

ADVANCES IN REAL-TIME PCR: APPLICATION TO CLINICAL LABORATORY DIAGNOSTICS

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1. Abstract

The polymerase chain reaction (PCR) has become one of the most important tools in molecular diagnostics, providing exquisite sensitivity and specificity for detection of nucleic acid targets. Real-time monitoring of PCR has simplified and accelerated PCR laboratory procedures and has increased information obtained from specimens including routine quantification and differentiation of amplification products. Clinical diagnostic applications and uses of real-time PCR are growing exponentially, real-time PCR is rapidly replacing traditional PCR, and new diagnostic uses likely will emerge. This review analyzes the scope of present and potential future clinical diagnostic applications of this powerful technique. Critical discussions focus on basic concepts, variations, data analysis, instrument platforms, signal detection formats, sample collection, assay design, and execution of real-time PCR.

2. Introduction

The invention of the polymerase chain reaction (PCR) by Kary B. Mullis in the 1980s [1–3] has been one of the formative events in molecular biology. Its influence is comparable to the discovery of the double-helix structure of DNA or to DNA cloning. Myriads of permutations of this elegant approach to sequence-specific and *in vitro* exponential DNA replication have opened new avenues of scientific inquiry in biology and have revolutionized molecular biology and medical diagnosis. Clinical laboratory diagnosis was one of the first fields to embrace PCR methodology, mainly because the exquisite sensitivity and specificity of nucleic acid detection by PCR allowed convenient detection of microorganisms that were difficult to cultivate [4, 5] or to confirm by analysis of their products [6, 7]. Subsequently, the concept of reverse transcription (RT) of RNA to cDNA prior to amplification was introduced as RT-PCR that allowed sensitive detection of RNA viruses and eukaryotic mRNA, greatly expanding the use of PCR. Furthermore, the advantage of facile sequence differentiation of amplified nucleic acids was used for genetic typing of microorganisms [8–10] as well as of mammalian gene polymorphisms such as those of histocompatibility antigens in transplantation medicine [11–13]. Next, drawbacks of the initial approaches of PCR were recognized such as cumbersome multistep procedures of amplification and subsequent product detection by electrophoresis. The strong operator influence in classic PCR methodology frequently gave rise to false-positive results by product carryover and to false-negative results by PCR inhibition, mostly due to incomplete nucleic acid purification [14]. The shortcomings prompted a flurry

of technological innovations of PCR chemistry and instrumentation, culminating in closed-tube methodologies with continuous detection of the amplification products, also termed real-time PCR. These mature technologies now allow quantification of any nucleic acid target in addition to detection and differentiation, because kinetic information about product accumulation is collected. A multitude of real-time PCR platform approaches is presently transforming nucleic acid-based diagnosis in the clinical laboratory. This review will focus on established and generally accepted standard methods of real-time PCR and on aspects that critically influence diagnostic results. Methods for detection of low nucleic acids target copy numbers (typically of microorganisms or mammalian mRNAs) will preferentially highlight the status of the field, since such methods generally are the most demanding applications of PCR. Nevertheless, such technical considerations apply equally to high-copy targets such as in PCRs that identify mammalian genetic polymorphisms. The challenge in these PCRs is the extraction of maximum information from a single specimen by simultaneous amplification of multiple target sequences, also called multiplex PCR [15, 16].

3. Real-time PCR: Concept, Variations, and Data Analysis

3.1. REAL-TIME PCR VERSUS TRADITIONAL PCR

Real-time PCR was first introduced by Higuchi *et al.* to analyze the kinetics of PCR by constructing a system to detect PCR products during the process of their amplification [17, 18]. In this real-time system, the fluorescent dye ethidium bromide intercalates preferentially into double-stranded amplification products, which increases its fluorescence. The progress of PCR amplification can be continuously monitored in real time by acquiring fluorescence signals in each amplification reaction cycle during segments in which double-stranded DNA (dsDNA) is present. Thus, quantitative information of the PCR process is obtained by plotting the intensity of the fluorescence signal versus cycle number. This is in contrast to the traditional PCR approach in which only qualitative information is obtained by detection of presence or absence of a specific dsDNA product by gel electrophoresis at the end of amplification.

Direct product monitoring in real-time PCR also has the advantage that reaction vessels do not need to be reopened after PCR. This minimizes post-amplification processing, and importantly, the potential for sample contamination with products of previous amplifications (product carryover [19]). Aside from the increased information that can be extracted from specimens, the ease of sample processing and prevention of false-positive results has

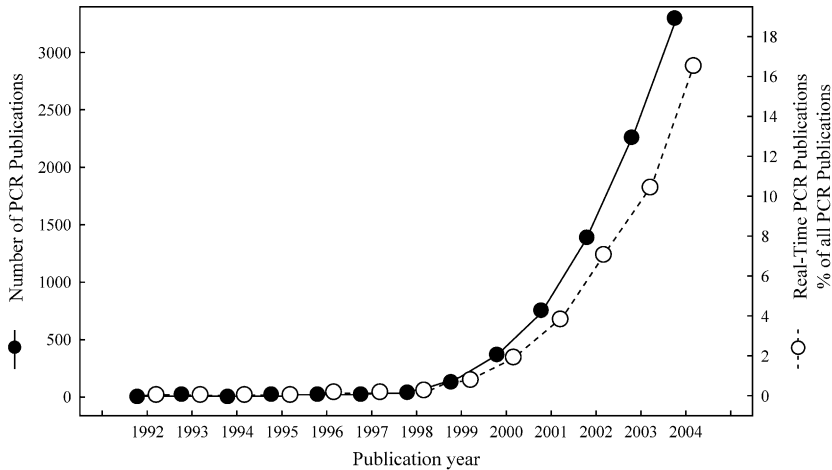


FIG. 1. Annual numbers of PCR publications (left ordinate) and percentage of real-time PCR publications among PCR publications (right ordinate). The numbers of publications citing PCR or real-time PCR in the title or in keywords were obtained by using the search terms “PCR” and “real-time PCR” in PubMed (Accessed December 2004).

been one of the main incentives for introduction of real-time PCR to clinical diagnostic laboratories [6]. Real-time PCR has been rapidly applied to many areas, such as detecting and quantifying DNA and RNA of microbes [5, 20, 21], genotyping microbes and parasites [8, 22, 23], monitoring the transcription of genes by quantifying mRNA concentrations relative to constantly expressed genes (housekeeping or reference genes) [24–26], and typing of genetic polymorphisms [11–13]. The profound impact of PCR and real-time PCR becomes obvious in the numbers of publications with the term “PCR” in the title or keywords. Similar to product accumulation in PCR, the number of PCR publications has increased exponentially since its invention, and among PCR publications the percentage of publications citing real-time PCR also shows a logarithmic increase (Fig. 1). These data suggest that real-time PCR methods will replace a large percentage of traditional PCR assays within the next five years.

3.2. VARIATIONS OF REAL-TIME PCR

A major variation between real-time PCR methods results from the amplification target, DNA versus RNA. All RNA targets require an initial reverse transcription step, using either random oligonucleotide, oligo(dT), or target sequence-specific priming [27]. Random oligonucleotide and oligo(dT)

priming occurs in a separate step prior to PCR, and multiple real-time PCRs are typically performed with such samples (two-step PCR). Target-specific priming of reverse transcription may occur as an initial thermal cycling step in the PCR vessel (one-step PCR) or prior to PCR (two-step PCR). Collectively, PCR methods for amplification from original RNA targets are termed real-time reverse transcription PCR (real-time RT-PCR).

Another difference between real-time PCR methods is the number of different target nucleic acids amplified and/or detected per reaction [15, 16]. Reactions that simultaneously amplify (or detect) two or more target fragments (or detect sequence polymorphisms within target fragments) are termed duplex and multiplex real-time PCR, respectively.

Detection of the amplification products represents yet another source of variation between real-time PCR methods. To date, only light emission (fluorescence) has been used to monitor the PCR process. The source of fluorescence may be a fluorescent dye such as ethidium bromide or SYBR[®] Green I that changes fluorescence emission upon binding to dsDNA [28, 29]. Another approach is the amplicon-specific detection of fluorescence, in which the light emission of a fluorescent dye coupled to an oligonucleotide changes after hybridization of the oligonucleotide to the amplification product [30–32].

