Drug Susceptibility of Brazilian Strains of Mycobacterium bovis Using Traditional and Molecular Techniques


Laboratório de Biologia Molecular Aplicada à Micobactérias, Departamento de Medicina Tropical, Instituto Oswaldo Cruz-Fiocruz, Av. Brasil 4365, 21040-900 Rio de Janeiro, RJ, Brasil *Escola de Veterinária, UFMG, Belo Horizonte, MG, Brasil **Laboratório Regional de Apoio Animal, Ministério da Agricultura, Pecuária e Abastecimento, Pedro Leopoldo, MG, Brasil ***Unidade de Pesquisa em Tuberculose, Instituto de Doenças de Tórax, HUCFF, UFRJ, Rio de Janeiro, RJ, Brasil

Transmission of Mycobacterium bovis from cattle to humans has been reported and can cause tuberculosis (Tb) and a problem in certain risk populations. Therefore, knowledge of resistance of M. bovis towards antibiotics used for therapy of human Tb could help avoiding care delay and treatment cost increase when dealing with drug resistant organisms. We therefore evaluated the susceptibility of M. bovis isolates towards streptomycin, isoniazide, rifampicin, ethambutol, and ethionamide, the first line antibiotics for human Tb. Therefore, 185 clinical samples from cattle with clinical signs of tuberculosis were processed and submitted to culturing and bacterial isolates to identification and drug susceptibility testing using the proportion method. Among 89 mycobacterial strains, 65 were identified as M. bovis and none were resistant to any of the antibiotics used. Confirmation of present results by future studies, enrolling a large number of isolates and designed to properly represent Brazilian regions, may favor the idea of using isoniazide preventive therapy as part of a Tb control strategy in special situations. Also, nucleic acids from bacterial isolates were submitted to rifoligotyping, a recently described reverse hybridization assay for detection of mutations causing resistance towards rifampicin. Concordance between the conventional and the molecular test was 100%, demonstrating the use of such methodology for rapid evaluation of drug susceptibility in M. bovis.

Key words: Mycobacterium bovis - drug susceptibility - isoniazide - rifoligotyping

Tuberculosis (Tb) is caused by infection with Mycobacterium tuberculosis and is one of most severe infectious diseases. During the last decade, control of Tb turned more complicated because of decreasing socio-economic conditions, the human immunodeficiency virus pandemic and the spread of drug (DR) and multi-drug resistant (MDR) M. tuberculosis strains. One third of the world population is infected and over 100,000 new Tb cases are reported each year in Brazil (WHO 1993). Infection with M. bovis followed by Tb has been reported mainly in cattle, wild and domestic pigs, primates (Nolte & Metchock 1995) and in humans, disease symptoms are indistinguishable from that caused by M. tuberculosis and is therefore generally treated in the same way. Bovine tuberculosis is much more frequent in developing than in developed countries, and although in most latter, the disease was considered under control (Ferreira Neto & Bernardi 1997, Mota & Lobato 1998), reports on resurgence of bovine tuberculosis have been demonstrated (Collins et al. 1994). In Brazil, the mean national prevalence of M. bovis infection in cattle Tb was estimated at 1.3% for the period 1989 to 1998 but this can increase to 15% in farms with some automated form of milk production (Belchior 2001).

Successful control of human tuberculosis should be accompanied by control of animal tuberculosis because transfer of M. bovis to humans has been reported, both through respiratory route and consumption of contaminated meat and dairy products. Especially farmers and veterinarians are at elevated risk to get infected with M. bovis while HIV-infection increases risk for Tb development after infection with M. tuberculosis or M. bovis (O’Reilly & Daborn 1995). According to WHO (1993), infection with M. bovis is responsible for about 5% of human Tb cases in Brazil, suggesting the importance of better control of transmission from cattle to man. M. bovis is naturally resistant to pyrazinamide (Cepanzo 1988) and usually susceptible to most antibiotics used to treat human Tb, caused either by infection with M. tuberculosis or M. bovis. The main control of bovine Tb is through “tuberculization test and slaughter” strategy but due to the financial burden of this procedure, uncontrolled, and unregistered use of antibiotics, mainly isoniazide, is frequently occurring for treatment of cattle with signs of Tb infection in Brazil (Mota 2003). Having in mind that monotherapy results quickly in the generation of drug-resistant bacilli and their possible transmission to humans, an evaluation of drug susceptibility of M. bovis strains is warranted to understand the magni-
tude of influence of the transmission of DR or MDR *M. bovis* strains on treatment of human Tb.

