Confirmation of the Species Prevotella intermedia and Prevotella nigrescens

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The elevation of the two genotypes of Prevotella intermedia to species rank as P. intermedia and Prevotella nigrescens has increased the need for reliable differentiation between the two taxa. In this study, 53 strains, including strains whose species affiliations were known as well as fresh dental plaque isolates, were subjected to a multilocus enzyme electrophoretic analysis, DNA analyses in which we used whole genomic DNA, rRNA-derived sequences, and an oligonucleotide specific for the former P. intermedia genotype II (P. nigrescens) as probes, and a sodium dodecyl sulfate-polyacrylamide gel electrophoresis analysis of soluble cellular proteins. All of these tests consistently separated the strains into the same two distinct groups corresponding to P. intermedia and P. nigrescens, confirming that the two species constitute two distinct populations of bacteria. Each of the tests used independently provided reliable identification to the species level. A previously reported heterogeneity in the pattern of human immunoglobulin A1 (IgA1) degradation was not confirmed. No differences between species were observed. All of the strains induced total degradation of IgA1 within 48 h, a property that may be a virulence factor in periodontal disease development. The enzymes responsible for IgA1 degradation were not inactivated by the proteinase inhibitors α2-macroglobulin and α1-proteinase inhibitor.

**Prevotella intermedia** (formerly *Bacteroides intermedium*) is a gram-negative, obligately anaerobic rod that is often isolated from subgingival plaque of humans with periodontal disease and from other oral and nonoral infections (for reviews see references 10, 18, and 26). The two previously recognized genotypes of *P. intermedia* (12) were recently elevated to species rank as *P. intermedia*, corresponding to genotype I, and *Prevotella nigrescens*, corresponding to genotype II (24). However, the suggested tests that distinguish the two species (24) have proved to be difficult to reproduce in other laboratories. The suggested association between *P. intermedia* (genotype I) and periodontal disease (7, 9, 18) has strengthened the need for reliable tests to differentiate the two species.

In addition to addition of levels of homology between whole-cell DNA (24), serotyping with monoclonal antibodies (8, 9) and hybridization to an oligonucleotide based on 16S rRNA sequences from the former *P. intermedia* genotype II (3) have been reported to be methods that can be used to distinguish the two genotypes of the former species *P. intermedia*. In addition, sodium dodecyl sulfate (SDS)-polyacrylamide gel electrophoresis (PAGE) of soluble cellular proteins has been used to differentiate the two taxa (7, 12).

In this study we used several methods, including multilocus enzyme electrophoresis, restriction fragment length polymorphism (RFLP) analyses in which we used whole genomic DNA as well as rRNA-derived sequences as probes for hybridization, and SDS-PAGE of soluble cellular proteins to evaluate the value of these methods for discriminating between *P. intermedia* and *P. nigrescens*. We performed tests with 53 strains, including the type strains, other strains assigned to one of the species according to the recent taxonomic revision (24), and fresh dental plaque isolates. The resulting data were combined with the results of an examination of the ability of each strain to degrade human immunoglobulin A1 (IgA1), a potential virulence factor in periodontal disease development, in order to determine whether the previously observed diverse pattern of IgA1 degradation (14) is species related.

