Real-time monitoring of rolling-circle amplification using a modified molecular beacon design

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ABSTRACT

We describe a method to monitor rolling-circle replication of circular oligonucleotides in dual-color and in real-time using molecular beacons. The method can be used to study the kinetics of the polymerization reaction and to amplify and quantify circularized oligonucleotide probes in a rolling-circle amplification (RCA) reaction. Modified molecular beacons were made of 2′-O-Me-RNA to prevent 3′-exonucleolytic degradation by the polymerase used. Moreover, the complement of one of the stem sequences of the molecular beacon was included in the RCA products to avoid fluorescence quenching due to inter-molecular hybridization of neighboring molecular beacons hybridizing to the concatemeric polymerization product. The method allows highly accurate quantification of circularized DNA over a broad concentration range by relating the signal from the test DNA circle to an internal reference DNA circle reporting in a distinct fluorescence color.

INTRODUCTION

Real-time measurements of the polymerase chain reaction (PCR) have enabled convenient quantification of target sequences over a wide concentration range by the use of homogeneous detection assays for the amplification products (1). Most commonly the assays are based on fluorescence resonance energy transfer (FRET), or fluorescence by compounds that intercalate in nucleic acids.

Hairpin-shaped hybridization probes, so-called molecular beacons, have been used to detect, distinguish or quantify amplification products of DNA or RNA target sequences (2). Molecular beacons are equipped with a fluorophore and a fluorescence quencher at either end, so that the fluorophore is brought close to the quencher when the free probes adopt a stem-loop structure. Upon hybridization to a target nucleic acid molecule, the fluorophore is extended away from the quencher and can emit fluorescence. The stem-loop structure can also be disrupted due to degradation or binding by proteins. This has been utilized to study the activity of single-strand nucleases (3) and single-strand binding proteins (4).

Rolling-circle amplification (RCA) has been applied for sensitive detection of nucleic acid sequences and of proteins (5,6). The method is based on the finding that also small DNA circles can be copied by a rolling-circle replication mechanism (5,7–9), known from the replication of phage genomes. RCA is particularly useful for signal amplification of padlock probes, i.e. linear DNA probes that become circularized upon recognition of a specific nucleic acid sequence (10), since only reacted probes are amplified. Padlock probes have been used to genotype samples of genomic DNA or RNA in solution (5,11–13), or for in situ genotyping of metaphase chromosomes and interphase nuclei (12,14,15). The combination of padlock probe circularization and amplification through RCA has proven useful for genetic analysis (5,16).

The course of RCA reactions has been studied using electrophoresis (5,7–9), or by hybridizing labeled oligonucleotides to RCA products attached to a glass slide (5,6,17,18). Both methods require laborious handling of gels or microscope slides, and the precision of quantification is limited. RCA reactions have also been analyzed using intercalating dyes (19). The major disadvantage of using intercalating dyes is that the nucleic acids present in the sample, as well as any non-specific polymerization products, contribute to the fluorescence. Moreover, only one RCA product can be analyzed per reaction.

Here we describe quantitative, dual-color, real-time monitoring of RCA using a modified molecular beacon design. We show that this isothermal linear signal amplification allows highly accurate quantification of circularized padlock probes.

MATERIALS AND METHODS

Oligonucleotides

All oligonucleotides were kindly provided by Eurogentec, Belgium. The padlock probes used were: pp90, P-CCTCCCATCATATTTAAAGGCTTCTCTATGTTAAGTGAC-CTACGACGATGCTGCTGCTGTACTACTCTTCCTAAGGCATTCTGCAAACAT; pp93, P-CCTCCATCATATTTAAAGGCTTCTCTATGTTAAGTGACCTACGACCTCAA-TGCTGCTGCTGACTACACTCTCTTCTGAAGGCCATTTCG-.

