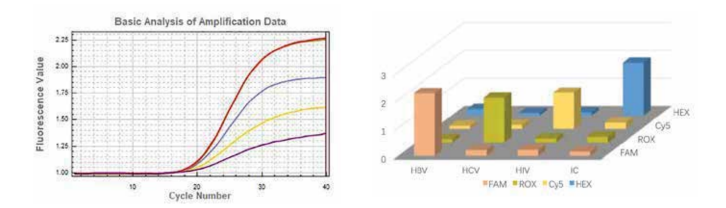
## -Effectively reduce multi-color crosstalk and edge effect, no ROX correction required

The multi-color crosstalk caused by the small sample spacing of 96 or 384-well plates has a great influence on the accuracy of the experimental results, especially in multiplex qPCR detection.

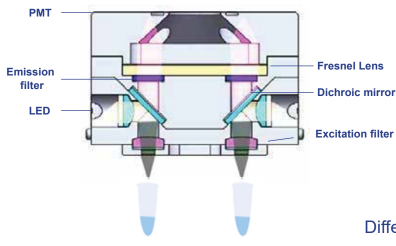
The new optical signal detection system and unique time-resolved scanning can reduce non-target sample optical signal collection. Thereby high repeatability of single fluorescent channel can be ensured.

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Four different target genes (2 repeats) of FAM/HEX/ROX/Cy5 were simultaneously detected in one reaction tube, and the results showed that there was almost no cross-interference between the different channels.

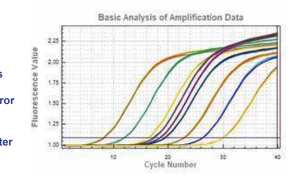


## New optical scanning detection system —High sensitivity /resolution

Sectional view of optical detection

• LED light source Efficient and mainte nance-free

• Fresnel Lens It greatly reduces the light collection of the nontarget area. And the relative position of the detector to the block hole ensures that one optical detection channel is aligned with one target to be tested at the bottom.



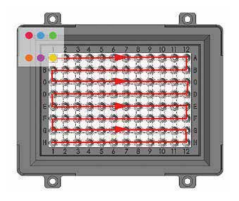
Different concentrations of plasmids were amplified by probe assay (concentration from left to right is  $5\mu g$ , 500 ng, 50ng, 20 ng, 10 ng, 5 ng, 500 pg, 50 pg, 5pg), three replicates for per concentration. The Ct values difference of the 10-fold dilution is exactly 3.3. The Ct values difference of the 2-fold dilution is exactly 1.



## Innovative scanning method and time-resolved signal separation technology —High accuracy

• Unique time-resolved scanning method The different fluorescence signals of the same sample are collected at different times.

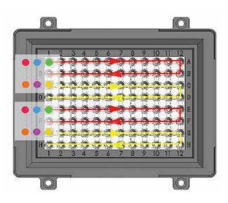
The high-speed stepping motor and the highly sensitive detector ensure that all signal acquisition of the entire sample plate is completed in a short time.



Multi-channel scanning for probe assay

• Innovative detection channel arrangement Interlaced arrangement of upper and lower channels further reduces inter-hole and multi-color fluo-rescence crosstalk.

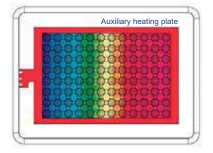
• Double FAM scanning for melting curve Scan time is shortened a lot.



**Double FAM scanning for melting curve** 

#### **>>** Technical Innovation 2

-Unique edge temperature compensation technology



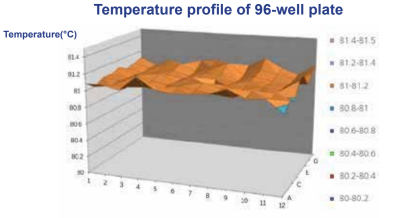


- Both temperature accuracy and uniformity are ± 0.2°C
- Module maximum ramp rate is 6°C/ sec
- The average sample ramp rate is 2.2°C / sec
- With a unique outframe protection design, the unit can achieve very even temperature across the whole plate. Effectively reduce the edge temperature variance.

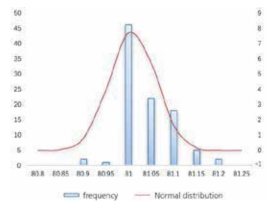


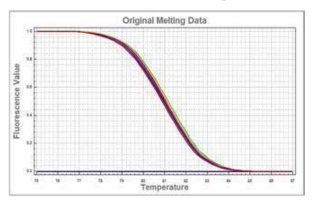


The Tm values of 96 replicate wells were detected. The results showed that the temperature uniformity difference was  $< \pm 0.2^{\circ}$ C@81°C.



