Performance of probe polymerization-conjunction-agarose gel electrophoresis in the rapid detection of KRAS gene mutation

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Abstract

This study aimed to develop a simple and rapid method to detect KRAS gene mutations for conventional clinical applications under laboratory conditions. The genotype of mutation sites was determined based on the occurrence of target bands in the corresponding lanes of the reaction tubes through polymerization-conjunction of the probes, probe purification and amplification, and agarose gel electrophoresis. Circulating DNA samples were obtained from the plasma of 72 patients with lung cancer, which were identified based on six mutation sites (G12S, G12R, G12C, G12D, G12A, and G12V) of codon 12 of the KRAS gene. The detection results were compared with direct sequencing data. The proposed detection method is characterized by simple operation, high specificity, and high sensitivity (2%). This method can detect the mutations of three samples at G12S, G12R, and G12A. In the direct sequencing spectra of these samples, the genotype could not be determined due to the lack of evident sequencing peaks that correspond to the basic group of mutations. In conclusion, a simple and rapid method was established based on probe polymerization-conjunction-agarose gel electrophoresis for detecting KRAS gene mutations. This method can be applied to the conventional mutation detection of inhomogeneous samples.

Keywords: Mutation, circulating DNA, polymerization-conjunction reaction, agarose gel electrophoresis, K-ras.

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Introduction

The KRAS gene is a common oncogene in humans, and encodes the 21-kDa RAS protein. The RAS protein exhibits intrinsic guanosine triphosphate (GTP) enzymatic activity, and is involved in cell proliferation, differentiation, and apoptosis by switching the mutual transformation regulatory signal system between guanosine diphosphate (GDP) and GTP. The KRAS gene has a high mutation rate in patients with colorectal cancer and lung cancer, and confers the resistance to epidermal growth factor receptor (EGFR), tyrosine kinase inhibitors (TKIs), and EGFR monoclonal antibody agents (Chen et al., 2013; Li et al., 2014; Leis et al., 2015; Hsu et al., 2016). Reports indicate that KRAS gene mutations in patients with non-small cell lung cancer are related to short survival time and poor prognosis (Li et al., 2016; Tao et al., 2016). The mutation rate of the KRAS gene significantly differs among various types of tumors and populations. For example, the mutation rate of the KRAS gene can reach as high as 15–30% in Caucasian patients with colorectal cancer, and only 4–10% in Asian patients with lung cancer (Mascaux et al., 2005; Lu et al., 2013; Kinugasa et al., 2015; Omidifar et al., 2015; Ohba et al., 2016). Approximately 80% of KRAS gene mutations occur at codon 12, and the remaining mutations take place mainly in codons 13 and 61 (Mascaux et al., 2005). Hence, the detection of KRAS gene mutations is important for the development of individualized treatments for patients with tumors, as well as to improve the effects of targeted clinical treatments and reduce treatment expenses.

KRAS gene mutations can be detected using various detection methods, including restriction fragment length polymorphism (RFLP) (Dobre et al., 2013), DNA sequencing (Kinugasa et al., 2015), capillary electrophoresis (Zhang et al., 2013), high-resolution melting (HRM) curve analysis (Heideman et al., 2009; Trujillo-Arribas et al., 2016), mass spectroscopy (Choi et al., 2014; Kriegsmann et al., 2015), pyrosequencing (Altimari et al., 2013; Vincenzi et al., 2015; Mack et al., 2016; Lee et al., 2016), and ARMS real-time PCR (Hamfjord et al., 2011; Zhang et al., 2015). DNA sequencing is a traditional detection method that has
low sensitivity. Pyrosequencing, HRM, and RT-PCR techniques exhibit high sensitivity, but also have high requirements in terms of experimental apparatus and conditions. These methods cannot be applied to conventional clinical detection under simple experimental conditions. In the present study, a simple, rapid, and sensitive method for detecting KRAS gene mutations was established based on probe polymerization-conjunction–agarose gel electrophoresis (PPC-AGE). This method was used to detect KRAS gene mutations at the six sites (G12S, G12R, G12C, G12D, G12A, and G12V) of codon 12.

Materials and Methods

Plasmid template and clinical samples
Plasmids that carry the wild and mutation alleles (G12S, G12R, G12D, and G12A) of codon 12 of the KRAS gene were constructed according to PCR-mediated site-specific mutagenesis technology. Plasma samples were collected from 72 patients with non-small cell lung cancer, who were admitted in the Affiliated Hospital of Hubei University of Arts and Science and Zaoyang Clinical College. Circulating DNA samples were extracted according to the instruction of the DNA extraction kit (QIAamp DNA Blood Mini Kit, Qiagen), and stored at -20°C for detection of KRAS gene mutations.