3.3. QUANTITATIVE ANALYSIS OF REAL-TIME PCR DATA

The kinetics of target copy numbers in a PCR, as visualized in a plot of fluorescence versus cycle number, follow a sigmoidal curve. The signal from early target amplification remains below the background noise. When enough amplicon is present to generate a signal exceeding the threshold baseline of background noise, an exponential increase in signal (and target numbers) ensues that typically lasts for 4–8 cycles [33]. This exponential phase of the amplification generates a line in a plot of the logarithm of fluorescence versus cycle number (Fig. 2A). As *Taq* DNA polymerase becomes the limiting factor and self reannealing of amplified DNA outcompetes primer annealing, the PCR efficiency falls from the almost 2-fold amplification in the exponential phase to 1, and product numbers and fluorescent signal reach a plateau phase without any more increase.

3.3.1. Absolute Quantification

Product detection in traditional PCR is limited to the end product in the plateau phase, which eventually is achieved in any positive PCR irrespective of the number of starting templates. Thus, quantitative information about input targets cannot be retrieved from traditional PCR. In contrast, in real-time PCR the fractional threshold cycle number (C_t) is linearly inversely related to the logarithm of starting templates [34]. Thus, the fewer starting

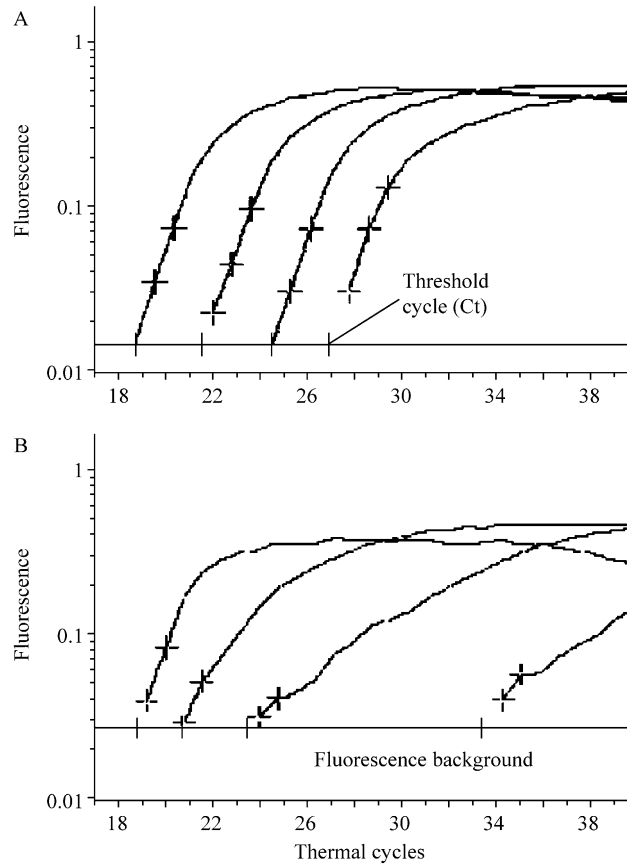


FIG. 2. The kinetics of real-time PCR. Panels A and B show the real-time PCR amplification of a segment of the 23s rRNA gene of *Chlamydia pneumoniae* from 7-fold template dilutions [76]. PCRs of panel A were performed at high-stringency annealing at 56 °C, those of panel B at low-stringency annealing at 48 °C. An amount of 10^5 starting templates was used at the highest concentration (leftmost curve) in both PCR series. A, at high stringency, amplification in the linear exponential phase is close to 2-fold per cycle (efficiency ≈ 2), and approximately 2.8 additional cycles are required to generate the threshold signal for each subsequent 1:7 template dilution ($2^{2.8} \approx 7$). Parallel slopes of the linear phase at different starting template numbers indicate equal efficiency of the PCRs. B, low stringency annealing caused extensive formation of aberrant amplification products and reduced efficiency during late PCR cycles at low copy numbers. This is evident in decreasing slopes of the linear phase for decreasing starting template numbers, and resulted in progressively increasing Ct values for higher dilutions of the template.

templates present, the more cycles are required for a fluorescent signal that exceeds in the linear phase of a log-linear fluorescence plot the background or an arbitrary cut-off fluorescence level (Fig. 2A).

For determination of absolute numbers of starting templates, the C_t values of individual unknown samples are compared to the C_t values of a series of quantitative standard PCRs. A linear regression equation of the logarithm of standard template numbers and the corresponding C_t values calculates the starting templates of an unknown sample from its C_t value [33–35]. This calculation assumes equal and constant efficiency of all standard and unknown sample PCRs. Figure 2B clearly demonstrates that poor adjustment of a single parameter such as annealing temperature can result in poor PCR performance and negate the requirement of equal efficiency as evident in different slopes of the linear portion of the fluorescence plot for different starting templates (linear portions are not parallel). Such unequal PCR efficiencies prevent correct quantitative interpretation of real-time PCR data [33, 36].

To remedy this problem, Ramakers *et al.* [37] present an algorithm for assumption-free analysis of real-time PCR data. They calculate the efficiency of each individual PCR from a “window-of-linearity” of the best linear fit and the steepest slope of the log (fluorescence) of data acquired in four consecutive cycles. Using individual C_t values, they can determine for each PCR the starting templates relative to an arbitrary copy number at threshold fluorescence without a need for calibration standards. Others have reported similar approaches to calculate PCR efficiency from the linear phase of the log-linear fluorescence plot [38–40].

However, the convex plots of the high-dilution samples in Fig. 2B suggest that the efficiency of a suboptimal PCR is not necessarily constant prior to the plateau phase, but declines even during the “linear,” exponential phase of amplification, that is, a “window-of-linearity” does not exist for all samples [41]. A nonlinear fit of fluorescence versus C_t would reflect the change in efficiency with cycle number, thus allowing more accurate deduction of efficiency at each PCR cycle, and potentially, more accurately calculating overall PCR efficiency in an assumption-free algorithm.

Such a nonlinear approach was pioneered by Liu and Saint [41] who showed that a sigmoid function modeled PCR amplification more effectively than the exponential model upon which the threshold method is based. Rutledge [42] most recently presented a refined sigmoidal approach in which an optimal cut-off cycle prior to the variable plateau phase was mathematically identified and the fluorescence data prior to this cut-off point were fitted with highest precision to a four-parameter sigmoid function (r^2 of individual reactions typically 0.99992). In this function, amplification efficiency decreases continuously throughout the amplification process and each cycle has a unique amplification efficiency. Relative quantification can thus be achieved without standard curves and prior knowledge of amplification

efficiency by modeling individual amplification reactions. A single calibration standard can convert such relative data into absolute copy numbers. This method has the potential to redefine analysis of real-time PCR by allowing for reliable, fully automated quantification of nucleic acids. This method would also be insensitive to variations in amplification efficiency between individual reactions. Such variations may, for instance, occur by chemical inhibition of PCRs, through competition in multiplex real-time PCRs, or in heat-block real-time PCR instruments.

3.3.2. *Relative Quantification of mRNAs*

Good precision in absolute quantification by real-time PCR is readily achievable if PCR protocols are well designed, PCR efficiencies are carefully monitored, and invariant controls are used [33]. In real-time RT-PCR quantification of analyte mRNAs relative to the mRNA of an internal reference gene, the problems are compounded by combining results of two real-time PCRs for each sample. These values are usually related to the corresponding values of a calibration sample or to an absolute standard. Levels of starting templates can again be expressed in absolute terms or relative to the reference mRNA in the same sample. Problems of the quantification accuracy of RT-PCR have been recognized [43, 44] and mathematical approaches have been published for calculations of relative mRNA levels. The methods assume either equal efficiencies of analyte and reference mRNA RT-PCR [45], or do not assume equal efficiencies, but require constant efficiency of analyte and reference RT-PCR [46]. However, it is important to keep in mind that Ct values, and typically used differences between Ct values of analyte and reference mRNA (ΔCt), are exponentially related to the numbers of target copies. Thus, small errors in ΔCt may become large errors in the ratios. In a simulation with different PCR efficiencies, a real 10-fold difference in mRNA levels was experimentally determined as any value between 0.7 and 210 [37]. These data underscore the extreme importance of tight control of all parameters and rigorous validation of real-time PCR protocols [33]. One-step duplex RT-PCR for simultaneous amplification of analyte and reference gene mRNA has the potential to optimally control variation in experimental parameters [26]. Deduction of starting templates by sigmoidal curve-fitting likely will maximize the accuracy of results [42] and use of a single target DNA standard (without reverse transcription [26]) in this approach likely will allow calculation of absolute starting targets in RT-PCRs.

