High levels of congenital transmission of *Toxoplasma gondii* in longitudinal and cross-sectional studies on sheep farms provides evidence of vertical transmission in ovine hosts

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SUMMARY

Recent research suggests that vertical transmission may play an important role in sustaining *Toxoplasma gondii* infection in some species. We report here that congenital transmission occurs at consistently high levels in pedigree Charollais and outbred sheep flocks sampled over a 3-year period. Overall rates of transmission per pregnancy determined by PCR based diagnosis, were consistent over time in a commercial sheep flock (69%) and in sympatric (60%) and allopatric (41%) populations of Charollais sheep. The result of this was that 53.7% of lambs were acquiring an infection prior to birth: 46.4% of live lambs and 90.0% of dead lambs (in agreement with the association made between *T. gondii* and abortion). No significant differences were observed between lamb sexes. Although we cannot distinguish between congenital transmission occurring due to primary infection at pregnancy or reactivation of chronic infection during pregnancy, our observations of consistently high levels of congenital transmission over successive lambings favour the latter.

Key words: Toxoplasma gondii, congenital transmission, ovine, abortion.

INTRODUCTION

The protozoan parasite Toxoplasma gondii is common to most warm-blooded organisms (Dubey & Beattie, 1988; Tenter, Heckeroth & Weiss, 2000). It is normally apathogenic and serious disease is restricted to those with a weakened or immature immune system as is the case with a developing foetus. Sheep are particularly susceptible and primary infection leads to congenital disease or abortion (Beverley & Watson, 1971; Buxton, 1990). Serological assessment suggests that 25-36% of sheep in the British Isles are infected (Leguia & Herbert, 1979; Samad & Clarkson, 1994) and the disease accounts for one third of all diagnosed ovine abortions (SAC Veterinary Science Division, 1999). It has long been held that the major route of transmission was via oocysts released in the faeces of the felid definitive host following sexual recombination (Hutchison, 1965) and this has been proposed as the main transmission route in sheep. In carnivores and omnivores, transmission of asexual bradyzoite stages by carnivory is also important and recent reports suggested that this favoured transmission route has led to the expansion and clonal population structure of the parasite (Su et al. 2003). Although transplacental transmission of the parasite is demonstrated in many species, it is usually considered in terms of disease rather than as a potential route of transmission.

Beverley was the first to suggest that serial vertical transmission might sustain the parasite (Beverley, 1959) and that this might be important in maintaining the parasite in some populations, however this has been a contentious issue (Johnson, 1997). Supportive evidence of vertical transmission over a single generation has been obtained in populations of experimentally infected mice (Apodemus sylvaticus and Mus domesticus) (Owen & Trees, 1998) and in natural populations of domestic mouse (M. domesticus)(Marshall et al. 2004). Experimentally infected rats exhibit high rates of parasite transmission to offspring when infected during pregnancy (Dubey & Shen, 1991; Dubey et al. 1997) although rates were found to differ depending on parasite strain, host infection route and parasite stage (Zenner et al. 1993). Furthermore, the situation in rats was different from that observed in mice and hamsters where T. gondiiinfected dams could produce several infected litters without re-infection (Beverley, 1959; Roever-Bonnet, 1969). More recently high rates of congenital transmission have been reported in epidemiological investigations in sheep (Duncanson et al. 2001). The mechanism by which congenital transmission in these various species is achieved is less clear. Two possible scenarios can be envisaged: firstly, primary

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exposure of the female during pregnancy followed by transmission to the foetus during pregnancy or secondly, reactivation of chronic infection in the female during pregnancy. There is currently contention as to the importance and relevance of congenital transmission in toxoplasmosis.

