MOLECULAR TYPING AND EPIDEMIOLOGY

Genomic DNA restriction site heterogeneity in bovine Pasteurella multocida serogroup A isolates detected with an rRNA probe

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A total of 81 Pasteurella multocida isolates from healthy and diseased dairy and beef cattle originating from various geographical locations was examined by rRNA gene restriction site polymorphism analysis (ribotyping), restriction endonuclease analysis (REA), SDS-PAGE analysis of whole-cell (WCP) and outer-membrane (OMP) proteins, and capsule and somatic serotyping. Bacterial strains were isolated from nose, lung and in one case testicle, of Holstein and cross-bred beef cattle. The isolates represented for the most part serogroup A3 (88%). Ribotyping was performed on DNA digested with HaeII, electrophoresed and then hybridised with 32P-labelled 16S-23S rRNA from Escherichia coli. Six ribotypes (R1-R6) and 10 REA types were found among the 81 isolates with similar discrimination index (DI) of c. 0.60. Protein profiles revealed reproducibility and high levels of polymorphisms among lung isolates. Isolates were compared according to their geographical habitat, their isolation from dairy or from beef cattle and from nasal cavities or lungs. No correlation was apparent between geographical locations and ribotypes. Overall, isolates obtained from dairy cattle were predominantly R1, whereas those obtained from beef cattle were equally distributed between R1 and R2. R1 was more representative of lung isolates. For some strains, particularly the single isolate ribotypes, good correlation was achieved between WCP analysis, REA types and ribotypes. For others, REA to some extent and WCP profiles were able to discriminate among isolates within ribotypes. The data suggest that a combination of ribotyping, REA and WCP analysis is useful for investigating the epidemiology of bovine P. multocida serogroup A.

Introduction

Pasteurella multocida, a pathogenic gram-negative bacterium with coccoid or short rod morphology, has a broad host range including most mammals, birds and man [1]. It contributes substantially to bovine pneumonic pasteurellosis, which causes major economic losses to beef cattle and dairy industries in the USA [2].

P. multocida comprises five capsular serogroups of 16 somatic serotypes [1, 3]. In cattle, P. multocida serogroups B and E are the pathogens of bacterial haemorrhagic septicaemia, and serogroup A is mainly associated with respiratory disease [1, 4, 5]. P. multocida A3 is the second most commonly isolated bacterium from fibrinous pneumonia in beef cattle (with prevalence of 21–34%) and the major cause of fibrinopurulent pneumonia in dairy calves (>65%) [6–9].

In epidemiology and other fields of applied microbiology, reliable methods for the identification and characterisation of bacterial strains within a particular genus and species are primary requirements. Several methods have been used to differentiate bacterial strains, including those of P. multocida [10–19]. Traditional phenotypic methods, such as outer-membrane-protein (OMP) subtyping, whole-cell-protein (WCP) analysis, serotyping and biotyping [11, 14, 18, 20–25] are not sensitive enough for strain differentiation. Restriction endonuclease analysis (REA) of whole genomic DNA has been useful in the genetic characterisation of bacterial strains [11, 16,
26-28, but its limitations have led to the use of *Escherichia coli* ribosomal RNA (rRNA) as a broad-spectrum probe (ribotyping) for both epidemiological and taxonomic purposes [29-37]. Recently, Townsend et al. [38] used ribotyping and field alteration gel electrophoresis (FAGE) to differentiate *P. multocida* isolates from haemorrhagic septicaemia. To our knowledge, the discriminatory power and application of molecular techniques have not been investigated in bovine *P. multocida* serogroup A isolates from the respiratory tract. Furthermore, there are no discriminating data available on such bovine *P. multocida* isolates in relation to animal group, organ or geographical habitat.

In this study, 81 bovine *P. multocida* isolates from healthy and diseased dairy and beef cattle originating from different geographical locations were characterised by five molecular and traditional typing methods – serotyping, SDS-PAGE of WCP and OMPs, REA of whole genomic DNA and ribotyping.

### Materials and methods

#### Bacterial strains and growth conditions

Eighty-one *P. multocida* isolates were examined (Table 1). They were isolated from nose (33), lung (47) and testicle (1) of Holstein (48) and cross-bred beef cattle (33). The isolates were collected from animals in Oklahoma (51), California (28), Kansas (1) and Missouri (1). They were grown on Bacto Brain-Heart Infusion (BHI) Agar (Difco Laboratories, Detroit, MI, USA), at 37°C in air with CO₂ 5% overnight [39]. For broth cultures, overnight seed cultures were used to inoculate BHI broth and grown to a mid-logarithmic phase at 37°C with shaking.