3.3.3. *Determination of T_M by Melting Curve Analysis*

The melting point of the amplification DNA fragment (T_M , temperature at which a given DNA fragment exists as 50% each double- and single-stranded DNA) can be determined after completion of real-time PCRs that use

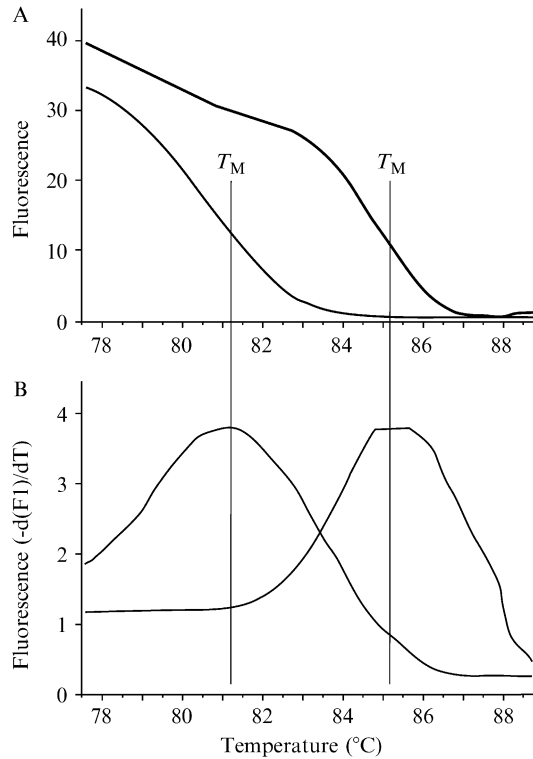


FIG. 3. Melting curve analysis for T_M determination. SYBR Green I real-time PCRs of fragments of the *ompA* and 23S rRNA genes of *C. pneumoniae* [5, 76]. A, after completion of the real-time PCRs, the reactions were cooled below the expected T_M of the dsDNAs and then slowly reheated while fluorescence was continuously measured. B, the negative first derivative of fluorescence over temperature produces a peak at the point of highest change in fluorescence, indicating the melting transition (T_M) of dsDNA. The lower T_M of the left peak reflects the lower GC content of the *ompA* fragment. In a similar approach for the T_M determination of oligonucleotide probes, shifts due to nucleotide mismatches are highly reproducible and can be effectively used for typing of nucleotide sequence polymorphisms.

dsDNA-binding dyes [47]. Fluorescence is continuously acquired during a slow temperature increase from $\sim 65^\circ\text{C}$ to 90°C . The negative first derivative of fluorescence over temperature shows a peak at the temperature of steepest decline in fluorescence (Fig. 3). The temperature at peak is equivalent to the T_M . Similarly, the T_M of oligonucleotide probes can be determined in an analogous way in PCRs using oligonucleotide, but not hydrolysis (*Taqman*[®]) probes. Hybridization of oligonucleotides is highly sensitive to destabilization by nucleotide mismatch and changes in the T_M as determined

by melting curve analysis can be very effectively used for typing of sequence polymorphisms [22, 48, 49].

4. Real-Time PCR Platforms

Market introduction of real-time PCR instrumentation was critical in enabling users to convert traditional PCR methods to real-time PCR, because this complex method can be performed only with sophisticated equipment. Three main concepts for temperature control of reaction vessels are currently in use for the critical thermal cycling component of real-time PCR instruments: (i) traditional heat blocks [50–54]; (ii) heated air [28, 32, 55–58]; and (iii) independent electronic temperature control for each individual reaction tube [59, 60]. Continuously updated information on the latest versions of real-time PCR platforms and many pertinent issues is available on the Gene Quantification website [61].

Heat block instruments are derived from classic PCR equipment with an added optical component for fluorescence acquisition. An advantage is that they accommodate samples in the 96-well format. Historically, heat block instruments tended to have a shortcoming in the uneven temperature distribution across the heat block [62, 63]. Uneven reaction temperature potentially presents a major problem because the accuracy of real-time PCR strongly depends on absolutely uniform temperature ramping and equilibrium temperature of all samples. Even minor temperature differences at each cycle may change PCR efficiency between samples and thus may result in widely divergent results [37, 63]. The profound effect of differences in annealing temperature on real-time PCR accuracy is clearly evident in Fig. 2B. However, the problem of temperature uniformity has been recognized and most current heat-block PCR instruments claim temperature uniformity within $\pm 0.5^\circ\text{C}$ that ensure reproducible and consistent results. Nevertheless, it is highly recommended to carefully evaluate the reproducibility of real-time PCR methods in a new instrument (63).

Real-time PCR instruments with heated air or independent electronic temperature control for reaction tubes are second generation instruments that address the problem of uneven temperature distribution very effectively, thus have the potential to yield highly accurate real-time PCR results. These real-time PCR platforms have the advantage of small reaction volumes and rapid temperature transfer and equilibration. The result is increased PCR specificity and short cycle times, therefore these systems are sometimes termed rapid-cycle PCRs.

Heated air instruments [55–57] rotate samples continuously at low speed to ensure temperature uniformity [63]. These instruments use special custom

reaction capillaries [56, 57] or standard polypropylene tubes [55] and allow more rapid cycling than heat block instruments. The principal disadvantage is that they do not seamlessly accept the 96-well format.

Novel concepts for tight temperature control and rapid cycling have been implemented in two real-time PCR platforms [59, 60]. The Cepheid Smart-Cycler[®] uses a heated fluorometer module for each individual reaction tube. Each module contains a solid state heating thermistor with forced-air cooling. The InSyte[®] thermal cycler uses composite tubes with integrated heating polymer and controls the temperature of individual reaction vessels by use of a noncontact infrared thermometer.

Virtually all instruments, particularly second generation platforms, provide single or multiple excitation wavelengths and support multiple emission wavelengths, and thus multiplex PCRs. Software formats differ between platforms, but all allow direct determination of Ct values, and offer algorithms for conversion of these values into starting copy numbers without exporting the data.

In the choice of a real-time PCR platform, a potential user needs to first ascertain the main use of the instrument and determine the requirements for accuracy of quantification, sensitivity of detection of low-copy targets, or typing of genetic polymorphisms of high-copy target DNA. Second, multi-color detection channels, anticipated sample throughput, and requirements for automation need to be considered. Careful evaluation of requirements for the real-time PCR assays and of advantages and disadvantages of each platform are important.

5. Signal Detection Formats

Presently only fluorescent light, with four main strategies, is used to detect PCR amplification products in real-time PCR. The initial and simplest real-time PCR method uses fluorescence dyes that specifically bind to dsDNA and that change their fluorescent behavior after dsDNA binding. The other three methods, hydrolysis probes, molecular beacon probes, and hybridization probes, rely on a change in fluorescence after the hybridization of fluorescently labeled oligonucleotides to the target amplicon. Detection with oligonucleotide probes adds another layer of specificity to the specificity of PCR primers. Well designed real-time PCR methods using probes to generate the fluorescent signal tend to be highly specific, with little or no background signal in negative samples. It is important to note that dual-labeled oligonucleotides such as hydrolysis probes, molecular beacons, or sunrise and scorpion primers are more expensive than single-labeled probes and require careful optimization of their hybridization properties. However,

once appropriate PCR conditions have been identified, dual-labeled probes provide reliable results.

5.1. DNA-BINDING FLUOROPHORES

The first dye used as DNA-binding fluorophore was ethidium bromide [17, 28], and other dyes such as YO-PRO-1 and BEBO have been also used [64–66]. SYBR Green I is now the most frequently used dsDNA-specific dye in real-time PCR. It is a cyanine dye and has an asymmetric structure related to the dsDNA-specific dyes YOYO-1 and TOTO-1. The binding affinity of SYBR Green I to dsDNA is more than 100 times higher than that of ethidium bromide [28, 29]. Intercalation of SYBR Green I does not affect the process of PCR amplification as ethidium bromide does, since SYBR Green I mainly binds to the minor groove of dsDNA. Unbound SYBR Green I exhibits little fluorescence in solution and fluorescence emitted by dsDNA-bound dye is more than 1000-fold higher than that of free dye, therefore this dye is ideally suited for real-time monitoring of PCR amplification. SYBR Green I can be excited at a wavelength of 480 nm and peak fluorescence emission is at 520 nm. In an optimized SYBR Green I real-time PCR, starting templates as low as a single target copy can be detected [5]. In some SYBR Green I real-time PCRs, the accumulation of primer dimers and other nonspecific amplification products may reduce specific amplification. High temperature fluorescence acquisition, just below the melting point of the specific amplification product, but above the melting point of primer dimers, may remedy this problem by collecting only specific signals.

