Nucleic acid sequence-based amplification (NASBA) for the identification of mycobacteria

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Nucleic acid sequence-based amplification (NASBA), an isothermal amplification technique for nucleic acids (NA), was investigated for the species-specific identification of mycobacteria. A set of primers was selected from a highly conserved region of the 16s rRNA sequence of mycobacteria sandwiching a variable sequence to perform amplification of mycobacterial RNA. Species-specific probes for the M. tuberculosis complex, M. avium-paratuberculosis, M. intracellulare and M. leprae were hybridized in-solution with the amplified nucleic acids of 10 pathogenic mycobacteria and 11 closely related bacteria, as well as with human-derived NA in an enzyme-linked gel assay (ELGA). Each probe was shown to hybridize specifically to the amplified single-stranded RNA of the corresponding species. Thirty-two clinical isolates of M. tuberculosis strains from different parts of the world were correctly identified by NASBA using the M. tuberculosis-complex-specific probe. In combination with the ELGA, NASBA could identify mycobacteria rapidly, i.e. in less than 6 h. The relative simplicity and rapidity of this technique makes it an attractive tool for species-specific identification of mycobacteria.

Introduction

The members of the genus Mycobacterium include some important human pathogens, such as Mycobacterium tuberculosis and M. leprae. Infection with mycobacteria can be detected by conventional acid-fast staining of clinical specimens, providing the number of bacteria present in the sample is more than 10^4 ml^-1 (Bates, 1979). However, identification of the species is required for clinical and epidemiological purposes.

Identification of mycobacteria can be based on phenotypic properties, such as biochemical characteristics or antigenic composition, or on genetic properties, as expressed in characteristic nucleotide sequences in the genome. Phenotypic determination is still routinely the most common strategy, but is time-consuming and laborious. Genotypic identification is a more rapid and reliable approach for identification of mycobacteria (Gonzalez & Hanna, 1987). The necessity of prior culture may be circumvented through the use of nucleic acid amplification techniques, which allow rapid and sensitive detection of species-specific nucleotide sequences (Meier et al., 1993).

16S rRNA is an appealing target for the purpose of genotypic characterization for two reasons (Woese, 1987). First, the 16S rRNA contains sequence information allowing identification of mycobacteria at the species level (Böddinghaus et al., 1990a, b; Rogall et al., 1990a, b; Stahl & Urbance, 1990; Teske et al., 1991). Secondly, an advantage is the high copy number of rRNA in each cell, which makes rRNA easier to detect than targets that occur in a low number of copies. Amplification of a genus-specific stretch of the 16S rRNA with the polymerase chain reaction (PCR), followed by sequence analysis of the PCR product (Rogall et al., 1990b) or hybridization with species-specific probes (Böddinghaus et al., 1990b) has been successfully applied for identification of mycobacteria. A drawback of this method is the need for sophisticated equipment, such as a thermocycler (Stoker, 1990; Telenti et al., 1993).

An alternative approach is the nucleic acid sequence-based amplification (NASBA) technique (Compton,
1991) to selectively amplify mycobacterial 16S rRNA. Isothermal nucleic acid amplification of RNA in NASBA is achieved through the concerted action of AMV reverse transcriptase (RT), T7 RNA polymerase and RNaase H. The reaction starts with a non-cyclic phase, in which a downstream primer containing a tail-sequence of the T7-promotor sequence, anneals to the single-stranded target sequence. Through the action of AMV-RT, cDNA is formed. The RNaase H hydrolyses the RNA from the RNA-DNA hybrid, which results in a single strand of DNA to which the upstream primer can anneal. The AMV-RT synthesizes, through its DNA polymerase activity, a second DNA strand producing a double-stranded DNA intermediate with a transcriptionally active promoter sequence. The T7 RNA polymerase generates from this intermediate single-stranded RNA copies (100–1000), which serve as a template in the nuclease phase of NASBA. Apart from the fact that NASBA can amplify nucleic acids a billionfold within 1–5 to 2 h, it has the advantage over other amplification techniques in that it is a continuous, isothermal process, without the need for specialized equipment. The entire procedure, from amplification to identification, can be performed within a day. NASBA has already been successfully applied to the detection of RNA viruses (Kievits et al., 1991). We have investigated NASBA followed by ‘in-solution’ hybridization in an enzymelinked gel assay (ELGA) to detect the amplified single-stranded RNA products with non-radioactive mycobacterial species-specific probes. We focused on M. tuberculosis, M. avium and M. leprae, important pathogens in immunocompetent patients and, in the case of the first two species, also in immunocompromised patients.

