The Pathology of Spontaneous Paratuberculosis in the North American Bison (Bison bison)

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Abstract. Gross and histopathologic examinations were performed on 70 North American bison (*Bison* bison) from a *Mycobacterium avium paratuberculosis* culture–positive herd. The bison examined were part of a breeding herd totaling 2,800 animals. Eight of 70 (11%) animals had gross findings of intestinal mucosal thickening, and 16 of 70 (23%) of the animals had enlarged mesenteric lymph nodes. Histologic lesions compatible with Johne's disease were diagnosed in 30 of 70 (43%) bison on the basis of the demonstration of noncaseating granulomatous inflammatory infiltrates and of one or more acid-fast bacilli characteristic of *Mycobacterium avium paratuberculosis*. A suspicious diagnosis of Johne's disease was obtained in 11 of 70 (16%) bison on the basis of the observation of noncaseating granulomatous inflammatory infiltrates granulomatous inflammatory infiltrates without demonstrable acid-fast bacteria. Twenty-nine of 70 (41%) animals were assessed as histologically paratuberculosis free. Histologic results were compared to Johne's disease tests such as culture, serology, and polymerase chain reaction, which were performed on some of the cohort animals.

Key words: Gross and microscopic pathology; Johne's disease; North American bison.

Paratuberculosis (Johne's disease), a chronic infectious disease of ruminants, has been described throughout the world since its discovery in 1895. The causative agent is Mycobacterium avium paratuberculosis, a facultative intracellular acid-fast bacillus.^{6,26} Paratuberculosis has been recognized in captive ruminants (artiodactylids) and free-range ruminants. Paratuberculosis has been reported as a spontaneous disease in a variety of nondomestic free-range ruminants,15 including moose,24 bighorn sheep, Rocky Mountain goats,²⁹ white-tailed deer,⁵ axis deer,²¹ Florida key deer,20 and tule elk.9 Paratuberculosis has also been documented in wild rabbits in Scotland.¹³ Other reports have dealt with outbreaks of paratuberculosis in farmed red deer around the world.10,11,16,23 Thus, a variety of free-range animal species that carry the bacilli may have a potential to spill over to domestic ruminants sharing the same pasture and habitat.

To our knowledge, paratuberculosis has been reported only recently in the North American bison (*Bison bison*) as a herd problem.²⁸ We describe the macroscopic and microscopic findings of paratuberculosis in 41 of 70 sacrificed bison from a ranch in the northern United States. The bison are from a herd proven

culture positive for *Mycobacterium avium paratuber*culosis.

Materials and Methods

Approximately 2,800 bison are grazed on 100,000 acres of a ranch in the foothills of the northern United States. The bison were originally acquired from various sources and were introduced to the ranch in 1989. The herd started with 100 bulls and 577 cows. Cattle occupied the ranch before the introduction of the first bison. Over the past 5 years, individual bison were noticed losing weight and failing to shed winter hair. The poor body condition was attributed to inadequate nutrition, endoparasites, and stress from herd social behavior. Management made adjustments to overcome the condition deficits, but these were to no avail. In July 1997, a thin adult female bison was subjected to a field necropsy and showed gross findings of mucosal corrugation in the distal small intestine and of enlarged mesenteric lymph nodes. These findings suggested Johne's disease. Histopathologic examination yielded a tentative diagnosis of Johne's disease. The bison tested negative on fecal culture for Mycobacterium avium paratuberculosis. Mycobacterial culture verification of herd disease was not obtained until bison 8 of the study cohort tested positive.

For the histologic Johne's disease assessment, multiple tissues were examined independently by pathologists from five different institutions. Each pathologist examined tissues of only a subset of the animals under study.