5.2. 5'-NUCLEASE HYDROLYSIS OLIGONUCLEOTIDE PROBES

A hydrolysis probe, also termed *Taqman*[®] probe, is an oligonucleotide labeled with a 5'-terminal reporter dye and a 3'-terminal fluorescence quencher [67]. In solution, unbound intact probes do not emit fluorescence because the emission of the reporter dye is suppressed by the 3'-terminal quencher. During PCR amplification, primers and probes hybridize to the cognate nucleotide sequence of the amplicon. When *Taq* DNA polymerase, extending the 3'-end of the primer, reaches the bound probe, the dsDNA-specific 5'-exonuclease activity of *Taq* polymerase will degrade the probe [30]. Degradation of the probe separates reporter dye and quencher and the reporter dye will emit unquenched fluorescence after excitation [68]. The strongest fluorescence signal is observed when the two labels are at the extreme terminals of the oligonucleotide probe, probably because a single hydrolytic

cleavage only of the 5'-nucleotide separates reporter and quencher and releases the signal. The T_M of hydrolysis probes is critical: binding to the target sequence needs to be tight in order to ensure hydrolysis of the probe rather than displacement by *Taq* polymerase. At a probe T_M of 70°C, performance of the PCR annealing and extension step at 60–62°C will ensure probe binding during primer extension and thus probe hydrolysis and generation of the fluorescent signal. The corollary of the mechanism of hydrolysis probes is that the signal generated is cumulative, because separation of reporter dye from quencher is irreversible. This implies that melting curve analysis is impossible with hydrolysis probes.

5.3. FLUORESCENCE RESONANCE ENERGY TRANSFER (FRET) OLIGONUCLEOTIDE PROBES

FRET probes measure the transfer of energy between two fluorophores that are attached to two oligonucleotide probes [69]. These probes are designed to hybridize to adjacent segments of a nucleotide sequence, and hybridization brings the fluorophore at the 3'-end of the donor probe in immediate proximity of the fluorophore at the 5'-end of the acceptor probe, the fundamental requirement for FRET. The donor probe typically is labeled with fluorescein or a similar fluorophore with an excitation peak around 480 nm wavelength. The acceptor probe may be labeled with one of a multitude of dyes with longer-wavelength peak excitation and emission such as cyanine dyes Cy3 and Cy5, TET (6-carboxy-4, 7, 2', 7'-tetrachloro-fluorescein), TAMRA (6-carboxy-N, N, N', N'-tetramethylrhodamine), or ROX (6-carboxyrhodamine), and must be blocked at its 3'-end to prevent extension during the PCR. When excited by light of 480 nm wavelength, fluorescein of the donor probe will emit green fluorescent light if the probe is in solution. However, if both donor and acceptor probe are hybridized simultaneously to the cognate nucleotide sequence on the amplification fragment, the activation energy of fluorescein will directly transfer to the acceptor dye, which will emit long-wave fluorescent light. Thus, during the PCR annealing step, FRET results in increased long-wave fluorescence proportional to the amount of PCR product. FRET probes ensure very high specificity of detection, because two probes must simultaneously hybridize to adjacent target sequences. In contrast to hydrolysis probes, FRET probes are not degraded and fluorescence is reversible, allowing the determination of probe T_M in melting curve analysis. Roche Applied Science has developed the fluorophores LightCycler RED-640 and LightCycler RED-705 specifically for use in FRET probes in the LightCycler [32]. However, other Cy5-like

fluorophores with similar spectral properties have been also successfully used [26, 70].

5.4. HAIRPIN OLIGONUCLEOTIDE PROBES: MOLECULAR BEACONS

Molecular beacons are DNA hybridization probes that form a stem-loop structure and are labeled on both ends. The loop portion is complementary to the target sequence [71, 72]. One arm is labeled with a fluorescent reporter and a quencher is attached to the other arm. The quencher is a nonfluorescent chromophore, which absorbs the emission energy from the fluorophore. In solution, quenching of fluorescence occurs because molecular beacons form a hairpin structure in which the stem keeps the arms together, and thus fluorophore and quencher in close proximity. During PCR annealing, the loop will hybridize to the specific target sequence, resulting in a conformational transition which separates the stem, and thus fluorophore and quencher [31]. This leads to the restoration of detectable fluorescence proportional to the amount of specific PCR amplification product. Similar to hydrolysis probes, molecular beacons require careful probe design. The stem must be optimized to ensure a complete stem-loop structure in solution, but effective unfolding and probe hybridization during PCR annealing. Suboptimal probe designs lead to a large background signal if the stem is not stable and to poor signal if probe annealing is unstable. Therefore, PCR annealing temperature and T_M of stem and loop of molecular beacons require careful optimization [71, 72].

5.5. LIGHT-UP PROBES

Light-up probes are peptide nucleic acids (PNAs). They use thiazole orange as the fluorophore, and no quencher is required [73]. PNA molecules have a backbone with peptide-like covalent bonds and exocyclic bases. When light-up probes hybridize with specific target DNA, the resulting duplex or triplex structures elicit increased fluorescence of the fluorophore [74].

5.6. SUNRISE PRIMERS

Sunrise primers have the dual function of PCR primer and generator of a specific fluorescent signal in PCR reactions [75]. They are labeled with a reporter fluorophore and a quencher at the ends, and have a 5'-terminal hairpin structure. The hairpin structure keeps the reporter and quencher in proximity when the primer is in solution. After annealing during PCR, sunrise primers are permanently extended to open the hairpin structure, which results in fluorescence emission.

5.7. SCORPION PRIMERS

Scorpion primers are similar to molecular beacons, but function as primers rather than as probes [76]. A specific probe sequence is held in a hairpin loop configuration by complementary stem sequences at the 5'- and 3'-sides of the probe. A fluorophore attached to the 5'-end is quenched by methyl red at the 3'-end of the stem-loop structure. The hairpin loop is linked to the 5'-end of a primer via a hexethylene glycol moiety which stops DNA synthesis. After extension of the primer during PCR the probe will hybridize to its complement on the same DNA strand. This stops quenching and results in fluorescence as long as the probe is bound in the PCR annealing phase. The hexethylene glycol moiety stops synthesis of a complementary strand beyond the primer, and thus prevents permanent opening of the hairpin loop. Therefore, scorpion primers, unlike sunrise primers, do not produce a fluorescent signal from primer dimers.

6. Design of Real-Time PCR Assays

For reliable quantitative results over a wide range of starting template concentrations, real-time PCR methods must be more robust than traditional PCR assays, that is, the efficiency of specific amplification should decline later than in traditional PCRs and a large amount of specific amplification fragment should be produced (the plateau should be high, Fig. 2A). In the design of such robust real-time PCR assays, it is important to keep in mind that any PCR is a finely tuned mixture of reagents and thermal reaction parameters and minor changes in any component may upset this balance. Components that have a desired effect in one reaction might not have the same effect in another reaction, or might even have a detrimental effect.

Real-time PCR with its strictly quantitative output lends itself to method optimization. Different strategies may be used to optimize a real-time PCR assay by changing the chemical composition as well as thermal cycling parameters. It is important to initially identify reaction conditions that yield reproducible results, typically at high starting template concentrations. An effective approach is then to define, by limiting dilution, the template concentration at which the PCR shows reduced performance (Fig. 2B), and to modify the PCR using this starting template concentration until the performance is reproducibly improved [5, 77]. This process should be repeated until the template becomes limiting, that is, when stochastic presence or absence of single templates (Poisson error) results in either strongly positive or unambiguously negative amplifications in multiple PCRs of the same template concentration.

Changes of different parameters may result in similar improvements of a PCR. Modification of the Mg^{2+} ion concentration, for instance, typically is used to adjust PCR efficiency and yield. For example, the inefficient real-time PCR of Fig. 2B can be substantially improved by reducing Mg^{2+} concentration from 4.5 mM to 3.0 mM. However, an even better improvement can be achieved at 4.5 mM Mg^{2+} by increasing the annealing temperature from 48 °C to 56 °C, as is evident in Fig. 1A. A highly effective approach is also touch-down or step-down thermal cycling, in which cycles of stringent, high-temperature annealing precede the actual real-time PCR with constant thermal conditions during fluorescent signal acquisition [26, 77–79]. Thus, two fundamental approaches may be used in improving a real-time PCR starting from successful, but inefficient amplification: preferential modification of either chemical or of physical (thermal cycling) reaction parameters. As in any scientific experiment, it is critically important to change only one parameter at a time, and to keep the overall reaction chemistry and protocol as simple as possible.