#### Capsule and somatic serotyping

Serogrouping was performed according to Rimler [19] by hyaluronidase, chondroitinase AC and heparinase III sensitivity tests. Serotyping was performed by agar gel precipitation with serotype specific antisera [5].

### SDS-PAGE of whole-cell and outer-membrane proteins

Bacterial envelopes were prepared by sonicaton [40]. The Sarkosyl insoluble (SI) method, which is based on the differential solubility of cytoplasmic (inner) and outer membranes in n-lauroyl sarcosine, was used as described previously [41]. Twelve *P. multocida* isolates, including a representative of each ribotype, were selected for OMP extraction. Whole-cell lysates were obtained and SDS-PAGE was done as described previously [42]. Protein bands were visualised after staining with Coomassie Brilliant Blue R250 (Fisher Scientific, Fair Lawn, NJ, USA) 0.1% w/v in acetic acid 10% v/v, methanol 40% v/v for 1 h at room temperature with gentle shaking and then de-staining in acetic acid 10% v/v, methanol 10% v/v.

### Genomic DNA preparation, REA and separation of DNA fragments

Genomic DNA of high mol.wt was prepared with a Qiagen DNA mini prep kit according to the manufacturer's instructions (Qiagen, Santa Clarita, CA, USA). Residual proteins were removed by phenol-chloroform extraction and DNA was precipitated with 2.5 volumes of cold absolute ethanol and 0.3 M ammonium acetate at -20°C overnight. The preparation was centrifuged at 12,000 rpm in a microcentrifuge at 4°C for 30 min. Pellets were washed with ethanol 70% v/v, dried, dissolved in 200 µl of TE buffer and stored at 4°C. DNA concentration was measured spectrophotometrically at 260 nm [43]. For ribotyping and REA, purified DNA (3 µg) was digested overnight at 37°C in 20-µl volumes containing the restriction endonucleases, according to manufacturer's instructions (Promega Corporation, Madison, WI, USA). Selected restriction endonuclease digests were checked continuously for reproducibility of the method. The digested DNA was electrophoresed at 24 V (constant voltage) overnight in agarose (Gibco-BRL, Grand Island, NY, USA) 0.7% w/v in TBE buffer (89 mM Tris base, 89 mM sodium acetate, 2.4 mM sodium EDTA, pH 8.0) and TAE buffer (40 mM Tris base, 1 mM EDTA, pH 8.0), respectively for ribotyping and REA.

### Radiolabelling of rRNA

Ribosomal 16S-23S RNA (3 µg) from *E. coli* (Boehringer Mannheim, Indianapolis, IN, USA) was alkaline hydrolysed with an equal volume of 100 mM sodium bicarbonate (NaHCO₃), pH 9, at 95°C for 20 min, precipitated with 2.5 volumes of absolute ethanol and 0.3 M NaOAc, dried and dissolved in 10 µl of sterile double-distilled water. The dissolved DNA was 5’ end-labelled with (γ-³²P)ATP with T4 polynucleotide kinase (United States Biochemical, Cleve-
land, OH, USA) according to the manufacturer's instructions.

Hybridisation in dried agarose gels

The general strategy for hybridisation in dried gels was adapted from Mather [44] and Lueders and Fewell [45]. Briefly, following electrophoretic separation of the digested DNA fragments, the agarose gel was placed on a piece of Whatman 3-mm paper, covered with plastic wrap and dried on a vacuum gel drier (Hoefer Scientific Instruments, San Francisco, CA, USA) for 30 min at room temperature and 30 min at 60°C followed by a 30-min cooling period. Before hybridisation, the dried gel was placed in a dish of de-ionised water for a few minutes to remove the paper backing. The gel was then treated with 0.5 M NaOH, 0.15 M NaCl for 45 min to denature the DNA, soaked in de-ionised water for 30 min, then neutralised in 0.5 M Tris-HCl, pH 7.5, 0.15 M NaCl for 45 min. The gel was washed for 30 min in de-ionised water then prehybridised for 3 h at 50°C with a solution containing 6× SSPE (1× SSPE is 20 mM NaH₂PO₄·H₂O, 180 mM NaCl, 8 mM NaOH, 1 mM EDTA, pH 7.0), 1× Denhardt's reagent, SDS 0.5% w/v, denatured sonicated salmon sperm DNA, 100 µg/ml and sodium pyrophosphate (NaPPi) 0.05%. Hybridisation was performed for 16 h at 50°C in the solution containing the radioactive rRNA probe. After hybridisation, gels were washed twice in 2× SSPE SDS 0.1% w/v at room temperature for 15 min each and once in 1× SSPE SDS 0.1% w/v at 60°C for 20 min. Gels were autoradiographed with Kodak X-OMAT film at −70°C with an intensifying screen.