Tissues from four animals that died in the field were randomly selected from a field necropsy, including the index case mentioned above (pilot animals 1-4, Group A), and were submitted to the Montana State Diagnostic Laboratory for a microscopic examination. A second group of tissues from 26 sick or dead bison (animals 1-26, Group B) were subjected to a complete field necropsy and were collected for various analyses, including microscopic examination. Tissue sections for microscopic examination included three sites in the jejunum, one site in the ileum, one site of jejunal and ileocecal lymph nodes, cecum, colon, heart, lung, spleen, kidney, liver, and abomasum. Alimentary tract tissues from Group B were microscopically examined by pathologists at the Montana State Diagnostic Laboratory, Cornell University, and the University of Wisconsin. A third group of 25 bison (animals 27-51, Group C) was sent to slaughter because of poor body condition and for Johne's disease surveillance. For these animals, a protocol was established to selectively take specimens from the lower jejunum, midjejunum, upper jejunum, associated mesenteric lymph nodes, ileum, ileocecal lymph node, cecum, colon, and other organs, as deemed necessary. Specimens from the Group C animals were microscopically assessed at the University of Florida and the University of Pennsylvania. Last, a subset of 15 bison (animals 52-66, Group D) was culled for reproductive failure, poor conformation, bad temperament, or Johne's disease surveillance. Ileum, mesenteric lymph nodes, and ileocecal lymph nodes were collected at slaughter. These tissues were microscopically examined at the University of Florida.

Tissues collected for histopathologic examination were fixed in 10% neutral buffered formalin, embedded in paraffin, and sectioned at 6 μ m. All sections were stained with hematoxylin and eosin, and selected sections were stained with an acid-fast Ziehl-Neelsen or a Brown-Brenn Gram stain.

Histomorphologic criteria were used to categorize each animal as positive, suspicious, or negative for paratuberculosis. The criteria for a definitive histomorphologic diagnosis of paratuberculosis required the presence of noncaseating granulomatous inflammation composed of Langhans' type giant cells and epithelioid macrophages and the identification of at least one intracellular small acid-fast rod compatible with *Mycobacterium avium paratuberculosis*. A case suspicious for paratuberculosis was categorized by the presence of epithelioid macrophages and especially of multinucleate giant cells of Langhans' type in either the intestinal mucosa or mesenteric lymph nodes, without demonstrable acid-fast bacilli. A case was considered negative for paratuberculosis on the basis of the absence of specific granulomatous cell infiltrates and stainable acid-fast bacilli.

The severity of the inflammation and the semiquantification of identifiable acid-fast bacilli were summarized by one investigator on the basis of the information rendered by other pathologists participating in the project. Lesions defined as noncaseating granulomatous inflammation were scored semiquantitatively by severity using a mild (1), moderate (2), and severe (3) scale of assessment. The category of acid-fast bacilli was semiquantitatively scored as paucibacillary or multibacillary. The bacterial score was subjectively compared with the inflammatory score.

The methodology of the cultural, serologic, and polymerase chain reaction (PCR) tests has been previously described.²⁸ Briefly, fecal and tissue cultures were cultured on Herrold's Egg Yolk, PCR was conducted by using primers of the IS900 sequence, and serum was tested on agar gel immunodiffusion (AGID) and with the CSL/IDEXX enzyme-linked immunosorbent assay (ELISA) Johne's kit.

Results

The cohort of animals, the necropsy and microscopic findings regarding paratuberculosis, the bacillary status based on the number of acid-fast bacilli stained, the host inflammatory response to the assumed presence of My-cobacterium avium paratuberculosis, and the results of antemortem tests are listed in Tables 1–3.

Most (68 in 70; 97%) of the animals examined were female. Only two males were included in the collection. The mean age of all bison sacrificed was 7.9 years (range 4-20 years). The mean age of the 30 bison considered positive for paratuberculosis was 8.2 years (range 4-20 years).

Of the examined bison, the gross changes consistent with paratuberculosis included weight loss due to skeletal muscle mass reduction, atrophy of body fat, or both, in 28 of 70 (40%) animals; thickening of the mucosa of the small intestine in 8 of 70 (11%) animals; and prominent mesenteric or ileocecal lymph nodes in 16 of 70 (23%) animals. One animal had distended serosal lymphatics.

In four of the bison with gross findings suggestive of paratuberculosis, three bison had intestinal ulcers covered by a laminated caseating exudate in the mucosa of the small intestine. The fourth bison had caseating necrosis in mesenteric lymph nodes. Culture of the intestinal and lymph glandular exudates resulted in the isolation of Rhodococcus equi. Similar gross foci of caseating necrosis were found in the lung, tracheobronchial lymph nodes, and liver of the same bison. Other gross findings included endoparasitism due to lungworms, gastrointestinal nematodes, or both in 10 animals; liver abscesses in eight bison in addition to the above mentioned three animals; other liver disorders in two animals; rumenal and pleural adhesions in three animals; lung abscesses in an additional three animals; retropharyngeal lymph node abscesses due to Arcanobacterium pyogenes (Actinomyces pyogenes) in three animals; and various genitourinary changes in five animals.