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AAACAT; ppWT, P-CTGCCATCTTAAACAAAACTTTCT-CCTATGATTATCTGACCTAGCCTCAATGCTG-CTGTGATCTACTCTCTCTGTGATCCACAGGGCT and
ppMUT, P-CTGCCATCTTAAACAAAACTTTCTCTCTATGATTACCTACCTGAATGCCTAG; tWt (for ppWT),
TTATGACCTACCTGACCTCAATGCACATGCTGAAAACAT; ppWT, P-CTGCCATCTTAACAAACCCTTTCTCTATGATTACCTACCTGAATGCCTAG; tMUT (for ppMUT), GTTTGTAAGATGGCAGGGCCCGGTAAATCG; and tMut
(see Figure 1), CTGCTGTACTACTCTTCTATGCATTGAGGTCGT.

**Padlock probe circularization**

Padlock probes (200 nM) were ligationed in 10 mM Tris±acetate (pH 7.5), 10 mM magnesiumacetate, 50 mM NaCl, 1 mM ATP, 1 µg/µl BSA, no dithiothreitol, 0.2 µg/µl T4 DNA ligase (Amersham Pharmacia Biotech, Uppsala, Sweden) at 37°C for 30 min in the presence of 600 nM ligation template.

**HPLC characterization of phi29-polymerase-dependent degradation of molecular beacons**

Three picomoles of molecular beacons from reactions with or without Φ29 DNA polymerase were analyzed using a LA-Chrom (Merck, Darmstadt, Germany) HPLC system with a Phenomenex Luna 5 µm C-18 column (250 × 2.0 mm). The samples were separated in a 30 min gradient from 13 to 57% Acetonitrile (AcN HPLC grade, Merck) in 0.1 M triethylammoniumacetate buffer. The column was washed for 5 min with 60% AcN and equilibrated for 5 min with 13% AcN before the next sample was injected. Fluorescence was measured at 490 nm excitation and 530 nm emission wavelengths.

**Rolling-circle amplification**

Polymerization reactions were performed in 50 mM Tris–HCl pH 7.5, 10 mM MgCl2, 20 mM (NH4)2SO4, 10 mM dithiothreitol, 0.2 µg/µl BSA, 0.25 mM dNTP and 2 ng/µl Φ29 DNA polymerase (kindly provided by Dr M. Salas) at 37°C, using the ligation template as primer for the polymerization. For real-time monitoring, the RCA was performed in the presence of 100 nM molecular beacon and 300 nM ROX dye. The reactions were followed in a ABI 7700 real-time PCR instrument. Fluorescence values are given as a ratio between the fluorescence emitted by the molecular beacon (FAM or HEX) and the ROX reference dye. The temperature profiles were obtained by sampling fluorescence after temperature increments of 1°C held for 30 s.

**Restriction digestion of RCA products**

A MseI restriction recognition sequence present once in every monomer of the RCA products was rendered double stranded and digested by adding 360 nM oligonucleotide (TCTATGTTAAGTGACC) and 73 mM/ml MseI (New England Biolabs) to the polymerization reaction. The restriction digestion was performed at 37°C for 30 min.

**Quantification of probe circles using an internal reference circle**

Polymerization reactions were performed and followed in real-time for 90 min as described except that 1 µg/µl BSA, no ROX reference and 50 mM iWT oligonucleotide was used as primer in a volume of 20 µl. RCA was measured for a serial dilution of circularized ppWT padlock probe, and 30 fmol ppMUT DNA circles as an internal reference. The RCA products from the two DNA circles were detected with the molecular beacons mbRNAfam and mbRNAhex, respectively.