Statistical analysis

The discrimination index (DI) of ribotyping and REA, i.e., the probability that two unrelated strains randomly selected from the test population would fall into different typing groups, was calculated by Simpson's index of diversity [46]. The percentage of similarity (%S) between ribotypes was calculated as follows: %S = [(number of matching bands × 2)/total number of bands] × 100.

Results

Serotyping

All 81 P. multocida isolates were confirmed as being capsular serogroup type A by the disk diffusion decapsulation test. Five serotypes were represented; these included one A2, 71 A3, three A4, one A7, four A10 and one untypable isolate. Serotype A3 represented 88% of total isolates, and 85% and 88% for beef and dairy cattle isolates, respectively. Serotype A10 isolates were all beef cattle isolates and represented 5% of total isolates and 12% of beef cattle isolates. In all, 4% of total isolates were serotype A4, which comprised one beef (3%) and two dairy (4%) cattle isolates. The single isolate serotypes A2 (dairy cattle), A7 (beef cattle) and the untypable isolate (dairy cattle) each comprised 1% of the total isolates.

Ribotyping

Preliminary experiments with different restriction endonucleases (REs) and RE combinations showed that HaeII was the most discriminating enzyme as judged by the number, range and variety of bands after hybridisation with the rRNA probe (data not shown). For that reason, HaeII was used in the ribotyping of the isolates under investigation. Ribotypes are defined as patterns of bands containing rRNA gene sequences and were assessed by including the preparations of DNA from isolates representative of known ribotypes in every Southern hybridisation experiment. Isolates were denoted as distinct when their hybridisation patterns differed by one or more bands. A total of six different ribotypes (designated R1–R6) was observed for the 81 isolates tested (Fig. 1) and these accounted for 57% (46), 21% (17), 14% (11), 6% (5), 1% (1) and 1% (1) of

Fig. 1. Representative ribotypes of P. multocida bovine isolates. Lane 1, 9040970; 2, T94289; 3, 95120769; 4, 95110872; 5, 96020298; 6, T931317. Hybridisation was performed with 32P-labelled DNA probe of 16+23S rRNA from E. coli. Note the similarity between R1, R2 and R5 and between R3, R4 and R6. Size markers in (kb) are indicated.
the total isolates, respectively. The numbers of bands in the six ribotypes varied from eight to nine with band sizes ranging from 1.6 to >12 kb. The six Haell ribotypes revealed four monomorphic bands (1.6, 2.0, 2.6 and 4.3 kb). The analysis of uncommon bands established 12 polymorphic Haell restriction sites for the six ribotypes. R1 and R2 had two polymorphic bands at the anode and one at the cathode. R3 and R4 differed only in one polymorphic band at the cathode. R5 and R6 were single isolate ribotypes.

**Restriction enzyme analysis (REA)**

For REA, the large number of bands generated by digestion of genomic DNA with various restriction endonucleases hindered a detailed visual assessment of the method. Haell was selected from the restriction endonucleases tested, because it displayed the most interpretable, consistent and discriminating restriction bands, the majority of which ranged between 1.6 and 12 kb. The enzyme revealed 10 REA types (Fig. 2) among the 81 *P. multocida* isolates. REA types 1, 3, and 2c accounted for 58% (47), 14% (11) and 12% (10) of the isolates, respectively. REA types 2a, 2b and 4a each accounted for 4% (3) of the isolates. The remaining REA types (4a, 4b, 5 and 6) each accounted for 1% (1) of the isolates. For the most part, the REA results correlated with ribotypes, with the exception of R2 and R4, which each exhibited three REA types. R1, R2 and the single isolate ribotypes R5 and R6 all exhibited different REA types.

