Real-time PCR for mRNA quantitation

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Real-time PCR has become one of the most widely used methods of gene quantitation because it has a large dynamic range, boasts tremendous sensitivity, can be highly sequence-specific, has little to no post-amplification processing, and is amenable to increasing sample throughput. However, optimal benefit from these advantages requires a clear understanding of the many options available for running a real-time PCR experiment. Starting with the theory behind real-time PCR, this review discusses the key components of a real-time PCR experiment, including one-step or two-step PCR, absolute versus relative quantitation, mathematical models available for relative quantitation and amplification efficiency calculations, types of normalization or data correction, and detection chemistries. In addition, the many causes of variation as well as methods to calculate intra- and inter-assay variation are addressed.

INTRODUCTION

The advent of real-time PCR and real-time reverse transcription PCR (real-time RT-PCR) has dramatically changed the field of measuring gene expression. Real-time PCR is the technique of collecting data throughout the PCR process as it occurs, thus combining amplification and detection into a single step. This is achieved using a variety of different fluorescent chemistries that correlate PCR product concentration to fluorescence intensity (1). Reactions are characterized by the point in time (or PCR cycle) where the target amplification is first detected. This value is usually referred to as cycle threshold ($C_T$), the time at which fluorescence intensity is greater than background fluorescence. Consequently, the greater the quantity of target DNA in the starting material, the faster a significant increase in fluorescent signal will appear, yielding a lower $C_T$ (2).

There are many benefits of using real-time PCR over other methods to quantify gene expression. It can produce quantitative data with an accurate dynamic range of 7 to 8 log orders of magnitude (3) and does not require post-amplification manipulation. Real-time PCR assays are 10,000- to 100,000-fold more sensitive than RNase protection assays (4), 1000-fold more sensitive than dot blot hybridization (5), and can even detect a single copy of a specific transcript (6). In addition, real-time PCR assays can reliably detect gene expression differences as small as 23% between samples (7) and have lower coefficients of variation (cv; SYBR® Green at 14.2%; TaqMan® at 24%) than end point assays such as band densitometry (44.9%) and probe hybridization (45.1%) (8). Real-time PCR can also discriminate between messenger RNAs (mRNAs) with almost identical sequences, requires much less RNA template than other methods of gene expression analysis, and can be relatively high-throughput given the proper equipment. The major disadvantage to real-time PCR is that it requires expensive equipment and reagents. In addition, due to its extremely high sensitivity, sound experimental design and an in-depth understanding of normalization techniques are imperative for accurate conclusions.

The general steps performed during a real-time PCR experiment, from RNA isolation to data analysis, are outlined in Figure 1. This review is intended to provide an overview of the many facets of real-time PCR, highlighting PCR theory, quantification methods and models, data normalization, types of detection chemistry, and causes of variation.

THEORY OF REAL-TIME PCR

PCR can be broken into four major phases (Figure 2): the linear ground phase, early exponential phase, log-linear (also known as exponential) phase, and plateau phase (9). During the linear ground phase (usually the first 10–15 cycles), PCR is just beginning, and fluorescence emission at each cycle has not yet risen above background. Baseline fluorescence is calculated at this time. At the early exponential phase, the amount of fluorescence has reached a threshold where it is significantly higher (usually 10 times the standard deviation of the baseline) than background levels. The cycle at which this occurs is known as $C_T$ in ABI Prism® literature (Applied Biosystems, Foster City, CA, USA) or crossing point (CP) in LightCycler® literature (Roche Applied Science, Indianapolis, IN, USA) (2,10). This value is representative of the starting copy number in the original template and is used to calculate experimental results (2). During the log-linear phase, PCR reaches its optimal amplification period with the PCR product doubling after every cycle in ideal reaction
REVIEW

One-Step Versus Two-Step Real-Time PCR

When quantifying mRNA, real-time PCR can be performed as either a one-step reaction, where the entire reaction from cDNA synthesis to PCR amplification is performed in a single tube, or as a two-step reaction, where reverse transcription and PCR amplification occur in separate tubes. There are several pros and cons associated with each method. One-step real-time PCR is thought to minimize experimental variation because both enzymatic reactions occur in a single tube. However, this method uses an RNA starting template, which is prone to rapid degradation if not handled properly. Therefore, a one-step reaction may not be suitable in situations where the same sample is assayed on several occasions over a period of time. One-step protocols are also reportedly less sensitive than two-step protocols (12).