Thirty of 70 (43%) bison were considered positive for Johne's disease on the basis of established histomorphologic criteria. A suspicious diagnosis was obtained in 11 of 70 (16%) bison. Twenty-nine of 70

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								Serol	ogy			
	Age		Gross	Granulo- matous	Bacillarv	Cult	ture	Agar Gel Immuno-	Enzyme- Linked Immuno- sorbent	Polymera Reac	e Chain ion	
Bison No.	(years)	Sex	Findings†	Infiltration [‡]	Category§	Feces	Tissue	diffusion	Assay	Feces	Tissue	Other Observations
Group A 1	∞	Ц	4	2	Mb	I	ND	I	I	ND	ND	Large intestine involved;
0	6<	Ц	NR	0	Mb	NR	I	I	I	ND	+	endoparasites
Group B												
1	20	Ц	4	7	Mb	+	+	Ι	S	+	+	Endoparasites
3	9	Ц	4	2	Pb	+	+	Ι	S	+	+	Ruminal adhesions
4	9	Ц	4	1	Pb	Ι	I	I	+	+	+	Liver abscess
5	5	Ц	NS	1	Pb	Ι	I	I	S	+	+	Liver abscess; endoparasites
9	4	Ц	NS	0	Pb	Ι	I	I	I	+	+	
7	8	Μ	3, 4	б	Pb	Ι	I	I	I	I	+	
8	9	Ц	1, 4	б	Mb	+	+	I	+	+	+	
6	9	Ц	3, 4	m	Mb	I	+	Ι	S	+	+	Rhodococcus equi
11	14	Ц	NR	m	Mb	I	+	Ι	+	+	+	ĸ
12	6	Ц	NR	б	Mb	+	+	Ι	+	+	+	
13	6	Ц	4	m	Mb	I	I	Ι	+	+	+	
14	6	Ц	NR	m	Mb	I	+	Ι	+	+	+	
15	6	Ц	NR	1	Mb	+	+	Ι	+	+	+	
17	13	Ц	1, 4	m	Mb	+	+	Ι	+	+	+	Rhodococcus equi
19	6	Ц	1, 3, 4	7	Mb		I	I	S	+	+	Liver abscess
21	9	Ц	4	7	Mb	+	+	I	+	I	+	Liver abscess; endoparasites
22	×	Ц	3, 4	б	Mb	+	+	I	+	+	+	Endoparasites
23	7	Ц	4	1	 +	Ι	+	Ι	+	Ι	+	Liver abscess; endoparasites;
č	c	Ľ	Ţ	-	_				۵		-	pleural adhesions
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		-		1					NTK T			
Group C												
30	UK	Ц	1, 3, 4	7	Pb	NR	NR	Ι	+	+	+	Arcanobacterium pyogenes
36	6	Ц	1	S	Mb	NR	NR	Ι	+	+	+	
38	6	Ц	1, 3	б	Mb	NR	NR	+	+	+	+	Fatty liver
39	×	Ц	1	0	Pb	NR	NR	I	+	+	+	
43	L	Ц	m	б	Pb	NR	NR	I	+	+	+	
47	UK	Ц	NS	0	Mb	NR	NR	I	+	I	+	
48	6	Ц	4	0	Pb	NR	NR	I	+	+	+	Liver abscess
49	UK	Ц	2, 4	С	Mb	NR	NR	Ι	+	+	+	Large intestine involved
* - = ne	gative resul	ts; + = pc	ositive results;	+- = weak	results; $S = s$	uspicious; N	$\sqrt{R} = not rec$	corded; NS =	none seen; U	K = unknov	vn; ND =	= not done.
+1 = mu	icosal corruș	$\frac{1}{2}$ = 2 =	= prominent ly	/mphatics; 3 =	 lymphadeno 	pathy; 4 =	weight loss.					
10 = 0	ne; $I = mult$	d; $2 = m0$	derate; $\beta = se$	vere.								
$s ro = p_{s}$	aucibacillary	v; Mb = n	nultibacillary.									