Detection probe

As shown in Figure 1, the mutations of codon 12 were detected using two pairs of oligonucleotide probes (P-12-1R/P-12-1L and P-12-2R/P-12-2L). P-12-1R and P-12-1L were used to detect mutations at G12S, G12R, and G12C, while P-12-2R and P-12-2L were used to test mutations at G12D, G12A, and G12V. Each probe pair comprises of five parts: homologous hybridization sequences (H1 and H2) with the template, general amplification primer sequences (Tag1 and Tag2), and the 5'-end biotin labeling of probes P-12-1L and P-12-2L. The oligonucleotide probe sequences were as follows:

P-12-1L: 5’-biotin-GGGTTCGTGGTAGAGCGTCGGAGTACTCTTCTGCTAGCCAC-3’;
P-12-1R: 5’-AGCTCCAACTACCACAAGTGGCTGCTATCTGCAGCGGTGTG-3’;
P-12-2L: 5’-biotin-GGGTTCGTGGTAGAGCGTCGGAGTAGCTCTTGCCTAGCCAC-3’;
P-12-2R: 5’-CAGCTCCTAATACCATCAGAGATGCAGCTGCTAGCCAC-3’.

Tag1 sequences and their sequences complementary with Tag2 (C Tag2) were as follows:

Tag1: 5’-GGGTTCGTGGTAGA GCGTCGGAGTACCTCTTGCCTAGCCAC-3’;
C Tag2: 5’-CCAGACGACACCGAGATAGCAGCCACTCTTGCCTAGCCAC-3’.

The oligonucleotide probes were synthesized by Shanghai Sangon Co., Ltd.

KRAS gene amplification

Exon 2 of the KRAS gene was amplified by PCR. The forward and reverse primers used were as follows: 5’-TAAGCGTCGATGGAGGAGTT-3’ and 5’-CATCATTGGACCCTGACATAC-3’, respectively. The amplification system comprised of 5 μL of the circulating DNA sample, 2.5 μL (10 pmol/μL) of each amplification primer, 5 μL of deionized water, and 15 μL of 2x Taq PCR Mastermix. The amplification conditions were as follows: 32 cycles of 95°C for five minutes, 95°C for 45 seconds, 54°C for 45 seconds, 72°C for 90 seconds, and 72°C for four minutes. The amplified DNA sequences were extracted (SanPrep Column PCR Product Purification Kit, Shanghai Sangon Biotech) and stored at -20°C. The amplification products were subjected to the direct sequencing method.

Polymerization and conjunction

P-12-1R and P-12-1L corresponded to a group of four reaction tubes (“G1,” “A1,” “C1,” and “T1”), and P-12-2R and P-12-2L to another group of four reaction tubes (“G2,” “A2,” “C2,” and “T2”). These eight reaction tubes were added with 28 μL of the reaction buffer (20 mM of Tris-HCl (pH 7.6), 25 mM of KAC, 10 mM of MgAC2, 10 mM of DTT, 1 mM of NAD, and 0.1% Trion-x100), 50 fmol of detection probes, and 5 fmol of template sequences. Next, the tubes were added with 1 μL of deoxyribonucleotide (25 μM) bases that are complementary with the labeling of different tubes (for example, dATP was added to tube T, and dCTP was added to tube G). These tubes were processed at 95°C for three minutes and at 49°C for five minutes. Subsequently, 1 μL of enzyme solution (1 unit Pu of DNA polymerase (Shanghai Sangon, Co., Ltd., China) and 2 units of...
Taq DNA ligase (New England BioLabs, USA) were added. Then, the tube was again processed at 49°C for five minutes and at 98°C for 10 minutes. All reaction tubes were cooled in ice bath.

Purification of the conjunction-dependent probe

Conjunction products in the different reaction tubes were purified through the magnetic particles of streptavidin (Xi’an GoldMag Nanobiotech Co. Ltd., China). These purified magnetic particles were washed by aqueous alkali (0.1 M of NaOH) to eliminate the non-specifically adsorbed nucleotide sequence. Finally, the magnetic particles were stored in Tris-HCl buffer solution (10 mM, pH 7.5).

