MOLECULAR IDENTIFICATION AND EPIDEMIOLOGY

Antigenic and genomic homogeneity of successive Mycoplasma hominis isolates

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Sixty Mycoplasma hominis isolates were obtained from the cervices of pregnant women and from the ears or pharynges of their newborn babies. The isolates were examined by SDS-PAGE and pulsed-field gel electrophoresis. Antigenic and genomic profiles were obtained for 16 series with two or more successive isolates. Both analyses led to the conclusion that isolates from the same woman were identical or nearly identical, while isolates from different women exhibited a high degree of variation with respect to both genomic and antigenic profiles.

Introduction

Mycoplasma hominis is one of >70 species comprising the class Mollicutes with a characteristically small genome size, an absence of cell wall and a generally low G+C content. Despite the limited coding capacity ranging from 580 kb (M. genitalium [1]) to 1300 kb (M. iowae [2]), mycoplasmas have adapted to a wide variety of hosts and are found to colonise mainly mucous tissues in man and animals. The lack of a cell wall exposes the plasma membrane directly to the environment, making the ability to alter surface proteins of critical importance for colonisation and survival of the micro-organism.

M. hominis is a sexually transmitted human pathogen and a frequent inhabitant of the genital tract. Despite years of intensive studies, the role of M. hominis in disease is not yet clear. However, there is an association with a number of urogenital diseases [3, 4]. Extragenital infections, particularly in immunosuppressed individuals, are being reported with increasing frequency [5, 6]. Assessment of the pathogenic potential of M. hominis is complicated by a high degree of genomic and antigenic heterogeneity within the species. The picture is further complicated by the fact that the micro-organism is isolated frequently from asymptomatic males and females [7].

Variability among different strains of M. hominis has been studied by several approaches. Ladefoged and Christiansen [8] constructed physical and genetic maps of the genomes of five M. hominis strains obtained from different individuals. Genome sizes obtained by pulsed-field gel electrophoresis ranged from 704 to 825 kb. Restriction patterns varied greatly; none of the strains had identical restriction patterns with any of the five restriction endonucleases used. However, the order of the mapped genes was conserved in the strains. The five strains were included previously in a study by Andersen et al. [9]. They compared the antigenic profiles of 14 M. hominis strains by SDS-PAGE, two-dimensional (2-D) gel electrophoresis and immunoblotting. It was possible to distinguish each of the 14 strains despite high similarities (76–99%) between the strains in SDS-PAGE. The electrophoretic 2-D protein patterns had lower similarities ranging from 41 to 72% between individual isolates. As 2-D gel electrophoresis separates proteins according to both mol.wt and iso-electric point, these lower similarity data seem to be more accurate than those obtained by SDS-PAGE. No two isolates were found to be identical by either of the two above-mentioned methods or by the immunoblotting analyses. Christiansen and Andersen [10] studied genetic variation of 26 M. hominis strains as reflected in differences in hybridisation patterns. DNA from the 26 strains was probed by Southern blotting with plasmids containing parts of the relatively conserved mycoplasma ribosomal RNA (rRNA) genes. Variability in restriction patterns was frequent outside the rRNA operons and was observed within the rRNA genes in three strains. The 26 strains were classified on the basis of RFLP

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patterns. Three classes contained six, three and two strains, respectively, with identical RFLP patterns. A fourth class contained the remaining 15 strains with unique restriction cleavage patterns. The finding that 15 of the 26 strains showed unique patterns was surprising, as genes encoding rRNA are among the most conserved genes in prokaryotic cells. In mycoplasmas, which harbour only one or two rRNA gene copies, it seems crucial to maintain functional rRNA gene operons. Therefore, the heterogeneity observed within these genes probably reflects a 'minimum of variation' detectable in *M. hominis*. The apparently identical strains with respect to RFLP patterns in rRNA genes were not the most related strains with respect to protein profiles as determined in the study by Andersen et al. [9].

The very heterogeneous character of *M. hominis* combined with a possibly minimal coding capacity, the lack of cell wall and the occasionally opportunistic behaviour make the organism an interesting subject in studies of not only *M. hominis* pathogenesis, but also with respect to the general development of bacterial mechanisms of surface variation.