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								Sero	logy			
	A oe		sour	Granulo- matous	Bacillary	Cult	ture	Agar Gel Immino-	Enzyme- Linked Immuno- sorbent	Polymera Reac	se Chain tion	
Bison No.	(years)	Sex	Findings†	Infiltration [‡]	Category	Feces	Tissue	diffusion	Assay	Feces	Tissue	Other Observations
Group B												
7	9	Ц	4	1	I	+	I	I	S	+	+	Endoparasites
16	6	Ч	4	2	Ι	Ι	Ι	Ι	+	+	+	Endoparasites
20	6	Ц	1, 4	б	I	I	I	I	S	+	+	Liver abscess;
												Rhodococcus equi
25	9	Ч	NR	1	Ι	ND	ŊŊ	I	NR	ND	ND	
Group C												
29	5	Н	3, 4	1	Ι	NR	NR	Ι	+	Ι	+	Lung abscess
31	5	Ц	1	2	I	NR	NR	Ι	I	Ι	Ι	
32	7	Ц	NS	1		NR	NR	Ι	+	+	Ι	
34	UK	Ч	б	1	I	NR	NR	Ι	S	+	+	Firm liver
40	6	Ч	NS	2	I	NR	NR	Ι	NR	+	+	
41	5	Ч	NS	1	I	NR	NR	I	+	+	+	
Group D												
61	5	Ц	NR	1	I	NR	NR	Ι	ND	ND	ND	
$\begin{array}{c} * - = ne \\ \dagger 1 = mu \\ \ddagger 0 = nor \end{array}$	gative results; cosal corrugati ne; 1 = mild; 2	+ = positive ion; 2 = prodera	ve results; + - ominent lymp ite; 3 = sever	 – = weak resublation <	lts; S = susp nphadenopatl	icious; NR = ıy; 4 = weig	not recorded ht loss.	l; NS = none	seen; UK = u	ıknown; ND	= not done	-

 Table 2.
 Signalment and summary of test results in paratuberculosis-suspicious bison.*

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Table 3.

								Serol	logy			
	Age		Gross	Granulo- matous	Bacillarv	Cult	ture	Agar Gel Immino-	Enzyme- Linked Immuno- sorbent	Polymera Reac	se Chain tion	
Bison No.	(years)	Sex	Findings†	Infiltration ‡	Category	Feces	Tissues	diffusion	Assay	Feces	Tissues	Other Observations
Group A												
ω ₹	4	Σι	4	0	I	Q Z	QN A	I	I	Q	Q Z	Endoparasites
4	UK	Ţ	N	0	I	UN	ND	I	I	ΠŊ	ΠN	Pleuritis; salmonellosis
Group B												
10	4	Ц	4	0		Ι	I	I	S	I	+	Pleuropneumonia
18	6	Ц	3, 4	0	I	Ι	I	I	S	+	 +	Endoparasites
Group C												
27	5	Ц	3, 4	0	Ι	NR	NR	Ι	+	Ι	+	
28	14	Ц	3, 4	0	I	NR	NR	Ι	+	Ι	+	Arcanobacterium pyogenes
33	5	Ц	ю	0		NR	NR	I	S	Ι	+	Abscesses; Rhodococcus equi
35	6	Ц	б	0	I	NR	NR	I	+	 +	+	Nephritis; Arcanobacterium
												pyogenes
37	UK	Ľ	NS	0	I	NR	NR	Ι	+	 +	 +	Glomerulonephritis; liver and
ć	Г	þ	¢	C		CIN	CIN		4		-	Photoscesses
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45	L	Ľ,	ŝ.	0	I	NK	NK	I	N	I	+	
46	6	Ľ	4	0	I	NR	NR	I	+	 +	 +	Nephritis; leptospirosis
50	UK	Ц I	NR	0	I	NR	NR	I	N i	+	+	
51	UK	Ц	NS	0	I	NR	NR	I	S	 +	+	
Group D												
52	UK	Ц	NR	0	I	NR	NR	Ι	ŊŊ	ND	ND	
53	5	Ц	NR	0	I	NR	NR	I	ND	ND	ŊŊ	
54	ŝ	Ц	NR	0	I	NR	NR	I	ND	ND	ND	
55	7	Ц	NR	0	I	NR	NR	I	ND	ND	ND	
56	6	Ц	NR	0	I	NR	NR	I	ND	QN	QN	
57	×	Ц	NR	0	I	NR	NR	I	ND	ŊŊ	QN	Ovarian teratoma
58	5	Ц	NR	0	I	NR	NR	I	ND	QN	QN	
59	9	Ц	NR	0	I	NR	NR	I	ND	QN	QN	
60	5	Ц	NR	0	Ι	NR	NR	Ι	ND	QN	QN	
62	7	Ц	NR	0	I	NR	NR	I	ND	ND	ND	
63	18	Ц	4	0	I	NR	NR	Ι	ND	QN	ND	
64	9	Ц	NR	0	Ι	NR	NR	I	ND	ŊŊ	ND	
65	12	Ц	NR	0	Ι	NR	NR	I	ND	ŊŊ	ND	Fetal mummification
66	6	Ц	NR	0	Ι	NR	NR	Ι	ŊŊ	ND	ŊŊ	
* - = ne + 1 = mu	icosal corrug	fs; + = pc sation; 2 =	esitive results; = prominent ly	+- = weak 1 /mphatics; $3 =$	results; S = s Iymphadeno	suspicious; l pathy; 4 =	NR = not rec weight loss.	corded; NS =	none seen; U	K = unknov	vn; ND =	not done.
1 = 10	ne; $I = mil($	$1; 2 = m_0$	derate; $3 = s_t$	evere.								

