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- 1 A case of transplacental transmission of *Theileria equi* in a foal in Trinidad
- 2

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17 Abstract

18	Equine piroplasmosis due to T. equi and B. caballi is endemic in Trinidad. A case
19	of equine piroplasmosis due to T. equi was diagnosed in a Thoroughbred foal at 10 hrs
20	post-partum. A high parasitaemia (63 %) of piroplasms was observed in a Wright-
21	Giemsa® stained thin blood smear from the foal. In addition, the 18S rRNA gene for
22	Babesia/Theileria was amplified from DNA extracted from the blood of the foal and the
23	mare. Amplified products were subjected to a reverse line blot hybridization assay
24	(RLB), which confirmed the presence of <i>T. equi</i> DNA in the foal. The mare was negative
25	by RLB but was positive for <i>T. equi</i> using a nested PCR and sequence analysis.
26	In areas where equine piroplasmosis is endemic, severe jaundice in a post-partum
27	foal may be easily misdiagnosed as neonatal isoerythrolysis. Foals with post-partum
28	jaundice should be screened for equine piroplasmosis, which may be confirmed using
29	molecular methods if available.
30	Key words: Neonatal foals, equine piroplasmosis, reverse line blot, Trinidad

31 Introduction

32	Equine piroplasmosis is a tick-transmitted haemoparasitic disease of equids
33	caused by the protozoa Theileria equi or Babesia caballi. Ixodid tick vectors of the
34	genera Rhipicephalus, Hyalomma and Dermacentor are known to transmit the disease (de
35	Waal, 1992). Both parasites are distributed world wide, however several countries restrict
36	the importation of horses that are serologically positive for the disease (Magnarelli et al.,
37	2000). The clinical signs of equine piroplasmosis are often variable and non-specific.
38	Acute infection results in fever (40°C), depression, reduced appetite, pallor, icterus,
39	dyspnoea, petechiation, sweating, colic, eyelid and distal limb oedema and incoordination
40	(Zobba et al., 2008). Massive destruction of erythrocytes results in haemoglobinuria.
41	Chronic infection tends to result in variable clinical signs (de Waal, 1992).
42	Equine piroplasmosis can also be transmitted iatrogenically via the common use
43	of blood contaminated syringes and needles and this route is thought to be responsible for
44	the 2008 outbreak of the disease in several horses in Florida (Florida Department of
45	Agriculture and Consumer Services Division of Animal Husbandry, 2009).
46	Transplacental transmission of T. equi has been reported and although suspected,
47	evidence of <i>B. caballi</i> infection via vertical transmission <i>in utero</i> is lacking (Allsopp et
48	al., 2007). Infection of foals by T. equi in utero can result in abortions, full term still
49	births or the birth of live foals with neonatal piroplasmosis (Allsopp et al., 2007; Phipps
50	and Otter, 2004). Equine piroplasmosis due to B. caballi and T. equi has been reported in
51	adult horses in Trinidad using molecular (Rampersad et al., 2003) and serological
52	(Asgarali et al., 2007) techniques. This is the first report of a case of transplacental
53	transmission of T. equi in Trinidad.

54 Case description:

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55	Diplomacy, a 12-year old thoroughbred mare had a normal pregnancy and
56	parturition. The foal was born weak and severely icteric (Figure 1) with blood tinged
57	urine. Blood from the foal was collected into an EDTA tube for routine haematological
58	analysis at 10 hrs postpartum. A thin blood smear was stained with Wright-Giemsa ® and
59	examined under light microscopy. Piroplasms in red blood cells (RBC) were evident
60	(Figure 2). A stress leukon of lymphopaenia with thrombocytopaenia was evident (Table
61	1). The foal died 24 hrs after birth. Microscopic examination of a Wright-Giemsa $\ensuremath{\mathbb{R}}$
62	stained peripheral blood smear from the mare was negative for piroplasms.
63	Materials and methods
64	Reverse line blot (RLB)
65	DNA was extracted from 100 μL of EDTA blood from the foal and the mare
66	using the DNeasy blood and tissue kit (Qiagen Sciences Maryland, USA) according to
67	the manufactures instructions. The 18S rRNA gene for Babesia /Theileria was amplified
68	using primers F2 and R2 as described previously (Gubbels et al., 1999). DNA of B. canis
69	rossi was used as a positive control and PCR grade water (Sigma St. Louis MO, USA)
70	was used as a negative control. The amplified products were then subjected to the reverse
71	line blot (RLB) hybridization process (Gubbels et al., 1999). The oligonucleotide probes
72	that were attached to the membrane for the RLB are listed in Table 2.(Matjila et al., 2005;
73	Nagore et al., 2004)
74	Nested PCR
75	DNA extracted from the blood of the mare was then shipped to the University of