**Materials and Methods**

**Bacterial strains.** Of the 53 strains included in this study, strains CCUG 24041 (=ATCC 25611T =type strain), MH3, MH6, MH7, MH14, MH15, MH16, MH18, OMZ248, and OMZ324 were assigned to *P. intermedia* and strains CCUG 9560 (=ATCC 33563T =ATCC 33563T), BEF3 (=NCTC 9361), NCTC 9398, MH11, MH20, OMZ251, OMZ265, OMZ328, OMZ310 (=TS88), and HG65 (=TS88) were assigned to *P. nigrescens* according to the recently published taxonomic revision of the species (24). The remaining 33 *P. intermedia* strains had not been reclassified after revision of the species. These strains included strains HG189, BN116, OMZ226, OMZ276, OMZ326, UB13-c, AB13a-f, and P11a-k and 25 strains freshly isolated from subgingival plaque on nonselective agar plates in our laboratory. The strains were received from the following sources: strains CCUG 24041 and CCUG 9560, Culture Collection of the University of Göteborg, Göteborg, Sweden; strain NCTC 9398, National Collection of Type Cultures, London, United Kingdom; strains MH3, MH6, MH7, MH14, MH15, MH16, MH20, and BN116, H. Shah, Eastman Dental Hospital, University of London, London, United Kingdom; strains OMZ248, OMZ251, OMZ266, OMZ276, OMZ310, OMZ324, OMZ326, and OMZ328, R. Gmur, University of Zürich, Zürich, Switzerland; strains HG65 and HG189, T. J. M. van Steenbergen, Vrije Universiteit, Amsterdam, The Netherlands; strain BEF3, J. Carlsson, University of Umeå, Umeå, Sweden; strains UBJ13-c, AB13a-f, and P11a-k, G. Sundqvist, University of Umeå, Umeå, Sweden; and strains MHSl-20, BK27, BK28, BK30, BK31, BK32, BK33, BK35, BK36, BK37, BK38, BK22, BK19, BK20, BK21, BK23, BK24, BK25, BK26, BK34, MHSl-18, MMJ1-22, MMJ1-23, and MMJ1-24, our laboratory. The 25 freshly isolated strains were assigned to the species *P. intermedia* as previously defined on the basis of the results obtained in the following tests: Gram staining; formation of black-pigmented colonies on blood-containing media; selective growth in an anaerobic atmosphere; production of indole (15), tryptophan (17), β-galactosidase, β-glucosidase, β-glucosaminidase, α-glucosidase, and α-fucosidase (13, 16); and fluorescence under UV light (25). The strains were grown on plate agar plates or in liquid plate medium for all analyses (4, 22).

**Multilocus enzyme electrophoretic analysis.** A 15-ml portion of a dense starter culture was used to inoculate 150 ml of medium, which was incubated anaerobically for 48 h. Bacteria were harvested by centrifugation, and the pellet was suspended in 2 ml of 50 mM Tris-HCl-5 mM EDTA (pH 7.5), frozen at −20°C for 2 h, thawed on ice, and sonicated intermittently for 10 min with a model B-12 Sonifier (Branson Sonic Power Co., Danbury, Conn.) equipped with a small tip at setting 4 (40 to 50 W). The sonicate was frozen again at −20°C for 2 h, thawed on ice, and centrifuged. The supernatant was stored at −70°C until it was used. Multilocus enzyme electrophoretic analysis was performed on horizontal starch gels as described previously (20, 23). Electromorphs (mobility variants) were determined by the following metabolic enzymes (the buffer systems used are indicated in parentheses): glutamate dehydrogenase (buffer system A), malate dehydrogenase (buffer system A), adenylate kinase (buffer system A), indole-3-acetamide deacetylase (buffer system A), nucleoside phosphorylase (buffer system A), glyceraldehyde-3-phosphate dehydrogenase (buffer system G), carbamate kinase (buffer system I), phosphoglucomutase (buffer system I), phosphoglucokinase
isoenzyme (buffer system I), g-glutamyl-2-phenylalanine peptidase (buffer system II), g-glutamyl-leucyl peptidase (buffer system II), phenylalanine-leucyl-glutamate isomerase; CDK, carbamyl phosphate synthetase; PGM, phosphoglucomutase; GP1, glycyl-phenylalanine peptidase; GP2, glycyl-phenylalanine peptidase; GP3, glycyl-leucyl peptidase; LGG, leucyl-glycyl-glycyl peptidase; PLP, phenylalanine-leucyl-glycyl peptidase.

where

\[ \text{probability that two randomly chosen strains have different alleles of the locus} \]

Operational taxonomic units (OTUs) were used to construct a dendrogram. The smallest genetic distance (above which dissimilar alleles were present), were used to construct the dendrogram shown in Fig. 1. The smallest genetic distance was 0.02. A dendrogram was constructed from the data obtained from a computerized cluster analysis performed by the average-linkage method by using a matrix of pairwise genetic distances between ETs and the program ETCLUS, which was kindly provided by T. S. Whittam, Department of Biology, Institute of Molecular Evolutionary Genetics, Pennsylvania State University, University Park. The absence of enzyme activity was treated as missing data.