(41%) animals were assessed as histologically paratuberculosis free. The majority of the paratuberculosispositive animals were from Group B; no animal was diagnosed as positive from Group D. Granulomatous infiltrations in the positive and suspicious paratuberculosis categories were graded as mild (1) in 12 of 70 (17%) bison, moderate (2) in 15 of 70 (21%) bison, and severe (3) in 14 of 70 (20%) bison. When there was a discrepancy of scoring between tissues from the same animal, the highest grade obtained from one tissue was applied to the final score.

Twelve (17%) of the 70 bison had mild cellular changes suggestive of paratuberculosis, but only two of these clearly had acid-fast bacilli indicative of *Mycobacterium avium paratuberculosis*. Individual Langhans' type giant cells were best detected within the paracortex of mesenteric lymph nodes; only on occasion were inflammatory giant cells present in the lamina propria of the intestinal mucosa (Fig. 1). Specific epithelioid macrophages infiltrating the intestinal lamina propria were difficult to identify from intestinal resident macrophages in this group. Because of the paucity of cellular infiltrates, the ratio of epithelioid cells to Langhans' type giant cells was difficult to establish in this category of animals: subjectively, Langhans' type cells were prominent.

Bison with moderate histologic lesions (15 of 70, 21%) had several small groups of epithelioid macrophages, often with several Langhans' type giant cells that were limited to the lamina propria of intestinal villi or within the paracortical zone of regional mesenteric lymph nodes. In eight of these bison, epithelioid macrophages were more numerous than giant cells; in four bison, the proportion of macrophages to giant cells favored giant cells; and in three other bison, only epithelioid macrophages were noted (Fig. 2). When specific sites of the lymph node selection were known (Group C), moderate granulomatous inflammation was present in the midjejunal, upper jejunal, and ileocecal lymph nodes, with the midjejunal lymph nodes showing characteristic changes in most of the cases.

Many macrophages and lesser numbers of Langhans' type giant cells had infiltrated the lamina propria and submucosa of various segments of the small intestine in bison with severe lesions (14 of 70; 20%) of paratuberculosis (Fig. 3). Some of these inflammatory cells had invaded the tunica muscularis and tunica serosa as well. Granulomatous cell infiltrates cuffed intramural intestinal lymphatics. Intestinal villi were short and distorted. Villous lacteals were prominent because of dilatation. The granulomatous inflammatory response extended to the adjacent mesenteric lymph nodes. The lymphoid cells within the paracortical zone were largely replaced by epithelioid macrophages and giant cells. Lymphoid follicles were spared. In two animals (I and 49), the cecum and spiral colon also had granulomatous inflammation. Seven bison in this group had inflammation composed exclusively of epithelioid macrophages, whereas one bison had predominantly giant cells present. Six other bison had a mixed granulomatous inflammation with primarily epithelioid macrophages.

