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A netlike rolling circle nucleic acid amplification technique

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A nucleic acid amplification technique termed as netlike rolling circle amplification is proposed by introducing a nicking enzyme into the existing hyperbranched rolling circle amplification system. Surprisingly dense and uniform network morphology is observed; and cubic amplification is achieved for the sensitive detection of a sequence from HIV.

Rolling circle amplification (RCA) is one of the mostly used isothermal amplification technology. It utilizes circular single-stranded DNA probes as everlasting templates for DNA polymerization to generate multiple single-stranded linear copies of the original DNA in a continuous head-to-tail series. The first generation of RCA also known as linear RCA (LRCA) results in linear growth of the products with up to several thousand fold amplification (Scheme 1a). Subsequently, several modified RCA techniques have been developed to improve the amplification efficiency. The most representative one is the hyperbranched rolling circle amplification (HRCA), also referred to as ramification amplification RCA. In addition to the linear extension around the circular probes, strand displacement and ramified extension proceed simultaneously in HRCA by adopting a second reverse primer (Scheme 1b). As a result, quadratic (also known as exponential) amplification is achieved with the amplification increasing to million fold, which may rival PCR.

As the growth in demand for the detection of ultra-trace level nucleic acids like microRNA, pathogenic DNA, and mutation in cancer cells, higher amplification is greatly and even urgently required. Here, by introducing a nicking enzyme into HRCA system, we have developed a novel netlike rolling circle amplification (NRCA) technique (Scheme 1c). Under the catalysis of a nicking enzyme, the tree structure of HRCA is scissored into branches (short ssDNA segments), which subsequently work as seeds (probes) for the growth of other HRCA trees. The process is similar to the plant breeding using cuttage method. Thus, a cascade of primer extension, strand displacement, and nicking reactions is integrated in a tube to achieve cubic amplification (Scheme 1c). In contrast to the linear amplification of LRCA and the quadratic amplification of HRCA, the cubic amplification of NRCA shows ultrahigh efficiency without sacrificing the time, labor, and cost required. The overlap of the NRCA product may finally result in a netlike or forest structure, while in the case of LRCA and HRCA, the corresponding structure is just like a trunk and a tree, respectively. The network of NRCA has been verified with atomic force microscopic imaging. As far as we know, it is the first report of dense and uniform network morphology from isothermal nucleic acid amplification. Sensitive detection of HIV-1 DNA and even single-nucleotide polymorphism (SNP) has also been achieved by adopting this NRCA technique.

Scheme 1 Schematic illustration of the principle of LRCA, HRCA, and NRCA. For NRCA, the tree structure of HRCA is scissored into branches (B1, B2, and etc.) under the catalysis of nicking enzyme. Each branch may work as a seed for the growth of another HRCA tree.

The experimental details including the sequences of the adopted oligonucleotides and the optimization of the experimental conditions have been shown in ESI† (Table S1, Fig. S1 and S2). Since amplification efficiency is the most important evaluation index of a
nucleic acid amplification technique, comparison of the amplification efficiency of NRCA with LRCA and HRCA is first conducted. In the case of a relatively high concentration of the primer DNA (primer 1, 10 nM ~ 200 nM), agarose gel electrophoretic results of the products of RCAs (NRCA and HRCA) have clearly shown that NRCA has higher amplification efficiency (Fig. S3, ES†). While the concentration is further increased to 1 μM, no remarkable difference is observed maybe owing to the saturation of both amplified products. Electrophoresis is visible but not sensitive enough. So, we have also conducted real-time monitoring of the fluorescent signals of NRCA as well as HRCA and LRCA by using a real-time PCR detection system. As are shown in Fig. 1a-c, the fluorescent signals increase along with the increasing concentration of the primer 1 in all the three cases. However, the magnitudes of the growth differ apparently with a gradient NRCA > HRCA > LRCA. Because the comparison of amplification efficiency under a low concentration is more meaningful for practical application, we then pick the curves of 1 pM of the primer 1 to show the differences (Fig. 1d). The stable fluorescent signals of NRCA, HRCA and LRCA for the amplification of 1 μM of the primer 1 reach 1848, 137, and a negligible quantity, respectively, showing a greatly different amplification efficiency. Certainly, the result of LRCA may not reflect the real amplification efficiency, since the products of LRCA are linear single-stranded DNA and the non-specific fluorescent dye SYBR Green I intercalates mostly with double-stranded DNA. Nevertheless, the huge gap between NRCA and HRCA still makes sense.

