

# New concepts for accelerated real-time PCR analysis



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## Introduction

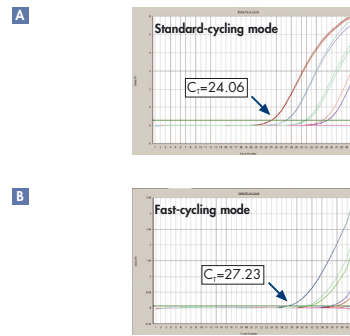
For researchers needing to increase their throughput or share a cyclor with other users, there is a strong demand for faster, real-time PCR.

**Fast, real-time PCR can be achieved by:**

- Reduced DNA polymerase activation time
- Shortened amplification cycles
- Combined annealing and extension steps
- Use of a dedicated fast-cycling instrument
- Shortened RT step in one-step RT-PCR

Until now, fast, real-time PCR using standard reaction chemistry has been hampered by reduced sensitivity and increased variability of quantification data (1). We demonstrate how the combination of a newly developed fast-cycling PCR buffer with a rapid-activating hot-start DNA polymerase allow significant reduction of PCR cycling times without sacrificing specificity and sensitivity.

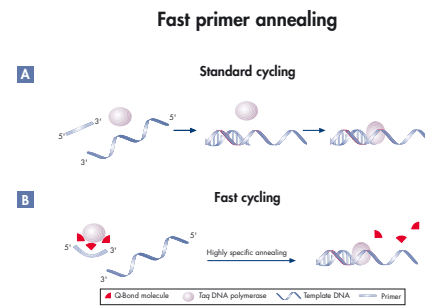
### Loss of sensitivity in fast cycling with standard reaction chemistry



Expression of MYC in human leukocytes was analyzed by real-time, two-step RT-PCR on the Applied Biosystems® 7500 Fast System. A kit for standard cycling from Supplier A was run (A) in recommended standard-cycling mode (reduced ramping rates, 15-second denaturation, 60-second annealing/extension) and (B) in fast-cycling mode (rapid ramping rates, 10-second denaturation, 30-second annealing/extension).

## New chemistries for fast-cycling PCR

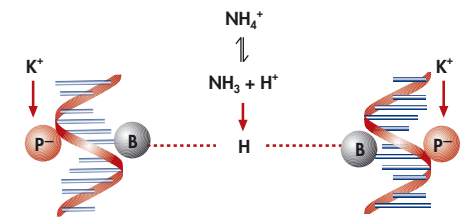
We have developed a fast-cycling PCR buffer that significantly reduces denaturation, annealing, and extension times. A novel additive, Q-Bond, dramatically increases the binding affinity of DNA polymerase to single-stranded DNA. This turns the 3-step process of template denaturation, primer annealing, and DNA polymerase binding in standard-cycling PCR (A) into a faster 2-step process (B).



(A) Without Q-Bond, the primer and DNA polymerase bind sequentially to the template, increasing primer annealing time. (B) Q-Bond increases the affinity of DNA polymerase for short single-stranded DNA, reducing primer annealing time to a few seconds.

High annealing specificity is maintained by a balanced combination of KCl and NH<sub>4</sub>Cl in the buffer. The binding of primers to imperfectly matching sequences on the template is suppressed.

### Specific primer annealing



The fast-cycling PCR buffer contains K<sup>+</sup> and NH<sub>4</sub><sup>+</sup> ions to ensure specific primer annealing and efficient extension. NH<sub>4</sub><sup>+</sup> destabilizes the weak hydrogen bonds at mismatched bases of nonspecifically bound primers.

A novel enzyme, HotStarTaq® Plus DNA Polymerase, is rapidly activated in 3 or 5 minutes by a 95°C incubation at the start of PCR.

## Ultrafast cycling for end-point PCR analysis

Although fast results in PCR can be achieved on cyclers with rapid ramping rates, even faster results are possible by reducing cycling times. The QIAGEN® Fast Cycling PCR Kit, which integrates the fast-cycling PCR buffer with HotStarTaq Plus DNA Polymerase, provides significant time savings of up to 78% in end-point PCR. Fast results can be accomplished on all cyclers, including cyclers not capable of fast ramping rates.

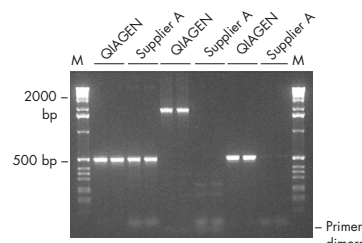
The PCR buffer minimizes amplification of nonspecific products, primer-dimer formation, and background smear in every PCR cycle. Q-Solution, an additive that enables efficient amplification of "difficult" (e.g., GC-rich) templates, is also provided with the kit.

### PCR cycling times calculated for different fragment lengths\*

Fragment length	QIAGEN Fast Cycling procedure (min)	Standard cycling procedure (min)	Time saving
200 bp	15	68	78%
500 bp	20	68	71%
1000 bp	29	85	66%
3000 bp	63	155	59%

\* Total time required for a PCR run of 35 cycles. The specified PCR cycling times do not include ramping times, which are cycler-dependent.

