Proliferative enteritis (PE) is an enteric disease that occurs primarily in weanling animals but is also seen in older animals. It has been reported in a variety of species, including the pig, hamster, ferret, fox, rat, rabbit (27), horse (5, 34), deer (4), ostrich (36), and emu (17). The disease has been most thoroughly studied in the pig and the hamster (27). It may be chronic causing weight loss and poor growth, or more acute, occasionally resulting in death. Antibiotics may be used to treat PE; however, many cases resolve spontaneously. The disease is widespread in swine. Poor weight gain and deaths during epizootics result in production loss and represent a significant cost to the swine industry (14, 27). The significance of PE in other species is not known. Because of the wide host range of the disease, it is likely that there are numerous reservoirs, and the disease may represent an equally serious threat to these species.

PE causes hyperplasia of the intestinal mucosa, specifically the crypt epithelial cells. Large numbers of small, curved intracellular bacteria are present within the proliferating cells (27). The disease may be diagnosed by observation of these lesions at necropsy, by detecting the intracellular organism in histologic sections with silver stain (27), or by various immunologic (15, 20) and molecular (8, 9, 12, 13, 19, 22) methods. Serologic diagnosis of PE has not proven to be useful (10, 16). The organism has been isolated in cell cultures from hamsters and pigs with PE but has not been grown on cell-free medium (15, 31). Studies have shown that this bacterium is the causative agent of PE in pigs (15, 23) and probably hamsters (31). Sequence analysis of the 16S rDNA of this bacterium shows that it is most closely related to the sulfate-reducing bacterium Desulfovibrio desulfuricans. The intracellular agents found in the lesions of different animal species are antigenically similar. In addition, strains from the pig, ferret, and hamster have been shown to be genetically similar. In this study we performed a partial 16S ribosomal DNA sequence analysis on the intracellular agent of proliferative enteritis from a hamster, a deer, and an ostrich, and compared these sequences to that of the porcine L. intracellularis isolate. Results of this study indicate that the intracellular agents from these species with proliferative enteritis have high sequence similarity, indicating that they are all in the genus Lawsonia and that they may also be the same species, L. intracellularis.

The intracellular agents of PE are morphologically identical in all species that the disease affects (4, 5, 17, 27, 34, 36). The ability of monoclonal antibodies developed from a pig isolate of the organism to detect the organism in tissues from other species demonstrates that the intracellular organisms seen in lesions from different species are antigenically similar (4, 6, 14, 24, 34, 36). Sequence analysis of the 16S rDNA from this organism from the pig (7, 10), the hamster (6, 26), and the ferret (6) suggests that the intracellular agents of PE in these species are also genetically similar.

In this study we performed a complete 16S rDNA sequence analysis of the intracellular agent of PE in the hamster and a partial 16S rDNA sequence analysis on the intracellular agent of PE extracted from formalin-fixed, paraffin-embedded tissues from a deer and an ostrich with PE. The sequence obtained demonstrated high similarity between the 16S rDNA from the intracellular agent of PE from these species and that of the pig isolate of L. intracellularis. These results suggest that the intracellular agents of PE in the hamster, deer, and ostrich are all isolates of L. intracellularis.

**MATERIALS AND METHODS**

**Samples.** The hamster intracellular agent of PE (strain auratus) was grown in INT 407 human embryonic intestinal epithelial cells, and the DNA was extracted for PCR and sequence analysis. The organism was fed to newly weaned hamsters (Mesocricetus auratus) according to protocols previously described (31). The animals were necropsied at 21 days, and the ilea were harvested. Ilea from one of these experimentally infected hamsters and from a deer (Odocoileus virginianus) and an ostrich (Struthio camelus D94-06196) with spontaneous PE were examined. PE was diagnosed by histopathologic examination of formalin-fixed, paraffin-embedded tissues stained with Warthin-Starry silver stain and by multiplex PCR (2). The hamster organism was kindly provided by Harold Stills, College of Veterinary Medicine, Ohio State University, Columbus. Deer tissues were kindly provided by Richard Drolet, Université de Montréal, Québec, Canada. Ostrich tissues were kindly provided by Leslie Woods, Veterinary Diagnostic Laboratory, University of California, Davis.) Tissues frozen at ~70°C or formalin-fixed, paraffin-embedded samples of the ilea from these animals were selected for DNA extraction, amplification, cloning, and sequencing.

