Isothermal Amplification of Rabies Virus Gene

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ABSTRACT. A sensitive and specific in situ amplification technique is needed to elucidate the dynamics of rabies virus in the body during the long incubation period after infection. To overcome the disadvantage of using the traditional reverse transcription (RT)-PCR in in situ studies, an isothermal nucleic acid sequence-based amplification (NASBA) technique was developed for detection of the rabies virus gene. The NASBA technique involves the use of 4 enzymatic activities to produce multiple RNA copies of the target sequence by means of double-strand cDNA intermediates under an isothermal condition without thermocycling. The amplified cDNA intermediates from the genomic RNA in the rabies virion and the total RNA in the infected cells in NASBA reaction were analyzed by Southern hybridization assays. The specific amplified products of the rabies viral gene with the expected length were detected after 8 hr of incubation in NASBA using both of the RNAs as templates. The NASBA system used in this study was less sensitive than the general RT-PCR technique. This may have been because we employed Southern hybridization for the amplified cDNA intermediates, not many RNA copies, to evaluate the NASBA results. In conclusion, we successfully amplified the rabies viral gene in the NASBA reaction under an isothermal condition. The unique character of this technique would make it particularly valuable for in situ studies not only on rabies virus but also on other RNA viruses.

KEY WORDS: isothermal amplification, NASBA, rabies virus, RT-PCR, Southern hybridization.


Rabies virus causes a fatal neurological disease in all mammals, including humans. Rabies virus belongs to the Lyssavirus genre of the family Rhabdoviridae. The genome is an unsegmented negative-sense RNA of about 12 kb, encoding five structural proteins: nucleoprotein (N), phosphoprotein (P), matrix protein, glycoprotein and large protein.

Rabies is probably the oldest recorded infection of mankind [5]. Humans and animals become infected with the rabies virus mainly by a bite from a rabid animal. The incubation period usually lasts several days to a few months, but periods of more than six years have been reported in exceptional cases [17]. A small number of rabies virus particles or viral genes is probably present in a living body during the incubation period. It is not clear what happens when the virus invades and replicates at the biting site and in the brain. This can only be determined by using a specific and sensitive method for detection of a single copy of the rabies gene in each cell. One of the most effective methods is in situ gene amplification [14]. In situ reverse transcription (RT)-PCR was developed for detection and localization of viral RNA in formalin-fixed tissue or cultured cells [1, 2, 13]. However, there are problems such as destruction of cell morphology, high background, and diffusion of the amplified product [13]. It is thought that the high temperature required for thermocycling in PCR often causes cell damage, creation of diffused products, and prevention of accurate localization in fixed tissue or cells [14].

An alternative method to RT-PCR is nucleic acid sequence-based amplification (NASBA), which is based on transcription using three enzymes, reverse transcriptase, T7 RNA polymerase and RNaseH, under an isothermal condition [3]. The NASBA technique has been successfully applied to the diagnosis of rabies [18]. Diagnosis of rabies by NASBA in clinical samples was performed using a commercially available electrochemiluminescence (ECL) system. It has also been used to detect a number of pathogens [4, 7, 8, 10–12, 20]. It has been reported that an in situ NASBA technique was useful for detecting viral RNA in measles-infected cell culture [9]. In this study, we focused on the development of an effective NASBA technique for amplification of the rabies virus gene in vitro prior to applying to in situ study.

MATERIALS AND METHODS

Cells and virus: BHK and CER [16] cells were maintained in Eagle’s minimal essential medium supplemented with 10% tryptose phosphate broth and 5% fetal calf serum. BHK cells were used for preparation of a stock of the CVS-CE strain of rabies virus. The CVS-CE strain was derived from the CVS strain after 84 passages in chick embryonic fibroblasts, as described previously [15].

Extraction of RNA: Genomic RNA of the CVS-CE strain was extracted from the stock virus using Isogen (Nippon Gene, Tokyo, Japan) and quantitated using a spectrophotometric method at a wavelength of 260 nm. Total RNA was also extracted from CER cells infected with the CVS-CE strain using Isogen.

PCR: To prepare template RHN29-T7 for NASBA, template TP8–10 for PCR, and probe RHN29 for Southern hybridization, the N gene was partially amplified by PCR with each pair of the primers using a plasmid containing the N gene of the RC-HL strain of rabies virus [6] (Fig. 1). These PCR products were purified in a low-melting-temperature agarose gel. PCR was also carried out to check the
synthesis of single- or double-stranded cDNA in the NASBA system.