**Southern blot analyses.** Total cellular DNA was prepared from a 10-mI 24- to 48-h liquid culture as described previously (21), except that lysozyme was omitted from the preparation. The quality and concentration of DNA were determined by agarose gel electrophoresis.

Approximately 2 mg of total cellular DNA was subjected to restriction endonuclease digestion with EcoRI or MspI (Boehringer Mannheim GmbH, Mannheim, Germany) according to the manufacturer's instructions. After digestion for 15 min at 37°C, digests were loaded on 1% agarose gels and analyzed by ethidium bromide staining. DNA fragments were cut out of the gels, recovered from the agarose by gel extraction, and rehydrated with water. Gel-extracted DNA was used as a probe stock. The DNA probe was labeled by nick translation with [\( ^{32} \)P]dCTP and used as a probe. The labeled DNA probe was hybridized to Southern blots to detect the presence of the IgAl substrate in the bacterial culture. The blots were hybridized in a solution containing 50% (vol/vol) formamide, 5× SSC, 0.5% SDS, 5× Denhardt's solution, 0.5 mg of denatured salmon sperm DNA per ml, 1 mg of [\( ^{32} \)P]dCTP per ml, and 0.05 μg of DNA from P. intermedia

The mean genetic diversity for all enzyme loci was 0.52.

### TABLE 1. Numbers of electrophoretic alleles and levels of genetic diversity per enzyme locus for 51 strains of *P. intermedia* and *P. nigrescens*

<table>
<thead>
<tr>
<th>Enzyme locus</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>6</th>
<th>7</th>
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<tr>
<td>ADK</td>
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<td>0.11</td>
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<td>0.02</td>
<td>0.09</td>
<td>0.04</td>
<td>0.006</td>
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<td>0.04</td>
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<td>0.02</td>
<td>0.002</td>
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<tr>
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<tr>
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a: Abbreviations: ADK, adenylate kinase; GDH, glutamate dehydrogenase; MDH, malate dehydrogenase; NSP, nucleoside phosphorylase; IPO, indophenol oxidase; GP1, glycerol-2-phosphate dehydrogenase; GP2, 2-phosphoglycerate phosphatase; GP3, glycyl-leucyl peptidase; LGG, leucyl-glycyl-glycyl peptidase; PLP, phenylalanine-leucyl-glycyl peptidase; GP1, g-glutamyl-2-phenylalanine peptidase; GP2, g-glutamyl-phenylalanine peptidase; GP3, g-glutamyl-leucyl peptidase; LGG, leucyl-glycyl-glycyl peptidase; PLP, phenylalanine-leucyl-glycyl peptidase.

b: The mean genetic diversity for all enzyme loci was 0.52.

g: The mean genetic diversity for all enzyme loci was 0.52.

### Results

Genetic diversity and relationships revealed by multilocus enzyme electrophoretic typing. A total of 51 of the strains were characterized by determining the electrophoretic mobilities of 14 intracellular metabolic enzymes (detailed data are available upon request). All 14 enzymes were polymorphic, with two to seven alleles per locus (Table 1). The mean number of alleles per locus was 3.9, and the mean level of genetic diversity per locus was 0.52. A total of 46 ETs, each characterized by a distinct combination of electrophoretic mobilities of the 14 enzymes, were identified. Six of the ETs were represented by two strains, whereas the remaining 39 ETs contained one strain (Fig. 1). The genetic distances between ETs, which were calculated by determining the proportions of the 14 enzyme loci at which dissimilar alleles were present, were used to construct the dendrogram shown in Fig. 1. The smallest genetic distance between ETs was 0.07, corresponding to a single difference in the 14 enzymes analyzed. The phylogenetic tree revealed two major lineages, designated divisions I and II, which separated at a mean genetic distance of 0.77 and included 21 and 30 strains, respectively (Fig. 1). The type strain of *P. intermedia*
with monoclonal antibodies (MAb) according to Gmiir and Guggenheim (8) and Gmiir and Wyss (9); degrees of hybridization