Georgia, USA and subjected to a nested PCR for the Babesia or Theileria 18S rRNA

77	gene using primary primers 5.1 and B (Medlin et al., 1989; Yabsley et al., 2005) followed
78	by a secondary reaction using primers F and R as described (Gubbels et al., 1999). To
79	prevent and detect contamination, DNA extraction, primary and secondary amplification,
80	and product analysis were done in separate dedicated areas. Two negative water controls
81	were included in each set of DNA extractions and 1 water control was included in each
82	set of primary and secondary PCR reactions. A blood sample from a white-tailed deer
83	(Odocoileus virginianus) naturally infected with Theileria cervi was used as a PCR-
84	positive control. To confirm identity, the positive sample was sequenced. The amplicon
85	was purified with a commercial gel-purification kit (QIAGEN, Valencia, CA) and
86	submitted to Clemson University Genomics Institute (Clemson, SC) for bi-directional
87	sequencing using the F and R primers.
88	Results
89	A high parasitaemia (63%) was noted in the thin blood smear from the foal. There
90	were on average 1-3 organisms per infected RBC. The RBC and haematocrit (HCT)
91	levels were on the low end of the reference interval(Weiss and Wardrop, 2010). Based on
92	the results of the RLB, the foal was infected with T. equi only (Figure 3). The mare tested
93	negative by RLB but an amplicon of ~ 550 bp was obtained by nested PCR analysis. The

nucleotide sequence (504 bp) obtained from the mare was 99.8% similar to *T. equi*

95 sequences from equines from Spain (AY 150062) and South Africa (EU 642508,

96 Z15105).

97 Discussion

98 Clinical signs of icterus and haemoglobinuria as well as reductions in the number
99 of red blood cells and platelets, reduction in haemoglobin concentration and leukopaenia

100 are common in equine babesiosis (de Waal, 1992). Hyperbilirubinaemia is due to 101 haemolytic anaemia. Haemolysis may be a result of mechanical damage to erythrocytes 102 by trophozoite intra-erythrocytic binary fission and toxic damage by haemolytic factors 103 produced by the parasite. The foal was anaemic, however the degree of true anaemia may 104 not be reflected on the foal's haemogramme because of the sharp increase in RBC 105 parameters at birth. During the first 12 hrs, transfusion of placental blood to the foal from 106 the mare increases the foals RBC parameters which then decline sharply over 12-24 hrs 107 (Weiss and Wardrop, 2010). Mild anisocytosis is also commonly observed.(Weiss and 108 Wardrop, 2010).

109 Our case has demonstrated that carrier mares can transmit *T. equi* to their foals 110 and such foals born at term can have an overwhelming parasitaemia. If icterus is not 111 observed immediately after birth, these cases of neonatal piroplasmosis could be 112 mistaken for neonatal isoerythrolysis.

113 Allsopp et al (2007) suggested that neonatal equine piroplasmosis may be acquired during days 40 - 150 of gestation which corresponds to the histotrophic when 114 115 the embryo is exposed to a mixture of uterine gland secretions, desquamated epithelial 116 cells and maternal erythrocytes (histotroph). A study which used medically induced 117 abortions in T. equi serologically positive mares, detected T. equi DNA in all aborted 118 foetuses. In that study, the earliest abortion was at day130 of gestation (Allsopp et al., 119 2007). However, microscopic examination of peripheral blood smears was not performed 120 in that study, hence the level of parasitaemia in the aborted foals and mares was 121 unknown. A report of transplacental transmission of T. equi has been described in 122 asymptomatic horses with no known natural exposure to the tick vector, thus indicating

123 that such carrier animals may act as reservoirs to establish equine piroplasmosis in areas 124 where competent tick vectors may be present (Phipps and Otter, 2004). The severely 125 icteric foal in our study had a 63% parasitaemia of T. equi, however, no piroplasms were 126 observed on light microscopy of Wright-Giemsa® stained peripheral blood smears of the 127 mare. The RLB was also not sensitive enough to detect T. equi DNA in the blood of the 128 mare but T. equi was detected using a nested PCR. The negative RLB result in the mare 129 in our study may be due to a very low level of parasitaemia which was below the limit of 130 detection of the RLB. The detection limit for the RLB for bovine piroplasms was estimated by Gubbels *et al* to be 10^{-6} % which corresponded to 3 parasites per µl of blood 131 132 (Gubbels et al., 1999). 133 The economic importance of transplacental transmission of equine piroplasmosis 134 on the thoroughbred industry in Trinidad needs further investigation as many cases may 135 be misdiagnosed. We therefore recommend that all icteric foals born in endemic areas 136 should be screened for neonatal piroplasmosis. This is the first reported case of neonatal 137 piroplasmosis in Trinidad.