**NASBA:** The NASBA method was performed as described by Guatelli et al. [7] with some modifications. The principle of the NASBA reaction is shown in Fig. 2. Basically, the target gene was brought up to a 50-µl final reaction volume containing 40 mM Tris-HCl (pH 8.0), 8 mM MgCl₂, 25 mM NaCl (or 50 mM KCl), 2 mM spermidine hydrochloride, 5 mM dithiothreitol, bovine serum albumin (BSA, 100 µg/ml), 1 mM (each) deoxyribonucleoside triphosphate, 2.5 mM (each) ribonucleoside-5-triphosphate, 31.3 U of reverse transcriptase RAV-2 (Takara, Shiga, Japan), 11.3 U of reverse transcriptase T7 (Takara, Shiga, Japan), and 4 µM each of two primers. RAV-2 has enzymatic activities of RNA- and DNA-dependent DNA polymerase and RNaseH. PCR product RHN29-T7, genomic RNA from the stock virus, or total RNA from virus-infected cells was used as a template in NASBA with each pair of the primers (Fig. 1). The reaction mixtures were incubated at 37°C for the times indicated, and the reactions were stopped by heating at 95°C for 5 min.

**Detection of amplification products:** Amplification products were detected in Southern hybridization assays using an ECL direct nucleotide labeling and detection system (Amersham Biosciences, NJ, U.S.A.) as described previously [19]. The products were separated by 1.2% agarose gel electrophoresis and were transferred to a Hybond N+ nylon membrane (Amersham Biosciences, NJ, U.S.A.). The membrane
was hybridized with probe RHN29 labeled with horseradish peroxidase. After Southern blot hybridization, the signal was detected using the ECL detection system.

RESULTS

Optimization of NASBA reaction: NASBA utilizes simultaneous reverse transcription and RNA transcription to produce multiple RNA copies by means of cDNA intermediates (Fig. 2). The double-stranded cDNAs from RNA are used to produce RNA copies. The target RNAs can further serve as templates for additional amplification of the target sequence. The synthesis of single- and double-stranded cDNAs in NASBA reaction with 25 mM NaCl or 50 mM KCl and with/without BSA (100 µg/ml) was confirmed by PCR. After 90-min incubation at 37°C under different conditions in NASBA reaction with two primers, TP8 and TP10, and 900 ng of rabies genomic RNA, PCRs with primers T7 and RHN2 and with T7 and RHN9 were carried out to confirm the presence of positivesense single- and double-stranded cDNAs, respectively (Fig. 3). In NASBA, positive-sense cDNA containing the T7 primer sequence at the 5’ end is first synthesized by reverse transcriptase from negative-sense genomic RNA using TP8 and then negative-sense RNA is removed by RNase H. This product can be amplified by PCR using primers T7 and RHN2, and the PCR product can be observed as a 536-bp band in agarose gel. As shown in Fig. 3, a single DNA product of 536 bp was detected in NASBA reaction with both NaCl and KCl and with/without BSA. The negative-sense cDNA with the T7 primer sequence is subsequently synthesized from the positive-sense cDNA using TP10. This negative-sense cDNA can be amplified by PCR using T7 and RHN9, and the PCR product can be observed as a 1,035-bp band in agarose gel. We found strong signals of 1,035 bp in NASBA-PCR reaction with BSA, a weak signal with NaCl and no BSA, and no signal with KCl and no BSA. Since the results indicated that BSA was essential for synthesis of double-stranded cDNA, we decided to use NASBA reaction buffer containing NaCl and BSA.

NASBA reaction was carried out using 50 pg of cDNA fragment RHN29-T7 as a template at 37°C for 2 to 6 hr. Those products were analyzed in Southern hybridization assays using probe RHN2–9 in the ECL system (Fig. 4). An intensified signal of 308 bp was detected after 2 hr, and the signal was strongest after 8 hr. These intensified signals indicated the existence of DNA, not RNA, because those signals did not disappear following treatment with RNase A (data not shown).

Amplification of rabies gene by NASBA: Genomic RNA extracted from the rabies virion was amplified by NASBA using primers TP8 and PN366. The NASBA reaction was carried out using 900 ng of genomic RNA for 30 min to 22 hr. In Southern hybridization assays, a signal of approximately 720 bp was detected after 2 hr, and the intensity of the signal increased with time (Fig. 5). The expected signal of approximately 1,220 bp appeared after 8 hr. A specific broad band of 720 to 1,220 bp was detected at 22 hr. These NASBA products were not detected in agarose gel stained with ethidium bromide (data not shown).

Total RNAs from rabies- and mock-infected CER cells were amplified by NASBA using TP8 and PN366 with incubation times of 2, 8 and 24 hr. The same template RNA was also used for RT-PCR with the primers RHN2 and RHN9.
These amplified cDNAs from 1,500 ng of total RNA in NASBA and RT-PCR were analyzed by Southern hybridization (Fig. 6). A weak signal of about 720 bp in length was detected after 2 hr of incubation in the NASBA reaction using rabies-infected cellular RNA and disappeared after 8 hr. The signals with expected length of 1,220 bp by the NASBA reaction were observed at and after 8 hr of incubation in rabies-infected CER cells. The signal intensity at 24 hr in NASBA was stronger than it at 8 hr. No signals were observed in NASBA and RT-PCR products using normal cellular RNA as a template by Southern hybridization analysis. The RT-PCR product with expected length of 283 bp from rabies-infected cellular RNA was detected in both Southern hybridization assays and ethidium bromide-staining gel, although the NASBA products were not directly detected at all in agarose gel (data not shown).