All bison with paratuberculous granulomatous inflammation had uniformly formed, well-differentiated epithelioid macrophages with dense eosinophilic cytoplasm, distinct cell boundaries, eccentric oval hyperchromatic nuclei, and several nucleoli of various sizes.

In the animal cells that stained positive with Ziehl-Neelsen stain, nine had a few (usually 1–3) acid-fast–positive bacilli within giant cells. This group of animals would fit the category of paucibacillary infection. Nineteen bison had a plethora of acid-fast bacilli, mainly within macrophages and typically arranged in clusters (Fig. 4). These animals would fit the category of multibacillary infection.

In four animals (9, 17, 20, and 42), the histologic changes of paratuberculosis were accompanied by necrotizing pyogranulomatous changes due to *Rhodococcus equi* infection. These changes were best seen in mesenteric lymph nodes, but they were also present in the intestine, liver, lung, and pulmonary lymph nodes. Microscopically, the necrotic cores had a laminated, hypereosinophilic appearance; epithelioid macrophages surrounded the necrotic cores and contained a multitude of small gram-positive bacterial coccoid rods in the cytoplasm.

The antemortem tests showed one (3%) of the 30 bison morphologically positive on AGID; 19 (63%) bison were positive on ELISA. With the fecal culture system, 8 of 20 morphologically positive bison were identified as shedders, whereas in the same group, 12 animals had a positive tissue culture. Fecal PCR identified 22 of 30 (73%) correctly but also had eight additional bison as positive; tissue PCR identified 100% of the morphologically positive animals but also 15 of the suspicious and negative bison.

Discussion

The distribution and character of pathologic changes of spontaneous bison paratuberculosis were similar to the lesions of bovine paratuberculosis.^{3,27} There was protracted weight loss leading to a state of emaciation. Gross intestinal lesions were most easily seen in the distal segments of the small intestine and there was gross evidence of enlargement of mesenteric lymph nodes of up to 3–4 times the normal size. Histologically, advanced cases of paratuberculosis had diffuse,



Fig. 1. Upper jejunum; bison 40. Individual Langhans' type giant cell adjacent to mucosal crypt is embedded among resident lymphocytes of the lamina propria. HE. Bar = $25 \mu m$.

Fig. 2. Mesenteric lymph node; bison 41. Moderate Johne's disease. Cluster of epithelioid macrophages present within the paracortex of the mesenteric lymph node. HE. Bar = $50 \mu m$.

Fig. 3. Small intestine; bison 50. Advanced Johne's disease. Epithelioid macrophages and individual Langhans' type giant cells have infiltrated the lamina propria and have replaced mucosal crypts. HE. Bar = $50 \mu m$.

Fig. 4. Small intestine; bison 50. Clusters of *Mycobacterium avium paratuberculosis* bacilli are present within epithelioid macrophages. Ziehl-Neelsen. Bar = $50 \mu m$.

severe, noncaseating granulomatous inflammation of the distal small intestine and associated mesenteric lymph nodes. Histologic changes of subclinical paratuberculosis were rather mild.

The granulomatous inflammation was noncaseating and composed of epithelioid macrophages with varying numbers of intermingled multinucleate Langhans' type giant cells. Animals with paucibacillary and multibacillary paratuberculosis were identified, the latter accompanied by severe granulomatous inflammation. Other animals had no visible acid-fast bacilli but had mild changes with individual giant cells, small clusters of epithelioid macrophages, or both, best seen in mesenteric lymph nodes.

The absence of demonstrable acid-fast bacilli in the inflammatory giant cells and epithelioid macrophages has prompted the creation of a morphologic category of paratuberculosis-suspicious animals if epithelioid macrophages, but, more typically, giant cells, are detected in target tissues despite the animal's exhibiting clinical signs suggestive of Johne's disease and its origin from a herd of established paratuberculosis-positive status. By applying this criterion to our case material, 11 bison would have been classified as suspicious.