**SDS-PAGE analysis of *P. multocida* isolates**

WCP profiles of the *P. multocida* isolates showed good reproducibility both within and between the gels, as indicated by the use of representative *P. multocida* WCP as reference profiles on SDS-PA gels (data not shown). The 81 isolates examined by SDS-PAGE gave 36 WCP profiles (Fig. 3) with a high level of polymorphisms within R1, R2, R3 and R4. WCP profile differences were based upon the presence, absence or intensity of protein bands. R1 alone had 17 different WCP profiles. However, the single isolate ribotypes R4 and R5 exhibited different WCP profiles and as such correlated with ribotype and REA results.

SDS-PAGE of OMP profiles revealed polymorphisms among the 12 *P. multocida* isolates investigated (data not shown). The OMP profiles of isolates representative of the six ribotypes (Fig. 4) revealed six distinct OMP profiles.

**Analysis of isolates**

The study examined the *P. multocida* lung isolates for any association between ribotypes and (i) animal
SUBTYPING OF BOVINE \textit{P. multocida} STRAINS

Fig. 4. Coomassie Blue-stained SDS 12.5% PA gel of OMPs of \textit{P. multocida} bovine isolates representative of the six ribotypes. Isolates in lanes 1–6 are as indicated in Fig. 1 for R1–R6. Broad-range (BioRad) size markers (in kDa) are indicated.

Isolates were grouped according to ribotype, WCP pattern, REA type and serotype with respect to whether they were nasal or lung, beef or Holstein cattle in origin (Table 3). For beef cattle lung isolates, R1 isolates contained five different WCP patterns among 10 isolates, one REA type and three different serotypes. Variations in WCP patterns and serotypes were also seen for R2 and R3. In contrast, Holstein cattle lung isolates of R1 contained eight different WCP patterns among 20 isolates, one REA type and a single serotype. WCP patterns varied for R2 and R3; however, REA types and serotypes were consistent for each of the ribotypes. Each beef cattle nasal isolate was of a single WCP pattern (except R2, which contained two different WCP patterns) and REA type for each of the five ribotypes. Each Holstein cattle nasal isolate was some variation among the R1 group as to WCP pattern and serotype; however, a single REA type and serotype was seen for both R1 and R4 and R4, respectively.

\textbf{Table 2.} State of origin of \textit{P. multocida} isolates with respect to ribotype

<table>
<thead>
<tr>
<th>Origin</th>
<th>Number of cattle</th>
<th>Number of isolates of ribotype</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beef cattle</td>
<td>33</td>
<td></td>
</tr>
<tr>
<td>Lung</td>
<td>15</td>
<td>7 3 3 - - -</td>
</tr>
<tr>
<td>Oklahoma</td>
<td>13</td>
<td>1 1 - - - -</td>
</tr>
<tr>
<td>Kansas</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Missouri</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Nose</td>
<td>18</td>
<td>3 10 1 3 1 -</td>
</tr>
<tr>
<td>Oklahoma</td>
<td>18</td>
<td></td>
</tr>
<tr>
<td>Holstein cattle</td>
<td>48</td>
<td></td>
</tr>
<tr>
<td>Lung</td>
<td>32</td>
<td>16 4 1 - - -</td>
</tr>
<tr>
<td>California</td>
<td>28</td>
<td>14 - - 1 - -</td>
</tr>
<tr>
<td>Oklahoma</td>
<td>4</td>
<td>1 1 - - - -</td>
</tr>
<tr>
<td>Nose</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>Oklahoma</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>Testicle</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Oklahoma</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>46</td>
<td>17 11 5 1 1</td>
</tr>
</tbody>
</table>

\textbf{Statistical analysis}

Simpson’s index of diversity [46] was used to evaluate the discrimination power of ribotyping and REA. The use of HaeII in both methods resulted in DI values of 0.60 and 0.63 for the six ribotypes and 10 REA types, respectively. This indicates that for both ribotyping and REA, when 2 isolates were randomly sampled from the tested population, they would fall into different types on at least 60% of the occasions. For ribotyping, the variation of the discrimination power with total lung isolates only (47) and isolates within diseased animal groups (32 and 15 for Holstein and beef cattle, respectively) was also evaluated. In all three cases the
isolates were discriminated with a DI of 0.60, similar to that reported above for the whole test population. The percentage similarity between ribotypes is shown in Table 4. The results suggest that R1, R2 and R5 (71–88% similarity) are related, as are R3, R4 and R6 (89% similarity).