The reactions were monitored in an ABI 7700 real-time PCR instrument. Instead of analyzing the data with the SDS v1.7 software package provided with the instrument, we used a simplified multi-component analysis to resolve the overlapping fluorescence spectra of the two fluorophores used. Individual fluorescence spectra were recorded from the two molecular beacons hybridizing to RCA products at 37°C to determine the relative contribution of the two fluorophores (IFAM and IHEX) at the optimal emission wavelength of respective fluorophore (FAM = λ1 and HEX = λ2). For each sample position in the ABI 7700 instrument a background spectrum was obtained in order to determine the background fraction at the two wavelengths (Bkg1,2 and Bkg1,3). To determine the signal from the two molecular beacons (SFAM and SHEX), we assumed the following relationships between the signals from the two molecular beacons and the measured fluorescence at respective wavelengths with 90 s interval (Fλ1 and Fλ2): Fλ1 = IFAMλ1 × SFAM + IHEXλ1 × SHEX + Bkgλ1, and Fλ2 = IFAMλ2 × SFAM + IHEXλ2 × SHEX + Bkgλ2. The ratio of the maximal slope of HEX and FAM signals as a function of time (ΔmaxHEX/ΔmaxFAM) was used as a measure of the amount of ppWT probe circles relative to the amount of reference ppMUT circles present in the RCA reaction. With this procedure we obtained more reproducible quantification, with minimal influence from well-to-well differences in background. To further minimize the contribution from instrument fluctuations the fluorescence values were averaged over scanning windows of five consecutive measurements. The maximum increase of fluorescence was similarly determined from averaged fluorescence of five consecutive changes in fluorescence.

**RESULTS**

Protection of molecular beacons from exonucleolytic degradation

In early attempts to detect RCA products using DNA molecular beacons, we observed a non-specific accumulation
of fluorescence. We suspected that this could be due to 3’ exonucleolytic degradation of the molecular beacon, separating the DABCYL quencher from the fluorophore at either end of the molecules. We therefore designed a molecular beacon composed of 2’-O-Me-RNA residues, to render the probes resistant to nuclease degradation (20). The nuclease resistance was tested in an experiment where 2’-O-Me-RNA molecular beacon mbRNAhex labeled with a HEX fluorophore (red) or DNA labeled FAM fluorophore (green) were added to the same test tube in the presence (circles) or absence (squares) of Φ29 DNA polymerase. The left portion of the graph shows the variation of fluorescence over time at 37°C. The right portion shows the variation of fluorescence in response to increasing temperature. This temperature profile gives an indication whether the stem parts of the molecular beacons are intact at the end of the 60 min incubation. The different molecular beacons have slightly different melting characteristics because of different hybrid stability of their respective stem sequences. (B) HPLC chromatograms showing the fluorescence from mbDNAfam (green line) and mbRNAfam (red line) molecular beacons before and after a 1 h treatment with Φ29 DNA polymerase at 37°C.

Figure 1. Non-specific accumulation of fluorescence due to exonucleolytic degradation of molecular beacons can be avoided by replacing all DNA residues of the molecular beacon with 2’-O-Me-RNA residues. (A) The DNA molecular beacon mbDNAfam labeled with a FAM fluorophore (green) and the 2’-O-Me-RNA molecular beacon mbRNAhex labeled with a HEX fluorophore (red) were added to the same test tube in the presence (circles) or absence (squares) of Φ29 DNA polymerase. The left portion of the graph shows the variation of fluorescence over time at 37°C. The right portion shows the variation of fluorescence in response to increasing temperature. This temperature profile gives an indication whether the stem parts of the molecular beacons are intact at the end of the 60 min incubation. The different molecular beacons have slightly different melting characteristics because of different hybrid stability of their respective stem sequences. (B) HPLC chromatograms showing the fluorescence from mbDNAfam (green line) and mbRNAfam (red line) molecular beacons before and after a 1 h treatment with Φ29 DNA polymerase at 37°C.
The samples were also analyzed with HPLC to investigate whether the Φ29 DNA polymerase-dependent accumulation of fluorescence was due to exonuclease degradation of the DNA molecular beacon (Fig. 1B). As expected from the temperature profiles, the HPLC traces of the Φ29 DNA polymerase treated and untreated 2′-O-Me-RNA beacons were identical, whereas additional peaks appeared from the DNA molecular beacon treated with Φ29 DNA polymerase compared with the untreated DNA molecular beacon. The new peaks are less hydrophobic than the original material, which is consistent with loss of the hydrophobic quencher and perhaps one or more nucleotides at the 3′ end. The degradation products emit more fluorescence than the corresponding decrease of fluorescence from the original material, which provides further evidence that the quencher has been physically separated from the fluorophore.