Past estimates of Toxoplasma infection in sheep suggest a median of 30% infection amongst ewe populations by serology (Blewett, 1983; Van der Puije et al. 2000). There are few reports on age-related prevalence; however, those that exist report a gradual increase in prevalence with approximately 24% seroconversion per annum (Waldeland et al. 1977). In contrast, the study of Duncanson et al. (2001), using PCR diagnosis to estimate disease transmission to newborn lambs found high transmission rates (61%). This study was based on a single lambing and could have been the result of an unusually high level of primary exposure to infection. Two important questions exist: are these levels of congenital transmission a consistent phenomenon across flocks or breeds and is congenital transmission a significant route of transmission in sheep? In this study we aim to address these questions by measuring rates of congenital transmission over time, between breeds and between farms.

MATERIALS AND METHODS

Commercial Suffolk-cross and pedigree Charollais sheep from a farm near Droitwich in Worcestershire were investigated over 5 subsequent lambings (March 1999–March 2001). Sampling was carried out over a period of 1–2 weeks during winter and spring lambings. Commercial sheep were lambed during both periods, whereas the Charollais flock were only lambed in winter. The two flocks were kept in separate housing and grazed in different areas allyear-round as required for the pedigree flock. This flock was maintained in house with replacement from home-grown stock rather than bought-in ewe lambs. Charollais sheep were also sampled on a second farm near Crewe in Cheshire for a single lambing period during the winter of 2001/02.

Collection of tissue samples was carried out as follows. For all samplings, a strict sterility regime was followed: tissues were removed using a fresh set of sterilized instruments for each tissue to prevent contamination across tissues and with external skin/ hair. Samples were kept in separate tubes and immediately frozen for storage. Instruments were thoroughly washed in Miltons solution and then sterilized using a portable steam sterilizer or by immersion in alcohol and flame treatment.

Aborted lambs

Lambs were dissected to reveal internal tissues (brain, heart, lung, liver, tongue) and tissues removed using the sterility protocol described above, taking care to use different, fresh, instruments for internal tissues following external dissection.

Live lambs

(a) During the Spring 1999 lambing, samples were taken from foetally-derived placental tissue or lamb cord immediately after birth as described previously (Duncanson et al. 2001). Sterile instruments were used throughout, care being taken to avoid crosscontamination and the sampler was present at all samplings. Samples were frozen immediately until DNA extraction could be undertaken. (b) All subsequent lambings (Winter 2000-Spring 2001; all Charollais samplings): umbilical cord tissue was taken aseptically immediately after birth using sterile instruments. The sampler was present at all samplings and the sample was not allowed to come into contact with other samples or with maternal tissue. Cord tissue was taken as close to the lamb as possible. Tissues were frozen immediately until DNA extraction could be performed.

All samples were transported to laboratories at Salford University for DNA extraction and PCR and these operations were conducted in such a way as to avoid any cross-contamination of samples. DNA was extracted from tissue samples as described previously (Duncanson et al. 2001; Terry et al. 2001) using a standard phenol-chloroform extraction process. For the detection of Toxoplasma gondii a nested polymerase chain reaction (PCR) was used to amplify the Surface Antigen Gene 1 (SAG1) (Savva et al. 1990), as described (Duncanson et al. 2001). Mammalian tubulin PCR was used as a positive PCR control for all samples as described previously (Terry et al. 2001. Products of amplification were identified by gel electrophoresis as described (Duncanson et al. 2001).

Reliability of PCR results was satisfied in a number of ways. Firstly, mice experimentally infected with reference isolates of Toxoplasma gondii (J. E. Smith, Leeds) showed SAG1 PCR amplification from internal tissues taken from infected mice while no amplification was achieved from uninfected mice. Secondly, the nested SAG1 PCR system was specific to Toxoplasma as amplification from DNA extracted from other related apicomplexans (Neospora, Sarcocystis, Hammondia) was not observed. Thirdly, the applicability of the procedure to field samples was judged by the successful SAG1 PCR amplification from sheep, fox, mouse, Apodemus, and human DNA spiked with low concentrations of Toxoplasma DNA but lack of amplification when they were spiked with any concentration of Neospora DNA. Finally, the reliability of the PCR analysis of lamb cord samples as indicators of PCR positivity of internal tissues, was judged by comparison in aborted lambs where both tissues could be sampled. In a sample of Table 1. Frequency of abortion and lamb losses in Suffolk cross and Charollais flocks