**DISCUSSION**

An *in situ* gene amplification technique is needed for the detection of a small number of rabies viral RNA in each cell during a long incubation period. The RT-PCR technique has been mainly used for amplification of the rabies virus gene. We propose a NASBA technique, an amplification approach that enables direct and isothermal amplification of viral RNA, as an alternative method to RT-PCR in *in situ* study. The relatively low isothermal temperature applied to NASBA allows it to be used for *in situ* amplification without disrupting the integrity of cells [3]. The NASBA technique was used in a previous study for rapid diagnosis of rabies [18]. However, since the diagnosis in that study requires use of a commercially available ECL kit, which contains standardized reagents for nucleic acid isolation, NASBA, and ECL detection, and automated reader, it is difficult to directly apply their system to *in situ* study. In the present study, we optimized the conditions of a NASBA reaction using rabies virus RNA prior to application to *in situ* study.

The amplification methodology involves the use of four enzymes, RNA-dependent DNA polymerase, RNaseH, DNA-dependent DNA polymerase and T7 RNA polymerase (Fig. 2). Two commercially available enzymes, T7 RNA polymerase and reverse transcriptase RAV-2, were used for NASBA in this study, because this reverse transcriptase has three enzymatic activities. Since there are differences in the compositions, NaCl, KCl or BSA, between the reaction buffer for each enzyme and the NASBA reaction buffer in the original report [7], different buffer conditions were tested to obtain double-stranded cDNA synthesized from the rabies viral genomic RNA. Negative-sense cDNA was synthesized only under the condition of the buffer containing BSA, while positive-sense cDNA from genomic rabies RNA was synthesized under all conditions (Fig. 3), suggesting that BSA is essential for the activity of RNaseH or DNA-dependent DNA polymerase in a NASBA reaction.

Under optimized conditions, double-stranded cDNA with a T7 promoter can produce RNA copies and subsequently proceed to the synthesis of new double-stranded cDNAs from their RNA copies in a NASBA reaction. In a preliminary test, the purified PCR product with a T7 promoter, RHN29-T7, was applied to the NASBA reaction as a template. The amplified double-stranded cDNA from RHN29-T7 in the NASBA reaction was detected in a Southern hybridization assay as shown in Fig. 4. This result suggested that new RNA copies and subsequent products, new RHN29-T7, were synthesized from the template RHN29-T7 by multienzymatic activities under the condition described above. Taken together, these results indicate that rabies viral genomic RNA can produce double-stranded cDNA with a T7 promoter and subsequent positive-sense single-stranded RNA in the NASBA reaction under the conditions used in this study.
A shorter-than-expected molecule of about 720 bp was detected after 2 hr of incubation in the NASBA reaction using both rabies viral genomic RNA and total rabies-infected cellular RNA as templates in Southern hybridization (Figs. 5 and 6). This molecule in NASBA using cellular RNA disappeared after 8 hr of incubation. In contrast, the amount of the expected molecule of 1,220 bp in length increased in the NASBA reaction using both RNAs after 8 hr. In the early stage of the reaction, double-stranded DNA with a T7 promoter that is shorter than the full-length DNA of 1,220 bp might be synthesized due to insufficient elongation of DNA by DNA polymerase and might produce the shorter-than-expected RNA and double-stranded DNA, although it is not clear why the length of the shorter DNA with T7 was 720 bp. The conversion of the shorter DNA into the full-length DNA might be due to sufficient elongation by DNA polymerase during the subsequent incubation. It has been reported that an amplified gene of less than 1,000 bp in size should be designed as a target for an effective NASBA reaction [3]. The insufficient elongation in this study may have been caused by the excessive length of a target gene. The design of a target gene of less than 1,000 bp may enable an amplified gene to be produced more effectively. These signals were thought to be specific for the rabies virus gene as well as for a previously reported amplified gene of hepatitis A virus in the NASBA reaction [11], because no signal was detected in normal CER cells.

Jean et al. [11] showed the superiority of the sensitivity of a NASBA system over RT-PCR for the amplification of viral target RNA. The NASBA system used in this study was much less sensitive than RT-PCR. A possible for this discrepancy in results is a difference in methods for analyzing NASBA results. For the analysis of NASBA products, Jean et al. employed Northern and dot hybridization methods for the amplified RNA, while we employed Southern hybridization for the amplified double-strand cDNA. It is thought that detection of the amplified RNA is needed to apply our system to diagnosis.

In summary, we successfully amplified the rabies viral gene in NASBA reaction under an isothermal condition. The unique character of this technique would make it particularly valuable for in situ study of low-level expression of rabies viral genes during the incubation period after infection. In addition, such a method could also be applied to the detection of other viral RNA genes.

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