Methods

Bacterial strains. Table 1 shows the bacterial strains that were used in this study. The cultivable mycobacteria were grown on Löwenstein-Jensen slants for 2–3 weeks. M. leprae was isolated from spleen tissue of an experimentally infected armadillo (Dasypus novemcinctus Linn.) as recommended by the World Health Organization (1980). Other bacteria which might be found in human samples or which are closely related to mycobacteria (see Table 1), were used for controls (courtesy of Dr R. J. van Ketel, Department of Medical Microbiology, Academic Medical Centre, Amsterdam, The Netherlands). For Actinomyces israelii lyophilized bacteria were used.

M. tuberculosis strains isolated from patients from Tanzania (n = 16), Thailand (n = 8) and The Netherlands (n = 8) were grown on Löwenstein-Jensen slants. All these strains were identified by conventional microbiological identification methods and characterized as M. tuberculosis. They were included in the study to explore the use of NASBA on clinical isolates from distinct geographical origins.

Nucleic acid (NA) isolation. All bacteria were harvested from a solid medium, except M. smegmatis, which was grown in Sauton medium. A suspension of the bacteria was made in 1 ml of phosphate-buffered saline (PBS) containing 0.1% (w/v) Tween 80 (Sambrook et al., 1989). The mycobacteria were centrifuged for 5 s at 13000 g in an Eppendorf centrifuge to remove large clumps.

All organisms were adjusted to a turbidity equivalent of a no. 4 McFarland barium sulphate nephelometer standard as described earlier by Verstijnen et al. (1991) and subsequently ten times diluted. A 50 µl vol. of the diluted samples contained approximately 10^3 viable mycobacteria as determined by counting the number of colony-forming units. A 50 µl vol. of the diluted samples was used for lysis and NA isolation according to ‘protocol Y/SC’ described by Boom et al. (1990). In brief, the bacteria were added to a guanidinium thiocyanate (GuSCN) lysis solution (5-25 M-GuSCN, 50 mM-Tris/HCl, pH 6.4, 20 mM-EDTA, 1.3%, w/v, Triton X-100) and rapid lysis was facilitated by two cycles of heat/cold shocks: heating for 5 min at 100 °C in boiling water and snap-freezing for 5 min at −170 °C in liquid nitrogen. Activated silica (50 µl silica (Sigma); 1 g ml^-1 suspension in 0.1 M-HCl) was added to bind the released NA. The NA–silica complex was washed twice with GuSCN washing solution (5-25 M-GuSCN; 50 mM-Tris/HCl, pH 6.4), twice with 70% (v/v) ethanol and once with acetone. After drying the silica at 56 °C, the NA were eluted from the silica with 50 µl RNAase-free H2O and stored at −20 °C.

Human NA were isolated in a similar fashion (kindly provided by Dr H. Smits, Department of Virology, Academic Medical Hospital, Amsterdam, The Netherlands). A 2 µl vol. containing 270 ng NA was tested. This amount corresponds to approximately 4 x 10^5 diploid human cells.