This study reports the variability among 60 *M. hominis* isolates randomly chosen from a recent study by Thorsen [11]. He evaluated the association between various micro-organisms, including *M. hominis*, isolated from the cervicis of 3178 pregnant women, and bacterial vaginosis. Among the 60 isolates, 16 series of two or more isolates from the same woman or her newborn baby, or both, were examined, and the variability among these successive isolates was compared with the variability among isolates from different women. Protein profiles were analysed by SDS-PAGE and genomic profiles were evaluated by pulsed-field gel electrophoresis (PFGE) of genomic DNA cleaved with two different restriction endonucleases.

**Materials and methods**

**Mycoplasma strains and cultivation**

Sixty *M. hominis* isolates were obtained from pregnant women and their newborn babies participating in a cross-sectional population-based study [11]. The numbers refer to the woman and the letters B, G, S, M and N refer to the relative time of recovery: B, G and S, indicating the first, second and third antenatal hospital visits, respectively; and M and N indicating samples taken from the ear and pharynx, respectively, of the newborn baby. Samples to be analysed for the presence of *M. hominis* were taken with sterile cotton-tipped swabs and inoculated directly into 1 ml of BE liquid growth medium [12], and frozen at −80°C for later cultivation. Cultures were grown in BEA medium [12] and filter cloned once according to the recommendations of the Subcommittee of the Taxonomy of Mollicutes [13].

**SDS-PAGE and immunoblotting**

Whole-cell preparations of *M. hominis* harvested in log-phase were dissolved in sample buffer containing 62.5 mM Tris HCl (pH 6.8), glycerol 10% v/v, SDS 2.3% w/v, β-mercaptoethanol 5% v/v and bromophenol blue 0.05% w/v. The samples were boiled for 5 min and subjected to electrophoresis in polyacrylamide 7.5% gels followed by Coomassie Blue staining as described previously [9].

**Construction of dendrogram**

Coomassie Blue-stained SDS-PAGE gels were analysed for the presence or absence of 35 distinct protein bands. A dendrogram based on an average linkage algorithm was constructed from these data by computer programmes (ETDIV and ETCLUS, v.2.3), kindly provided by T. S. Whittam (Institute of Molecular Evolutionary Genetics, University of Pennsylvania, USA).

**Preparation, restriction endonuclease digestion and PFGE of DNA in agarose blocks**

Six ml of *M. hominis* cultures in log-phase were harvested by centrifugation at 15 000 rpm for 40 min and washed twice with phosphate-buffered saline (PBS; 20 mM sodium phosphate, 250 mM NaCl, pH 7.4). The pellets were resuspended in 150 μl of PBS, mixed with 150 μl of low-melting-point agarose (Incert agarose; FMC Bioproducts, Rockland, MN, USA) 1.8% in PBS, previously melted and equilibrated to 43°C. The suspension was immediately transferred to a plastic mould (20 × 9 × 1.2 mm) (BioRad Laboratories, Richmond, CA, USA) which had been cooled on ice to ensure instant setting of the gel. The agarose plugs were transferred to tubes containing lysing buffer (sarcosyl 1%, 0.5 M EDTA, 10 mM Tris-HCl, pH 9.5) with Proteinase K (Boehringer GmbH, Mannheim, Germany) 200 μg/L. After incubation for 2 h at 50°C, the lysis buffer was changed and the plugs were incubated overnight at 50°C. The plugs were washed with lysis buffer without proteinase K for 2 h and stored in 100 mM EDTA. Before endonuclease digestion, small blocks (3 × 4.5 × 1.2 mm) were washed for 1 h with several changes of TE buffer (10 mM Tris-HCl, 1 mM EDTA, pH 8.0), and equilibrated in 200 μl of the appropriate restriction endonuclease buffer at room temperature. Digestion was carried out for 2 h in 120 μl restriction endonuclease buffer, 20 units of enzyme and bovine serum albumin (Boehringer Mannheim) 12 μg at temperatures specified by the supplier.

Before electrophoresis the blocks were washed in TE buffer for 30 min, followed by equilibration in 0.5 × TBE buffer (44.5 mM Tris-borate, 44.5 mM boric acid, 1 mM EDTA, pH 8.0). Agarose 1% gels in 0.5 × TBE buffer were run for 20 h in 0.5 × TBE buffer at 14°C and 200 V in the CHEF-DR II system (BioRad) and the sub-cell electrophoresis cell (Bio-
Results

**SDS-PAGE and antigenic cluster analysis**

For each isolate 30–40 protein bands were observed, of which c. 20 were consistently present (Fig. 1). The variability in banding pattern was most pronounced for the high mol. wt bands, while smaller proteins were more consistently observed. Despite the often pronounced similarities, a marked heterogeneity between isolates from different women was evident. Isolates originating from the same woman were identical or differed only by the presence of one or two protein bands (Fig. 1).