The discriminatory power of WCP patterns expressed as the number of different WCP profiles/number of isolates revealed a high frequency of polymorphism within R1, R2 and R3 for both Holstein and beef cattle lung isolates as compared with nasal isolates (Table 3). Only R1 Holstein and R2 beef cattle nasal isolates exhibited some WCP polymorphisms.

### Discussion

Radiolabelled *E. coli* rRNA was used successfully as a broad-spectrum probe to characterise 81 bovine *P. multocida* isolates from healthy and diseased dairy and beef cattle originating from different geographical locations. The isolates were also subtyped by serotyping, REA of whole genomic DNA, and WCP and OMP profiles. Characterisation of the isolates by serotyping turned out to be of limited value as reported previously [35,47]; >88% of the isolates were of the same serotype, A3, consistent with previous reports [6–9].

The 81 *P. multocida* isolates were discriminated into six ribotypes with HaeII restriction endonuclease. Townsend *et al.* [38] reported seven distinct ribotypes for both *EcoRI* and *PstI*-digested DNA among 19 haemorrhagic septicaemia (HS)-associated and non-associated *P. multocida* isolates from Asia, Africa and North America. There was a high degree of homogeneity among the *P. multocida* isolates within geographical locations with the exception of the African *P. multocida* isolates, which exhibited similarities with various unrelated isolates. In this study, the *HaeII* enzyme alone produced easily interpretable rRNA banding patterns with a high discriminatory power of 0.60. This DI was identical to those calculated for *P. multocida* lung isolates and for isolates within each diseased animal group. R1 alone and R2, R3 and R4 together made up 57% and 41% of the isolates, respectively. The remaining 2% included single isolate ribotypes 5 and 6.

Differences between R1, R2 and R5 and between R3, R4 and R6 were small when based upon the percentage similarity and the number of polymorphic bands. These *HaeII* ribotype differences clearly grouped the bovine *P. multocida* isolates into two major clusters. As indicated by Townsend *et al.* [38], DNA macrosection differences of up to three fragments are probably representative of genotypic variants of the same isolate. Therefore, the above ribotype differences were probably caused by small genomic re-arrangements, such as insertion or deletion around the *HaeII* restriction site, primarily between the rRNA transcriptional units.

The large number of bands resulting from REA is an obstacle when comparing large numbers of isolates. Nevertheless, REA of genomic DNA, performed by digestion with *HaeII*, revealed 10 REA types with a DI of 0.60. In general, REA results correlated with

### Table 3. Grouping of *P. multocida* isolates by ribotypes, WCP patterns, REA types and serotypes

<table>
<thead>
<tr>
<th>Origin</th>
<th>Ribotype</th>
<th>WCP pattern*</th>
<th>REA type</th>
<th>Serotype†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beef cattle</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lung isolates</td>
<td>R1</td>
<td>5/10</td>
<td>1</td>
<td>A10(2), A3(7), A4(1)</td>
</tr>
<tr>
<td></td>
<td>R2</td>
<td>3/3</td>
<td>2a</td>
<td>A3(3)</td>
</tr>
<tr>
<td></td>
<td>R3</td>
<td>3/3</td>
<td>3</td>
<td>A10(2), A3(1)</td>
</tr>
<tr>
<td>Nasal isolates</td>
<td>R1</td>
<td>1/3</td>
<td>1</td>
<td>A3(3)</td>
</tr>
<tr>
<td></td>
<td>R2</td>
<td>2/10</td>
<td>2c</td>
<td>A3(9), A7(1)</td>
</tr>
<tr>
<td></td>
<td>R3</td>
<td>1/1</td>
<td>3</td>
<td>A3(1)</td>
</tr>
<tr>
<td></td>
<td>R4</td>
<td>1/3</td>
<td>4a</td>
<td>A3(3)</td>
</tr>
<tr>
<td></td>
<td>R5</td>
<td>1/1</td>
<td>5</td>
<td>A3(1)</td>
</tr>
<tr>
<td>Holstein cattle</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lung isolates</td>
<td>R1</td>
<td>8/20</td>
<td>1</td>
<td>A3(20)</td>
</tr>
<tr>
<td></td>
<td>R2</td>
<td>3/3</td>
<td>2b</td>
<td>A3(3)</td>
</tr>
<tr>
<td></td>
<td>R3</td>
<td>2/7</td>
<td>3</td>
<td>A3(7)</td>
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<tr>
<td></td>
<td>R4</td>
<td>1/1</td>
<td>4</td>
<td>A4(1)</td>
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<tr>
<td></td>
<td>R6</td>
<td>1/1</td>
<td>6</td>
<td>UT(1)</td>
</tr>
<tr>
<td>Nasal isolates</td>
<td>R1</td>
<td>3/14</td>
<td>1</td>
<td>A3(12), A4(1), A2(1)</td>
</tr>
<tr>
<td></td>
<td>R4</td>
<td>1/1</td>
<td>4c</td>
<td>A3(1)</td>
</tr>
</tbody>
</table>