Total RNA isolation. The procedure for total RNA isolation was adapted from Chirgwin et al. (1979). M. tuberculosis was harvested and resuspended in 2-5 ml of GTC (4 M-GuSCN, 25 mM-sodium citrate, pH 7.0, 0.5% Sarkosyl, 0.1 M-β-mercaptoethanol). Lysis of the mycobacteria was facilitated by two series of alternating heat/cold shocks (see above). The GTC suspension was loaded onto a cushion of 1:5 ml 5 M-CsCl in a polyallomer centrifuge tube. The tube was filled up with GTC (final volume about 6 ml), and centrifuged in a swing-out rotor at 150000 g for 15–18 h at 25 °C. The supernatant was removed leaving a

Table 1. Bacterial species and strains

<table>
<thead>
<tr>
<th>No.</th>
<th>Strain/source</th>
<th>Species</th>
<th>Origin*</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5544</td>
<td>Mycobacterium africanum</td>
<td>RIVM</td>
</tr>
<tr>
<td>2</td>
<td>3875 (serovar 2)</td>
<td>Mycobacterium avium</td>
<td>RIVM</td>
</tr>
<tr>
<td>3</td>
<td>8316</td>
<td>Mycobacterium bovis</td>
<td>RIVM</td>
</tr>
<tr>
<td>5</td>
<td>IWGMT3 (serovar 4)</td>
<td>Mycobacterium intracellularis</td>
<td>RIVM</td>
</tr>
<tr>
<td>6</td>
<td>1012</td>
<td>Mycobacterium kansasii</td>
<td>RIVM</td>
</tr>
<tr>
<td>7</td>
<td>Armadillo isolate</td>
<td>Mycobacterium leprae</td>
<td>KIT</td>
</tr>
<tr>
<td>8</td>
<td>138601-24</td>
<td>Mycobacterium paratuberculosis</td>
<td>CDI</td>
</tr>
<tr>
<td>9</td>
<td>3442</td>
<td>Mycobacterium scrofulaceum</td>
<td>RIVM</td>
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<tr>
<td>10</td>
<td>4514</td>
<td>Mycobacterium tuberculosis</td>
<td>RIVM</td>
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<td>11</td>
<td>103.62</td>
<td>Actinomyces israelii</td>
<td>CBS</td>
</tr>
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<td>12</td>
<td>Clinical isolate</td>
<td>Corynebacterium belfi</td>
<td>AMC</td>
</tr>
<tr>
<td>13</td>
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<td>Corynebacterium J.K.</td>
<td>AMC</td>
</tr>
<tr>
<td>14</td>
<td>Clinical isolate</td>
<td>Escherichia coli</td>
<td>AMC</td>
</tr>
<tr>
<td>15</td>
<td>Clinical isolate</td>
<td>Haemophilus influenzae</td>
<td>AMC</td>
</tr>
<tr>
<td>16</td>
<td>Clinical isolate</td>
<td>Klebsiella pneumoniae</td>
<td>AMC</td>
</tr>
<tr>
<td>17</td>
<td>Clinical isolate</td>
<td>Legionella pneumophila</td>
<td>AMC</td>
</tr>
<tr>
<td>18</td>
<td>Clinical isolate</td>
<td>Nocardia asteroides</td>
<td>AMC</td>
</tr>
<tr>
<td>19</td>
<td>Clinical isolate</td>
<td>Pseudomonas aeruginosa</td>
<td>AMC</td>
</tr>
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<td>20</td>
<td>Clinical isolate</td>
<td>Staphylococcus aureus</td>
<td>AMC</td>
</tr>
<tr>
<td>21</td>
<td>Clinical isolate</td>
<td>Streptococcus pneumoniae</td>
<td>AMC</td>
</tr>
</tbody>
</table>