**Sample digestion.** Formalin-fixed, paraffin-embedded samples were digested by methods similar to the procedures of Isola et al. (11). Briefly, 10 to 20 5-mm thick slices were cut from each block, deparaffinized once in 1 ml of xylene, and rinsed twice in 100% ethanol. Samples were dried at room temperature and resuspended in 1 ml of DNA extraction buffer (0.3 mg of proteinase K/ml, 100

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**Comparison of the 16S Ribosomal DNA Sequences from the Intracellular Agents of Proliferative Enteritis in a Hamster, Deer, and Ostrich with the Sequence of a Porcine Isolate of Lawsonia intracellularis**

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mNaCl, 10 mM Tris-HCl [pH 8.3], 25 mM EDTA [pH 8.0], 0.5% sodium dodecyl sulfate). Samples were incubated at 56°C for 3 days with 0.2 mg of proteinase K added for 24 and 48 h. After 72 h, the samples were cooled to 4°C prior to DNA extraction.

**DNA extraction.** DNA was extracted from cultured cells with a phenol-chloroform (26). DNA was extracted from tissue samples with diatomaceous earth and guanidine thiocyanate (GuSCN) as described by Blot et al. (13). Briefly, 0.5 ml of a sample was mixed with 0.9 ml of lysis buffer (6 mM GuSCN, 0.1 M Tris-HCl [pH 6.4], 0.65% Triton X-100) and 40 μl of diatomaceous-earth suspension (200 mg/ml of 0.17 M HCl). The sample was vortexed, incubated at 20°C for 30 min, and pelleted at 14,000 × g for 5 min. The pellet was resuspended in 200 μl of DNA solubilizer buffer (20 mM Tris-HCl, pH 6.4) and twice in 200 μl of 70% ethanol, with vortexing and pelleting of the sample with each rinse. The pellet was dried at 20°C overnight and resuspended in 6.41 ml of water and subsequently 200 μl of water.

Briefly, 0.5 ml of a sample was mixed with 0.9 ml of lysis buffer (60 mM Tris-HCl [pH 8.3], 50 mM KCl, 1.5 mM MgCl₂, 50 mM NaCl, 1 mM bovine serum albumin/ml, 70 mM p-mercaptoethanol, 1 μM ATP, 20 mM dithiothreitol, and 1 μM of extracted sample DNA (−1 ng) or 1 μl of sterile distilled water as a negative control. The mixture was heated to 95°C for 5 min and cooled to 4°C, and 1 U of Taq polymerase (Boehringer Mannheim, Indianapolis, Ind.) was added. The sample was subjected to 25 cycles of PCR (DNA thermal cycle 480; Perkin-Elmer, Foster City, Calif.) with an amplification efficiency of 30% for 30 s, 45°C for 1 min, and 72°C for 1 min. PCR products from mucosal samples and plasmids purified from clones, described in the next section, were analyzed by agarose gel electrophoresis. Briefly, 10 μl of the PCR product or of a 1:10 dilution of plasmid was combined with a 0.5% agarose gel loading solution (40% sucrose, 0.1 M EDTA [pH 8.0], 0.5% sodium lauryl sulfate, and 0.05% bromophenol blue; Sigma, St. Louis, Mo.) and subjected to agarose gel electrophoresis on 2% agarose gel in TAE buffer (0.04 M Tris-acetate, 0.001 M EDTA with 0.8 μg of ethidium bromide/ml) for 1.5 h at 60 V and photographed under UV illumination.