* Expressed as number of different WCP patterns/number of isolates.
† Numbers in parentheses are the numbers of isolates of the serotype.

### Table 4. Percentage similarity between ribotypes

<table>
<thead>
<tr>
<th>Ribotype</th>
<th>R1</th>
<th>R2</th>
<th>R3</th>
<th>R4</th>
<th>R5</th>
<th>R6</th>
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<tr>
<td>R1</td>
<td>82</td>
<td>56</td>
<td>56</td>
<td>71</td>
<td>42</td>
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</tr>
<tr>
<td>R2</td>
<td>71</td>
<td>71</td>
<td>88</td>
<td>71</td>
<td>42</td>
<td></td>
</tr>
<tr>
<td>R3</td>
<td>89</td>
<td>89</td>
<td>89</td>
<td>89</td>
<td>89</td>
<td></td>
</tr>
<tr>
<td>R4</td>
<td>82</td>
<td>89</td>
<td>89</td>
<td>89</td>
<td>89</td>
<td></td>
</tr>
<tr>
<td>R5</td>
<td>71</td>
<td></td>
<td></td>
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</tbody>
</table>
ribotypes. In a few cases where ribotyping was found to be less discriminatory than the REA of genomic DNA, it is possible that the maximal level of sensitivity of the ribotyping may not have been reached by the use of a single HaeII restriction endonuclease. Previous studies indicated that increasing the number of enzymes used in ribotyping is usually followed by an increase in discriminatory power [30, 47].

To our knowledge, no discriminating data are available for ribotyping of bovine Pasteurella multocida serogroup A isolates from the respiratory tract, on the basis of animal group, site of isolation or geographical location. The data from the present study convincingly showed no association between geographical locations and ribotypes as indicated by the predominance of R1 in all geographical locations. However, R1 seemed to be most representative of isolates from Holstein calves (74%) and lung (63%). The preponderance of R1 in particular, and to some extent R2 and R3, in all geographical locations suggest a spread of infection resulting from the same source or a single clone. This may have occurred over time as infected cattle were moved from one area to another within and between states. In contrast, Townsend et al. [38] reported some correlation between ribotype and geographical origin of HS-causing Pasteurella multocida isolates. However, the geographical locations varied greatly in their study and included Asia, Africa and North America. Chelasius-Dancla et al. [31], in their ribotyping study of Pasteurella species isolated from animals, reported limited polymorphism in bovine Pasteurella haemolytica strains from independent breeding herds and high diversity in Pasteurella multocida strains from rabbit farms. They hypothesized that high selective pressure by antibiotics may have introduced an evolutionary constraint on bovine Pasteurella haemolytica with drastic consequences on their genetic polymorphism. There was no indication of such high selective pressure by antibiotics on the bovine Pasteurella multocida isolates in this study. A more complete ribotype study with isolates from outbreaks of pneumonia within dairy herds would be necessary to determine consistency of various ribotypes in those situations.

The isolates were also subtyped by WCP and OMP. Unlike serotyping and REA, WCP exhibited high frequency of polymorphisms within and between ribotypes. Most polymorphisms seen in WCP were in lung isolates. Although a definitive correlation between ribotypes and WCP profiles was not observed, the single isolate ribotypes exhibited different WCP profiles. OMP subtyping was able to discriminate between isolates representative of the six ribotypes.

In conclusion, ribotyping was clearly discriminating and six ribotypes were observed for the 81 Pasteurella multocida serogroup A bovine isolates. The six HaeII ribotypes may have originated from two Pasteurella multocida genetic variants through small genomic re-arrangements such as deletion or insertion around the HaeII restriction site. REA, OMP and WCP analysis, with all their reported limitations, can be useful complements to ribotyping in epidemiological and taxonomic studies of Pasteurella multocida isolates from cattle.

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References