Two-step real-time PCR separates the reverse transcription reaction from the real-time PCR assay, allowing several different real-time PCR assays on dilutions of a single cDNA. Because the process of reverse transcription is notorious for its highly variable reaction efficiency (13), using dilutions from the same cDNA template ensures that reactions from subsequent assays have the same amount of template as those assayed earlier. Data from two-step real-time PCR is quite reproducible with Pearson correlation coefficients ranging from 0.974 to 0.988 (14). A two-step protocol may be preferred when using a DNA binding dye (such as SYBR Green I) because it is easier to eliminate primer-dimers through the manipulation of melting temperatures ($T_m$) (14). However, two-step protocols allow for increased opportunities of DNA contamination in real-time PCR.

TYPES OF REAL-TIME QUANTIFICATION

Absolute Quantitation

Absolute quantitation uses serially diluted standards of known concentrations to generate a standard curve. The standard curve produces a linear relationship between $C_t$ and initial amounts of total RNA or cDNA, allowing the determination of the concentration of unknowns based on their $C_t$ values (2). This method assumes all standards and samples have approximately equal amplification efficiencies (15). In addition, the concentration of serial dilutions should encompass the levels in the experimental samples and stay within the range of accurately quantifiable and detectable levels specific for both the real-time PCR machine and assay.

The PCR standard is a fragment of double-stranded DNA (dsDNA), single-stranded DNA (ssDNA), or cRNA bearing the target sequence. A simple protocol for constructing a cRNA standard for one-step PCR can be found in Fronhoffs et al. (16), while a DNA standard for two-step real-time PCR can be synthesized by cloning the target sequence into a plasmid (17), purifying a conventional PCR product (18), or directly synthesizing the target nucleic acid. The standard used must be a pure species. DNA standards have
been shown to have a larger quantification range and greater sensitivity, reproducibility, and stability than RNA standards (19). However, a DNA standard cannot be used for a one-step real-time RT-PCR due to the absence of a control for the reverse transcription efficiency (20).

### Relative Quantitation

During relative quantitation, changes in sample gene expression are measured based on either an external standard or a reference sample, also known as a calibrator (21). When using a calibrator, the results are expressed as a target/reference ratio. There are numerous mathematical models available to calculate the mean normalized gene expression from relative quantitation assays. Depending on the method employed, these can yield different results and thus discrepant measures of standard error (22,23). Table 1 shows a comparison of the different methods, with an explanation of each method to follow.

<table>
<thead>
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<th>Table 1. Characteristics of Relative Quantitation Methods</th>
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<td><strong>Methods (Reference)</strong></td>
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<td>Standard Curve (31)</td>
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<sup>a</sup>www.gene-quantification.info  
<sup>b</sup>www.BioTechniques.com  
<sup>c</sup>nar.oupjournals.org/cgi/content/full/31/14/e73/DC1

Amplification efficiency. Amplification efficiency of the reaction is an important consideration when performing relative quantitation. Past methods of calculating gene expression have assumed the amplification efficiency of the reaction is ideal, or 1, meaning the PCR product concentration doubles during every cycle within the exponential phase of the reaction (24). However, many PCRs do not have ideal amplification efficiencies, and calculations without an appropriate correction factor may overestimate starting concentration (22). Current mathematical models make assumptions of reaction kinetics and usually require its accurate measurement (7,21,22,25,26).

Traditionally, amplification efficiency of a reaction is calculated using data collected from a standard curve with the following formula (27):

$$\text{Efficiency} = 10^{(-1/slope)}$$  

**[Eq. 1]**

Exponential amplification = \(10^{(-1/slope)}\)

The amplification efficiency of the reaction varies from being relatively stable in the early exponential phase and gradually declining to zero (22). This decay is due to the depletion of PCR components, the decline of polymerase activity, and competition with PCR products. Calculation of amplification efficiency using a standard curve is not reflective of this changing efficiency (28) and may overestimate efficiencies (9). Because PCR results are based on C<sub>i</sub>, which are determined very early in the exponential phase of the reaction, these differences in amplification efficiency usually generate only minor differences in C<sub>i</sub> value (20). Nonetheless, after 26 cycles, a 5% difference in amplification efficiency can result in a 2-fold difference of PCR product concentration (29).