**Inter-molecular quenching of molecular beacons**

Despite the chemical modification of the molecular beacons we were still unable to obtain reliable fluorescence signals from RCA products using a conventional molecular beacon design. We hypothesized that this could be due to hybridization between stem segments of neighboring molecular beacons when hybridizing to an RCA product. Thereby the fluorescence quencher and the fluorophore of the molecular beacons would be brought close together by inter- rather than intra-molecular hybridization (Fig. 2). We therefore redesigned the original pp90 padlock probe to a new pp93 version that generated an RCA product which included the complement of one of the stem sequences of the molecular beacon. With this modification the molecular beacons hybridize to the RCA product with one of the stem arms in addition to the loop part. We expected that this would prevent inter-molecular molecular beacon quenching.

Figure 3A (left) demonstrates the results of an experiment where the mbRNAhex molecular beacon was used to follow in real-time RCA reactions of either of the two padlock probe versions. The molecular beacon signal was ~6-fold stronger in the RCA reaction with the modified pp93 padlock probe compared with the original pp90 version. The temperature profiles of the two RCA products support the notion that inter-molecular stems were formed between neighboring molecular beacons hybridizing to the RCA product. The quenched inter-molecular structures formed in the original version of the padlock probe melt at a temperature ~35°C lower than that required to melt the intra-molecular hairpins of molecular beacons that do not hybridize to a target sequence (Fig. 3A, right). The lower $T_m$ of the hairpin structure formed by the RCA products compared with the one formed by free molecular beacons is probably a consequence of increased entropy due to the larger loops formed by the RCA products.

To further verify that the reduced signal from molecular beacons hybridizing to the original RCA product design was due to inter-molecular quenching of the molecular beacons rather than differences in the amount of RCA product, the RCA products were monomerized by adding an oligonucleotide complementary to a $Mse$ restriction enzyme recognition sequence in the RCA product, along with the corresponding restriction enzyme. Fluorescence temperature profiles of the restriction-digested products were recorded after a 30 min incubation at 37°C (Fig. 3B). After restriction digest, the molecular beacons hybridizing to the quenched version of the RCA product emitted 10-fold more fluorescence at 37°C compared with when hybridizing to the non-digested product, while the fluorescence from molecular beacons hybridizing to the non-quenched version of the RCA product was only marginally affected by the restriction digestion.

Notably, the restriction-digested originally quenched product, fluoresce 2-fold stronger than the corresponding non-quenched product at 50°C. This phenomenon could not be explained by differences in restriction digestion efficiencies or in molar amounts of the two products, as judged by gel electrophoresis analysis of the restriction-digested products (data not shown). The molecular beacon hybridizes to the two RCA products in a slightly different manner. In the original design both the quencher and the fluorophore are connected to stem sequences that do not hybridize to the target sequence, while in the non-quenched version, the stem sequence that carries the fluorophore hybridizes to the target sequence (Fig. 2). To investigate whether this physical difference could explain the difference in fluorescence, we synthesized three oligonucleotide hybridization targets to which the molecular beacon would hybridize in three different ways. To the first target the molecular beacon could hybridize only with the loop segment, as a traditional molecular beacon. To the other two targets, either the quencher stem or the fluorophore stem could hybridize in addition to the loop sequence. Figure 3C shows the fluorescence temperature profiles of the mbRNAHex molecular beacon hybridizing to the three different targets. Evidently the target that hybridizes to the fluorophore-carrying stem yields less fluorescence than the one hybridizing to the quencher-carrying stem. The decreased fluorescence from molecular beacons with hybridized fluorophore stem may be due to quenching caused by the bases in the target sequence (21).
Quantitative analysis of circularized padlock probes