The columns on the right show the ET types; strain designations; species affiliations (PL

proteins as determined by an SDS-PAGE analysis (see Fig. 4).

we observed faint bands having mobilities identical to the mo-

in addition to the

and one slowly migrating band in division II strains. The dis-

we observed faint bands having mobilities identical to the mo-

characteristic of the other species in addition to the

strongly stained band.

Genetic diversity and relationships revealed by RFLP.

Whole-cell DNAs were prepared from 46 of the 53 strains

included in this study. Only one strain, BK22, contained visible plasmid DNA. To examine the levels of genomic similarity

among strains assigned to the two divisions, Southern blots of

were hybridized with total genomic DNA from P. intermedia CCUG 24041 (= ATCC 25611) (P.i.) DNA and whole genomic P. nigrescens CCUG 9560 (= NCTC 9336) (P.n.) DNA were used as probes (+, ++, and +++ indicate intensities of hybridization, as determined from Fig. 2); degrees of hybridization to EcoRI digests when rRNA sequences were used as the probe or when oligonucleotide probe Bi31 specific for the foresort P. intermedia genotype II (P. nigrescens) was used; and protein profiles of soluble cellular proteins as determined by an SDS-PAGE analysis (see Fig. 4).

(CCUG 24041 [= ATCC 25611] in ET 15) and all of the strains
designated P. intermedia except one clustered in division I.

Strain MH3, which was received as P. intermedia, clustered
with division II (ET 26) together with the type strain of P.
nigrescens (BEF3 [= NCTC 9336 = ATCC 33563] in ET 19)
and all of the strains received as P. nigrescens.

For every enzyme tested except glutamate dehydrogenase there was at least one allele that was shared by strains belonging
to the two divisions. For glutamate dehydrogenase we ob-

erved one of two fast-migrating bands in all division I strains

and one slowly migrating band in division II strains. The dis-

ances between the three bands were small, and in some strains

we observed faint bands having mobilities identical to the mo-

bility characteristic of the other species in addition to the strongly stained band.

Genetic diversity and relationships revealed by RFLP.

Whole-cell DNAs were prepared from 46 of the 53 strains

included in this study. Only one strain, BK22, contained visible plasmid DNA. To examine the levels of genomic similarity among strains assigned to the two divisions, Southern blots of Mspl-digested whole-cell DNAs were hybridized with total genomic DNAs from the two type strains (Fig. 2). Mspl was chosen for digestion because it resulted in an equal distribution of DNA fragments in the gel. A comparison of the relative intensities of the hybridizing bands divided the strains into two groups that were identical to the two divisions based on multilocus enzyme electrophoresis data (Fig. 1 and 2). As this type of hybridization is very sensitive to the amount of DNA applied to the gel, the same filter was used for hybridization with each of the two probes, and the results were based on the relative hybridization intensity. For unknown reasons, when genomic DNA from P. intermedia CCUG 24041 (= ATCC 25611) was used as the hybridization probe, it gave clearer results than DNA from P. nigrescens CCUG 9560 (= NCTC 9336 =

FIG. 1. Genetic relationships among 51 strains based on ET typing data. Two major lineages (ET divisions) that separated at a genetic distance of 0.77 are indicated on the left. The columns on the right show the ET types; strain designations; species affilations (P.I., P. intermedia; P.n., P. nigrescens) if defined and serological reactions with monoclonal antibodies (MAb) according to Gmiir and Guggenheim (8) and Gmiir and Wyss (9); degrees of hybridization to EcoRI digests when whole genomic P. intermedia CCUG 24041 (= ATCC 25611) (P.i.) DNA and whole genomic P. nigrescens CCUG 9560 (= NCTC 9336) (P.n.) DNA were used as probes (+, ++, and +++ indicate intensities of hybridization, as determined from Fig. 2); degrees of hybridization to EcoRI digests when rRNA sequences were used as the probe or when oligonucleotide probe Bi31 specific for the foresort P. intermedia genotype II (P. nigrescens) was used; and protein profiles of soluble cellular proteins as determined by an SDS-PAGE analysis (see Fig. 4).