6.1. CHEMICAL COMPOSITION OF REAL-TIME PCR ASSAYS

The basic composition of a real-time PCR is similar to a traditional PCR and consists of a thermostable DNA polymerase, oligonucleotide primers, dsDNA-binding fluorophores or fluorescently labeled oligonucleotide probes, nucleotides, $MgCl_2$, KCl, and Tris-HCl [2, 5, 43, 77]. Amplification facilitators such as bovine serum albumin or nonionic detergents are sometimes added to enhance amplification efficiency. Frequently, uracil-N-glycosylase is added to prevent product carryover contamination of a PCR by degrading products of previous PCRs in which TTP was substituted with dUTP [19, 80, 81].

6.1.1. *Thermostable DNA Polymerases*

The introduction of *Taq* DNA polymerase, the thermostable DNA polymerase from *Thermus aquaticus*, greatly simplified PCR and facilitated its success [2]. *Taq* DNA polymerase and its derivatives are still the most widely used DNA polymerases for traditional as well as real-time PCR. Numerous DNA polymerases from other organisms are also commercially available, such as *Tth* and *rTth* DNA polymerase from *Thermus thermophilus*, or DyNzyme from *T. brockianus*. These polymerases have different resistance to PCR-inhibitory substances and they exhibit different performance depending on sample type [14, 82, 83]. For example, *Taq* DNA and Ampli-Taq[®] Gold polymerases were completely inhibited in the presence of 0.004 % (v/v) blood in the PCR master mixture, while Hot *Tub*, *Pwo*, *rTth*, and *Tfl*

DNA polymerases provided fully efficient amplification in the presence of 20% (v/v) blood [14, 84]. *Tth* and *rTth* polymerases were 10- to 100-fold more effective than *Taq* polymerase in detecting influenza A virus in nasopharyngeal swab samples [85], *Helicobacter hepaticus* in mouse feces [86], or *Clostridium botulinum* in pig feces [87].

At room temperature, *Taq* polymerase shows low activity, but nevertheless extends spuriously paired DNA of primers, palindromes, or loop sequences [88]. Such mispriming is particularly detrimental in RT-PCRs, because cDNA is derived from RNA which has a high frequency of self-complementary, self-folding sequence structures. Inhibition of amplification during the assembly phase of a PCR is highly effective in reducing nonspecific amplification. This approach is termed hot-start PCR, because amplification ensues only after the first heating step. Hot-start PCR has traditionally been achieved by a multitude of approaches. Most frequently, wax has been layered on incompletely assembled reactions, followed by addition of the remaining reaction components [88]. The first heating step then melted the wax, resulting in mixing of all reaction components and initiation of amplification. This cumbersome approach has been virtually completely replaced in real-time PCR by initial inactivation of the DNA polymerase in fully assembled reaction. Inactivation is typically achieved via antibody directed against the active site of the enzyme [89] or via use of a recombinant *Taq* DNA polymerase with a sequence deletion that renders the enzyme inactive without exposure to high temperature [90]. These hot-start methods are highly effective and real-time PCR is mostly performed as hot-start reaction.

6.1.2. PCR Facilitators

Almost all PCR chemistries include one or more reagents that improve amplification efficiency. Five groups of such facilitators have been described [91]: (i) proteins such as bovine serum albumin (BSA) and single-stranded DNA-binding proteins; (ii) nonionic detergents such as Tween 20, Triton-X, or NP-40; (iii) organic solvents such as dimethyl sulfoxide (DMSO); (iv) biologically compatible solutes such as betaine and glycerol; and (v) polymers such as polyethylene glycol (PEG) and dextran. Amplification facilitators affect amplification by modifying the thermal stability of the template or affecting the accuracy of DNA polymerase, and influence overall PCR performance and frequently alleviate inhibition of amplification [82]. For example, glass capillaries used in LightCycler platforms require higher levels of bovine serum albumin than polypropylene reaction tubes [5], presumably to block surface binding of reaction components.

6.2. PRIMERS AND OLIGONUCLEOTIDE PROBES

A large array of computer software is available for primer or probe design. Typically, producers of oligonucleotides and of real-time PCR platforms offer proprietary software that supports design of oligonucleotides for real-time PCR. All programs implement algorithms that identify oligonucleotides with matched T_M , and that avoid incorporation of 3'-complementary sequences and palindromes. Nevertheless, it is important to remember that not all problems with PCRs stem from poor primer design. In many cases, internal sequences of amplification fragments lead to recombination during PCR that causes aberrant products independently of primer influence [26, 92]. Also, limitations in the choice of target sequences may impose restrictions on the design of primers and probes.

The thermodynamic rules for oligonucleotide probe design follow those for primers [93]. Design of FRET probes is simple and requires essentially only the observation of a specific T_M range. Design and optimization of dual-labeled, self-folding probes or primers should follow the suggestions and use software provided by industry, but may be tedious and time consuming.

The optimal amplicon length for real-time PCR is between 100 to 200 base pairs and amplicons ideally should have 50–60% GC content. Amplification of short fragments is more efficient than that of longer fragments and the probability of broken template strands is also lower for short fragments. The 3' end of primers should avoid stretches of 3 or more GC nucleotides, which may cause unspecific priming. To ensure specificity as well as options for adjustment of annealing temperature, primers should be between 20–30 bp, with a calculated melting temperature of at least 60°C, and as high as 72°C if possible. Such primers allow for high, stringent annealing temperatures which typically result in highly specific and efficient real-time PCR assays. Primers and probes for eukaryotic mRNA used in RT-PCR that bridge exons will anneal only to cDNA, but not to genomic DNA. However, in special circumstances, design of exon-bridging oligonucleotides is not feasible such as for single exon genes, genes for which processed pseudogenes exist, or for prokaryotic genes.

6.3. REAL-TIME RT-PCR

Real-time RT-PCR is the gold standard for quantification of low-abundance mRNAs and its wide use is partly responsible for the current exponential growth in the application of real-time PCR. The first step is reverse transcription of RNA to cDNA, followed by the real-time PCR. The reverse transcription may be performed separately from PCR as part of a two-step RT-PCR, or preceding the PCR in a one-step reaction in the PCR

vessel. The advantage of the two-step system is that a pool of cDNA can be created that can be used for quantification of multiple target mRNAs. Conversely, one-step RT-PCR minimizes the number of steps in a protocol and thus the potential for inaccuracies. Two-step real-time PCR typically utilizes random oligonucleotides or oligo(dT), one step RT-PCR predominantly uses the specific PCR primers for priming reverse transcription. Total RNA is used as a template for reverse transcription, but use of poly(A)⁺ RNA increases sensitivity. DNase treatment of extracted nucleic acids is required if the RT-PCR does not discriminate between target genomic DNA and cDNA. The mostly frequently used reverse transcriptases are avian myeloblastosis virus reverse transcriptase (AMV-RT) and moloney murine leukemia virus reverse transcriptase (MMLV-RT) and recombinant engineered versions of these enzymes [94, 95]. MMLV-RT is less robust than AMV-RT, but has lower RNase H activity than AMV-RT. Since RNase H activity interferes with the synthesis of long cDNAs, MMLV-RT is preferred for generating full-length cDNA molecules.

Concentrations of mRNAs found by real-time PCR are generally related to the levels of reference transcripts of housekeeping genes in the same sample. An ideal internal standard mRNA should have constant expression among different samples, or between individuals of the same species. In duplex or multiplex RT-PCR, the concentration of the reference mRNA should be less than 100-fold different from the analyte mRNA, in order to avoid competitive PCR inhibition of the lower copy mRNA. The expression of these internal controls should also be independent of normal physiological and pathological changes. The three most frequently used reference transcripts are ribosomal RNAs (rRNAs), β -actin (ACTB), and glyceraldehyde-3-phosphate dehydrogenase (GAPDH). However, none of these targets is universally suited as reference transcript. rRNAs are abundant, stably expressed, but can be used only in total RNA preparations and the expression levels are very high in comparison to most mRNAs [43, 96, 97]. ACTB is highly expressed, but less abundant than rRNAs, varying expression levels have been reported and the existence of pseudogenes needs to be considered in the design of oligonucleotides [24, 98]. GAPDH is the most problematic of the reference RNAs, with inconstant expression reported [43, 99, 100]. Transcripts of numerous other genes have been used as internal reference transcripts, and Vandesompele *et al.* [25] explore in depth the relevant criteria for their use.