### Specific and reliable results

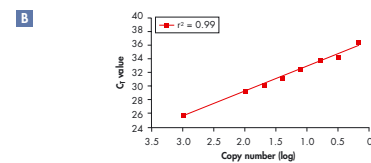
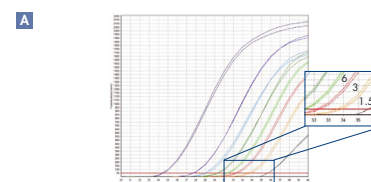


Three different PCR assays (IL1R, PKC, and AGRT2) were performed using the QIAGEN Fast Cycling PCR Kit (QIAGEN) and a fast-cycling PCR solution from another supplier (Supplier A) according to the manufacturer's instructions. Reactions were performed on a fast cycler from Supplier A. The QIAGEN Fast Cycling PCR Kit provided highly specific results for each assay, whereas results using Supplier A were unpredictable with high drop-out rates and unspecific results (e.g., primer-dimers). M: marker.

## Fast SYBR Green quantification with high specificity

In biological systems, minute changes in transcript abundance often lead to strong biological effects. Therefore, a method for reliable and reproducible discrimination between similar copy numbers is critical. With QuantiFast™ SYBR Green Kits, even small differences in the amount of low-copy targets can be clearly distinguished.

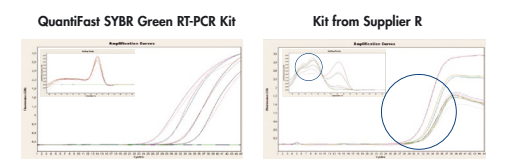
### Resolution of small differences in copy number



The QuantiFast SYBR Green PCR Kit and Mastercycler® ep realplex were used to detect the single-copy gene SRY in a genomic DNA sample. (A) Curves for 1000 down to 1.5 copies can be clearly distinguished from each other. (B) A plot of copy number (log) versus C<sub>t</sub> value demonstrates high linearity.

Achieving high specificity with SYBR Green based real-time PCR is important, since SYBR Green binds all double-stranded DNA. QuantiFast SYBR Green Kits promote specific primer annealing and provide a stringent hot start, preventing the formation of nonspecific PCR products that can affect SYBR Green fluorescence. Quantification is specific and sensitive, even with low template amounts.

### Specific one-step RT-PCR

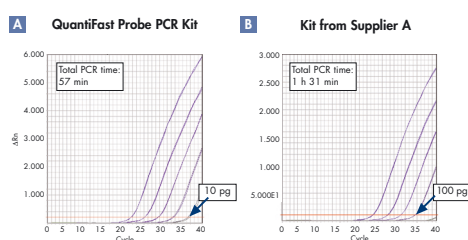


Expression of BCL2 in HeLa cells was analyzed on the LightCycler® 2.0. Unlike the instrument-dedicated kit from Supplier R (which was used according to the fast-cycling protocol), the QuantiFast SYBR Green RT-PCR Kit provided specific amplification and sensitive quantification.

## Speed and sensitivity in probe-based detection

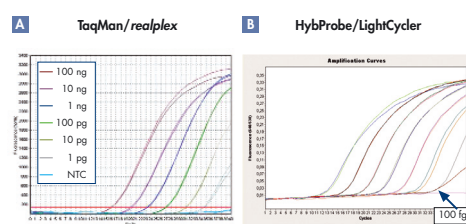
Reducing the duration of each PCR cycle leads to faster PCR, but can impair PCR performance. The use of our fast-cycling technology by QuantiFast Probe Kits enables faster results in probe-based real-time PCR without compromising sensitivity. Fast and sensitive quantification is possible on all available cyclers and with different types of sequence-specific probe, such as TaqMan® and FRET probes.

### Faster results without compromising sensitivity



Reactions were run on the ABI PRISM® 7900 using a TaqMan gene expression assay for IL1RN (a cytokine). The QuantiFast Probe PCR Kit was 40% faster than the standard-cycling kit from Supplier A, and also more sensitive: C<sub>t</sub> values were much lower and transcript could be quantified from down to 10 pg cDNA.

### Sensitive detection independent of cycler and probe



The QuantiFast Probe PCR +ROX Vial Kit provided accurate gene expression analysis from high to low amounts of human leukocyte cDNA with PCR efficiencies of greater than 90%. (A) Reactions run on the Mastercycler ep realplex (block cycler) using a TaqMan assay for ubiquitin. (B) Reactions run on the LightCycler 2.0 (capillary cycler) using a FRET assay for β-2 microglobulin. Both cyclers were run using maximum ramping rates and short cycling steps.

## Summary

- A patent-pending, fast-cycling PCR buffer containing Q-Bond significantly reduces denaturation, annealing, and extension times.
- HotStarTaq Plus DNA Polymerase possesses no enzyme activity prior to PCR, and is rapidly heat-activated in 3 or 5 minutes.
- The fast-cycling PCR conditions provide significant time savings of up to 78% in end-point PCR and up to 60% in real-time PCR without compromising specificity and sensitivity.
- New QIAGEN chemistries enable fast-cycling on all cyclers, including those not capable of achieving rapid ramping rates.

### References

- Hilscher, C., Vahrson, W., and Dittmer, D.P. (2005) Faster quantitative real-time PCR protocols may lose sensitivity and show increased variability. *Nucleic Acids Res.* 33:e182.

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