(Lambing losses in the 2 separate sheep flocks at the Droitwich Farm during the time period investigated. Differences in mean losses between commercial and Charollais sheep were found to be statistically significant. (Lamb losses: $\chi^2 = 27.6$; D.F. = 1; P < 0.001; Pregnancy losses: $\chi^2 = 23.9$; D.F. = 1; P < 0.001).)

Flock	Lambing	Spring 1999	Winter 2000	Spring 2000	Winter 2001	Spring 2001	Total
Commercial	No. aborted lambs	61	46	26	34	30	197
	Total no. lambs	619	505	454	386	443	2407
	% Lambs aborted	9.9%	9.1%	5.7%	8.8%	6.8%	8·2%
	No. unsuccessful pregnancies	39	36	20	23	20	138
	Total no. pregnancies	298	296	242	240	243	1319
	% Pregnancies	$13 \cdot 1\%$	$12 \cdot 2\%$	8.3%	9.6%	8·2%	10.5%
Charollais 1	No. aborted lambs	N/A	21	N/A	20	N/A	41
(Sympatric)	Total no. lambs	N/A	102	Ň/A	115	N/A	217
	% Lambs aborted	N/A	20.6%	N/A	17.4%	N/A	18.9%
	No. unsuccessful pregnancies	N/A	14	N/A	16	N/A	30
	Total no. pregnancies	Ń/A	55	Ň/A	62	Ń/A	117
	% Pregnancies	N/A	25.5%	N/A	25.8%	N/A	25.6%
Charollais 2	No. aborted lambs	N/A	N/A	N/A	2	N/A	2
(Allopatric)	Total no. lambs	N/A	N/A	N/A	44	N/A	44
	% Lambs aborted	Ń/A	Ň/A	Ň/A	4.5%	Ń/A	4.5%
	No. unsuccessful pregnancies	N/A	N/A	N/A	2	N/A	2
	Total no. pregnancies	Ń/A	Ň/A	N/A	24	Ń/A	24
	% Pregnancies	Ń/A	Ń/A	Ń/A	8.3%	Ń/A	8.3%

42 aborted lambs, 36 (86%) were either positive (n=29) or negative (n=7) for both lamb cord and the internal tissue while the remaining 6 were only positive in the internal tissues leading us to conclude that PCR amplification from lamb cord is a good indicator of the PCR status of internal tissue and, if anything, underestimates PCR positivity in internal lamb tissues. Good agreement was also found when comparing a range of internal tissues.

Statistical analyses of lambing results were carried out using the Chi² test.

RESULTS

The aim of this study was to compare rates of *Toxoplasma* transmission and abortion over time, between breeds and between farms. Rates of congenital transmission were assessed in 5 sequential lambings over a period of 2 years in a flock of commercial sheep. In addition two Charollais sheep flocks were followed over the same period-first a sympatric and the second an allopatric population. In the commercial flocks between 8 and 13% of ewes aborted resulting in lamb losses of between 5.7 and 9.9% (Table 1). In the sympatric Charollais flock these levels were higher with one in four ewes suffering abortion and lamb losses of 17–20% (Table 1). In the allopatric population abortion occurred in 8.3% of ewes with lamb losses of 4.5% (Table 1).