* RIVM, National Institute of Public Health and Environmental Protection, Bilthoven, The Netherlands; SSI, Statens Serum Institut, Copenhagen, Denmark; KIT, Royal Tropical Institute, Amsterdam, The Netherlands; CDI, Central Veterinary Institute, Lelystad, The Netherlands; CBS, Centraalbureau voor Schimmelcultures, Baarn, The Netherlands; AMC, Academic Medical Centre, Department of Medical Microbiology, Amsterdam, The Netherlands.
layer of 1–1.5 ml CsCl and the walls of the tube were rinsed with GTC,
keeping the RNA pellet covered with a CsCl layer. After decanting
the remaining CsCl, the bottom of the tube containing the RNA pellet
was cut off. The pellet was rinsed twice for 3 min with 100 μl of ice-cold
70% ethanol. The RNA was resuspended in 450 μl of RNAase-free
H2O. This 450 μl vol. of RNA suspension was transferred to a new tube
and was precipitated with 3 m- sodium acetate, pH 5-2, and 70%
ethanol. After washing the RNA pellet with 70% ethanol, it was dried
and resuspended in 100 μl of RNAase-free H2O. The concentration of
RNA was determined spectrophotometrically.

Selection of primers and probes. The primers and probes used in this
study are listed in Table 2. Primers OT 727 and OT 737 were chosen
from a highly conserved region of the nucleotide sequence of the
mycobacterial 16S rRNA genes (Rogall et al., 1990a; Liesack et al.,
1990). These primers facilitate the amplification of 16S rRNA when
there is sufficient homology between the primers and target. From the
published sequence data these primers were calculated to be approxi-
mately 200 nucleotides apart depending on the species of mycobacteria
(Rogall et al., 1990a). This primer set sandwiched a variable sequence
in mycobacteria; stretches of sequence that occur in this region are
specific for certain species. On the basis of these characteristic sequences
probes were constructed for identification (Böddinghaus et al., 1990a; Stahl & Urbance, 1990; Teske et al., 1991). Previously described
species-specific probes for the detection of NA derived from M.
tuberculosis (OT 999), M. avium-paratuberculosis (OT 998), M. intra-
cellularare (OT 979) and M. leprae (OT 852) were used in this study
(Böddinghaus et al., 1990a; Teske et al., 1991). A species-specific probe
for M. smegmatis (OT 851) was chosen by us from the same variable
region.

NASBA. The NASBA reactions were performed as described by
Kievits et al. (1991) with some modifications. The final volume of the
reaction mixture was 25 μl. First, a 21 μl vol. prereaction mixture was
assembled so that the final concentration for 25 μl would be: 40 mM-
Tris/HCl, pH 8.5, 12 mM-MgCl2, 42 mM-KCl, 5 mM-DTT, 1 mM
of each dNTP, 2 mM of each NTP, 15 % (v/v) DMSO, 12 U RNA guard
(Pharmacia), 0.2 μM-Primer 1 (OT 737, Table 2) and 0.2 μM-Primer 2
(OT 727, Table 2). After addition of 2 μl of target RNA the tubes were
incubated for 4 min at 65 °C in order to uncoil the tertiary and
secondary structures of the 16S rRNA. The reaction mixtures were then
directly transferred to 41 °C. After 2 min, 2 μl of an enzyme mix was
added to the tubes, resulting in a final volume of 25 μl, containing
0.1 μg BSA μl-1, 38 U T7 RNA polymerase (Pharmacia), 8 U AMV-
RT (Seikagaku), 0.1 U RNAase H (Pharmacia). Isothermal amplifi-
cation of the RNA target was performed by incubation of these samples
at 41 °C for 2 h. Reaction mixtures containing no target NA served as negative controls. The amplification products were either
directly processed for detection analysis by Southern blotting or ELGA
(see below) or stored for 2-4 weeks at -20 °C.

Blotting procedure. For detection of the amplification products, 7 μl
of each reaction mixture was loaded onto a 2% agarose ( Pronarose,
SpheroQ, The Netherlands) gel containing ethidium bromide (Sam-
brook et al., 1989). The separated products were blotted onto Zeta-
Probe (Bio-Rad) membrane using a vacuum blot apparatus (Pharmacia) and hybridized with a 32P-labelled oligonucleotide
(OT 999 and OT 851, Table 2) (Sambrook et al., 1989). The membranes were
exposed to X-ray film (Kodak) for 4 h.

