New Developments in Quantitative Real-time Polymerase Chain Reaction Technology

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Abstract
Real time-quantitative PCR (RT-qPCR) technology has revolutionized the detection landscape in every area of molecular biology. The fundamental basis of this technology has remained unchanged since its inception, however various modifications have enhanced the overall performance of this highly versatile technology. These improvements have ranged from changes in the individual components of the enzymatic reaction cocktail (polymerizing enzymes, reaction buffers, probes, etc.) to the detection system itself (instrumentation, software, etc.). The RT-qPCR technology currently available to researchers is more sensitive, faster and affordable than when this technology was first introduced. In this article, we summarize the developments of the last few years in RT-qPCR technology and nucleic acid amplification.

Introduction
Since its introduction in the late 1990s, real time-quantitative PCR (RT-qPCR) technology has revolutionized the detection landscape in every aspect of molecular biology (Filion, 2012). The fundamental basis of this technology has remained unchanged since its inception, which is fluorescence based detection of its target, amplified via classical multistep PCR. A steady modification(s) of individual components which makeup the RT-qPCR platform has however enhanced the overall performance of this highly versatile technology. These improvements have ranged from changes in the individual components of the enzymatic reaction cocktail (polymerizing enzymes, reaction buffers, probes, etc.) to the detection system itself (instrumentation, software, etc.). As a result, the RT-qPCR technology currently available to researchers is more sensitive, faster and most importantly, affordable for laboratories on moderate/modest budgets, than when this technology was first introduced (Filion, 2012).

In this article, we seek to briefly summarize some of the developments which have happen within the last few years in the area of RT-qPCR technology and nucleic acid amplification (e.g. classical PCR). In our opinion, these developments might have a direct relevance for researchers working in the area of environmental transcriptomics. Some of described examples in this article have been validated on RT-qPCR platforms, making it relatively easy to adapt in their current state, while others have only been implemented on a conventional end-point detection platform.

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Isothermal detection methods
The current RT-qPCR technology is based on the classical three-step thermal cycling process which is, template denaturation, followed by primer/probe annealing and finally, extension/detection of the fluorescence signal, to amplify and detect the target transcripts all under real-time conditions (Gelfand, 1992). A very commonly observed phenomenon in this multistep thermo-cycling amplification system is the generation of spurious fluorescence signal due to mispriming of primer/probes. To overcome such limitations, detection platforms have been proposed which amplify the target exponentially like PCR, but under isothermal conditions, i.e. at a fixed, user-defined temperature (reviewed by Gill and Ghaemi, 2008).

The helicase-dependent (HDA) amplification system is one such novel ‘non-PCR’ system for amplifying target DNA (Vincent et al., 2004) and RNA (Goldmeyer et al., 2007), under isothermal conditions. This revolutionary amplification system makes use of a novel enzymatic cocktail which does not require the DNA to be cycled between different temperatures, like that done for reactions based on Taq DNA polymerase amplification or any of its variants. In lieu of a standard denaturation step, the HDA system uses the helicase enzyme to unwind the double-stranded DNA and with the aid of other polymerizing enzymes, an exponential amplification is achieved (Vincent et al., 2004). All these steps are performed at a fixed, user defined temperature. Though extremely novel when first introduced, the HDA system suffers from one major limitation- its inability to amplify DNA targets greater than 200bp (Vincent et al., 2004). As a result, in its present state, it is seriously unable to challenge and act as a viable alternative to the highly versatile PCR, or any amplification system based on it.

Despite this technical limitation, HDA-based amplification is increasingly being viewed as an attractive alternative in any amplification system, where length of the amplicon is less than 200bp. One excellent example of such short amplicon size detection system is RT-qPCR. It is a well known fact that, for an efficient amplification, the primer/probe parameters of the RT-qPCR detection system is usually within the 50–150bp size range. It is therefore not impossible to develop a RT-qHDA detection system, where the classical Taq DNA polymerase amplification cocktail is substituted by an enzymatic system, which amplifies its target at a fixed (isothermal) temperature.

Recently, Tong et al. (2008) showed that such enzymatic substitution is indeed feasible. In their work, primer/probes designed for TaqMan RT-qPCR fluorescence system were used to develop a novel RT-qHDA system to detect various biothreat microorganisms. Since pre-existing TaqMan primer/probes were used, the RT-qHDA system was optimized to work with the 60–65°C temperature range and detection sensitivity, similar to what one would...
expect from a classical RT-qPCR system. Though very preliminary, it is tempting to conclude that the RT-qHDA system, at least in theory, is ‘backward compatible’ with a pre-existing RT-qPCR system. In other words, the user does not have to separately invest in a new detection platform (i.e. instrumentation) or custom synthesize the fluorogenic probes, both of which could be prohibitively expensive. The only modification one would have to make is to substitute the Taq DNA polymerase based reaction components, with a HDA amplification cocktail, which is now commercially available in various formats from Biohelix Corp., USA. Interestingly enough, isothermal amplification systems themselves are also constantly being refined to enhance fidelity (amplicon length) and robustness of the amplification process. Protocols which make use of novel enzymatic cocktails which are more efficient in unwinding and amplifying the target double helix are constantly being reported (Kiesling et al., 2007; Tan et al., 2008; Schaerli et al., 2010; Joneja and Huang, 2011).