6.4. MULTIPLEX REAL-TIME PCR

Duplex and multiplex real-time PCRs are reactions in which two or more sets of primers and probes simultaneously amplify and detect two or more target fragments. These reactions are possible because probes labeled with fluorescent

dyes with different excitation and emission spectra can be used and discriminated concurrently in real-time PCR platforms. Multiple photodetectors and electronic compensation of overlapping emission spectra ensure specificity and accuracy of data [101]. In theory, multiplex PCR saves time and cost, and reduce the required sample volume. However, multiplex real-time PCR is burdened with a number of inherent technical challenges: (i) all amplifications must be designed for identical reaction parameters; (ii) the parallel amplifications compete with each other; and (iii) the probability for undesirable interaction between amplifications increases with the number of targets in a multiplex PCR. For these reasons, successful multiplex approaches have almost exclusively been used for typing of genetic polymorphisms in real-time PCR with high starting template numbers [4, 102, 103]. Wang *et al.* [26] established a duplex real-time PCR approach on the LightCycler platform for simultaneous quantification of transcripts of a reference and analyte gene in a one-step RT-PCR. This approach is accurate and sensitive, and uses hot-start and step-down cycling and high-temperature annealing between 63°C to 72°C after reverse transcription at 55°C for 20 min.

7. Execution of Real-Time PCR Assays

A major challenge in successful routine use of real-time PCR is the consistent output of high quality results. The previous chapters make it abundantly clear that it is not sufficient to obtain positive amplification results. Vaerman *et al.* [33] conclude that it is today's challenge "to design experimental protocols that are rigorously validated. Each step of any real-time PCR-based assay must be controlled, from sampling to PCR, including manipulations like extraction and reverse transcription. The evaluation of analytical parameters such as linearity, precision, accuracy, specificity, recovery, limit of detection, robustness, reference interval (and so on) helps us to achieve this goal. Statistical tools such as control charts allow us to study, in the real-time PCR process, the impact of various factors that change over time."

In order to obtain consistent and contamination-free real-time PCR results, separate laboratories should be designated for sample processing and nucleic acid extraction, for assembly of real-time PCRs, for the real-time PCR instrument, and for contamination-free disposal of closed PCR vessels.

7.1. SPECIMEN COLLECTION

Specimen collection and processing for real-time PCR falls largely into two categories: (i) specimens for high sensitivity real-time PCR of low-copy targets such as microorganisms, for which specimen quantity may be limiting,

or (ii) specimens for genetic typing for which specimen quantity is not limiting. Specimens for low-copy targets require careful preservation of nucleic acids, removal of PCR inhibitors, and highest recovery during nucleic acid extraction [91]. Frequently, the limiting factor in detection of low-copy targets is target preservation and recovery in nucleic acid extraction, not the real-time PCR assay [77]. For genetic typing of high-copy targets, it is important that PCR inhibitors are quantitatively removed during nucleic acid extraction while sample preservation and recovery rates are of less concern.

It is a misconception that immediate freezing is the best approach for nucleic acid preservation. In fact, nucleic acids in frozen samples become exposed to nucleases during thawing, and low-copy targets may easily be lost. It is imperative to collect low-copy samples specifically for PCR analysis in nucleic acid preservatives such as guanidinium isothiocyanate buffer before freezing [77]. In most cases, it will make good sense to collect any sample specifically for real-time PCR. This requires good communication and clear instructions between laboratory and clinical personnel [104]. Other factors that affect the stability and property of samples should also be considered. This may include anticoagulants, stabilizing agents, temperature, timing before initial processing, sterility, and even the endogenous degrading properties of specific sample types [104].

7.2. NUCLEIC ACID EXTRACTION

In real-time PCR assays, optimal extraction of nucleic acids from a wide range of clinical samples is one of the most underappreciated, but nevertheless challenging and important steps. Proper extraction must efficiently release nucleic acids from samples, remove PCR inhibitors, avert the degradation of nucleic acids, and ensure adequate concentration of the nucleic acids after extraction. This is relatively easily achievable for viral and mammalian amplification targets, but much more difficult and inconsistent for nucleic acid targets of bacteria, fungi, protozoa, and parasites. These pathogens frequently possess highly resistant outer membranes and specific methods for disruption of these membranes are necessary for quantitative recovery of nucleic acids from these organisms. In some cases, concentration of target organisms by physical means increases assays sensitivity, but it is important to ascertain the enrichment effect [77]. Such methods include immunomagnetic, centrifugal, or filter concentration [91, 105, 106].

The presence of inhibitory substances in the samples may strongly influence PCR performance and detection sensitivity. PCR inhibitors typically interfere with the action of DNA polymerase [14], but may also degrade nucleic acids, or interfere with the cell lysis procedure [107]. Bile salts and

complex polysaccharides in feces [108, 109], heme, immunoglobulin G, and lactoferrin in blood [82, 84], collagen in food samples [110], and humic substances in soil [111] have been identified as PCR-inhibitors.

Nucleic acid preparations used as samples in real-time PCR assays include total nucleic acids, purified DNA, total RNA, and poly(A)⁺ RNA. An abundance of quality-assured commercial kits are available for extraction of nucleic acids and it is strongly advisable to use such kits in the clinical laboratory. It is imperative, however, to validate recovery and removal of PCR inhibitors of any nucleic acid extraction method prior to routine use [91]. Also, compatibility of the nucleic acid storage buffer with the real-time PCR and reproducibility of results after freeze-thaw cycles of extracted nucleic acids should be evaluated [77].

7.3. AUTOMATION

The universal laboratory need for nucleic acid extraction and the analysis of large sample numbers in high-throughput instruments has given rise to major efforts in automation of these procedures. Automation has the potential to reduce the variability in sample preparation the probability of contamination [112]. Automated fluidic devices are available from many producers for extraction, purification, and concentration of nucleic acids [113–120]. These devices use mainly chemical and thermal lysis to release nucleic acids from samples. However, extraction of nucleic acids from organisms with rigid membranes still requires specialized methods such as mechanical disruption that have not yet been automated.

The limitations for the present formats of automated instruments are (i) that these instruments are expensive, both in acquisition and in operation; (ii) that recovery of nucleic acids from samples with low-copy targets needs to be validated; and (iii) that this equipment functions properly only if continuously operated and maintained by highly trained personnel.

8. Applications of Real-Time PCR

Presently, clinical diagnosis and differentiation of pathogens still largely relies on phenotypic identification of the agents. However, the polymerase chain reaction has been accepted as the gold standard for detection of many of these pathogens. Conversion of cumbersome traditional PCR assays into real-time PCR methods and development of new methods will lead to a rapid expansion of the applications of real-time PCR. Over the next decade, real-time PCR may well replace many of the present phenotypic pathogen detection methods in the clinical laboratory. In addition, genetic differentiation of

amplification products such as rapid genetic identification of antibiotic resistance may well open new clinical uses of laboratory diagnostic information.

The limiting factor in all real-time PCR methods is nucleic acid extraction from specimens. Thus, sample collection and processing deserves considerably more attention than it has received. However, once high quality purified nucleic acid is available, multiple real-time PCRs can rapidly extract a wealth of information from a single specimen. Another principal advantage of the widespread use of real-time PCR will be a rapid sample turnaround and accelerated distribution of diagnostic information, which in many cases may be available within 24 hours. These advantages will, and frequently already do, apply similarly to cancer and prenatal diagnostics, and identify genetic polymorphisms by real-time PCR. The following paragraphs cannot cover the advances in depth, but will try to highlight paradigms of current developments in real-time PCR clinical diagnostics.

8.1. DETECTION, QUANTIFICATION, AND GENETIC TYPING OF MICROORGANISMS

8.1.1. *Bacterial Agents*

Results of real-time PCR assays for detection bacteria can, in principle, inform the clinician or veterinarian of the infection status of the patient more rapidly than culture methods, allowing more specific and timely administration of appropriate therapy. This will increase the treatment efficiency, reduce the time of hospital stay, avoid the inappropriate use of antibiotics, and may lower the probability of emerging antibiotic-resistant bacterial strains.

PCR has long been the preferred method for diagnosis of bacteria that are difficult to cultivate, and many real-time PCR assays have already been developed to replace standard PCR methods for bacteria such as *Listeria monocytogenes* [121], *Legionella pneumophila* [122], *Mycobacterium* spp. [123–126], *Propionibacterium* spp. [127], *Borrelia burgdorferi* [128], and *Leptospira* genospecies [129]. DeGraves *et al.* [5] established an efficient real-time PCR platform to detect single *Chlamydia* spp. organisms in the LightCycler by SYBR Green fluorescence and by FRET hybridization probes. This sensitivity is approximately equivalent to detection of a single target in a 10^{12} -fold background of unrelated DNA. The number of chlamydial genomes detected by FRET PCR correlated well with cell culture determination of inclusion forming units [5]. Using optimized nucleic acid extraction methods [77], these PCR methods are at least 150-fold, but typically about 1000-fold more sensitive than the cell culture quantitative assay. Detection of exotic bacteria such as the methanotropic bioremediating *Methylocystis* spp. has also been improved by real-time PCR [130].