There are several alternate methods of calculating amplification efficiency based on raw data collected during PCR (7,9,22,25,30). During the exponential phase, the absolute fluorescence increase at each PCR cycle for each individual sample reflects the true reaction kinetics of that sample. Consequently, data collected during the exponential phase can be log-transformed and plotted with the slope of the regression line representing the sample’s amplification efficiency. In the Liu and Saint (22) method, the individual researcher designates which cycles have exponential characteristics, while the method proposed by Tichopad et al. (9) uses a statistical calculation to define the period of exponential growth. Amplification efficiency calculated from raw data analysis is reportedly more accurate than when derived from a standard curve (9,25).

**Standard curve method for relative quantification.** The quantity of each experimental sample is first determined using a standard curve and then expressed relative to a single calibrator sample (31). The calibrator is designated as 1-fold, with all experimentally derived quantities reported as an n-fold difference relative to the calibrator. Because sample quantity is divided...
by calibrator quantity, standard curve units are eliminated, requiring only the relative dilution factors of the standards for quantification. This method is often applied when the amplification efficiencies of the reference and target genes are unequal (22). It is also the simplest method of quantification because it requires no preparation of exogenous standards, no quantification of calibrator samples, and is not based on complex mathematics. However, because this method does not incorporate an endogenous control (usually a housekeeping gene), results must still be normalized.

**Comparative Ct** \(2^{-\Delta C_{t}}\) method. The comparative \(C_{t}\) method is a mathematical model that calculates changes in gene expression as a relative fold difference between an experimental and calibrator sample. While this method includes a correction for nonideal amplification efficiencies (i.e., not 1; Reference 21), the amplification kinetics of the target gene and reference gene assays must be approximately equal (32) because different efficiencies will generate errors when using this method (22). Consequently, a validation assay must be performed where serial dilutions are assayed for the target and reference gene and the results plotted with the log input concentration for each dilution on the x-axis, and the difference in \(C_{t}\) (target-reference) for each dilution on the y-axis. If the absolute value of the slope of the line is less than 0.1, the comparative \(C_{t}\) method may be used (21). The PCR product size should be kept small (less than 150 bp) and the reaction rigorously optimized (25). Because the comparative \(C_{t}\) method does not require a standard curve, it is useful when assaying a large number of samples since all reaction wells are filled with sample reactions rather than standards.

**Pfaffl model.** The Pfaffl model (26) combines gene quantification and normalization into a single calculation. This model incorporates the amplification efficiencies of the target and reference (normalization) genes to correct for differences between the two assays. The relative expression software tool (REST\textsuperscript{®}), which runs in Microsoft\textsuperscript{®} Excel, automates data analysis using this model (33). REST uses the Pairwise Fixed Reallocation Randomization Test\textsuperscript{®} to calculate result significance and will indicate if the reference gene used is suitable for normalization.

**Q-Gene.** Q-Gene is a fully comprehensive Microsoft Excel-based software application that aids in the entire process of a real-time PCR experiment, from experimental planning and setup through data analysis and graphical presentation (23). Q-Gene calculates the mean normalized gene expression with standard errors using two different mathematical models, both correcting for amplification efficiencies. The calculated expression values are then compared between two matched groups to determine the expression of a sample relative to a calibrator. The program also includes several statistical tests such as the paired or unpaired Student's t-test, a Mann-Whitney U-test, Wilcoxon signed-rank test, together with Pearson’s correlation analysis to fully assess the significance of experimental results. When running large or complex real-time PCR experiments, having an organized and automated method such as Q-Gene can significantly expedite data processing and management.

**Gentle et al.** Gentle et al. (7) designed one of the first models in which both fold changes between samples and amplification efficiencies of experimental versus control samples are calculated without the use of standard curves. Linear regression analyses of the mean of the raw log fluorescence data collected during the exponential phase of the PCR are used to calculate the amplification efficiency of each sample. By graphing the control and experimental samples together, they show that the vertical distance between the control and experimental lines is the log of the fold difference between the two, with the slopes of the lines representing the log of their amplification efficiencies (7).