ELGA. A rapid non-radioactive, 'in-solution' hybridization assay
(ELGA) was developed to identify the NASBA products with species-
specific horseradish peroxidase (HRP) 5'-labelled oligonucleotide
probes (ELGA probes). Hybridization was performed without the need
for prior denaturation of the NASBA products, since the product of
NASBA is mainly single-stranded RNA. Stringent hybridization
conditions are compulsory to distinguish the mismatches in a target by
the ELGA probe. After hybridization, excess non-hybridized ELGA
probes were separated from the homologous hybridized product by gel
electrophoresis. The free ELGA probes and the hybridized products
could be directly visualized in the acrylamide gel by incubating the gel
directly in substrate solution for HRP. Due to its lower mobility the
homologous hybridized product migrates in the gel above the free
ELGA probe.

Table 2 shows the species-specific synthetic oligonucleotides that
were labelled on the 5' end with HRP and used as probes in the ELGA.
An ELGA hybridization was performed by mixing 1 μl of the NASBA
reaction product with 4 μl hybridization solution [final concentration
of this reaction mix was 1 × SSC (0.15 M-NaCl; 0.015 M-sodium citrate),
5%, v/v, glycerol, 2 mM-sodium phosphate buffer, pH 7.0, 0.01%
bromphenol blue, 0.01% xylene cyanol and 105 molecules of HRP-
labelled probe]. The hybridization reaction mixtures were incubated for
15 min at 45 °C. These stringency conditions, which were sufficient
to distinguish at least two mismatches with the oligonucleotide, were used
for all ELGA probes tested. Hybridization reaction mixtures (2.5 μl)
were then directly applied onto a 7 % acrylamide gel, containing 0.04 %
(w/v) dextran sulphate. After electrophoresis, the hybridized amplifi-
cation products and the ELGA probes in the gel were visualized by
staining with 80 μl substrate solution (125 μg 3,3',5,5'-tetrathienylbenzidine ml-1 and 0003 % H2O2 in 01 mM-sodium citrate
buffer, pH 5.2). The gel was incubated in the substrate solution for
approximately 10 min at room temperature before blue-stained bands
appeared. The gel was fixed with incubation in 50% (v/v) methanol
overnight at room temperature.

Results and Discussion

Specificity of NASBA

The NASBA primers (OT 727 and OT 737; Table 2) directed the amplification of an approximately 200 nt sequence of the 16S rRNA containing a region which is variable in mycobacteria. Previously described species-specific probes (Table 2) were used for identification of the NASBA products derived from ten different myco-
bacteria and eleven other relevant bacteria. Identification of the NASBA products was performed using the ELGA technique, an ‘in-solution’ hybridization assay. Electrophoresis of the hybridization reaction mixtures is
necessary to discriminate between free ELGA probe (fast mobility) and ELGA probe that has specifically hybrid-
ized to the NASBA product (slow mobility). Both products could be visualized directly in an acrylamide gel
(Fig. 1).

The specificity of the NASBA in combination with ELGA is shown in Fig. 1. Probe OT 999 hybridized with amplified RNA from M. africanaum, M. bovis, M. bovis BCG and M. tuberculosis (Fig. 1a, lanes 1, 3, 4 and 10,
respectively). Probe OT 999 was shown previously to be specific for M. tuberculosis (Böddinghaus et al., 1990a). As expected, hybridization occurred between amplifi-
cation products derived from other members of the M. tuberculosis complex due to their identical 16S rRNA
sequences (Rogall et al., 1990b).
Table 2. Sequences of primers and probes used in amplification of 16S rRNA