Real-time PCR has now become an important tool in the rapid detection of potential use of bacterial pathogens as biological weapons. Conventional assays for such agents usually require more than 24 hours. Real-time PCR is much more rapid and in addition allows for immediate discrimination of weaponized pathogens from the less harmful wild-type strains. A good example is the successful application of real-time PCR for screening for the presence of *Bacillus anthracis* spores and for differentiation based on virulence-encoding plasmids and chromosomal markers [131].

Results of real-time PCR may also inform the clinical use of antimicrobials. Conventional microbiological diagnosis by culture is too slow to provide relevant information for physician and veterinarian for the initial use of antibiotics in infectious disease. The reduced time for real-time PCR diagnosis and the information about pathogen load and genetic properties have the potential to revolutionize clinical medicine by insuring better management of patients, reducing health care costs, and slowing the problem of antibiotic resistance. Real-time PCR can detect and monitor the antibiotic resistance of clinical isolates of bacteria such as *S. aureus*, *S. epidermitis*, *Helicobacter pylori*, *Enterococcus faecalis*, and *Enterococcus faecium* [132]. For example, melting curve analysis distinguished multiple rifampin resistance mutations and high-level isoniazid resistance mutations in *M. tuberculosis* [22], and a real-time PCR was applied successfully to screen for methicillin-resistant *S. aureus* from nasal swabs [133].

8.1.2. *Viral Agents*

Of all fields in clinical microbiology, real-time PCR has had the strongest impact on virology. A number of factors have contributed to this impact: (i) in terms of genetic complexity, viruses are the simplest pathogens; (ii) isolation of nucleic acids from viruses is not complicated by the need to break up rigid membranes of the agents; and (iii) the effort of standard microbiological diagnosis by culture is greater than for many other pathogens, thus real-time PCR diagnosis compares favorably with standard virological methods [134]. Real-time PCR helps to identify co-infections in epidemiological studies, discriminates a multiplicity of viral genotypes within a single reaction vessel, gives accurate quantitative results of viral infection load and its change after antiviral therapy, and demonstrates the epidemiological links between unique viral sequences and clinical signs of disease [20, 135–138]. Therefore, real-time PCR has been used widely for detection of viruses in humans, animals, and plants, and in specimens such as cells, tissue cultures, swabs, serum, plasma, saliva, urine, and cerebrospinal fluid.

One example is the use of real-time PCR for diagnosis of infections with avian Newcastle disease virus, which has two radically different pathogenic phenotypes caused by small nucleotide changes. Differential diagnosis

between diseases induced by these two strains had been difficult with standard methods, but these pathotypes are now rapidly and accurately genetically differentiated by fluorescence melting curve analysis in real-time PCR [8].

A significant number of real-time PCR assays have been developed and applied for detection, quantification, and differentiation of viruses inducing a wide range of human and animal diseases such as coronavirus [139], flaviviruses [140–142], gyrovirus [21], hepadnaviruses [143, 144], herpesviruses [145–147], orthomyxoviruses [148], parvoviruses [149], papovaviruses [150], paramyxoviruses [8], picornaviruses [151, 152], poxviruses [153], retroviruses [154, 155], and rhabdoviruses [156].

8.1.3. Parasites, Fungi, and Protozoa

Over the past 20 years, dramatic advancements have occurred in molecular approaches to the investigation of parasites and parasitic diseases. Among these approaches, real-time PCR methodology has been efficiently used to detect, differentiate, and diagnose parasites of human and animals. Specific FRET real-time PCRs were developed to diagnose *Toxoplasma gondii*, *Trypanosoma cruzi*, and *Cryptosporidium parvum* [157]. The traditional culture method, mouse inoculation, and serological detection are time-consuming and insensitive, while real-time PCR vastly improved speed and sensitivity of diagnosis of parasites such as *Toxoplasma gondii*. *C. parvum* oocysts and *Encephalitozoon* spp. have been genotyped by use of real-time PCR, which was highly advantageous over conventional laboratory microscopic diagnosis for low-concentration samples [158, 159]. Real-time PCR assays were also successfully applied to the diagnosis of diseases caused by *Aspergillus fumigatus* and *Aspergillus flavus* [6, 160]. Similar methods were utilized to investigate the presence of potentially harmful concentrations of toxic fungal spores and conidia produced by *Stachybotrys chartarum* [7].

8.2. CANCER DIAGNOSTICS

Progress in our basic understanding of the molecular causes of many forms of cancer provides an ever increasing array of molecular targets for cancer diagnosis, treatment, and prognosis. Both expression analysis of genes as well as differentiation of gene polymorphisms are required and real-time PCR technology has rapidly been adopted for these diagnostic needs. Thus, real-time PCR methodology is being used to translate recent discoveries in oncology into clinical practice, by routine use in the clinical laboratory for tumor profiling and prognosis [161]. Quantitative real-time PCR can determine gene duplications, deletions, fusions, and overexpression, and melting curve analysis of PCR products can be used to identify mutations, down to single base changes.

8.2.1. *Quantitative Transcript Analysis*

The human epidermal growth factor receptor 2 (HER-2) and DNA topoisomerase II α (topo II α) genes are important in the prognosis and treatment of breast cancer [162, 163]. The HER-2 proto-oncogene is expressed by, and involved in the growth of, 20–30% of metastatic breast cancers [164, 165]. Positive HER-2 status may indicate a poor prognosis and aggressive tumor transformation [166]. The topo II α gene is located near HER-2 within the chromosome band region 17q12–q21, an area that is often mutated in breast tumors. Immunohistochemistry is the conventional HER-2 testing method interpreted by anatomic pathologists, but this technique is subjective and error prone, and it is difficult to differentiate between weakly positive and background staining [167]. In contrast, real-time PCR provides a rapid and sensitive quantitative analysis of the HER-2 gene and its expression [168]. Duplex real-time PCR can be used to detect DNA amplification of HER-2 concomitant with topo II α alterations in a single reaction, and changes in topo II α mRNA copy number indicate the chemotherapy response to topo II α inhibitor drugs [169].

A three-color multiplex PCR reaction was established to simultaneously detect HER-2 and topo II α mRNA normalized to the reference albumin mRNA [161]. In this system, the cross points for all three targets in a wild-type sample are very similar. However, in a tumor sample, both HER-2 and topo II α have cross points three cycles earlier than albumin. This indicates approximately 8-fold mRNA over-expression of HER-2 and topo II α .

8.2.2. *Point Mutation Analysis*

Mutations in DNA may be small alterations such as point mutations and base insertions/deletions, as well as large rearrangements including translocations, inversions, gene fusions, and gene amplification. Small mutations occur frequently in carcinomas and somatic mutations may accumulate during the evolution of many tumors and may significantly impact the course of the malignancy [170]. The *p53* tumor suppressor gene is the most frequently mutated gene in human cancers. Initially, *p53* was thought to function as oncogene, because over-expression of *p53* appeared to cause oncogenic transformation of cells [171]. However, several critical discoveries defined the normal function of wild-type *p53* as anti-oncogenic, and screening of DNA from colon cancer patients revealed that *p53* mutations occur with unusually high frequency in tumor tissue, an observation that was extended to most of the other major forms of human cancer. Most mutations in *p53* are seen in the DNA-binding domain as single base changes [172]. Using real-time PCR, overlapping fluorescein-labeled probes were used to scan colon

cancers for mutations in exons encoding the DNA-binding domain of *p53*, and somatic mutations were identified by melting curve analysis. Compared to the *p53* nonmutated wild-type profile, mutated *p53* induces additional peaks or creates changes in peak areas in the melting curve analysis [161].

Screening for genetic polymorphisms is in its approach similar to mutation analysis. With increasing knowledge about the association of specific single nucleotide polymorphisms (SNPs) with genetic susceptibility/resistance against diseases, screening for such polymorphisms will become a major medical need in the near future that can be met by real-time PCR. Determination of the histocompatibility type is already a major need in transplantation medicine, and real-time PCR methodology for MHC typing has been described [11–13].