**Liu and Saint.** Liu and Saint (22) developed a sigmoidal mathematical model to quantitate and normalize gene expression. Similar to Gentle et al. (7), this method calculates amplification efficiencies from the actual slope of the amplification plot rather than a standard curve. The authors found this method was more accurate than the comparative \(C_{t}\) method with regard to the varying amplification efficiency throughout the PCR because the user defines which amplification efficiency is used for the calculation (22).

**Amplification plot method.** The amplification plot method uses a simple algorithm to calculate the amplification efficiency of experimental versus control samples. This method is based on the assumption that the amplification of the target gene is linear and that the standard curve is linear. The amplification plot method calculates the amplification efficiency by calibrator quantity, standard curve units are eliminated, requiring only the relative dilution factors of the standards for quantification. This method is often applied when the amplification efficiencies of the reference and target genes are unequal (22). It is also the simplest method of quantification because it requires no preparation of exogenous standards, no quantification of calibrator samples, and is not based on complex mathematics. However, because this method does not incorporate an endogenous control (usually a housekeeping gene), results must still be normalized.

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efficiency of every sample individually within the real-time PCR assay. These data are then used in the calculation for expression quantitation (30). To ease data handling, Peirson et al. (30) have developed a Microsoft Excel workbook entitled Data Analysis for Real-Time PCR (DART-PCR) that quickly calculates all results from raw data.

**Absolute or Relative Quantitation: Pros and Cons**

Absolute quantitation is considered to be more labor-intensive than relative quantitation because of the necessity to create reliable standards for quantitation and include these standards in every PCR (19). However, when performing relative quantitation, the data \( (C_T) \) used for comparison are arbitrary values and only applicable to the samples run within the same PCR. To compare samples between two different PCRs, it is necessary to include a reference control in every plate or run. In cases where data compared are assayed on different days or in different laboratories, absolute quantitation may be preferred because results are based on a constant. In terms of fold-change data, absolute and relative quantitation methods produce comparable results (30).

**Controls**

There are several types of controls that ensure the integrity of every step of the real-time PCR process. DNA contamination in the sample may be accounted for with a minus reverse transcription control. However, when one has numerous samples, an alternate method to prevent the detection of genomic DNA is to design the target PCR product to span an exon/exon boundary. Variation in the efficiency of the reverse transcriptase as well as the amount of RNA added into the reaction can be accounted for using an endogenous control, which is a nucleic acid already present in an individual sample. The use of endogenous controls is discussed in detail in the section entitled Normalization. PCR master mix volume has been shown to be a factor in PCR amplification efficiency such that differences in master mix volume in reactions using the same amount of starting template have different amplification efficiencies (22). A passive reference dye (such as ROX) is often included in the master mix to account for subtle differences in PCR master mix volumes as well as non-PCR-related fluctuations in

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**Figure 3. Real-time PCR detection chemistries.** Probe sequences are shown in blue while target DNA sequences are shown in black. Primers are indicated by horizontal arrowheads. Not all unlabeled PCR primers are shown. Oligo, oligonucleotide.
fluorescence signal. Problems with the PCR master mix itself can be accounted for using an exogenous control, which is a synthesized construct of characterized RNA or DNA spiked into each reaction (34).

Normalization

Normalization of gene expression data is used to correct sample-to-sample variation. Starting material obtained from different individuals usually varies in tissue mass or cell number, RNA integrity or quantity, or experimental treatment. Under ideal conditions, mRNA levels can be standardized to cell number, but when using whole tissue samples, this type of normalization is impossible (35). Therefore, real-time PCR results are usually normalized against a control gene that may also serve as a positive control for the reaction (36).

Housekeeping genes (mRNA). Traditionally, genes thought to have stable expression have been employed as controls in gene expression assays. Due to the increased sensitivity and dynamic range of real-time PCR over traditional quantitation techniques, many of the well-known housekeeping genes such as GAPDH and β-actin have been shown to be affected by different treatments, biological processes, and even different tissues or cell types (reviewed in depth in Reference 11). Consequently, normalization with a single housekeeping gene can falsely bias results. When using a housekeeping gene for normalization, it is absolutely imperative to validate its expression stability with one’s own samples rather than relying on previously published materials.