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- 140 Figure 1 Icteric sclera and conjunctiva of neonatal foal at 10 h post- partum
- 141 Figure 2 Red blood cells (RBC) of foal infected with *T. equi* piroplasms. There were 126
- 142 infected RBCs per high power field (HPF). There are approximately 200 RBC per HPF
- 143 (100x oil objective).
- 144 Figure 3 RLB hyperfilm result for foal diplomacy (b) (1, *Babesia/Theileria* genera probe;
- 145 2, Babesia canis rossi; 3, Babesia caballi; 4, Theileria equi). a, b, c (equine samples), d
- 146 positive control DNA containing *B. canis rossi*, e (negative control)

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Thoroughbred foals at 24 hrs ¹		
RBC (x 10^{12} / L) – (9.1 – 11.9)	7.4	
Hgb (g/L) – (129.0-155.0)	107.0	
HCT(L/L) – (0.38 - 0.46)	0.34	
MCV (fl) - (36.0 - 44.0)	46.0	
MCH(pg) - (13.0 – 15.0)	14.4	
Anisocytosis	2+	
MCHC (g/L) - (320.0 - 360.0)	312.0	
WBC (x 10 ⁹ / L) – (6.0 – 12.0)	5.0	
Neutrophils (%) – (58.0-79.0) (x 10^9 / L) – (4.0-9.5)	74.0 3.7	
Bands (%) – (0 – 3.0) (x 10^9 / L) – (0 - 0.1)	4.0 0.2	
Metamyelocytes (%) – (0) (x 10^9 / L) - (0)	1.0 0.05	
Lymphocytes (%) – (15.0 – 35.0) (x 10^9 / L) – (1.3-3.1)	10.0 0.5	
Monocytes (%) – (1.0-7.0) (x 10^9 / L) – (0 -0.6)	9.0 0.45	
Eosinophils (%)- (1.0-10.0) (x 10 ⁹ / L)- (0 - 0.8)	0	
Platelets (x 10^9 / L) – (100.0 – 600.0)	58.0	
Protein (g/L) – (53.0 – 71.0)	68.0	
Fibrinogen (g/L) – (2.1- 3.3)	5.0	

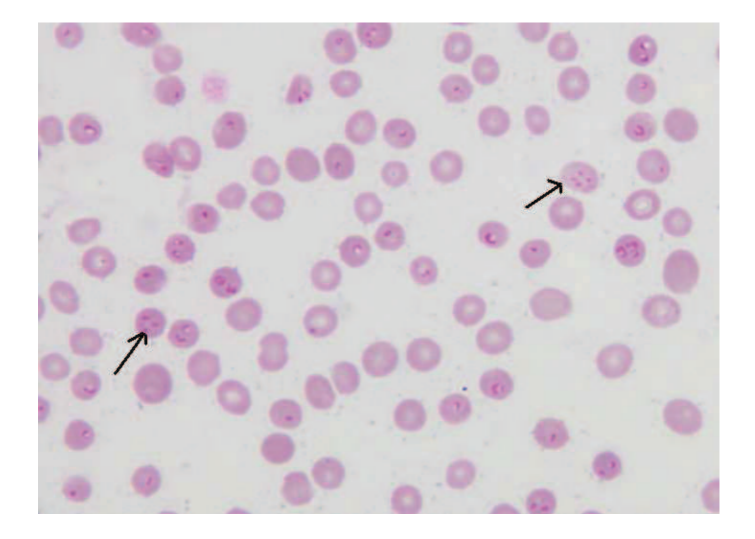
Table 1 The foal's haematological parameters at 10 hours post- partum Variable – Reference interval for Thoroughbred foals at 24 hrs¹

2

RBC = red blood cell, Hgb = haemoglobin, HCT = haematocrit, MCV = mean corpuscular volume, MCHC = mean corpuscular haemoglobin concentration, WBC = white blood cell. ¹ Adapted from Schalm's Veterinary Hematology, Sixth Edition (ed3

Weiss and Wardrop, 2010) 4





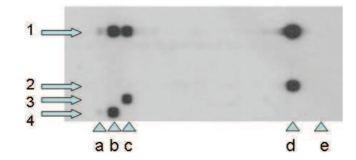


Table 2. 5'-3' sequence of oligor Oligonucleotide	nucleotide probes used in this study ²	ed in this study ²	
probe	5'-3' sequence of oligonucleotide probe	Reference	
Babesia/Theileria			
genera	TAATGGTTAATAGGARCRGTTG	(Gubbels et al., 1999)	
Babesia caballi	CGGGTTATTGACTTCGCTTTTTCTT	(Nagore et al., 2004)	
Babesia canis rossi	CGGTTTGTTGCCTTTGTG	(Matjila et al., 2005)	
Theileria equi	TCTGCTGTTTCGTTGACTG	GenBank Accession no. Z15105, AY150062	



 2 R = A or G