<table>
<thead>
<tr>
<th>Oligo</th>
<th>Sequence</th>
<th>Positions*</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>OT 727</td>
<td>5' AAC ACG TGG GTG-ATC TGC CCT GCA 3'</td>
<td>77–100</td>
<td>Upstream primer, highly conserved region of mycobacterial 16S rRNA†</td>
</tr>
<tr>
<td>OT 737</td>
<td>5' AAT TCT AAT ACG-ACG-GCT TGT GGC CTT CCA TCT CA 3'</td>
<td>262–284</td>
<td>Downstream primer, highly conserved region of mycobacterial 16S rRNA†</td>
</tr>
<tr>
<td>OT 851</td>
<td>5' CTG GTG GCA TGG-CCT GGT AG 3'</td>
<td>148–168</td>
<td>M. smegmatis probe†</td>
</tr>
<tr>
<td>OT 852</td>
<td>5' AAG CTT TTT GCG-GTG CAG GAT G 3'</td>
<td>172–196</td>
<td>M. leprae probe‡</td>
</tr>
<tr>
<td>OT 997</td>
<td>5' TTA GGC GCA TGT-CTT TAG GT 3'</td>
<td>148–168</td>
<td>M. intracellulare probe§</td>
</tr>
<tr>
<td>OT 998</td>
<td>5' CAA GAC GCA TGT-CTT CTG GT 3'</td>
<td>148–168</td>
<td>M. avium-paratuberculosis probe§</td>
</tr>
<tr>
<td>OT 999</td>
<td>5' CGG GAT GCA TGT-CTT GTG GT 3'</td>
<td>148–168</td>
<td>M. tuberculosis complex probe§</td>
</tr>
</tbody>
</table>

* Rogall et al. (1990a).
† This study.
‡ Teske et al. (1991).
§ Böddinghaus et al. (1990a).

Fig. 1. Specificity of NASBA in combination with ELGA probes: (a) OT 999, M. tuberculosis probe; (b) OT 998, M. avium-paratuberculosis probe; (c) OT 997, M. intracellulare probe; and (d) OT 852, M. leprae probe. NASBA was performed using primers OT 727 and OT 737 on NA from bacterial species and strains listed in Table 1. Lane 22 contained human NA. The negative control samples used in this assay contained no target. After ‘in-solution’ hybridization (ELGA) of the NASBA product with the species-specific ELGA probes, the hybridization reaction mixtures were run in a 7% acrylamide gel, to separate the free ELGA probe (fast mobility) from the homologous hybridized NASBA product (slow mobility).
Identification of Mycobacterium spp. with NASBA

Fig. 2. Sensitivity of the NASBA. NASBA and Southern blotting were performed as described in Methods. (a) Analysis of NASBA reactions containing total RNA purified from M. tuberculosis. The amount of total RNA added per NASBA reaction was 500 pg (lane 1), 50 pg (lane 2), 5 pg (lane 3), 500 fg (lane 4), and 50 fg (lane 5). (b) Analysis of NASBA reactions performed on NA isolated from a suspension of M. tuberculosis. The number of bacteria per sample was as follows: 2 x 10^4 (lane 1), 2 x 10^3 (lane 2), 2 x 10^2 (lane 3), 2 x 10^1 (lane 4) and 2 x 10^0 (lane 5).

(?) Alzheimer disease (AD) is a neurodegenerative disorder characterized by the accumulation of amyloid-beta (Aβ) peptides and neurofibrillary tangles (NFTs) in the brain. The pathogenesis of AD is believed to be multifactorial, involving both genetic and environmental factors.

Recently, the sequences of the 16S rRNA of different serovars belonging to the M. avium–M. intracellulare complex have become available (Böddinghaus et al., 1990b). The 16S rRNA sequences of M. intracellulare serovars 16 and 21–28 are still not known. Alignment of the known 16S rRNA sequences of the members of the M. avium–M. intracellulare complex with probes OT 998 and OT 997 suggests that probe OT 998 has the potential to detect all three M. avium serovars and M. intracellulare serovars 4–6 and 8–11; probe OT 997 detects M. intracellulare serovars 12–15 and 17/19/20. Since M. intracellulare serovars 7 and 18 are not identified by these two probes, an additional probe would be necessary to include these serovars also.