8.2.3. Exon Deletion and Gene Amplification Screening

Deletions or duplications of single or multiple exons, entire genes, or larger chromosomal regions are a frequent cause of human genetic disorders, including tumors. No fast and efficient methods were available to detect exon/gene deletions before the successful application of quantitative PCR. Hoebeeck *et al.* [173] successfully developed and applied real-time quantitative PCR as a fast, sensitive, and high-resolution strategy to detect single exon or larger deletions in the von Hippel-Lindau (VHL) gene. This VHL exon quantification approach is based on SYBR Green I detection which is normalized by using two reference genes, ZNF80 and GPR15, with a normal copy number.

Many types of cancer are characterized by increases in the copy numbers of oncogenes and such gene amplifications are typically associated with increased transcription and overexpression of these genes. Conventional methods such as Southern blot and fluorescence in situ hybridization have their intrinsic limitations for detection of gene copy number, such as low sensitivity and high laboratory work load requirement. Real-time PCR has been used very effectively at quantifying such gene amplifications at the chromosomal DNA level in numerous types of tumors [174–176]. Zatkova *et al.* [177] found 8q amplification in hepatoblastomas by a real-time, PCR-based genomic copy assay, and subsequently evaluated the transcript levels of four genes and expressed sequence tags within this chromosomal region. The PLAG1 oncogene, a transcriptional activator of IGF2 in other tumor types, was found highly overexpressed relative to normal liver tissue. These results suggested that PLAG1 overexpression may be responsible for the frequently observed up-regulation of IGF2 in hepatoblastoma and may be implicated in the molecular pathogenesis of this childhood neoplasia.

8.2.4. Fusion Gene Analysis

Many types of leukemia are characterized by chromosomal rearrangements that result in the fusion of genes. Such fusion genes are typically highly expressed by transformed cells and can serve both as markers of the type as well as of the progression of a tumor. Guerrasio *et al.* [178] detected and quantified real-time RT-PCR of the CBFbeta/MYH11 fusion transcript that characterizes the FAB M4Eo subtype of acute myeloid leukemia. The CBFbeta/MYH11 transcript was detected in all patients tested and highest copy numbers of the fusion gene relative to the reference ABL copies indicated patients destined to relapse. A 2–3 log decline in fusion transcripts was observed after induction/consolidation therapy. After therapy, minimal residual disease was continuously monitored by repeated real-time RT-PCR. In patients with continuous complete remission (CCR), fusion transcripts remained undetectable; while in relapsing patients, the fusion transcript copy number during complete remission (CR) never declined below the detection threshold.

Similarly, real-time RT-PCR quantification of SIL-TAL1 fusion gene transcripts allowed the earliest detection of tumor cells in human T-cell lymphocytic leukemia [179]. Quantification of BCR-ABL fusion transcripts after hematopoietic stem cell transplantation allowed sensitive monitoring for minimal residual disease in patients with chronic myeloid leukemia [180]. Osomi *et al.* [181] established a multiplex real-time PCR method that simultaneously detects 10 different fusion transcripts that characterize frequent forms of acute leukemia. This method enables rapid characterization of the leukemia as well as post-therapeutic monitoring for minimal residual disease.

8.3. PRENATAL DIAGNOSIS

Since the early 1970s, prenatal diagnosis has been applied to identify fetal genetic disorders, predominantly trisomy of chromosomes 21, 13, 18, and sex chromosome aneuploidies [182]. Ultrasound and biochemical tests have become the standard procedures to screen for fetal chromosomal abnormalities, but require confirmation by cytogenetic analysis of amniotic fetal cells. Two standard methods to obtain fetal cells, amniocentesis usually performed at 15 weeks of gestation and chorionic villus sampling at 9–11 weeks of gestation, are invasive and have the risk of induced abortion. To alleviate parental anxiety, accelerated molecular diagnosis of aneuploidy is highly preferable over time-consuming karyotyping of cultured fetal cells [182]. Real-time PCR is currently replacing many of these rapid molecular techniques for uncultured fetal cells that can deliver results in 1–2 days. For instance, in a duplex real-time PCR assay for trisomy 21 (Down syndrome),

the most common cytogenetic anomaly, simultaneous amplification from amniocytes or chorionic mesenchyme of the chromosome 12 locus GADPH and the amyloid gene locus on chromosome 21 allows the determination of chromosomal ploidy, and thus diagnosis of trisomy 21 [183, 184]. Other prenatal diagnoses by real-time PCR from amniocentesis or chorionic villus sampling include the most common beta-thalassemia mutations [185], achondroplasia and hypochondroplasia [186], and even an infectious disease, congenital toxoplasmosis [187].

Recently, the presence of cell-free fetal DNA in the maternal circulation was recognized (~ 30 fetal genome equivalents/ml maternal plasma), and extraction of fetal DNA from maternal plasma has opened the possibility of risk-free, noninvasive prenatal real-time PCR genetic diagnosis [188]. Such assays are robust, with sensitivity of approximately 95–100% and specificity near 100%. Zhong *et al.* [189] established a duplex real-time PCR method for simultaneous amplification of the Rhesus D gene and the SRY locus on the Y chromosome. This approach allows the identification of fetal Rhesus D status along with sex determination. Quantitative real-time PCR determination found that 2–3 times more fetal genome equivalents are present in maternal plasma in pregnancies with trisomy 21 and other fetal aneuploidies than in pregnancies with euploid fetus [190, 191]. Noninvasive genetic screening by real-time PCR has thus the potential to replace most invasive cytogenetic and molecular prenatal tests.

9. Conclusions

Real-time PCR is rapidly becoming the “gold standard” of nucleic acid sequence detection and quantification. The strengths of this still developing technology are manifold: ease and speed of assay execution for large batches of samples; highest sensitivity (single targets can be detected in a $\sim 10^{12}$ -fold excess of unrelated nucleic acids); wide dynamic range (at least six orders of magnitude) of detection and quantification with a linear relation between log target to detection threshold cycle; highest specificity approaching 100% in well designed assays with hybridization probes; and differentiation of detected nucleic acid sequences. These advantages ensure that real-time PCR will continue to replace ever more diagnostic assays in the clinical laboratory and that new assays will be developed that would not have been possible without real-time PCR. Current trends clearly indicate that, within the next decade, real-time PCR will become the dominant method in genetics-based diagnostics.

Many of the early shortcomings of have been corrected. For instance, use of dUTP-uracil glycosylase reaction chemistry reliably prevents product

carryover and has become standard in real-time PCR. False-positive results may also be caused by contamination during sample handling and nucleic acid extraction. Careful sample handling, dedicated laboratories and equipment, well-trained personnel, and a well-planned and strictly forward-moving workflow are essential to control systematic contamination problems [104]. However, some problems still impede wider use of real-time PCR technology in the clinical diagnostic laboratory [33]. The main problem is quality assurance and consistency of results, divided into two main areas: (i) false-negative results and (ii) inaccurate quantification and differentiation. Certain aspects of real-time PCR, as highlighted below, disproportionately contribute to inconsistent results.

The importance of false-negative results is substantially underestimated in PCR diagnostics, including real-time PCR. Inefficient preservation, removal of PCR inhibitors, and extraction of low-copy nucleic acids, and their insufficient concentration into the low sample volume of a real-time PCR assay, is probably the most underappreciated cause of false-negative results [77]. Of all aspects of real-time PCR, pre-PCR nucleic acid extraction has the greatest need for technological improvements and standardization.

Another main cause of false-negative results is lack of robustness of the real-time PCR assay, resulting also in poor reproducibility. Poorly designed real-time PCR assays can easily be recognized in a limiting standard template dilution series by the rapidly declining slopes of the linear phase of semi-log amplification plots. To ensure high PCR efficiency, the prerequisite for robustness, careful optimization of real-time PCR assays in a limiting template dilution series is vital. If many different PCR assays are performed, it is easier to modify thermal reaction parameters for optimum robustness than to modify reaction chemistry. One of the most effective ways to improve PCR efficiency is the use of a step-down thermal protocol prior to fluorescence acquisition cycling.

Lack of robustness may also contribute significantly to inaccuracies and imprecision in target quantification and differentiation, and improved quantification and differentiation is a "fringe benefit" of robust real-time PCR assays. Carefully constructed duplex one-step RT-PCR assays for simultaneous amplification of analyte and reference mRNA minimize errors in relative quantification of transcripts [26].

The theoretical underpinnings of quantification in real-time PCR have only recently become the subject of in-depth mathematical modeling. Four-parameter sigmoidal curve fitting of individual reactions has the potential to become the method of choice for objective, highly accurate quantification [42]. By modeling individual reactions, this approach would remove subjectivity in threshold cycle quantification with standard series, would require only a single calibrator standard for absolute quantification,

would be insensitive to differences in PCR efficiency between samples and in multiplex PCRs, and would allow automation.

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