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FIG. 2. RFLP analysis of chromosomal DNAs from *P. intermedia* and *P. nigrescens* strains digested with *MspI* and hybridized with total genomic DNA from *P. intermedia* CCUG 24041 (= ATCC 25611T) (A) and total genomic DNA from *P. nigrescens* CCUG 9560 (= NCTC 9336T = ATCC 35837T) (B). Lane 1, CCUG 9560T (= NCTC 9336 = ACIT 33563T); lane 2, CCUG 24041 (= ATCC 25611T); lane 3, BK38; lane 4, BK37; lane 5, BK36; lane 6, BK35; lane 7, BK33; lane 8, BK32; lane 9, BK31; lane 10, BK30; lane 11, BK29; lane 12, BK27. The positions of size markers (in kilobases) are indicated on the right. When we took the amount of DNA applied to each lane into account and compared the intensities of hybridization in panels A and B as described in the text, lanes 2, 5, and 7 were scored +++ in panel A and ++ in panel B. The remaining lanes were scored + in panel A and +++ in panel B.

ATCC 33563T) (Fig. 2). Strain MH3 reacted like *P. nigrescens* strains, which is consistent with the results of the multilocus enzyme electrophoretic analysis (Fig. 1 and 2).

To further estimate the levels of genetic diversity and relationships at the DNA level, we performed ribotyping experiments with *EcoRI*-digested genomic DNAs (Fig. 1 and 3A). Hybridization with the rRNA probe from *E. coli* revealed 10 different patterns among the 15 division I strains tested. A total of 17 distinct patterns were detected among the 29 division I1 strains tested; a single predominant type was represented by 11 strains, which were scattered throughout division I1 in the dendrogram based on multilocus enzyme electrophoresis data. No ribotype was shared by members of the two divisions. Both strains belonging to five of the six ETs containing two isolates were included in the ribotyping experiments. In all five ETs the two strains exhibited different ribotypes, indicating that they represent different clones.

Oligonucleotide probe Bi31, which originated from the rRNA genes and was previously found to be specific for the former *P. intermedia* genotype I1 (*P. nigrescens*), hybridized to all division I1 strains, including strain MH3 (Fig. 1 and 3), whereas no hybridization to the division I strains was observed.

**Protein profiles.** SDS-PAGE of soluble cellular proteins of all of the strains included in this study except strain MH18, which was lost and could not be replaced, resulted in overall very similar patterns for the strains belonging to the two species (Fig. 4). However, the location of a single predominant band with a molecular weight of approximately 20,000 consistently separated the strains, which was consistent with the results obtained with the other methods (Fig. 1 and 4).

### Degradation of IgA1

After 24 h of incubation with IgA1 each strain produced immunoelectrophoretic patterns for IgA1 degradation products identical to those described previously (10): two precipitation lines (Fig. 5, experiments B1 and C1) previously identified as Fab and Fc fragments (10); one precipitation line with an electrophoretic mobility significantly different from that of intact IgA1 (Fig. 5, experiments B4 and C4) and previously identified as an Fc fragment; or no precipitation lines, which was interpreted as total degradation of the IgA1 substrate (Fig. 5, experiments B24 and C24). After prolonged incubation for 48 h with a heavy inoculum of bacteria, all strains completely degraded IgA1, as revealed by the total absence of precipitation lines (Fig. 5).