To examine whether the isolates could be grouped in antigenically related clusters, a dendrogram was constructed on the basis of protein profiles obtained by SDS-PAGE. Coomassie Blue-stained gels were analysed for the presence or absence of 35 distinct protein bands (indicated at the top, Fig. 1). The quantity of individual polypeptides was not taken into account. The data were processed by the computer programmes ETDIV and ETCLUS and a dendrogram of relatedness was constructed (Fig. 2). The isolates were evenly distributed and no clustering could be observed. Isolates originating from the same woman were identical or closely localised on the dendrogram, reflecting a high degree of homogeneity. The five isolates from series 1720 had identical protein profiles as did the series 1697, 2462, 2641 and 2509. With three isolates in series 2867, the first (B) and the second (G) differed by a single protein, while the isolate from the newborn baby (N) was identical to the second isolate from the mother. Isolate 1572B differed from the two identical isolates from the newborn baby (M and N) by a single band. Isolate 1762B was identical to one of the isolates from the baby (1762M), but differed by three bands from the other isolate (1762N). Several series with two isolates differed by a single mismatch (3052, 2763, 2457 and 2032), two mismatches (2347 and 2682) or three mismatches (2223 and 1935) in protein profiles. Isolates from different women were more distant in the dendrogram; however, some isolates differed by only three mismatches (305B and 327, 519B and the series 2641, 549B and the series 1572). Randomised data input resulted in dendrograms in which identical isolates or isolates with one or two mismatches were always closely localised. Whenever more than two mismatches occurred the localisation in the dendrogram must be considered random.

**Genomic analysis of isolates**

All isolates were digested with the restriction endonucleases *XhoI* and *SmaI*. Restriction patterns of the series and some individual isolates digested with *SmaI* are shown in Fig. 3a and b, respectively. Genome sizes calculated on the basis of the band patterns varied from 650 to 800 kb. No isolates from different women were identical in both digestions (Fig. 3a and b), in contrast to isolates originating from the same woman. Ten series (1720, 2867, 1697, 1762, 2457, 2462, 2509, 2641, 2682 and 3052) contained identical isolates in both digestions (Fig. 3a). Two series (2223 and 1935) showed variations in both the *XhoI* and *SmaI* digestions (Figs. 3a and 4). In addition to the bands seen in 2223B, 2223S exhibited additional bands corresponding to an increased size of 300 kb. As these additional bands were of lower intensity, 2223S was considered to be a mixed culture (Fig. 3a). The isolates from series 1935 differed markedly in the intensity of one band (175 kb) and the presence of another (40 kb) in the *SmaI* digestion consistent with a deletion of >200 kb in 1935S. Digestion with *XhoI* showed replacement of bands of 325 and 275 kb with a band of 300 kb in 1935S compared with 1935B (Fig. 4a). However, the 300-kb band could be a double band resulting from co-migration of similarly sized fragments. If not, an apparent deletion of >200 kb would result in a genome size of 550 kb, which is even smaller than that of *M. genitalium* which possesses the smallest genome (580 kb) reported so far. Therefore, the endonucleases *ApaI*, *SalI* and *BamHI* were used for further analysis of the two isolates (Fig. 4a). While *ApaI* did not cut either of the genomes, *SalI* digestion showed variations of band sizes yet adding up to approximately identical genome sizes. *BamHI* showed variation of two bands. Considering the 35-kb band of 1935B as a double band, the genome sizes were thus identical, and the major deletion indicated by the *SmaI* digestion could be excluded by these findings. The isolates in series 1935 demonstrate the difficulty of interpreting intricate band patterns, mainly caused by co-migration of similar sized fragments and, furthermore, emphasise the need to use different endonucleases for comparison of
isolates. Four series (1572, 2763, 2032, 2347) showed small variations in only one of the digestions. Isolate 1572B from the mother was identical to isolate 1572M from the pharynx of the newborn baby. Isolate 1572N from the ear of the newborn baby, differed in the Smal digestion from these two isolates in the sizes of two DNA fragments because of an insertion of 25 kb (Fig. 3a). The insertion was probably not recognisable in the XhoI digestion because of the relatively small size. Series 2763 showed band patterns indicating a deletion. Compared to the band pattern of the maternal isolate 2763B, the baby isolate 2763N lacked a 125-kb fragment and had an additional 75-kb fragment in the Smal digestion (Fig. 3a). Neither of the isolates had an XhoI restriction site. Series 2032 gave identical isolates in the Smal digestion. In the BglI digestion an additional DNA fragment of 125 kb was seen in 2032B (Fig. 4b). The band was of low intensity and could have been due a small subpopulation of mycoplasmas exhibiting a frequently found XhoI restriction pattern. It should be kept in mind that the cultures were subcloned only once before use. Series 2347 appeared...
VARIABILITY OF SUCCESSIVE \textit{M. HOMINIS} ISOLATES