While in use, RCA technique is usually combined with a padlock probe, i.e. a linear DNA probe that becomes circularized upon recognition of a target nucleic acid sequence. The target-binding region of the padlock probe is equally split into two segments placed in opposite orientation at 5′- and 3′-ends. When hybridized head-to-tail with the target DNA, the ends are positioned adjacent and are sealed by DNA ligase, resulting in a circular probe with two ends connected. Non-circularized probe as well as the target DNA is removed by exonuclease(Exo) treatment, while the circularized probe may participate in the RCA system subsequently. It is also reported that the step of Exo treatment can be skipped, since the non-circularized probe cannot serve as the template for continuous extension. Here, the padlock probe is combined with our NRCA system to detect a 21-mer ssDNA from the HIV-1 U5 long terminal repeat (LTR) sequence. The schematic illustration of the well-designed padlock probe is shown in Scheme 2. Strategies either with or without Exo treatment are conducted, which have also been illustrated in Scheme 2.

**Scheme 2** Schematic illustration of (a) the functional regions of the padlock probe, (b) the principle of the padlock probe-based NRCA for the detection of the target. Fluorescence spectrum analysis, which is more sensitive than real-time PCR in the acquisition of fluorescent signals, is adopted as the detection tool. As is shown in Fig.3a and 3c, no matter Exotreatment is conducted or not, the fluorescence intensities of the dye-
intercalated NRCA products increase remarkably with the increase of the concentrations of the target DNA. In both cases, the relationship between the fluorescence intensities and the logarithm of the concentrations of the target DNA can be linearly fitted (Fig. 3b and 3d). The fitting equations are shown as following:

\[
Y = 12777 + 5610X (R^2 = 0.992) \quad \text{(no Exo treatment)}
\]

\[
Y = 10438 + 5122X (R^2 = 0.996) \quad \text{(Exo treatment)}
\]

The detection limits reach 5.4 aM (ca. 65 molecules, S/N = 3, no Exo treatment) and 9.3 aM (ca. 112 molecules, S/N = 3, Exo treatment), respectively. The little variation with and without Exo treatment may be ascribed to the fact that Exo treatment reduces the circular probe-independent polymerization (e.g. polymerization using linear probe as the template), and consequently results in the slight depression of the fluorescent signals and the sensitivity. The detection ranges both expand from 10 aM to 1 μM, a range across twelve orders of magnitude. Thereby, successful detection of the target DNA with outstanding performance is achieved.

![Fig. 3 Fluorescent detection of HIV-1 DNA by using the padlock probe-based NRCA. (a) The emission spectra of the dye-intercalated products of the padlock probe-based NRCA in the presence of different concentrations of the target HIV-1 DNA. The excitation wavelength was fixed at 470 nm. No Exo treatment was performed. (b) The relationship between the fluorescence intensity at 520 nm of (a) and the corresponding concentration of the target DNA. (c) The same as (a) with the exception that Exo treatment was performed. (d) The relationship between the fluorescence intensity at 520 nm of (c) and the corresponding concentration of the target DNA.](image)

Non-specific amplification is one major problem of the currently used amplification techniques. It is very interesting that the specificity of our NRCA technique is much better than that of HRCA. While in the detection of HIV-1 DNA, a clear discrimination of even SNP is achieved.

Fig. 4a shows the comparison of NRCA with HRCA by using the primer 1 as well as its mutant variants. Taking a 5-bases mutation of the primer 1 (mp-5) for instance, no ladder-like band is observed for NRCA whereas shallow bands still exist in the case of HRCA. We have also studied the differences in details through the image analysis of the gray value of the bands using Photoshop and ImageJ software. For NRCA, the gray values of 1-base, 3-bases, and 5-bases mutations of the primer 1 (mp-1, mp-3, and mp-5) decrease by 7%, 75%, and 100%, respectively, in comparison with that of the primer 1 (the gray value of the background has been subtracted) (Red columns in Fig. 4d). As for HRCA, the decline is 3%, 51%, and 83%, respectively (Green columns in Fig. 4d). The enhanced specificity of NRCA may be owing to the superimposition of the specificity of both the polymerase and the nicking enzyme.