Cloning. The PCR products were cloned into a 3.9-kb plasmid vector (TA cloning kit; Invitrogen, San Diego, Calif.) according to the manufacturer’s directions. Briefly, 7 μl (−7 ng) of PCR product was added to 50 ng of pCRII vector and 50 ng of pGEM5 easy vector (Promega, Madison, Wis.) in 250 μl of ligase buffer (CRII [pH 8.0], 50 mM NaCl, 1 mg of bovine serum albumin/ml, 70 mM β-mercaptoethanol, 1 mM ATP, 20 mM dithiothreitol, and 10 mM spermidine) and incubated at 12°C for 16 h. One microliter of the ligation mixture was added to 500 μl of fresh SOC medium (Boucheicher Mannheim, Indianapolis, Ind.) and added. The sample was subjected to 35 cycles of PCR (DNA thermal cycle 480; Perkin-Elmer, Foster City, Calif.) with an amplification efficiency of 93% for 30 s, 45°C for 1 min, and 72°C for 1 min. PCR products from mucosal samples and plasmids purified from clones, described in the next section, were analyzed by agarose gel electrophoresis. Briefly, 10 μl of the PCR product or of a 1:10 dilution of plasmid was combined with a 0.5% agarose gel loading solution (40% sucrose, 0.1 M EDTA [pH 8.0], 0.5% sodium lauryl sulfate, and 0.05% bromophenol blue; Sigma, St. Louis, Mo.) and subjected to agarose gel electrophoresis on 2% agarose gel in TAE buffer (0.04 M Tris-acetate, 0.001 M EDTA with 0.8 μg of ethidium bromide/ml) for 1.5 h at 60 V and photographed under UV illumination.

Sequence analysis. Sequences determined by the above-described methods were aligned manually with the GDE multiple sequence editor (distributed by the Ribosomal Database Project [18]) and compared with the 16s RNA sequence for L. intracellularis, the swine intracellular agent of PE (GenBank accession no. U03147), with that for the intracellular agent of ferret proliferative colitis (GenBank accession no. U07570) (6), with that for another isolate of the intracellular agent of hamster PE (GenBank accession no. U07569) (6), and with another report of the sequence for this strain (auratus) of the intracellular agent of hamster PE (GenBank accession no. U084623) (26). To also determine the relationships between these organisms and their nearest taxonomic relative with pathogenic potential, L. intracellularis was compared with a Desulfovibrio isolate from a human liver abscess (GenBank accession no. U42221) (32) by the same methods.

Nucleotide sequence accession numbers. The sequences of the cloned 16s rDNAs from each species have been submitted to GenBank with the following accession numbers: hamster, U65995; deer, U65996; ostrich, U65997.

**RESULTS**

Diagnosis of PE. Ileum from a hamster, an ostrich, and a deer, all with PE, were fixed, stained with hematoxylin and eosin and with Warthin-Starry silver stains, and examined by light microscopy. There was hyperplasia of the ileal crypt epithelium with silver-stained organisms present within the apical cytoplasm of crypt enterocytes in the hamster, deer, and ostrich with PE (Fig. 1).

Sequence analysis. Individual recombinant clones, each containing the partial 16s rDNA sequence from the intracellular agent of PE from the hamster, the ostrich, or the deer, were selected for plasmid purification and cycle sequencing with fluorescence-labeled nucleotides and automated sequence determination. These sequences were compared with the sequence of the L. intracellularis type strain (an isolate of ileal sibomint intracellularis, NCTC 12656, grown in cell culture; GenBank accession no. U30147) (21) and with sequences determined for strains of the intracellular agent of proliferative colitis in the ferret and of the agent of PE in the hamster. A 1,109-bp sequence corresponding to bp 358 to 1465 of L. intracellularis was determined from the hamster sample. Sequences of 255-bp corresponding to bp 815 to 1070 of L. intracellularis were determined from the ostrich and deer samples. There was 100% sequence similarity between the 255-bp sequence from L. intracellularis and those of the intracellular agents of PE in the hamster, deer, and ostrich (Fig. 2). There was 100% similarity over the entire 1,109-bp sequence from the hamster and the corresponding sequence from L. intracellularis (data not shown). The sequences of the cloned 16s rDNAs from each species have been submitted to GenBank (see above for accession numbers). The sequences were 100% similar to the sequence of the ferret intracellular organism over this region. Our hamster sequence had 98.4% similarity over the entire 16s rDNA with another sequence (GenBank accession no. U06423) (26), of the same strain (auratus) but only 97.7% similarity with the partial sequences we determined for the deer and ostrich isolates. Another isolate of the intracellular agent of hamster PE (GenBank accession no. U07569) (6) had 98.8% 16s rDNA sequence similarity with our hamster isolate over a partial sequence that did not include the hypervariable region. L. intracellularis had 91% 16s rDNA sequence
FIG. 1. Intestinal crypts from a hamster (A), a deer (B), and an ostrich (C) with PE stained with Warthin-Starry silver stain. Numerous elongate, curved organisms are present within the apical cytoplasm of the crypt enterocytes (arrows). Magnification, ×660.