In our opinion, adapting such ‘non-PCR’ amplification technologies could in the near future lead to detection platforms (Mahalanabis et al., 2010) which are more robust and would not suffer from the inherent drawbacks, for example spurious amplification, cycling parameter standardization, typically associated with the classical three-stage PCR system.

Development of novel normalization controls

The first step towards analysing microbial gene expression requires a quantitative extraction of RNA. This step has proven to be highly problematic for environmental matrices, due to compounded inefficiencies in individual steps which include, but not limited to, incomplete cell lysis, RNA degradation by ubiquitous RNases, co-extraction of inhibitors and their interaction with the enzymes used (Lorenz and Wackernagel, 1987; Moran et al., 1993, Alm and Stahl, 2000). One straightforward approach to quantify such losses and apply the necessary correction is to include an internal amplification control (IAC), so as to make the final gene expression meaningful and reproducible. An IAC is essentially a non-target DNA/RNA sequence co-amplified, preferably in the same reaction tube, under the same reaction conditions (Hoorfar et al., 2003, 2004; Nolte, 2004; van Doorn et al., 2009). While attempts to develop IAC’s have met with some success for experimental systems which are highly controlled (Okonomou et al., 2008; Scipioni et al., 2008), developing such controls have proven to be highly problematic for certain experimental set-ups, for example complex environmental matrices. The main difficulty in these cases has been in our inability to identify an inert IAC which is able to (a) withstand the harsh nucleic acid extraction procedures usually employed for environmental matrices, and if such a sequence is indeed developed/identified (b) designing a primer/probe combination which would not cross-react with other non-target (nucleic acids) components of the matrices. While few potential IAC based solutions have been proposed, for example the Biotrove OpenArray platform (van Doorn et al., 2009), high costs and proprietary issues of some IAC sequences (Nordstrom et al., 2007) have served as a deterrent for researchers who are seriously interested in rigorously implementing this external normalization strategy.

Recently, Gonzalez-Escalona et al. (2009) proposed the implementation of a highly versatile RNA based IAC, called the myIC system. The myIC is essentially a 200 bp artificial sequence, with no known homology to any nucleic acid accession in the GENBANK (NCBI) database. As a result, primer/probes designed for this sequence are not expected to cross-react with any non-target nucleic acids, typically expected to be present in high amounts in RNA/DNA extracted from complex environmental matrices. Recently, its applicability was successfully demonstrated in quantifying bacterial gene transcripts from natural soil samples spiked with a defined amount of the bacterium Pseudomonas sp. LBUM300 (DeCoste et al., 2011). This success should spur researchers to implement the myIC system to quantify microbial gene expression in other environmental matrices, but were unable due to lack of a suitable IAC. The added advantage of its non-proprietary usage and its applicability in both DNA/RNA formats (Gonzalez-Escalona et al., 2009; Deer et al., 2010), makes myIC a very attractive IAC to implement in various RT-qPCR based detection systems.

LNA/Zip nucleic acids

One critical factor for any PCR based detection platform, including RT-qPCR, is the invariant ability of the primer/probe to discern its target from a heterogeneous mixture and bind it with extreme specificity. In RT-qPCR, the design of these primers/probes is governed by strict guidelines (hybridization/sequence parameters) which, in turn, are based on the fluorescence chemistry used for that particular assay. For example, the hybridization parameters for TaqMan assays are different from the Molecular beacon based fluorogenic assays. Empirically, these primers/probes are designed with the help of specific software’s (e.g. PrimerExpress™ for TaqMan detection system) which are proprietary and usually supplied with the detection platform. Since these software use specific built-in algorithms, with a narrow set of highly defined binding parameters, it is not very uncommon to find these design software unable to identify optimal primer/probe combinations for a particular user input sequence.

While these design limitations have not exactly impeded our ability to vigorously implement the RT-qPCR technique, there has always been an ongoing endeavour to increase the sensitivity of an assay by enhancing the ‘specificity and recognition’ capacity of a primer/probe. The most conventional approach to achieve this goal has been to manipulate various components of the amplification reaction, for example use of reaction buffer which are highly optimized, increased level of divalent ions (Mg²⁺) and use of specific additives which are known to prevent primer mispriming. Early research has however showed that it is also possible to enhance the specificity and recognition aspects of an oligonucleotide, by another novel approach-chemical modification of the oligonucleotide backbone itself. As a result of this modification, the oligonucleotide is able to bind to its target with extreme specificity and enhanced accuracy.