Congenital transmission of *Toxoplasma* parasites was determined during a 2-week sampling window in each lambing period of pregnancies (more sheep in Table 1) by SAG1 PCR amplification from internal tissues of dead lambs, lamb umbilical cord from live

born lambs or occasionally placental tissue, which is foetally derived (see Materials and Methods section for details). These data show sustained high levels of congenital transmission over the study period (Table 2). Levels of congenital transmission remained high throughout the study with 47-78% of pregnancies affected and an overall mean for the 5 lambings of 69%. Transmission rates in unsuccessful pregnancies (scored as those where one or more of the lambs was aborted, stillborn or died shortly after birth) were consistently high, with a mean of 91% lambs infected overall. In contrast, levels of congenital transmission in successful pregnancies had an overall mean of 65%. There was a significant difference in the rate of maternal transmission between successful and unsuccessful pregnancies ($\chi^2 = 15.8$; D.F. = 1; P < 0.001).

To examine whether the high transmission rate was peculiar to the commercial sheep flock we conducted a parallel analysis of Charollais flocks on sympatric and allopatric farms (Table 3). Congenital transmission rates in the sympatric Charollais flock were found to be similar to the overall levels found in the commercial flock (60% of pregnancies). Again this level was significantly higher in the unsuccessful pregnancies (96%) than in the successful ones (43%) $(\chi^2 = 18.7; D.F. = 1; P < 0.001)$. T. gondii transmission rates were also high (41% of pregnancies) in a second Charollais flock on a separate geographically distant farm (Table 4). Infection rates here were not significantly higher in unsuccessful pregnancies (100%) than in the successful ones (36%) ($\chi^2 = 3.27$; D.F. =1; P = 0.07; however, lack of statistical significance may be due to the small sample size.

Table 2. Toxoplasma infection in a Suffolk-cross flock (Droitwich)

(Congenital transmission of *Toxoplasma gondii* in a flock of commercial sheep over a period of 2 years. (Note: Spring 1999 data taken from Duncanson *et al.*, 2001.) For cumulative data, differences in infection rates of lambs between successful and unsuccessful pregnancies were found to be significant ($\chi^2 = 15.8$; D.F. = 1; P < 0.001).)

Lambing		Outcome of pr		
	Data	Successful	Unsuccessful	Total
Spring 1999	No. of pregnancies	70	18	88
	No. of PCR-positive pregnancies	37	17	54
	Total	42%	94%	61%
Winter 2000	No. of pregnancies	57	15	72
	No. of PCR-positive pregnancies	41	12	53
	Total	72%	80%	74%
Spring 2000	No. of pregnancies	113	18	131
	No. of PCR-positive pregnancies	83	18	101
	Total	73%	100%	77%
Winter 2001	No. of pregnancies	44	7	51
	No. of PCR-positive pregnancies	18	6	24
	Total	41%	86%	47%
Spring 2001	No. of pregnancies	50	0	50
	No. of PCR-positive pregnancies	39	0	39
	Total	78%	N/A	78%
Cumulative	No. of pregnancies	334	58	392
	No. of PCR-positive pregnancies	218	53	271
	Total	65%	91%	69%

Table 3. *Toxoplasma* infection in Charollais flock 1 (sympatric population)

(Congenital transmission of *Toxoplasma gondii* in a flock of Charollais pedigree sheep over a period of 3 lambings (Winter- 2000, 2001 and 2002). Differences in lamb infection rates between successful and unsuccessful pregnancies were found to be significant ($\chi^2 = 18.7$; D.F.=1; P < 0.001).)

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	Outcome of				
Data	Successful	Unsuccessful	Total		
No. of pregnancies	49	24	73		
No. of PCR-positive pregnancies	21	23	44		
Total	43%	96%	60%		

Table 4. Toxoplasma infection in Charollais flock 2(allopatric population)

(Congenital transmission of *Toxoplasma gondii* over a single lambing period in the Cheshire farm flock of Charollais pedigree sheep. Differences in lamb infection rates between successful and unsuccessful pregnancies were not found to be significant using the chi squared test (χ^2 =3·27; D.F.=1; P=0·07).)