The molecular basis of resistance has been clarified to a different extent for the antituberculosis drugs. In the case of rifampicin, detection of mutations in a small fragment of the rpoB gene predicts drug susceptibility in about 95% of the *M. tuberculosis* strains and although the mechanism of resistance seams identical in *M. bovis*, no data are available on distribution of mutation frequency in this species. Recently, a reverse hybridization-based assay for detection of these mutations was shown to be highly sensitive and specific for human isolates of *M. tuberculosis* (Morcillo et al. 2002) but again, no *M. bovis* isolates have been included in the study.

We therefore evaluated susceptibility of isolates of *M. bovis* of bovine origin towards the five mostly used first line antituberculous drugs. Also, the efficiency of the rifoligotyping assay was evaluated in a retrospective way on these isolates of *M. bovis*.

**MATERIALS AND METHODS**

**Collection of sample and processing for culture** - During the period of 2001-2002, we performed an exploratory, transversal study by collecting tuberculous caseum lesions from 185 different animals of cattle, mainly from the state of Minas Gerais and some other regions of Brazil, including Santa Catarina, Mato Grosso do Sul, Amazonas, Goiás, Paraíba, and São Paulo. Sample collection occurred under supervision of the Federal Inspection of the state of Minas Gerais and the material sent frozen or in ice in a plastic recipient to the Tuberculosis Laboratory of the Regional Laboratory of the Ministry of Agriculture. Among these, 56 had been isolated in the field and 129 from slaughterhouses; no information was available regarding PPD skin test evaluation. The material was decontaminated according to Mota (1985), in brief, by incubating in 6% sulfuric acid during 30 min and followed by centrifugation at 500 g for 15 min. The pellet was washed twice in 0.85% saline solution and distributed in four tubes containing Stonebrink medium. Cultures were incubated at 37°C and observed weekly until being discarded after 4 weeks.

**Identification and drug susceptibility testing** - These tests were performed at same laboratory according to procedures were according to Cepanzo (1988). In summary, 65 cultures were obtained and evaluated for production of niacin, nitrate reduction capacity, susceptibility to 100 mg/ml pyrazinamide in Löwenstein-Jensen medium, hydrolysis of Tween and catalase activity at room temperature and at 68°C. Drug susceptibility was performed in Stonebrink medium containing sodium glutamate and without pyruvate. Antibiotics (Sigma Chemicals Co., US) were added separately in different concentrations (4 µg/ml streptomycin, 40 µg/ml rifampicin, 2 µg/ml ethambutol, 20 µg/ml ethionamide, and 0.2 µg/ml isoniazide) and considered resistant when either 10% (streptomycin and ethionamide) or 1% (others) of the amount of colonies observed in the sample without drug was counted in the antibiotic containing vial. The drug susceptible strain *M. tuberculosis* H37Rv was included as a control for antibiotic activity.