Ribosomal RNA (rRNA). rRNA is another possible reference gene for normalization. Of the two main rRNAs, 28S and 18S, 28S is considered more representative of mRNA integrity because 18S may remain intact in samples with degraded mRNA (37). There are several problems with using 28S rRNA to normalize mRNA gene measurements. rRNAs are transcribed with a different polymerase than mRNAs, so changes in polymerase activity may not affect both types of RNA expression equally (38). This is likely reflected in the fact that rRNA expression tends to be less affected by treatments that significantly alter mRNA expression (39). Varying ratios of rRNA to mRNA have been reported (40) and, given the extreme abundance of 28S rRNA in a total RNA sample [in a 10-µg total RNA sample, on average 2 µg are 18S rRNA and 5.5 µg are 28S rRNA (Technical Bulletin #151, www.ambion.com/techlib/tb/tb_151.html; Reference 40a)], it may be impossible to accurately measure both 28S and a rare transcript in the same RNA or cDNA dilution. Lastly, rRNA, which lacks a poly(A) tail, cannot be measured if an oligo(dT) or gene-specific primer has been used for reverse transcription.

<table>
<thead>
<tr>
<th>Detection Chemistries</th>
<th>Specificity</th>
<th>Multiplex Capability</th>
<th>Specific Oligonucleotide Required</th>
<th>Allelic Discrimination</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA Binding Dyes</td>
<td>Two PCR primers</td>
<td>No</td>
<td>No</td>
<td>No</td>
<td>$</td>
</tr>
<tr>
<td>Hybridization Probe Four Oligonucleotide Method</td>
<td>Two PCR primers; two specific probes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>$$$</td>
</tr>
<tr>
<td>Hybridization Probe Three Oligonucleotide Method</td>
<td>Two PCR primers; one specific probe</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>$$$</td>
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<tr>
<td>Hydrolysis Probes</td>
<td>Two PCR primers; one specific probe</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>$$$</td>
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<td>Molecular Beacons</td>
<td>Two PCR primers; one specific probe</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td>Scorpions</td>
<td>One PCR primer; one primer/probe</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
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<tr>
<td>Sunrise Primers</td>
<td>Two PCR primers</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>$$$</td>
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<tr>
<td>LUX Primers</td>
<td>Two PCR primers</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
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$, very expensive; $\$, moderately expensive; $\$, inexpensive. LUX, light upon extension.
DNA binding dyes emit fluorescence when bound to dsDNA (Figure 3A). As the double-stranded PCR product accumulates during cycling, more dye can bind and emit fluorescence. Thus, the fluorescence intensity increases proportionally to dsDNA concentration (43). This technique is very flexible because one dye can be used for different gene assays. Consequently, multiplexing reactions is not possible. Because DNA binding dyes do not bind in a sequence-specific manner, these assays are prone to false positives (44). Accurate results demand a specific PCR, which can be confirmed via dissociation curve analysis, where the presence of different PCR products is reflected in the number of first-derivative melting peaks (45) or gel analysis (46). A protocol for SYBR Green I PCR master mix can be found in Ramos-Payen et al. (47).

Hybridization Probes

Hybridization probes can be utilized in either a four or three oligonucleotide manner (for a short review, see Reference 48) (Figure 3B). The four oligonucleotide method consists of two PCR primers and two sequence-specific probes that bind adjacent to each other in a head-to-tail arrangement. The upstream probe is labeled with an acceptor dye on the 3' end, and the downstream probe with a donor dye on the 5' end (49), allowing the donor and acceptor fluorophores to experience an increase in fluorescence resonance energy transfer (FRET) when bound (48). The three oligonucleotide method is similar to the four oligonucleotide method, except that the upstream PCR primer is labeled with an acceptor dye on the 3' end, and thus replaces the function of one of the probes from the four oligonucleotide method.

In both cases, the downstream probe can be designed to cover a mutation site and discriminate between known alleles and detect new alleles simultaneously (50). Alleles are identified and differentiated via dissociation curve (48). A single melting curve can distinguish up to four different T_m values, and six differentially labeled probes may be multiplexed, theoretically allowing a run of 24 assays in a single tube (48). While multiplex reactions are theoretically a simple way to increase the efficiency of data collection, in reality it is a very technically challenging process that requires extensive optimization to ensure that reactions do not compete with each other (34).