	Outcome of		
Data	Successful	Unsuccessful	Total
No. of pregnancies	22	2	24
No. of PCR-positive pregnancies	8	2	10
Total	36%	100%	41%

During later commercial lambings and for the two Charollais flocks, lamb cord was also sampled. This allowed rates of individual lamb infection to be determined, as well as maternal transmission. By analysis of this cumulative data (Table 5), it was shown that transmission of the parasite in these sheep flocks reaches 53.7% of lambs. This rate of transmission was significantly higher in dead lambs (90%) than in live ones (46.4%) ($\chi^2 = 43.1$; D.F.=1; P < 0.001).

Given the high rates of congenital transmission occurring within these groups, we investigated possible patterns in the way the parasite was transmitted. No significant differences were seen with regard to lamb sex ($\chi^2 = 1.92$; D.F. = 1; P = 0.17),

although infection rates and mean losses were slightly higher in males. No significant differences was seen between the winter and spring lambings of the commercial flock ($\chi^2 = 1.2$; D.F. = 1; P = 0.27).

Certain patterns did, however, emerge by grouping litter types together (i.e. singletons, twins, triplets) (Table 6). Firstly, there is a trend towards a higher level of transmission with an increased number of offspring, although this variation is not significant ($\chi^2 = 2.49$; D.F.=2; P=0.29). Secondly, as the number of offspring increases there is a significant increase in the risk of abortion ($\chi^2 = 16.2$; D.F.=2; P < 0.001). This can be explained by the increased risk of mortality of infected lambs, which varies significantly with offspring number ($\chi^2 = 14.6$; Table 5. Cumulative individual lamb infection data (including all available data from the 3 flocks)

(Cumulative infection data in individual lambs. Differences in infection rates between live and dead lambs were found to be significant using the chi squared test ($\chi^2 = 43.1$; D.F. = 1; P < 0.001).)

Lamb Status	Alive	Dead	Total
Total no. of lambs infected Total no. of lambs	163 351	63 70	226 421
Total	46.4%	90.0%	53.7%

Table 6. Lamb abortion and *Toxoplasma* infection rates by number of offspring (cumulative data for the 3 flocks)

(Abortion and lamb infection rates by number of offspring. (*Total values includes a single quadruplet litter group that was excluded from the table due lack of statistical significance.) Variation in % aborted with increased offspring number is significant ($\chi^2 = 16 \cdot 2$; D.F. = 2; P = 0.0003); variation in % infected with increased offspring number is not significant ($\chi^2 = 2 \cdot 49$; D.F. = 2; P = 0.29); variation in % mortality of infected lambs with increased offspring number is significant ($\chi^2 = 14 \cdot 6$; D.F. = 2; P = 0.0007); variation in % mortality of uninfected lambs with increased offspring number is not significant (Fisher's exact test; P > 0.05).)

Litter type	% Aborted lambs	% Lambs infected	Mortality in infected lambs	Mortality in uninfected lambs
Singletons	17·9%	50·7%	31·3%	3·0%
	(12/67)	(34/67)	(11/34)	(1/33)
Twins	12·2%	52·2%	20%	3·8%
	(34/278)	(145/278)	(29/145)	(5/133)
Triplets	31·9%	62·5%	48·9%	3·7%
	(23/72)	(45/72)	(22/45)	(1/27)
Total*	16·6%	53·7%	31·9%	3·6%
	(70/421)	(226/421)	(63/226)	(7/195)

D.F. = 2; P < 0.001). There is no significant increase in uninfected offspring (Fisher's exact test; P > 0.05).

DISCUSSION

At the outset of study we raised the question of whether the elevated level of vertical transmission reported in sheep (Duncanson et al. 2001) was caused by unusual circumstances such as a high level of primary infection. Abortion storms, presumed to be associated with a wave of primary infection with Toxoplasma, have been known to occur (Hartley & Marshall, 1957) and it is suggested that these outbreaks can be attributed to contact between the flock and infected cats during gestation. By using PCR positivity of the Toxoplasma SAG 1 gene as a measure of infection with Toxoplasma, our data clearly show that vertical transmission occurs in 60-70% of all pregnancies and that this figure remains consistent through time, and with season, breed and geographical area. The result of this is that approximately 46.4% of live lambs are born infected. This figure is much higher than the 7.6–16.2% predicted from serological assessment (Skjerve et al.