As the RCA is a linear signal amplification reaction, it should be possible to measure accurately the amount of circles present in a given sample. To investigate this we made a dilution series of preformed ppWT probe circles, and used these as templates in an RCA reaction. Each reaction also contained a fixed amount of circularized ppMUT probe serving as an internal quantification reference to standardize quantifications in separate experiments. The RCA of the ppWT was monitored for HEX fluorescence using the mbRNAhex molecular beacon, while the reference probe circle templates an RCA that was monitored using the mbRNAfam molecular beacon labeled with a separate fluorophore (FAM). The increase in HEX and FAM fluorescence was followed in real-time and the maximal slope from the two molecular beacons ($D_{\text{max,HEX}}$ and $D_{\text{max,FAM}}$) was determined in each reaction. Figure 4A shows the increase in HEX fluorescence from a subset of these reactions. The slope of the fluorescence increase was steeper for RCA reactions templated by higher concentrations of probe circles. As demonstrated in Figure 4B the $D_{\text{max,HEX}}$ signal increases in proportion to the number of probe circles in the reaction. By normalizing the $D_{\text{max,HEX}}$ signal to the $D_{\text{max,FAM}}$ signal in the same reaction, the proportionality in signal response to the

Figure 3. Comparison of real-time RCA of circularized padlock probes that generate quenching or non-quenching products, and demonstration of the intramolecular quenching structure by varying the temperature or by restriction digestion of the RCA products. (A) A real-time fluorescence measurement (left) was followed by a temperature gradient to study the temperature dependence of the fluorescence (right). Analysis of the fluorescence temperature profile demonstrates the inter-molecular quenching structure. RCA was performed of padlock probes subjected to ligation (filled symbols) or no ligation (open symbols). The RCA reactions were templated by 10 fmol of either the pp90 (blue squares) or the pp93 (red circles) padlock probes, producing quenching and non-quenching RCA products, respectively. In the left portion of the graph the fluorescence from the mbRNAhex molecular beacon was followed in real-time at 37°C. The right portion shows the fluorescence temperature profile of the products at the end of the RCA. (B) Temperature profiles of the products generated in (A) after restriction digestion. (C) Temperature profiles of the mbRNAhex molecular beacon in presence of no hybridization target (black rectangles), or hybridization targets that in addition to the loop sequence also hybridizes to the quencher-carrying stem sequence (QUENCH, green circles), the fluorophore-carrying stem sequence (FLUO, red circles), or none of the stem sequences (NONE, blue squares).
number of probe circles in the reaction improved, and the experimental variability was reduced. Figure 4C shows that the $\Delta_{\text{max}}\text{HEX}/\Delta_{\text{max}}\text{FAM}$ signal ratio is proportional to the amount of probe circles in the reaction over a $>100$-fold concentration range with an average CV of 1.7% (based on 12 triplicate measurements with a range of 0.27–3%).

**DISCUSSION**

We show herein that molecular beacons can be used to monitor RCA reactions in real-time, allowing highly quantitative measurements of the number of circles templating the RCA. However, the molecular beacon design had to be modified in two ways to obtain reliable results.

First, the molecular beacon had to be resistant to the 3’ exonucleolytic activity of the polymerase. In this study we used the $\Phi 29$ DNA polymerase as it is a very efficient RCA polymerase (9), but it also has a very strong 3’ exonucleolytic activity (22). We found that by replacing all DNA residues in the molecular beacon with 2’-O-Me-RNA residues, the degradation of molecular beacons was avoided. More recently, we have found that it is sufficient to replace the three 3’-most residues in a 5 bp stem with 2’-O-Me-RNA to obtain full protection of molecular beacons. We have further found that also DNA molecular beacons with a 5 bp stem is degraded by the exonucleolytic activity of the polymerase, so the protection of the molecular beacons seems to be due to the 2’-O-Me-RNA residues as such, and not due to the shorter stem used in the modified molecular beacons compared with the DNA molecular beacons (F. Dahl, unpublished results).