Hydrolysis Probes

Hydrolysis probes, exemplified by the TaqMan chemistry, also known as 5' nuclease assay, fluoresce upon probe hydrolysis to detect PCR product accumulation (Figure 3C). The sequence-specific probe is labeled with a reporter dye on the 5' end and a quencher dye on the 3' end (24), which allows the quencher to reduce the reporter fluorescence intensity by FRET when the probe is intact (51). While both hydrolysis and hybridization probes rely on FRET to alter the intensity of fluorescence emission, the energy transfer works in opposite manners in these two chemistries. FRET reduces fluorescence intensity in hydrolysis probes and increases intensity in hybridization probes.

When annealed to the target sequence, the bound and quenched probe will be degraded by the DNA polymerase's 5' nuclease activity during the extension step of the PCR. Probe degradation allows for separation of the reporter from the quencher dye, resulting in increased fluorescence emission (2,24).

Minor groove binders (MGBs), such as dihydrocyclopentpyrroloindole tripeptide (DPI), may be added to these probes to increase their T_m and allow the use of a shorter probe (52). These probes are not only less expensive to produce but have reduced background fluorescence and a larger dynamic range due to increased efficiency of reporter quenching (52).

Hairpin Probes

Molecular beacons. Consisting of a sequence-specific region (loop region) flanked by two inverted repeats, molecular beacons are the simplest hairpin probe (Figure 3D) (53). Reporter and quencher dyes are attached to each end of the molecule, causing a reduction in fluorescence emission via contact quenching (FRET) when the beacon is in hairpin formation (free in solution). When bound to
the target, the quencher and reporter are separated, allowing reporter emission. Hairpin probes tend to have greater specificity than linear probes because the probe-target complex must be thermodynamically more stable than the hairpin structure itself (54), a property often exploited for allele discrimination (55). To increase fluorescence emission, “wavelength-shifting molecular beacons” have been developed, which fluoresce in a number of colors from a single monochromatic light source (56).

**Scorpions.** Scorpions combine the detection probe with the upstream PCR primer (Figure 3E) (57) and consist of a fluorophore on the 5’ end, followed by a complementary stem-loop structure (also containing the specific probe sequence), quencher dye, DNA polymerase blocker (a nonamplifiable monomer that prevents DNA polymerase extension), and finally a PCR primer on the 3’ end. The probe sequence contained within the hairpin allows the scorpion to anneal to the template strand, which separates the quencher for the fluorophore and results in increased fluorescence. Because sequence-specific priming and probing is a unimolecular event, scorpions perform better than bimolecular methods under conditions of rapid cycling such as the LightCycler (58). Cycling is performed at a temperature optimal for DNA polymerase activity instead of the reduced temperature necessary for the 5’ nuclease assay. Scorpions are specific enough for allele discrimination and may be multiplexed easily (58).

The scorpion chemistry has been improved with the creation of duplex scorpions in which the reporter dye/probe and quencher fragment are on separate complementary molecules (59). The duplex scorpions still bind in a unimolecular event, but because the reporter and quenchers are on separate molecules, they yield greater signal intensity because the reporter and quencher can separate completely.

**Sunrise™ primers.** Created by Oncor (Gaithersburg, MD, USA), Sunrise primers are similar to scorpions in that they combine both the PCR primer and detection mechanism in the same molecule (Figure 3F) (60). These probes consist of a dual-labeled (reporter and quencher fluorophores) hairpin loop on the 5’ end, with the 3’ end acting as the PCR primer. When unbound, the hairpin is intact, causing reporter quenching via FRET. Upon integration into the newly formed PCR product, the reporter and quencher are held far enough apart to allow reporter emission.

**LUX™ fluorogenic primers.** Light upon extension (LUX) primers (Invitrogen, Carlsbad, CA, USA) are self-quenched single-fluorophore labeled primers almost identical to Sunrise primers (Figure 3G). However, rather than using a quencher fluorophore, the secondary structure of the 3′ end reduces initial fluorescence to a minimal amount (61). Because this chemistry does not require a quencher dye, it is much less expensive than dual-labeled probes. While this system relies on only two oligonucleotides for specificity, unlike the SYBR Green I platform in which a dissociation curve is used to detect erroneous amplification, no such convenient detection exists for the LUX platform. Agarose gels must be run to ensure the presence of a single PCR product, a step that is extremely important not only for the LUX primers but also for the Sunrise primers and scorpions because PCR priming and probe binding are not independent in these chemistries.