**Mutation detection using the RIFO-assay** - The procedure was performed at the Leprosy Laboratory of the Oswaldo Cruz Institute at the Oswaldo Cruz Foundation (Fiocruz) in Rio de Janeiro, Brazil. Initially, nucleic acid extraction was performed on a fraction of the culture as described by Van Embden et al. (1993). Basically, a loop full of bacteria were submitted to digestion with lysozyme and proteinase K, followed by treatment with CTAB and chloroform/isoamylalcohol and precipitation with isopropanol. For detection of the rpoB genotype, rifoligotyping was performed basically as described by Morcillo et al. (2002). Therefore, a fragment from 1497 to 1653 of the rpoB gene was amplified using 100 ng of primers TR110A (5’-CGC CGC GAT CAA GGA GT-3’) and biotinylated TR111A (5’-ACGTGCGGGACCCTCA-3’) in a 25 µl reaction volume containing 1 U of Taq polymerase (Gibco BRL, US) and buffer, 0.2 mM of each dNTP and 10 ng of target DNA. Cycling was performed by incubating for 3 min at 95°C; followed by a touch-down PCR (2°C per step) from two cycles of 20 s 95°C, 30 s 65°C, 30 s 72°C to two cycles of 20 s 95°C, 30 s 57°C, 30 s 72°C, and more 25 steps of 20 s 95°C, 30 s 55°C, 30 s 72°C. Amplification was verified on a 2% agarose gel and product used in the reverse line blot hybridization assay basically as described by Morcillo et al. (2002) except that we included an oligonucleotide (5’-G CTG TTG GGG TTG ACC-3’) for hybridization with the mutant allele at codon 522 (TGG → TTG). Hence, a membrane (Biodyne C membrane, Pall Biosupport) was activated by incubation with 16% (wt/vol) 1-ethyl-3-(3-dimethylaminopropyl) carbodiimide (Sigma) for 15 min. The oligonucleotides with aminolink were diluted in 0.5 M NaHCO₃ (pH 8.4) at a concentration ranging from 12.5 to 200 pmol/150 µl (Morcillo et al. 2002), applied on the membrane using a Miniblotter MN45 (Isogen, the Netherlands) and after 1 min incubation at RT, the membrane was inactivated with 100 mM NaOH for 10 min and washed in 2 x SSPE containing 0.1% BDS for 10 min at 50°C. For hybridization, 10 ml of amplicon was diluted in 150 µl 2 x SSPE/0.1% BDS, heat denatured at 100°C for 10 min, chilled on ice and applied on the membrane using the miniblottor, in a perpendicular direction to the immobilized oligonucleotides. Hybridization occurred for 45 min at 50°C and the membrane was washed twice in 2 x SSPE/0.5% BDS for 10 min at 5°C, followed by incubation in 10 ml 2 x SSPE/0.5% BDS containing 1.25 U streptavidin-peroxidase conjugate (Amersham Pharmacia Biotech) for 30 min at 42°C. This was followed by washing in 2X SSPE/0.5% BDS, twice for 10 min at 42°C and twice for 5 min at RT. Hybridization to the membrane was visualized by enhanced chemiluminescence (ECL; Amersham Pharmacia Biotech) and exposure to a light sensitive film (Hyperfilm ECL, Amersham). As controls for the rifoligotyping procedure, one drug sensitive and six rifampicin resistant (one 531-TGG and five 531 TTG strains) of *M. tuberculosis* were included in an experiment. Interpretation of the hybridization patterns was performed by two persons and yielded the same conclusion.

**RESULTS**

From 185 clinical samples, 89 mycobacterial isolates were obtained of which 24 were rapidly growing organisms
not belonging to the *M. tuberculosis* Complex. No information on identity of these isolates was available at the time of this study. Biochemical characteristics, including nitrate reduction, absence of niacin, hydrolysis of Tween and catalase activity at 68°C and most importantly, natural resistance to pyrazinamide, demonstrated that 65 isolates were *M. bovis*. This observation was confirmed by the generation of *M. bovis* genotypes after submitting the isolates to spoligotyping (data not shown).

All isolates of *M. bovis* were successfully submitted to conventional susceptibility testing for the five antibiotics; none was resistant against any of the drugs tested.

Of the 65 isolates of *M. bovis*, 50 could be submitted to rifoligotyping and as partly demonstrated in the Figure, all generated the wild-type *rpoB* hybridization pattern. Two independent persons performed the interpretation of the genotypes. The rifampicin resistant and sensitive *M. tuberculosis* control strains generated the expected genotype patterns, demonstrating that the rifoligotyping experiments were executed under appropriate conditions and that the assay is accurate (kappa = 1).

**DISCUSSION**

There exists an increasing concern in the scientific community about the increase in both human and bovine Tb and the more frequent observation of drug resistant isolates of *M. bovis* and *M. tuberculosis* (Langenegger et al. 1981, 1991). In Brazil, three among 200 (1.5%) isolates from Tb patients were MDR *M. bovis* strains (Corrêa & Corrêa 1974) and recently, Leite and Lage (1999) reported to evaluate better the problem of drug resistant *M. bovis* strains in bovine Tb. Although none of the 65 isolates identified as *M. bovis* were resistant to any of the five anti-Tb drugs tested, we are aware that our study has limited statistical power. Indeed, estimating prevalence as 1.5%, a greater number of isolates (n = 536) would be tested to achieve a 95% confidence interval. Therefore, future studies are needed to definitively clarify the question of *M. bovis* anti-Tb drug resistance in Brazil.