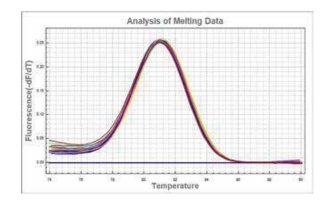
## Tm value normal distribution map of 96-well plate





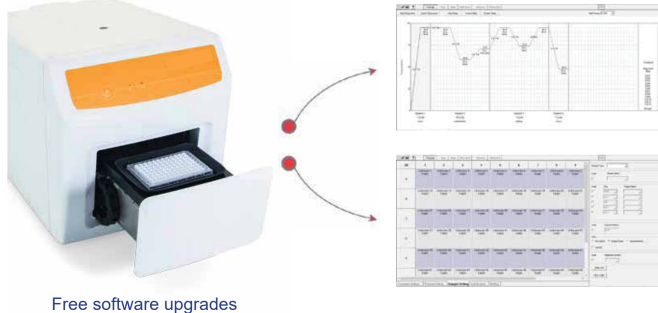
#### The melting curves of 48 replicate samples

05





#### **bb** User-friendly software



The software can provide you with a full range of solutions for sample detection and result analysis.



### One-stop process and solution



#### **Classic Examples-Swine fever virus ASFV detection**

QPCR was used to detect the swine fever virus DNA in vitro for clinical diagnosis of suspected diseased pigs.

**Kit**: Swine fever virus (ASFV) nucleic acid detection kit (PCR probe assay), DNA extraction kit (Spin column method)

**Method**: 5 mL of blood was extract from a live pig syringe to be examined. DNA was extracted by DNA extraction kit. According to the ASFV detection kit operation method, the extracted DNA, the positive control substance, and the negative control substance is separately added to the PCR reaction solution and the enzyme. Then the mixture is centrifuged and tested on real-time qPCR. The reporter and quencher of the TaqMan probe is FAM and TAMRA.

**Judgment basis**: Positive: Ct≤35, and the amplification curve has a significant exponential growth curve.Negative:Ct>37, or no significant amplification of the curve. Recommended retest: 35<Ct≤37.

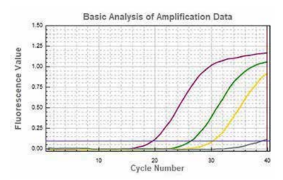
**Data analysis**: The Ct value of the positive control is 20, and the negative control product has no obvious amplification curve. The Ct values of the sample 1 and 2 to be tested is 26 and 30 respectively. According to the judgment basis of the kit, the samples to be tested are ASFV positive.

07

#### **Program:**

Step	Temperature	Time	Cycle	End point
15	95°C	10min	.1	No
) +++++++++++++++++++++++++++++++++++++			100000000000000000000000000000000000000	
2	.94°C	15580	40	No
	55°C	30sec		Yes

#### Amplification curve:





#### **>>** Technical Parameters

## **Temperature control system**

Sample capacity	0.1ml PCR tubes×96, 8×12 PCR plate or 96 well plate×1	
Reaction volume	10-50µl	
Thermal cycle technology	Peltier	
Max. Heating/Cooling rate	6.0°C/s	
Heating temperature range	4 – 100°C	
Temperature accuracy	±0.2°C	
Temperature uniformity	±0.2°C@60, ±0.2°C@95°C	
Temperature gradient		
setting range	30–100°C	
Temperature gradient		
difference setting range	1 – 36°C	

## **Detection system**

Excitation light source	4/6 monochrome high efficiency LEDs		
Detection device	PMT		
Time-resolved signal	Detection mode		
Separating technology			
Excitation/detection wavelength	455-650nm/510-715nm		
range			
Fluorescent channels	4/6 channels		
Supported dye	FAM/SYBR Green, VIC/JOE/HEX/TET, ABY/NED/		
	TAMRA/Cy3, JUN, ROX/Texas Red,		
	Mustang Purple, Cy5/LIZ		
Sensitivity	Single copy gene		
Resolution	1.33 folds copy number difference can be		
	distinguished in single-plex qPCR		
Dynamic range	10 orders of magnitude copies		

08



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HPLC Servicing :HPLC Servicing : We have team of service engineers who can attend to any make of HPLC promptly @the most affordable cost. Trainings :We also take up preventive Maintenace to reduce downtime of HPLC's Trainings.

AMC's/CMC :AMC's/CMC :We offer user training both in-House and at customer sites on HPLC principles, operations, troubleshooting.

Validations :Validations :We have protocols for carrying out periodic Validations as per GLP/GMP/USFDA norms.

:Instruments :We offer instruments/Renting Services Modules like pumps, detector etc. on Rent. Instruments

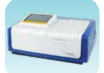




### About Analytical Technologies

Analytical Technologies is synonymous for offering technologies for doing analysis and is the Fastest Growing Global Brand having presence in at least 96 countries across the global. Analytical Technologies Limited is an ISO:9001 Certified Company engaged in Designing, Manufaturing, Marketing & providing Services for the Analytical, Chromatography, Spectroscopy, Bio Technology, Bio Medical, Clinical Diagnostics, Material Science & General Laboratory Instrumentation. Analytical Technologies, India has across the Country operations with at least 4 Regional Offices, 6 Branch Offices & Service Centers. Distributors & Channel partners worldwide.

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Atomic Absorption

Spectrophotometer

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NOVA-2100 Chemistry Analyzer

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TOC Analyzer

Laser Particle Size Analyzer

Micro Plate

Reader/Washer

Ion Chromatograph

Water purification system



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### **Regulatory compliances**



### **Corporate Social Responsibility**

Analytical Foundation is a nonprofit organization (NGO) found for the purpose of:



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