**Causes of Variation**

In theory, PCR is quite robust and predictable, but in actuality, minor variations in reaction components, thermal cycling conditions, and mispriming events during the early stages of the reaction can lead to large changes in the overall amount of amplified product (11,62). Due to the high sensitivity of the real-time PCR assay and the numerous steps that may introduce experimental error, awareness of the causes of variation help produce the most accurate data possible.

Whether using a one- or two-step process, cDNA synthesis can greatly affect the overall real-time PCR results. Both reverse transcriptase enzyme and dithiothreitol (DTT) are PCR inhibitors that may affect reaction kinetics in a one-step process or when carried over during a two-step reaction (18,46). In addition, many samples from complex biological sources often have other PCR inhibitors that may be carried over during sample preparation (63). Inhibitor carryover can be avoided using a cDNA precipitation protocol (18), while DTT may be omitted from the reaction (24).

The oligonucleotides used for reverse transcription priming affect overall cDNA levels. Gene-specific primers yield the most efficient reaction, oligo(dT) primers have an intermediate efficiency, and random hexamers are the least efficient (46). Gene-specific priming is often not ideal because one cannot assay both a target and a normalization gene from the same cDNA template, while with oligo(dT) priming, one may not effectively transcribe the 5′ end of long transcripts. The use of random and specific hexamers has been reported to overestimate mRNA copy number up to 19-fold and 4-fold, respectively, in comparison to 22-mer gene-specific primers (64). Consequently, one solution is to use a mixture of both oligo(dT) and random hexamer primers during the reverse transcription reaction.

The structure and concentration of the RNA template and the reverse transcriptase enzyme itself are other sources of variation during cDNA synthesis. RNA secondary structure and protein complexes present on the target RNA can interfere with the reaction by causing enzyme pausing, dissociation, or skipping over looped regions (18). Raising reaction temperature above 47°C may minimize this problem (65). Different reverse transcriptase enzymes have differing abilities to read through secondary structure (66). For example, SuperScript™ RT II (Invitrogen) has greater efficiency and accuracy than Sensiscript® (Qiagen, Valencia, CA, USA) (34).

As mentioned in the Normalization section, the biological sample itself is often a source of much variation. In cases where whole tissue is assayed, measuring several different cell types within a single sample yields an average expression value of the different cell types. Techniques such as laser-capture microdissection (LCM) may be utilized to extract a pure subpopulation of cells from a heterogeneous source (67).

Variation during PCR can be
incurred from several sources including assay design, PCR reagents, PCR equipment, and human error. Assay design, particularly primer stability and specificity as well as PCR product size, is crucial for an accurate result because amplification efficiency can greatly affect overall results (22). When using a block thermal cycler versus capillary tubes, it is important to measure any positional effects because slight variations in temperature when measuring fluorescence can lead to a variation in the amount detected, especially when using a DNA binding dye. If a service contract is used to maintain the real-time PCR machine, these effects are usually monitored as part of the routine maintenance. Variation in annealing temperature can also affect the enzymatic ability of the polymerase, primer binding, and formation or melting of secondary structure, etc., all of which have compounding effects on the overall PCR.

Variation can occur from the PCR reagents even when using premade master mixes from the same manufacturer. Bustin (34) reported a significant Ct value difference from a single template assayed with two different batches of the TaqMan EZ RT-PCR system (a one enzyme/tube system; Applied Biosystems) master mix that translated into a 2.5-fold difference in median mRNA copy number. Different probe lots synthesized within 6 months of each other also generated significant differences in Ct value, resulting in a 7-fold difference in mRNA copy number (34). Probes manufactured from different sources vary in stability. Bustin (34) reported that Applied Biosystems produces the most stable probes.