Amplification and detection

The conjunction-dependent probes that bound to the streptavidin magnetic beads were subjected to PCR amplification. A 20-μL reaction mix contained 5 μL of the magnetic particle template, 2.5 pmol of Tag1 and CTag2 primers, and 10 μL of 2 × Taq PCR MasterMix (Tiangen Co. Ltd., China). The reaction conditions were as follows: 95 °C for 30 s followed by 28 cycles of 95 °C for 25 s, 60 °C for 45 s, and 72 °C for 20 s, and 72 °C for 1 min. The PCR products were placed in different tubes and used for 3.5% agarose gel electrophoresis (AGE). The mutation detection results could be judged by the appearance of targeted detection fragments in the corresponding reaction tubes. For example, if the targeted band only appears in lanes corresponding to “G1” and “G2,” it is considered a wild template (the base sequence of codon 12 is GGT). If the targeted band appears in lanes corresponding to “G1,” “G2,” and “C1,” the template of codon 12 has the heterozygosis mutation of G12R (the base sequence of codon 12 is CGT/GGT). If the targeted band only appears in lanes corresponding to “C1”, the template of codon 12 has the homogenous mutation of G12R (the base sequence of codon 12 is CGT). The rest could be judged in the same manner.

Results

Detection flow

As shown in Figure 2, the entire detection process includes polymerization/conjunction reaction (PC reaction), purification, amplification, and detection. During the PC reaction, each pair of detection probes corresponds to the C, T, G, and A reaction tubes. After all reaction systems were heated and denatured to the annealing temperature, gaps corresponding to the detected mutation base were formed among different probe pairs. Each deoxyribonucleotide was correspondingly added to different reaction tubes (for example, dATP was added to tube T and dCTP was added to tube G). When the added bases were complementary to the detection templates, the corresponding pair of detection probes was connected to the conjunction-dependent probes under the effect of DNA polymerase and ligase. However, these detection probes were not connected when the added base mismatched with the bases at the mutation sites of the template. After the PC reaction, the probes were purified using streptavidin magnetic particles. The conjunction-dependent probes were purified on the magnetic particle surface, and were used as a template. All purified products were amplified by PCR using Tag1 and CTag2. Then, the amplification products were subjected to 3.5% AGE. The mutation type was judged by the appearance of targeted bands in the corresponding lanes of the reaction tubes.

Optimization of hybridization temperature

The annealing temperatures of P-12-1L (P-12-2L), P-12-1R (P-12-2R), and of the template ranged from 49 °C to 51 °C. The annealing temperature of the PC reaction was optimized using three temperature gradients: 44 °C, 49 °C, and 54 °C. The wild plasmid and mutation plasmid templates of G12R were used as detection objects. The AGE results are shown in Figure 3. At 44 °C and 49 °C, the amplification bands were clearly observed in lanes corresponding to tube C; the remaining lanes did not exhibit amplification bands. When the annealing temperature increased to 54 °C, all lanes had no amplification bands. This finding shows that the annealing ability of probes on the detection template is negatively correlated with annealing temperature. The detection probes could be annealed to the template from 44 °C to 49 °C, in order to obtain the conjunction reaction. However, these probes could not be annealed to the template at temperatures higher than 54 °C.
due to the absence of targeted amplification bands. Hence, 49 °C was selected as the optimal temperature to achieve an effective annealing of probes on the template and a specificity of the PC reaction.

**Specificity of detection**

The specificity of the proposed detection method was tested by determining the cycle number in the amplification reaction. Equal proportions of wild plasmid template and mutation plasmid template (G12S or G12D) were mixed, and the cycle number of the amplification reaction was set to 25 and 35. Approximately 5 μL of the amplification products were collected from each reaction tube and used for 3.5% AGE (Figure 4). Under two circulation conditions, the wild-type and mutation plasmids of G12S, P-12-1L, and P-12-1R corresponded to G1, A1, C1, and T1 reactions. Obvious bands were detected in lanes corresponding to G1 and A1. In addition, P-12-2L and P-12-2R corresponded to the G2, A2, C2, and T2 reactions, and an obvious band was detected in the lane corresponding to G2. Concerning wild and G12D mutations, an obvious band was detected in lanes corresponding to G1, G2, and A2. Meanwhile, nonspecific bands in other lanes were absent for 35 cycles. Hence, the proposed detection method exhibits high specificity and amplification volume.

**Sensitivity test**

Different proportions of wild plasmid and mutation plasmid templates (G12S) were mixed. The proportions of the mutation plasmid template in the total detection template were set to 50, 10, 5, and 2% (Figure 5). Under all proportions of the mutation plasmid templates, amplification

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**Figure 3** - Optimization of hybridization temperature. G, A, C and T represent the four reactions corresponding to each pair of detection probes.

**Figure 4** - Specificity detection with the PCR cycle numbers. G, A, C and T represent the four reactions corresponding to each pair of detection probes.