1998; Van der Puije et al. 2000). These differences in the sensitivity of PCR and serological assay are known to exist. PCR detects the presence of parasite DNA in tissue samples while serological methods measure exposure to the parasite. In a comparative study of PCR with mouse inoculation and serological methods it was found that PCR was at least as sensitive as mouse inoculation and better than some serological methods as a diagnostic tool for ovine toxoplasmosis (Owen, Clarkson & Trees, 1997). Several authors report that PCR-based assays are more sensitive in detection of the parasite than serology (Owen & Trees, 1998; Hafid et al. 2001) and in one study it was reported to be more accurate than the modified agglutination test (MAT) especially in the diagnosis of vertical transmission (Owen & Trees, 1998). In mice, it has been suggested that this latter phenomenon may be explained by compromised antibody responsiveness of young mice born to chronically infected dams (Suzuki & Kobayashi, 1990). There clearly remains a discrepancy between levels of serological and PCR positivity in newborn lambs in the literature, and further investigation is needed to understand the immunological interactions

occurring at this stage. We do not know the consequences of infection in live born lambs, whether they go on to develop systemic disease and chronic parasitaemia or whether parasite infection fails to establish. However, this early exposure to the parasite could compromise the development of anti-parasite immunity enhancing the likelihood that serial vertical transmission could occur. Vertical transmission is consistent with the data presented in this study.

Vertical transmission of *T. gondii* over one generation has been demonstrated unequivocally in experimentally infected mice (Owen & Trees, 1998). In the closely related parasite *Neospora caninum*, vertical transmission is known to occur in several host animals including mice (Cole *et al.* 1995), dogs (Barber & Trees, 1998), foxes (Schares *et al.* 2001) and in cattle, where serial vertical transmission is very efficient (Davison, Otter & Trees, 1999) and is often associated with foetal pathology (Bjorkman *et al.* 1996).

The overall rate of 53.7% vertical transmission for *Toxoplasma* in sheep reported in this study may not be sufficient to sustain parasite infection within sheep populations and it is likely that the parasite has the capacity for both vertical and horizontal transmission and the use of these routes might vary according to host, parasite strain and prevailing environmental conditions. In support of a role for vertical transmission, Terry *et al.* (2001), carrying out an initial genotyping study on this farm, found that *Toxoplasma* isolates taken from this flock had identical MGE genotypes.

We found a strong relationship between infection and abortion with 90% of aborted animals being Toxoplasma positive compared to 46.4% of live-born individuals. Abortion rates were higher in Charollais raising the possibility that pathogenesis varies with host breed. In addition to breed, other factors may also contribute to abortion; one of the most interesting of these is the positive relationship between the number of offspring and rates of infection and abortion. In humans, it has been proposed that reactivation of Toxoplasma infection and consequent abortion might result from pregnancy-induced stress and that that would be more severe in triplets than in twins and singletons (Avelino & Campos, 2002). If this were also the case in sheep, our data would be consistent with such a concept.

The importance of vertical transmission of *Toxoplasma* requires closer evaluation. In particular, a crucial question which needs to be addressed, is whether this transmission is due to primary infection during pregnancy or reactivation from chronic infection. If the latter is the case then animal husbandry, control methods and management techniques may need to be re-evaluated; vaccine strategies may need to focus on preventing transmission rather than reducing levels of foetal abortion. Further work is necessary to address these important questions.

Future epidemiological studies will be needed to determine whether vertical transmission of *Toxoplasma* is a common feature in herbivores and if so whether it might contribute to the clonal 'asexual' expansion and parasite population structure seen across Europe and North America (Su *et al.* 2003).

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