FIG. 2. DNA sequences of a portion of the 16S rDNA of the intracellular agents of PE derived from the hamster (strain auratus), deer (strain 93-3669), and ostrich (strain D94-06196) compared to the consensus sequences of a pig isolate of L. intracellularis (NCTC 12656; GenBank accession no. U30147) and of D. desulfuricans (GenBank accession no. M34113), the closest relative of L. intracellularis. Underlined regions correspond to 878F, the forward primer for the PCR amplification of the intracellular organism 16S rDNA from lesions of PE to 1100R, the reverse primer, and to a hypervariable region that distinguishes L. intracellularis from D. desulfuricans by nine bases. Dashes indicate gaps introduced to facilitate comparisons.

similarity with a Desulfovibrio isolate from a human liver abscess.

**DISCUSSION**

The results of this study show that the 16S rDNA sequences of the intracellular agents of PE from the pig, hamster, deer, and ostrich have high similarity, suggesting that they are all isolates of L. intracellularis. rDNA sequence analysis is a powerful technique that is used to identify the taxonomic position of prokaryotes (35). Partial 16S rDNA analysis has been used to determine the relationships between species of Desulfovibrio (3), the genus most closely related to L. intracellularis (7, 21). D. desulfuricans and L. intracellularis have 91% 16S rDNA sequence similarity (7). The region of the 16S rDNA of the intracellular agents of PE from the hamster, deer, and ostrich that was sequenced includes a hypervariable region that differentiates L. intracellularis from D. desulfuricans, its closest relative (Fig. 2) (7). There is 96.1% similarity between L. intracellularis and D. desulfuricans across this region. There was 100% sequence similarity across this hypervariable region (and for the entire partial 16S rDNA sequence we determined) between L. intracellularis and the intracellular agents of PE in the hamster, deer, and ostrich.

These results closely parallel those of other investigators who have shown that the intracellular agents of proliferative colitis in the ferret (6) and of PE in the hamster (6, 26) are closely related to L. intracellularis, based on 16S rDNA sequence similarity. We obtained a slightly different sequence for the same strain (auratus, isolated from a hamster) from that of Peace et al. (26), with only 98.4% similarity overall and 97.7% over the partial sequence we determined for the deer and
The organisms in the genus Lawsonia are clearly different from Desulfovibrio ATCC 27774 (the type strain) (7) and from a Desulfovibrio isolate from a human liver abscess (32), based on only 91% 16S rDNA sequence similarity. In addition to the differences in 16S rDNA sequences, these organisms differ from the desulfovibrios in several important areas. They are obligate intracellular parasites (7, 15, 21, 31), whereas the desulfovibrios can be grown on cell-free media (3, 32). In addition, the presence of the intracellular organism is always associated with lesions, and the pathogenesis of the disease (4–7, 17, 23, 24, 27, 31) is clearly distinct from the diseases that have been associated with the normally nonpathogenic desulfovibrios (32).

We have previously discussed the amplification of L. intracellularis DNA from formalin-fixed, paraffin-embedded tissues by PCR (2). It was necessary to use prolonged digestion with proteinase K and subsequent extraction with diatomaceous earth in the presence of GuSCN to purify DNA for successful amplification from these samples. While the reason for this is not known, we hypothesize that either there was very little target DNA, the target DNA was extensively cross-linked by formalin fixation and inaccessible to the DNA polymerase, the bacterial cell wall was resistant to degradation, or there were PCR inhibitors present in the fixation preparation. Preparation of human or viral nucleic acids from archival tissues for PCR amplification does not require prolonged digestion (11, 29), while preparation of bacterial DNA may (25).

This study demonstrates that the causative agent of PE in pigs, L. intracellularis, is closely related or identical to the intracellular agents of PE in the hamster, deer, and ostrich. PE is emerging as a worldwide disease that occurs in many species of animals. The causative agent, L. intracellularis, appears to be a versatile pathogen with a wide host range. An understanding of the relationships between different isolates of this organism will greatly facilitate the study of the epidemiology of the disease.