The Langhans' type giant cells may be the first to form within the lamina propria or, together with small clusters of epithelioid macrophages, within the paracortex of the mesenteric lymph nodes in the cases of mild multifocal lesions. Langhans's type giant cells are thought to arise by fusion of mature macrophages.¹

Epithelioid cells have been described as the most differentiated and the end stage of stimulated phagocytic macrophages. They are derived from circulating monocytes of bone marrow origin.¹ Macrophages turn into epithelioid cells when they become immobilized at the site of inflammation. Epithelioid cells are active in the phagocytosis and degradation of ingested bacilli. They have a secretory function, which may partially explain the calcification of cardiac endocardium and the aorta in some bovine animals infected with *My*-*cobacterium avium paratuberculosis*.² Such calcification of bison.

The epithelioid macrophages of mild paratuberculosis should not be confused with enlarged resident macrophages of the intestinal tract and associated lymph nodes that contain yellow granules or crystalline material in their cytoplasm.¹⁸ The latter are readily distinguished from epithelioid cells and giant cells because of the pigment and because of the observation that they have not been demonstrated to contain acidfast organisms. The cytoplasm of these resident macrophages constantly stained deep blue with acid-fast stain, suggesting metabolic activity.¹⁸

It is probably the combined capability of activated epithelioid macrophages and giant cells that destroys phagocytized mycobacterial microbes in the early phases of infection with *Mycobacterium avium paratuberculosis* such that they become nonstainable with acid-fast stains. Alternatively, the bacilli may transform to a status of spheroplasts or become completely degraded. Thus, the diagnosis of paratuberculosis will not be confirmed in many of the mild cases, and instead such infected animals will be categorized as suspicious. To overcome such a diagnostic situation, immunohistochemical staining by specific polyclonal or monoclonal antibodies has been suggested.¹⁹ The immunohistochemical techniques are not only of value for the detection of intact viable organisms, but also for soluble *Mycobacterium avium paratuberculosis* antigens, cell fragments, and altered microorganisms with defective cell walls.¹⁹ The technique has the potential to be more sensitive and will thus help to reduce the number of animals in the suspicious category and to increase the number of animals in the positive category. The immunostaining technique was not employed in this study because of a lack of specific reagents for bison.

In four animals, the demonstration of gram-positive intracellular small coccoid rods suggested the etiologic diagnosis of *Rhodococcus equi* infection. The diagnosis of alimentary tract rhodococcal infection was confirmed by cultural isolation of the organisms.

Rhodococcus equi are soil-associated bacteria that are usually isolated from pneumonic lungs of foals, with half of all foals developing ulcerative enterocolitis, mesenteric lymphadenitis, or both, along with the pneumonia.²⁵ In swine and cattle, cervical lymphadenitis due to Rhodococcus equi may mimic tuberculosis.¹⁴ Of interest was the observation in three bison of retropharyngeal abscesses thought to be the result of infection with Arcanobacterium pyogenes (Actinomyces pyogenes). Indeed, these could also have been due to Rhodococcus equi, as cultural isolation was not attempted from these tissues. There are reports in the literature of the isolation of Rhodococcus equi from liver abscesses in goats;¹² some of the liver abscesses in our bison might have been also due to Rhodococcus equi. Stress or infection with Mycobacterium paratuberculosis might have predisposed the bison to the development of rhodococcal infection. It has been reported that infection with Mycobacterium avium paratuberculosis may predispose cattle to salmonellosis, bovine viral diarrhea, and bacterial mastitis.²⁷ Predisposition to infection with Rhodococcus equi may be more frequent than is presently recognized in paratuberculosis-positive ruminants. Routine aerobic bacterial isolation tests should be performed on all ruminants that exhibit laminated abscesses in internal organs.

Infection with lungworms (*Dictyocaulus* sp.) and gastrointestinal nematodes (*Trichostrongylus* sp., and *Ostertagia* sp., and *Oesophagostomum* sp.) in 10 bison from Groups A and B is of concern because of worming frequency. Infection with *Salmonella* was not a

problem, as only one animal had a positive fecal isolation. Most of the animals were serologically negative for brucellosis, bluetongue, anaplasmosis, infectious bovine rhinotracheitis, and leptospirosis. One bison had nephritic leptospirosis.