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Fig. 3. Ethidium bromide-stained PFGE gels of \textit{SmaI}-
cleaved chromosomal DNA of: \textbf{a}, series isolates; \textbf{b},
individual isolates.

to contain identical isolates in the \textit{SmaI} digestion. A 150-kb
deletion, including an \textit{XhoI} restriction site, could account
for the absence of one band seen in the \textit{XhoI} digestion in isolate
2347G (Fig. 4c). In the \textit{SmaI} digestion the band sizes were identical;
however, in isolate 2347B, high intensity staining indicated double
bands of 90 and 60 kb which confirmed the finding of a deletion.

\textit{SmaI} restriction sites were found in all isolates,
whereas four isolates were resistant to \textit{XhoI} digestion,
possibly because of modification/methylation of the
recognition site.

Similar band patterns were produced in several
isolates when digested with \textit{XhoI} (data not shown).
However, when these isolates were electrophoresed
under different conditions, small size differences were
seen. The variability was confirmed since different
band patterns were observed when the restriction
endonuclease \textit{SmaI} was used. This indicates that the
\textit{XhoI} restriction sites, in contrast to the \textit{SmaI}
restriction sites, are located in relatively conserved
gene regions, and emphasises the importance of using
more than one restriction endonuclease for comparison
of isolates.

\textbf{Comparison of antigenic and genomic profiles}

Genomic differences were found in series 1572, 2223,
1935, 2032, 2347 and 2763 which could have resulted
from mixed cultures. These series also showed
differences in antigenic profiles. The major deletion
found in isolate 2347G was antigenically reflected in
the absence of two protein bands in SDS-PAGE.
However, none of the deletions (series 2347 and
2763) affected the growth of the isolates. Differences
in antigenic profiles were further seen in series 3052,
2682, 2457, 2867 and 1762. The remaining five series
– 1697, 2462, 2641, 2509 and 1720 – showed identity
in both genomic and antigenic profiles. While more
than half of the series exhibited antigenic or genomic
variations, or both, these were small compared with the
overall variability as reflected among the series and the
individual isolates.

\textbf{Discussion}

This study investigated the genomic and antigenic
variation of 60 \textit{M. hominis} isolates, comprising 16
series of successive isolates. The isolates did not form
antigenically related clusters on the basis of SDS-PAGE
when analysed with the computer programs ETDIV and
ETCLUS. Isolates from the same woman were identical
substantial genomic variation was revealed within
isolates originating from different women. In contrast,
isolates originating from the same woman had identical
genomic profiles in 10 of the 16 series. The isolates
from series 2032 and 2223 exhibited differences. As
the intensity and presence of bands varied between the
isolates, and as the isolates were filter-cloned only once before use, these differences were considered to result from mixed cultures. Differences between the isolates from series 2763 and 2347 probably resulted from genomic deletions, while an insertion most probably accounted for the differences in band pattern observed for series 1572. Series 1935 showed intricate band patterns when digested with five different enzymes, probably due to several double bands. Despite varying band patterns, genome sizes in four digestions were identical, suggesting that substitutions at cleavage sites and inversions accounted for the observed variations. The differences for some series were seen in only one of the digestions. This may be because the relatively small differences of fragment sizes could not be distinguished in the high mol. wt bands.