**Figure 5** - Sensitivity detection with agarose gel electrophoresis. The proportions of mutated alleles in the total plasmid DNA were 50, 10, 5 and 2%. G, A, C and T represent the four reactions corresponding to each pair of detection probes.
bands were found in lanes corresponding to C and T. With the reduction of the mutation allele (G12S), amplification bands in lanes corresponding to T darkened. However, amplification bands were still detected in lanes corresponding to T when the proportion of the mutation allele was lower than 2%. This finding shows that the proposed method exhibits high sensitivity, and that it can be applied to detect mutation alleles in inhomogeneous samples.

Sample detection

The mutations of codon 12 of the KRAS gene in 72 patients were detected using the proposed method. The detection results were compared with the direct sequencing data (Figure 6A,B). Mutations were detected in three samples: the G12S (GGT/AGT) mutation of the #17 sample, the G12R (GGT/CGT) mutation of the #51 sample, and the G12A (GGT/GCT) mutation of the #38 sample. However, these direct sequencing results did not exhibit obvious sequencing peaks corresponding to mutation bases at codon 12 of the KRAS gene. Impure peaks corresponding to mutation bases were found in #17, #51, and #38 samples, and almost overlapped with the background peak. Therefore, determining the genotype according to direct sequencing results alone is difficult. These results confirm that the proposed method is superior to direct sequencing in terms of its sensitivity and applicability for detecting inhomogeneous samples.

Discussion

The present study established a simple and fast method for KRAS gene mutation detection based on the method for detecting EGFR gene mutations (Tang et al., 2014). The proposed detection method can detect six kinds of mutation behaviors of codon 12 through eight reactions, as well as the mutation behaviors of codons 13 and 16 using the same principle. This method is simpler and does not need expensive experimental instruments or reagents, such as fluorescent probes and high-quality detection samples, when compared to methods like fluorescent quantitative PCR, mass spectrometry, or DNA direct sequencing. The probe length is less than 50 bp, and it can detect mutation alleles using ordinary PCR instruments and AGE. Compared with the RFLP method, ours has no limitations on the enzyme cutting site and is more applicable for mutation detection under simple and conventional experimental conditions.

The high specificity of the proposed method can be maintained by: (1) using one pair of probes and four similar reaction systems, when detecting one single-base mutation site. Each reaction system was added with one dNTP to ensure that the PC reaction could be accomplished only upon complementation between the added dNTP and the base at the mutation site of the detection template. In the four reaction systems, the PCR amplification results of two systems could be used to determine the genotype of the mutation site, and the remaining reaction systems could be used as a control group to avoid non-specific results. (2) The PC re-

![Figure 6 - Mutation detection of the clinical samples. (A) The results of agarose gel electrophoresis for codon 12 of the K-ras gene. (B) Results of the direct sequencing. G, A, C and T represent the four reactions corresponding to each pair of detection probes.](image-url)
action can only be accomplished under the dual specificity of DNA polymerase and DNA ligase. (3) The Tm values of H1 and H2 of the detection probe differed by 11 °C compared with that of Tag1 and Tag2 in the amplification primers, thereby reducing the mutual influence between the connection reaction and the follow-up amplification reaction.

This method exhibits high detection sensitivity and can detect as low as 2% mutation alleles in the mixed plasmid template where wild/mutation sites are located. Therefore, the proposed method can be used to detect mutations in inhomogeneous samples, design Tag1 and Tag2 on the probe sequence, and implement the secondary amplification of products after PC reaction in order to increase their numbers, resulting in a fast detection under the AGE level. Although DNA sequencing is the gold standard of mutation detection, it exhibits low sensitivity (only 20–50%; Garcia et al., 2000; Dufort et al., 2009), which is inadequate for detecting gene mutations in tumor tissues, circulating DNA samples, and other inhomogeneous samples. In the present study, KRAS mutations in the samples could not be judged based on the sequencing spectra alone, because the sequencing peaks that corresponded to the mutation sites in the direct sequencing spectra could not be separated from the background signal. Pyrosequencing, RT-PCR, and mass spectrum detection methods exhibit sensitivity as high as 1% (Jarry et al., 2004; Dufort et al., 2009), but require an expensive detection platform, equipment, and reagents. Hence, these techniques cannot be applied under conventional laboratory conditions.

In conclusion, a method for detecting KRAS gene mutations was established based on PPC-AGE. This method can be easily operated, and is applicable for inhomogeneous samples under simple experimental conditions (e.g., ordinary nucleic acid electrophoresis and PCR equipment). Furthermore, this method does not need expensive experimental apparatuses and reagents. However, it cannot detect unknown mutation sites, and the detection sensitivity of AGE is limited. Therefore, this method cannot be applied when the total proportion of mutation alleles in inhomogeneous samples is lower than 2%. The proposed method has limited detection capabilities.

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