As for the detection of the target HIV-1 DNA using padlock probe, the specificity of the padlock probe-based NRCA (P-NRCA) system is further improved. It has been known from Fig. 4a (ESI†) that the target-dependent circularization of the padlock probe (i.e. the ligation-based ligation reaction) has certain specificity. No circular probe can be formed when a 5-bases mutation of the target DNA (mT-5) or a totally unmatched variant (uT) instead of the target DNA is adopted (Fig. S5, ESI†). However, the specificity is not enough for SNP discrimination, since that a little amount of circular probe still forms in the case of 1-base or 3-bases mutations of the target DNA (Fig. S5, ESI†). While the circularization of the padlock probe is combined with our NRCA, it is expected that the specificity of each part can be integrated to present a better performance. As is shown in Fig. 4b, regardless of the Exo treatment, clear ladder-like band is observed only in the case of the target DNA, suggesting an outstanding specificity for SNP discrimination. The gray value of the 1-base mutation of the target decreases by 93% and 96% for the absence and presence of Exo, respectively (Fig. 4d). As comparison, if the circularization of the padlock probe is combined with HRCA, the improvement of the specificity is limited. The 3-bases mutation of the target DNA still cannot be well discriminated (Fig. 4c). Detection of single-nucleotide polymorphism has been widely reported. Nevertheless, the performances differ a lot, mainly in the discrimination of the signals of the target and the 1-base mutation. In our results, the signal intensity of the target is over 14 times stronger than that of the 1-base mutation (Fig. 4), suggesting a high performance over many reports.

![Fig. 4 Comparison of the specificity of NRCA with HRCA. (a) Agarose gel electrophoretic patterns of the products of NRCA and HRCA. Primer 1 and its mutant variants were adopted. “N”, “P-1”, “mp-1”, “-3”, “-5”, and “uP” stand for no primer 1, primer 1, 1, 3, 5-base(s) mutant and unmatched variants of primer 1, respectively. (b) Agarose gel electrophoretic patterns of the products of P-NRCA. The target DNA and its mutant variants were adopted. The meaning of the abbreviations is the same as that in Fig. 4a. Lane 2-7: without Exo treatment; lane 9-14: with Exo treatment. (c) The same as (b) with the exception that P-HRCA was performed here. (d) Image analysis of the relative gray value of the ladder-like bands from (a)-(c). The whole ladder-like bands of a single lane were selected for the statistics of the gray value. The background gray value of the gel was subtracted and set as 0, while the gray values of all the bands from different lanes were relative to that of the target DNA in percentage terms.](image)
usability. 2) Unlike some other cascade amplification strategies which need step-by-step procedure, the three amplification reactions of NRCA (i.e. primer extension, strand displacement, and nicking reactions) proceed simultaneously in a tube. Thus, it also has the advantages of time-saving and labor-saving. 3) It is noted that the group of Komiyama once developed a primer generation-rolling circle amplification (PG-RCA) technique. In their scheme, a nicking enzyme is also introduced. Nevertheless, nicking enzyme is used on the basis of LRCA, so only primer extension and nicking reactions, without strand displacement, are integrated, thus quadratic amplification instead of cubic amplification can be achieved. So, the sensitivity of the PG-RCA technique is only comparable to HRCA. Comparing our NRCA with PG-RCA proposed by Komiyama et al., besides the gap between cubic amplification and quadratic amplification, NRCA also has the advantage of the production of dsDNA, which may facilitate the sensitive real-time fluorescent detection. 4) Besides the efforts on the development of various RCA techniques to improve amplification efficiency, the improvement of specificity is also of great importance. Our results have shown that NRCA may provide a better specificity than HRCA. Moreover, when combined with the padlock probe for the practical detection of a target DNA, the specificity of the whole P-NRCA system is further improved. Detection of SNP can be achieved with 25-fold difference in the amplification products between the target and the 1-base mutant variant. To our knowledge, few existing strategies for gene assays may reach such high amplification efficiency while achieving the outstanding specificity at the same time.

The RCA technology has been widely used in the amplified detection of miRNA gene sequences, DNA methylation, and the targets of nucleic acid aptamers. Various detection methods including fluorescence spectrometry, UV-vis spectrometry, and electrochemistry have also been adopted. Benefited from the high performance and easy-operation of NRCA, rational design further will no doubt contribute to a wide application of this novel RCA technique. On the other hand, it is noted that there are still some problems to be solved. For example, the mechanism of the effect of the buffer solutions is still unknown; whether other nicking enzymes can be adopted should be explored further; it is also unknown why the specificity of P-NRCA can be improved so much. So, a lot of work still lies ahead to make the mechanism clear and to expand the application.

Conclusions

In summary, we have proposed a NRCA technique in this work. Simply upon addition of a nicking enzyme into a well-established HRCA system, amplification of nucleic acid with super-high efficiency and outstanding specificity is achieved. It is the first time that RCA is upgraded from linear amplification (LRCA in 1995) and quadratic amplification (HRCA in 1998) to cubic amplification. In comparison with LRCA, HRCA, and some other existing DNA amplification strategies, the NRCA technique shows some obvious advantages. It has also been successfully applied for the detection of a HIV-1 DNA with a detection limit lowered to 5.4 aM and a detection range extended from 10 aM to 1 μM. Moreover, dense and uniform network morphology of RCA products is observed for the first time, which validates the “netlike” and also does favour to the understanding of RCA technology.

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