Studies of variability have previously been performed on isolates of miscellaneous origin [14–16]. However, the use of consecutive isolates to examine antigenic variation might provide valuable information on possible mechanisms used by the pathogen to change antigenically over time. Only a single report has described antigenic variation in successive *M. hominis* isolates [17]. Olson et al. examined the expression of surface antigens of 14 consecutive isolates obtained from the synovial fluid of a patient with chronic septic arthritis. Monoclonal antibodies (MAbs) were raised against the primary *M. hominis* isolate and subsequently used in immunoblotting with all isolates. The isolates differed markedly in protein profiles, while restriction and hybridisation patterns were virtually identical when fragments of the conserved rRNA operon and elongation factor Tu gene were used as probes. When these probes were used in hybridisation studies with isolates obtained from different sources, a high degree of variation in hybridisation patterns was obtained. This supports the assumption that, in the study by Olson et al. [17], antigenic variation rather than re-infection had occurred.

A high degree of heterogeneity as demonstrated for several mycoplasma species reflects a rapid evolution. This could be explained by either a general genomic instability or by evolution caused by selective pressure. The results of the present study indicate the latter, as isolates from different women exhibited substantial variation, whereas isolates originating from the same woman were found to be nearly identical, suggesting adaptation to the host environment. This confirms a previous study involving characterisation of three antigenic variants of *M. hominis* PG21 selected by culture with MAb 552 added to the growth medium [18]. To establish whether addition of a MAb to the growth medium could mediate major changes in DNA or protein profiles, the type strain *M. hominis* PG21 was cultured for a 3-month period in the presence of MAb 552, which reacts with epitopes encoded by 471-bp long tandemly repeated sequences identified in the PG21 *Imp* genes [19,20]. Southern and Western blot analyses were performed on three occasions during culture. The cultures with MAb 552 added to the medium showed deletions of a number of the 471-bp repeats. The genetic deletions were
accompanied by a decrease in size of the proteins. Control cultures without antibody added to the medium showed no alterations. It was concluded that the observed antigenic variation resulted from in-vitro antibody-mediated selection.

Variation among species is generally studied with respect to either antigenic profiles with MAbs, SDS-PAGE and 2-D gel electrophoresis [9, 21-25] or genomic and genetic profiles by DNA-DNA hybridisations, PCR and restriction endonuclease cleavage analysis [14, 16, 26]. In the present study the examination of both antigenic and genomic variability resulted in a limited correlation between genomic RFLP and antigenic properties. Some antigenic differences were not reflected in the PFGE analysis as illustrated by series 2867 and 2682, in which the isolates exhibited identical PFGE patterns. On the other hand, series that showed different PFGE profiles always exhibited minor antigenic differences as reflected in SDS-PAGE. Antigenic variations could result from a number of genomic changes not detectable in PFGE, such as minor insertions or deletions, point mutations, on-off phase shifts and recombination events, or to erroneous transcription or translation. Alternatively, the variation found in the PFGE analysis may only reflect either major chromosomal re-arrangements such as large deletions or insertions, or point mutations specific to the endonuclease restriction sites. Therefore, not surprisingly, the correlation between antigenic and genomic variability was limited to the detection of a general heterogeneity among isolates from different women and homogeneity among isolates from the same women in the present study. Lack of correlation between genomic and antigenic variability was previously reported by Poumarat et al. [27] who examined genomic and antigenic variability of 37 M. bovis strains. The isolates were classified into 13 different genomic groups based on restriction endonuclease analysis with three enzymes. Antigenic profiles were investigated by immunoblotting with a polyclonal calf serum and three MAbs. An equally pronounced antigenic variability was observed whether the isolates came from the same or different genomic groups.

Variability in relation to time, geography and anatomical site of isolation is poorly understood. Frey et al. [28] found that the chromosomal heterogeneity as demonstrated by PFGE of M. hyopneumoniae strains from different countries was markedly larger than between strains isolated from the same laboratory. Solsona et al. [29] also found that the antigenic variability of 31 M. agalactiae strains was related to the geographic origin of the strains. In a study by Barle et al. [30] strains isolated from the same sources were found to form clusters of strains having very high genomic homologies. However, these observations could not be confirmed when antigenic variation among 14 M. hominis isolates from different sources and 12 genital isolates was compared. The genital isolates were found to vary as much as the 14 isolates from different sources [31].

The isolates used in the present study were collected within a rather small geographical area and over a period of 18 months. All samples originated from the cervixes of pregnant women or from newborn babies. A high degree of heterogeneity was found with respect to both antigenic and genomic variation despite the uniformity in time, geography and anatomical origin of the isolates. Successive isolates in general remained genomically stable over the sampling period, suggesting that the colonisation site and local host immune response might be of considerable importance for the micro-organism to accomplish antigenic variation.

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