To examine the kinetics of IgA1 degradation, aliquots of IgA1 and buffer were incubated with equal amounts of five representative strains grown in liquid culture to the late log phase. Each reaction mixture was subsequently reduced and subjected to SDS-PAGE and Western blot analyses (Fig. 6). Overall similar patterns of IgA1 heavy chain degradation were observed for all five strains tested. After 1 h, intense staining with antibodies against the α chain occurred, with bands corresponding to the intact monomeric α chain (apparent molecular weight, 61,000), a fragment with an apparent molecular weight of 46,500, and a fragment with an apparent molecular weight of 20,000.
FIG. 3. RFLP analysis of chromosomal DNAs from \textit{P. intermedia} and \textit{P. nigrescens} strains digested with EcoRI and hybridized with an rRNA probe from \textit{E. coli} (A) and oligonucleotide probe Bi31 specific for the former \textit{P. intermedia} genotype II (\textit{P. nigrescens}) (B). Lane 1 HG65; lane 2, CCUG 9560 (= NCTC 9336$^{T}$ = ATCC 33563$^{T}$); lane 3, NCTC 9538; lane 4, OMZ265; lane 5, OMZ256; lane 6, MH20; lane 7, MH11; lane 8, MH7; lane 9, MH6; lane 10, MH3; lane 11, OMZ328; lane 12, OMZ310; lane 13, OMZ251; lane 14, CCUG 24041 (= ATCC 25611$^{T}$). The positions of size markers (in kilobases) are indicated on the right.

weight of 35,500. Only a faint band with an apparent molecular weight of 35,500 was observed after incubation for 48 h (Fig. 6A). The 46.5-kDa fragment may reflect proteolytic degradation of a single domain, while the 35.5-kDa fragment may reflect proteolytic degradation of two domains. The $\lambda$ light chain of the substrate IgAl (apparent molecular weight, 30,000) was also degraded, and a breakdown product with an apparent molecular weight of 14,000 was observed; this product was also degraded after incubation for 48 h (Fig. 6B).

To examine if all or part of the proteolytic activity was influenced by protease inhibitors, five representative strains were tested for their ability to degrade IgAl after preincubation with the human physiological proteinase inhibitors $\alpha_{2}$-macroglobulin and $\alpha_{1}$-proteinase inhibitor. This experiment revealed that the degradation of IgAl was not influenced by these protease inhibitors (e.g., Fig. 5).

**DISCUSSION**

A total of 53 strains, including the type strains of \textit{P. intermedia} and \textit{P. nigrescens}, as well as reference collection strains and fresh dental plaque isolates, were subjected to various tests

FIG. 4. SDS-PAGE of soluble cellular proteins from reference strains of \textit{P. intermedia} and \textit{P. nigrescens} and wild strains that clustered with one of the two species. Lanes 1 and 10, high-molecular-weight and low-molecular-weight standards (Bio-Rad Laboratories, Richmond, Calif.), respectively; lane 2, \textit{P. nigrescens} CCUG 9560 (= NCTC 9336$^{T}$ = ATCC 33563$^{T}$); lane 3, \textit{P. intermedia} CCUG 24041 (= ATCC 25611$^{T}$); lane 4, BK36; lane 5, BK28; lane 6, \textit{P. intermedia} OMZ324; lane 7, HG189; lane 8, MH51-20; lane 9, MH3. The band that differentiates species is indicated by an asterisk. Protein profile I is represented by the \textit{P. intermedia} strains in lanes 3 through 6. Protein profile II is represented by the \textit{P. nigrescens} strains in lanes 2 and 7 through 9.

FIG. 5. Immunoelectrophoretic patterns of IgAl after incubation with buffer (A), \textit{P. nigrescens} CCUG 9560 (= NCTC 9336$^{T}$ = ATCC 33563$^{T}$) cells preincubated for 1.5 h with buffer (B), and \textit{P. nigrescens} CCUG 9560 (= NCTC 9336$^{T}$ = ATCC 33563$^{T}$) cells preincubated with $\alpha_{2}$-macroglobulin at a final concentration of 1 mM (C). Preparations were incubated at 37°C for 1, 4, 24, and 48 h, as shown on the figure.
in order to determine the value of the tests for discriminating between the two species.