The bison with confirmed paratuberculosis originated from other paratuberculosis-positive bison herds, so it is likely that the disease was introduced to the ranch as a preexisting, subclinical condition at the time of purchase of the original bison rather than from cattle previously inhabiting and grazing the ranch or from free-range wildlife. At least six different bison herds supplied original animals to the ranch. A strain fingerprinting by restriction-length fragment polymorphisms (RLFP) analysis resulted in the identification of B-C1,CC-1, and CC-18 strain types in the bison herd as evidence of multiple strains having infected the herd.^{17,28} These strains have also been identified in cattle.¹⁷ The fingerprinting method has been proposed to identify the identity of Mycobacterium avium paratuberculosis strains isolated from different world-wide geographic locations and from different ruminant animal hosts.¹⁷ The possibility of cross infection from free-range wildlife, especially elk, was disregarded, as no elk from the same region were reported to be positive for Johne's disease.22 Historical information relative to the possible avenue of transmission from cattle to bison was not available.

A restraining aspect for Johne's disease control of the herd was the dilemma associated with the diagnostic deficiency of the tests applied to the bison.²⁸ The 70 bison of the different subsets were subjected to one or several different Johne's disease tests (fecal and tissue culture methods, AGID, ELISA, tissue, and fecal PCR) commonly used for cattle.

The AGID test identified only one (3%) of the 30 mophologically positive bison, whereas the ELISA test, after some modification, identified 19 (63%) of the morphologically positive bison. Only 25 bison were subjected to fecal culture, and 8 of the 20 (40%)of the morphologically positive bison were correctly identified. One bison morphologically classified as suspicious had a positive fecal culture. The same 25 bison were also subjected to tissue culture, and 12 of 20 (60%) of them were correctly identified. Finally, 49 bison were subjected to a PCR test on feces, and 50 were subjected to a tissue PCR test. The fecal PCR identified 22 of 30 (73%) of the morphologically positive bison correctly, but it also identified eight additional bison as positive when the other tests, including the histologic tests, were negative in these animals. The tissue PCR identified 100% of the positive bison, but it also identified 15 of the suspicious and negative bison as positive.

From the comparison of these results, we concluded that the incidence of paratuberculosis infection in bison was higher when using PCR testing than histopathology or other tests. Some of the tests correlated with each other, whereas others did not. In some circumstances, acid-fast bacteria were microscopically detected in the tissue of bison, but the organisms could not be cultured from feces or tissues, and specific antibodies could not be detected from serum samples. This suggests that tests that work for cattle need to be modified to make them work reliably for bison. The discrepancy between tests prompted the overall conclusion that the results of attempts to diagnose paratuberculosis in bison with the test methods usually applied to cattle do not follow the pattern of sensitivity and specificity expected from these tests in cattle.

Prevention of paratuberculosis infection and the establishment of paratuberculosis-free populations is a management goal. This goal is extremely important when livestock mix with free-range wildlife or when one species of free-range wildlife mixes with another.15 There are presently no treatment modalities or effective vaccines available for paratuberculosis control. Control of infection and disease is recommended through test-and-cull procedures combined with management and hygiene adjustments.7 Vigorous removal of animals with poor body condition should be practiced with routine monitoring for Johne's disease by gross examination and histopathology at the time of culling. Of the 28 bison with poor body condition, 17 (61%) had a final positive diagnosis of paratuberculosis after microscopic examination. The observation of failing body condition created a simple, inexpensive, presumptive first antemortem test for paratuberculosis infection on the ranch once the presence of Johne's disease was originally established. Animals identified in such a way were then subjected to various Johne's disease tests.

Finally, it is very important to establish and adhere to morphologic criteria for the confirmatory diagnosis of paratuberculosis, as histologic evaluation, next to fecal or tissue culture, is presently considered the gold standard for paratuberculosis diagnosis.²⁷ Comparison among the various institutions involved in the histologic assessment of tissue material obtained from the bison originally showed disparity for the diagnosis of paratuberculosis status in a few animals among pathologists because of different morphologic definitions used. The disparity was resolved, and reclassification was done by adhering to a uniform definition of a conPathologists should not be divided as to morphologic criteria used for a positive diagnosis of paratuberculosis. Only in some countries, such as Australia, have standard recommendations for histopathologic assessment and reporting of results been implemented as a quality assurance program.^{4,8} Specific recommendations have been drafted, including how many tissues, which specific tissues, how many sections, and for what time duration these sections should be examined under the microscope.

There should be global uniformity of interpreting and reporting pathologic findings in paratuberculosis. This uniformity will ultimately benefit the efficacy of herd surveillance programs for Johne's disease control and eradication.

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