Sensitivity of NASBA

The sensitivity of the NASBA technique was investigated with M. tuberculosis as a model system. This was done by adding a 10-fold serial dilution of purified total RNA derived from M. tuberculosis in amounts ranging from 5 ng to 50 fg to the reactions. A 500 fg amount of total RNA could be amplified reproducibly to give a clearly detectable signal on a Southern blot using 32P-labelled probe OT 999 (Fig. 2a). In addition, 10-fold serial dilutions were made from a suspension of M. tuberculosis ranging from 10^5 to 10 bacteria per NA extraction. Only a part of the extracted NA was subjected to the NASBA reaction: the final amount of NA derived from 2 x 10^4 to 2 bacteria was added. The addition of NA derived from 200 bacteria resulted in a visible band on a Southern blot using 32P-labelled probe OT 999 (Fig. 2b). A similar experiment using M. smegmatis and probe OT 851 revealed a detection level of approximately 20 bacteria (results not shown).

The effect of using the high copy number of rRNA on the sensitivity of the NASBA assay was not apparent. However, NASBA has been shown previously to be able to detect as little as 100 molecules of RNA (Kievits et al., 1991). Using the protocol described here we did not reach such high sensitivity. However, in our investigation emphasis was put on specificity rather than on sensitivity. Optimization of NASBA for the detection of low numbers of mycobacterial target rRNA molecules is now in progress.

A 10-fold difference in sensitivity of the NASBA method was observed using M. tuberculosis and M. smegmatis NA derived from whole bacteria as targets. Two reasons can be put forward to explain this observation. First, fast-growing mycobacteria such as M. smegmatis may contain a larger number of ribosomes than slow-growing mycobacteria such as M. tuberculosis. Thus, more rRNA per bacterium can be isolated from rapid growing mycobacteria (Ratledge, 1982). The higher...
content of rRNA per cell in fast-growing mycobacterial species may lead to a more sensitive detection limit using the NASBA technique. Secondly, the apparent reduced sensitivity of NASBA when using *M. tuberculosis* compared to *M. smegmatis* might be due to different culture conditions. *M. tuberculosis* was grown on a solid medium and *M. smegmatis* in a liquid medium. A colony grown on a solid medium comprises a larger proportion of non-viable bacteria than the same amount of bacteria grown in a liquid medium (Winder, 1982). With both explanations, it is assumed that the apparent difference in sensitivity of NASBA using different mycobacteria reflects a difference in 16S rRNA content of the bacteria. The content of 16S rRNA may be related to metabolic activity. If this is true, the NASBA technique has the potential to discriminate between dead and living bacteria by detecting the different RNA levels of the cells.

**Identification of different *M. tuberculosis* isolates**

Probe OT 999 was based on the 16S rRNA sequence of a single *M. tuberculosis* strain (Böddinghaus et al., 1990a). Therefore, different *M. tuberculosis* isolates were tested in this study. NASBA in combination with ELGA using probe OT 999 allowed identification of the 32 *M. tuberculosis* strains from clinical isolates from different parts of the world (16 from Tanzania, 8 from Thailand and 8 from The Netherlands). The correct identification by NASBA of these *M. tuberculosis* isolates from various sources shows the potential general applicability of this method. Further studies are necessary to evaluate whether this method can be used for the identification and possibly also detection of mycobacteria in biological specimens.

**Conclusion**

NASBA in combination with ELGA offers a less complicated and more rapid alternative to routine biochemical identification, as it can be used to identify cultured mycobacteria in less than 6 h. The fact that NASBA does not require specialized equipment makes it an attractive method for simple settings. Additionally, NASBA has the potential to be used for detection and identification of mycobacteria (and other organisms) in biological specimens, since it meets the requirements of sensitivity, specificity and rapidity. This would have definite advantages over identification of phenotypic characteristics of cultured mycobacteria.

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**References**