The results of our multilocus enzyme electrophoretic analysis based on 14 enzymes showed that *P. intermedia* and *P. nigrescens* constitute two genetically distinct populations and support the results of a quantitative DNA homology analysis (24).

Additional heterogeneity corresponding to a third serotype (8, 9) was not observed, which is in agreement with previously reported results (4, 24). Our results confirmed the combined results of Devine et al. (4), Gmur and Guggenheim (8), Gmur and Wyss (9), and Shah and Gharbia (24) which indicated that serotype 1 corresponds to *P. intermedia* (formerly genotype I of *P. intermedia*; our division I [Fig. 1]), while serotypes 2 and 3 correspond to *P. nigrescens* (formerly genotype II of *P. intermedia*; our division II [Fig. 1]).

It has been suggested that differences in the mobilities of the enzymes malate dehydrogenase and glutamate dehydrogenase can be used as differentiating characteristics for the two species (24). In this study, malate dehydrogenase was detected in seven alleles, some of which were shared by strains of both species. Glutamate dehydrogenase was detected in only three alleles, in overall agreement with the two different mobilities reported previously (24). Thus, the *P. nigrescens* strains used in our study produced only one slowly migrating band, while the *P. intermedia* strains produced either of two fast-migrating bands. However, some strains produced faint bands with mobilities identical to the mobility of the band in the other species in addition to the strongly stained band. In addition, a single mutation may influence the electrophoretic mobility of an enzyme. For these reasons, species differentiation based on the electrophoretic mobilities of only these two enzymes is not advisable. All of the other enzymes which we examined had at least one allele that occurred in both divisions, and many of the enzymes were difficult to score because of double or triple bands, which presumably were due to the presence of conformational isozymes or degradation products (20). Thus, we concluded that species differentiation by multilocus enzyme electrophoretic analysis should be based on several enzymes.

The RFLP analysis also confirmed that *P. intermedia* and *P. nigrescens* are two genetically distinct populations. In addition, the reliability of probe Bi31, which is specific for the former *P. intermedia* genotype II (*P. nigrescens*) (3), was confirmed. The very diverse ribotype patterns of *P. intermedia* and *P. nigrescens* make this method especially suitable for epidemiological studies.

In this study we found that the location of a single predominant band in SDS-PAGE gels containing soluble cellular proteins provided a way to reproducibly differentiate the two species. The protein profiles of five strains representing the two genotypes of the former species *P. intermedia* were presented by Johnson and Holdeman (12), but the usefulness of these profiles for discrimination was not commented upon. Gharbia et al. reported that a 31-kDa band was present in *P. nigrescens* but absent in *P. intermedia* (7). However, inspection of the illustration of these authors revealed a band with a molecular weight of approximately 29,000 in *P. intermedia*. Since the electrophoretic conditions which Gharbia et al. used differed from our conditions, we cannot eliminate the possibility that we observed the same phenomenon.

All of the strains of both species were capable of totally degrading human IgA1. Thus, the different degradation patterns observed previously (14) were presumably due to variations in the amounts and conditions of the bacteria in the reaction mixtures. Western blotting (Fig. 6) showed the presence of a 35.5-kDa degradation fragment, which on the basis of its size has previously been identified as the Fc fragment of IgA1 after cleavage with specific IgA1 proteases (6, 19). However, it is not clear on the basis of our results whether a specific IgA1 protease was involved in IgA1 degradation by the two *Prevotella* species, since Fab fragments with molecular weights of approximately 26,000 were not observed. The inability of physiological human proteinase inhibitors to prevent part or all of the proteolytic degradation of IgA1 could be due to the resistance of the IgA1-degrading enzymes to the protease inhibitors or to degradation of the inhibitors (2). Since the IgA1-degrading enzymes were not influenced by physiological protease inhibitors, they might be expected to be active in vivo in periodontal pockets.

In conclusion, in this study we confirmed that *P. intermedia* and *P. nigrescens* represent genetically distinct populations of bacteria. The methods which we employed allow clear separation of the two species.

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REFERENCES