Of the several modifications developed till date (Hyrup and Neilson, 1996; Schulz and Gryaznov, 1996; VanAerschot et al., 1996; Hendrix et al., 1997; Nielson and Haaima, 1997), the locked nucleic acids (LNA) are probably seen as the most promising in terms of design simplicity and superior functionality, i.e. its unique ability to recognize
and bind to its target with an extreme precision (Obika et al., 1998; Singh et al., 1998). The high specificity of LNAs have made it possible to apply it in single-nucleotide polymorphism (SNP) genotyping and detection of difficult templates which, otherwise is not feasible with conventional primer/probes (Simeonov and Nikiforov, 2002; Latorra et al., 2003; Johnson et al., 2004; Ugozzoli et al., 2004). The recent introduction of zip nucleic acids (ZNA), which have similar functional properties of LNAs but, based on a different modification chemistry (Voirin et al., 2007), is also an exciting development. The ZNA oligonucleotides have a spermine residues conjugated to the basic oligonucleotide backbone. This modification results in reduced electrostatic repulsion brought about by the polyanionic nature of nucleic acids. As a result, an enhanced sensitivity is routinely observed in detecting its target during both RT-qPCR and reverse transcription assay (Moreau et al., 2009).

In our opinion, these ‘enhanced specificity’ oligonucleotides like LNAs, could be further exploited to design RT-qPCR primer/probes used to detect microbial gene expression from environmental matrices, especially complex types. As analysing gene expression from environmental samples entails discerning the target from a heterogeneous background, an enhanced specificity primer/probe would result in a very high signal-to-noise ratio and hence a more robust, accurate and reproducible detection of the target transcripts. The commercial availability of both the LNA and ZIP oligonucleotides should make it easier for researchers to develop specific applications to accurately detect its target from environmental matrices.

Use of polymerases engineered to resist inhibitors
As gene expression analysis using RT-qPCR entails the synthesis of cDNA by the reverse transcriptase (RT) enzyme, followed by an exponential amplification of the template by DNA polymerase, any chemical moiety, which is able to alter the fidelity (negatively) of these enzymes, has a direct bearing on the end results (Suslov and Steindler, 2005). The net effect of such interferences, at least in RT-qPCR, is the invariable quenching of the fluorescent signal, leading to sub-optimal or irreproducible gene expression data. Consequently, solutions are constantly being sought which could in some way, alleviate the effects of these interfering moieties on the normal functioning of these enzymes. Some of the commonly used inhibitor mitigation strategies include, use of RNA extraction protocols specifically designed to eliminate co-extracted inhibitors, and addition of specific additives in the RT-qPCR reaction which are known to bind the inhibitor moieties (e.g. BSA, T4 gene 32 protein). One attractive solution which could be an ‘add-on’ to these aforementioned strategies is the use of enzymes (reverse transcriptase and DNA polymerase enzymes) which are inherently ‘resistant’ to any of the signal quenching inhibitors.

The first step in RT-qPCR based gene expression analysis is the faithful conversion of RNA to cDNA by reverse transcription. Therefore, the first step would be the development of an RT enzyme which is able to resist a wide spectrum or a specific class of environmental inhibitors. While most RTs are highly sensitive to inhibitors, the recent report of a mutant Moloney murine leukemia virus (MMLV)-RT enzyme, resistant to few common known inhibitors (Arezi et al., 2010), is an exciting development. In the second step of RT-qPCR, the DNA polymerases are the most important component used for the exponential amplification of the target. While certain variants of DNA polymerases are capable of tolerating chemical inhibitors up to certain extent (Al-Soud and Rådström, 1998), the development of a DNA polymerase specifically engineered to resist inhibitors, has not been attempted till recently. Kermekchiev et al. (2009) using a site directed mutagenesis strategy, whereby a ‘designer’ DNA polymerase capable of resisting two recalcitrant inhibitors namely humic/fulvic acid (commonly found in soil) and haemoglobin (common in meat based matrices) was developed. Recently, Baar et al. (2011) showed through molecular breeding, development of a novel DNA polymerase called 2D9, which has sequence elements from four different DNA polymerases. This ‘chimeric’ polymerase exhibits a striking resistance to a broad spectrum of complex inhibitors including humic acid, bone dust, coprolite, peat extract, clay-rich soil, cave sediment and tar.

Taking both these development into consideration, it could be possible in the near future to develop RT-qPCR protocols which are specifically tailored for environmental microbiology. This modified approach would allow the end user to perform a more reproducible detection and quantification of gene expression from environmental matrices. Commercial availability, unit cost and proprietary issues are some of the factors which one would have to consider prior to implementing this approach.

Conclusion
The application of the RT-qPCR technology has contributed immensely to our understanding of gene expression in various biological systems; however in certain areas of research, for example applied microbiology, application of this technique has not progressed as one would have liked to expect. This application gap could at best be attributed to the extreme difficulties in extracting nucleic acids from environmental samples and the high sensitivity of the RT-qPCR system towards chemical components inherently co-extracted from environmental samples. Development of a more robust RT-qPCR platform is one possible solution to overcome this problem. We believe that a cross-adaption of some new developments in amplification and enzymatic technology would alleviate some of the drawbacks inherent to the RT-qPCR technology so that it's potentially is fully realized in areas like applied microbiology (Filion, 2012).

References
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