Secondly, one of the stem sequences should be designed to hybridize to the RCA product in order to avoid inter-molecular quenching between neighboring probes (Fig. 2). Preferably the fluorophore should be connected to the non-target complementary stem sequence to avoid interactions between the fluorophore and the target sequence (Fig. 3C). In this paper the fluorophore was attached to the target-complementary stem sequence, thus the results would probably improve by using the other configuration. We are currently successfully using molecular beacons of both configurations (F. Dahl et al., unpublished results). The extent of inter-molecular quenching between neighboring molecular beacons of the traditional design probably varies with the stability of the stem sequences, the distance between the molecular beacons, and the temperature during the polymerization reaction. One could probably find conditions under which traditional molecular beacons are not quenched when hybridizing to the RCA products. However, by using the present design, the inter-molecular quenching can be avoided completely, without optimizations.

Precise quantification and specific recognition are important for accurate measurements and comparisons of mRNA expression levels. Therefore, the combination of the specificity obtained using padlock probe circularization and the precise quantitative detection described herein should prove useful for quantification of cDNA or mRNA sequences. The coefficient of variation in signal (1.7%) in our experiments corresponds to a variation in circularized probes of 2.7% (range 0.45–5.2%). The quantitative precision was greatly increased by introducing an internal reference circle that reports in a separate color. Such an active reference can

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**Figure 4.** Real-time quantification of probe circles using RCA and molecular beacons. (A) HEX generated fluorescence from mbRNAhex molecular beacons plotted against time from RCA reactions containing 0 (orange), 0.87 (green), 8 (red), 50 (blue) and 100 (black) fmol ppWT probe circles, respectively. (B) $\Delta_{\text{max}}\text{HEX}$ (red circles) and $\Delta_{\text{max}}\text{FAM}$ (green triangles) signal values from the mbRNAhex and mbRNAfam molecular beacons, respectively, plotted against the amount of ppWT probe circles present in triplicate RCA reactions containing 30 fmol ppMUT reference circles. The corresponding $\Delta_{\text{max}}\text{HEX}/\Delta_{\text{max}}\text{FAM}$ signal ratios were calculated for the individual reactions and plotted as blue diamonds. (C) The $\Delta_{\text{max}}\text{HEX}/\Delta_{\text{max}}\text{FAM}$ signal ratio is plotted against the amount of ppWT probe circles (HEX signal) present in RCA reactions containing 30 fmol ppMUT reference circles (FAM signal). The data points are from reactions containing 0.87, 1, 7, 8, 33, 38, 44, 50, 87, 100, 130 and 150 fmol of ppWT probe circles. The error bars represent the standard deviation of triplicate reactions at each concentration of ppWT probe circles.
normalize for differences in reaction and detection conditions among reaction wells. We noticed further that the linear range of quantification increased with the use of an internal control. This effect was due to that at the highest concentrations of HEX probe circles, the relatively small increase of $\Delta_{\max}$HEX signal was compensated by a decrease of the corresponding $\Delta_{\max}$FAM signal, resulting in an enhanced increase of the $\Delta_{\max}$HEX/$\Delta_{\max}$FAM signal ratio (Fig. 4B). For quantitative studies of specific nucleic acid sequences, a reference circle can be added to a sample in a known amount for absolute measurements, or a control probe can be circularized by detecting a reference nucleic acid sequence for relative measurements.

As concentration differences of 15% can be reliably distinguished (Fig. 4), the precision in quantification compares advantageously with that of the Taqman assay, currently the most widely used quantitative RT–PCR method. Interestingly, the precision of quantification we report is similar to that of the recently described quantitative invasive cleavage assay (23), which also represents a linear amplification technique, in contrast to the exponential amplification obtained in the Taqman assay. Therefore, the limited precision of quantification in the Taqman assay may be a consequence of the exponential mode of amplification. The sensitivity and dynamic range of the Taqman assay is however superior to the method described here. We are currently developing means to greatly increase the sensitivity of the RCA with preserved quantification accuracy (F.Dahl et al., in preparation).

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