Nevertheless, the most likely source of variation is the person performing the experiment (34). Three different people used the same micropipets, master mix, laboratory, template, and machine (ABI Prism 7700; Applied Biosystems) to quantify the same target and found initial copy numbers ranging from $8.7 \times 10^3$ to $2.7 \times 10^3$. Even the most careful pipetting technique may have a 1% relative error. With a 10-fold dilution, this original error will result in a 1% error in amplification efficiency (30). Consequently, precision pipetting and pipet calibration are also essential for preventing cumulative error. Running a standard curve during every reaction can help alleviate this issue because the standard will be affected to the same extent as the unknowns. Using the same batch of enzymes, buffers, master mixes, pipets, and especially the same person will all help reduce experimental variability.

**Calculating Variation**

Because experimental variation is unavoidable, it is important to validate assay results by measuring intra- and inter-assay variation. Variation should not be calculated using Ct values because these are logarithmic units and will misrepresent true variability (8). Therefore, data used for calculation must be a linear value (such as copy number) to obtain accurate measurements of coefficients of variation.

Intra-assay variation quantifies the amount of error seen within a single assay when the same template is run multiple times on the same plate with the same reagents. Intra-assay variation can be calculated for every single sample of every reaction if the real-time PCR experiments are performed in triplicate, with a pooled variance for all sets of PCR triplicates representing statistical power (41). This variation is thought to be both primer and template dependent, with lower concentrations of starting template tending to have higher intra-assay variability. PCR reproducibility is influenced by distribution statistics and stochastic effects (Poisson’s Law; Reference 25). However, several reports have found no correlation between initial template copy number and overall variability (7).

Inter-assay variation should be quantified in cases where comparisons are made of results from two separate assays run on either the same or different days. Variation can be measured by running the same sample on every plate used during a single experiment. This calculation may often be performed using data from either a calibrator or standard sample because these are often already included on all plates.

**CONCLUSION**

Given the number of choices available for every aspect of real-time PCR, it may be difficult to determine what detection chemistry, quantitation method, normalization gene, etc., to use. Although every experimental situation is unique and requires specialized consideration, some general guidelines can be suggested. In terms of quantitation method (absolute versus relative), the majority of users will not require absolute data such as copy number of transcripts or nanograms of DNA, and therefore, relative quantitation will suffice. As discussed, there are many mathematical models available for relative quantitation. Larger projects would benefit greatly by using a method with an associated Excel worksheet such as Pfaffl (26), QGene (23), or DART-PCR (30). While amplification efficiency may be more efficiently calculated from raw fluorescence data instead of a standard curve, using a set of serial dilutions is recommended not only to check the dynamic range of the assay but also to ensure the accuracy of the quantitation. In addition, inclusion of a standard curve would allow results to be calculated using any of the relative quantitation methods available.

The choice of detection chemistry is highly dependent on the characteristics of an individual experiment. During the validation of microarray results, which tends to have only a few samples and several target genes, it is reasonable to use a DNA binding dye. However, in situations where it may be difficult to design a specific PCR (perhaps due to the presence of processed pseudogenes), a sequence-specific probe-based method would have increased reaction specificity. Of the many probe-based techniques available, a well-established system such as the hybridization TaqMan probes may be the best choice. This system has very well-written guidelines and protocols and is fairly error-proof when designed and run according to protocol.

In terms of normalization, the use of multiple housekeeping genes is the most accurate method. Nevertheless, when one has only a few genes to assay or a sample set with low diversity (such
as cell culture), it may not be feasible to run multiple housekeeping genes. If a single gene is used, its stability should be validated in an assay similar to the one used to rank gene stability in geNorm.

Because real-time PCR is now a common method for measuring gene expression, it is increasingly important for users to be aware of the numerous choices available in all aspects of this technology. Unlike traditional PCR, there are many complexities with real-time PCR that can affect overall results. However, with a well-designed experiment performed with the proper controls, real-time PCR can be one of the most sensitive, efficient, fast, and reproducible methods of measuring gene expression.

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COMPETING INTERESTS STATEMENT

The authors declare no competing interests.

REFERENCES

real-time PCR. Nucleic Acids Res. 30:e36.
40a. Technical Bulletin #151. Use of Internal and External Standards or Reference RNAs for Accurate Quantitation of RNA Levels. Ambion, Austin, TX, USA.
46. Lekanne Deprez, R.H., A.C. Fijnvandraat, J.M. Ruljter, and A.F. Moorman. 2002. Sensitivity and accuracy of quantitative real-time polymerase chain reaction using SYBR green I depends on cDNA synthesis condi-

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