The need to monitor incidence and transmission of drug resistant Tb has been suggested by WHO since 1993, when the disease was considered again a serious treat to human health, partly because of the increase in HIV and development of MDR strains of *M. tuberculosis* (Blázquez et al. 1997). Multi-drug resistant Tb can also be caused by infection with MDR *M. bovis* strains in HIV positive individuals (Blázquez et al. 1997) and AIDS increases Tb development, also after infection with *M. bovis* (O’Reilly & Daborn 1995). This increases concern about transmission of MDR *M. bovis* strains in a human risk population and transfer of *M. bovis* strains from cattle to human is occurring with a certain frequency among individuals with more intense contact (Mota 2003). Furthermore, the presence of viable *M. bovis* was recently demonstrated in livestock specimens and milk obtained in Brazil (Leite et al. 2003). Although transmission of MDR strains from cattle to man has to our knowledge not being reported, evaluation of drug susceptibility of *M. bovis* strains is important, not only for estimating such risk but also to understand the effect of uncontrolled use of antibiotics in cattle. Especially in the states of Minas Gerais and São Paulo, use of isoniazide for treatment of dairy cattle has become routine, principally because of the collaboration between farmers and large corporations for the purchase and distribution of the drug on a large scale (Belchior 2001).

Such lack of resistance, particularly towards isoniazide, orients health professionals in relation to correct treatment of humans infected by *M. bovis* and on the risk-free use of isoniazide for treatment of infected animals, when performed under strict supervision of veterinary official of Ministry of Agriculture. There is an ongoing discussion in Brazil on the use of antituberculous drugs for cattle treatment: arguments against treatment are the possible
increase of transmission from animals that are treated but not cured when “test and slaughter” is replaced by general-ized treatment; also, selection of drug resistant strains by use of monotherapy or inadequate treatment is an arg-ument. In favor of treatment are those who believe that “test and slaughter” is not popular in regions where Tb prevalence is high, stimulating frauds and omission of information. Fortunately, studies on treatment results in Brazil are promising as studies by Langenegger et al. (1981, 1991) demonstrated cure rates using isoniazid in highly infected farms was 93% using as a cure criterion rever-sion of sensibility to tuberculin testing during a period of more than two years. Diminished or even lack of tubercu-lin reaction after isoniazide treatment had also been de-scribed by Leite and Lage (1999). More recently, Mota (2003) demonstrated, using microbiologic examination and allergy desensibilization, a cure rate of 99% among natu rally infected animals in the region of Zona da Mata, Minas Gerais. Besides that, this study demonstrated a consider-able increase in milk production among treated animals, being one of the contributing factors for decreasing the concentration of isoniazid in milk for human consump-tion (Leite & Lage 1999). Considering cure rates, lack of circulation of isoniazid resistant M. bovis strains and decreased uptake of isoniazid in milk by pasteurization, isoniazide treatment under strict supervision seems an alternative for “test and slaughter”.

In this study, we also used a recently developed re-verse hybridization based assay, rifilogotyping, for de-tection of resistance towards rifampicin in M. tuberculo-sis (Morcillo et al. 2002). This assay is based on the de-tection of point mutations present in a small region (hot spot) of the rpoB gene and has not been evaluated on M. bovis strains. All 50 isolates that were characterized as sensitive towards rifampicin gave the sensitive rpoB geno-type after submission to the hybridization procedure. This is no surprise because it has been described that the mechanism of resistance in M. tuberculosis is shared by M. tuberculosis, M. africanaum, and M. leprae (Williams et al. 1994) and that the rpoB sequence is identical for the organisms belonging to the M. tuberculosis Complex (Kim et al. 1999), even when considering a highly polymorphic region of this gene (Lee et al. 2003). Although no rifampici-r resistant M. bovis isolates were included so we cannot conclude on the sensitivity of rifilogotyping in this species, the sequence data from literature and the correct hybridization of the sensitive strains with all five probes for the wild rpoB sequence strongly support the idea that rifilogotyping will be a quick assay for large scale suscepti-bility typing in M. bovis.

REFERENCES


