Contagious epididymitis due to *Brucella ovis*: relationship between sexual function, serology and bacterial shedding in semen

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**Abstract**

**Background:** Contagious Epididymitis (CE) due to *Brucella ovis* (*B. ovis*) is a contagious disease that impairs rams’ fertility due to epididymis, testicle and accessory sexual gland alterations. An increased incidence of CE has been observed in South Eastern France ("PACA" region) since the Rev.1 vaccination against *B. melitensis* has been stopped in 2008. The objective of this study was to evaluate the relationship between the infection by *B. ovis* and the sexual function of rams.

Two-hundred eighteen sexually-mature rams, from 11 seropositive flocks, were submitted to a clinical examination of the genital tract, a semen collection by electro-ejaculation for spermogram and culture, and a serological examination for anti-*B. ovis* antibodies by complement fixation test (CFT) and indirect ELISA (I-ELISA). The relationships between clinical, seminal, bacteriological and serological parameters were studied using the Fisher exact test and a logistic regression model (binomial logit).

**Results:** *B. ovis* shedding in semen was significantly associated with seropositivity (CFT and I-ELISA; \( p < 0.001 \) and 0.01 respectively), genital tract alterations \( (p < 0.05) \) and poor semen quality \( (p < 0.001) \). Seropositive rams presented significantly more genital tract alterations \( (p < 0.001) \) and a poor seminal score \( (p < 0.001) \) than seronegative rams.

**Conclusions:** Since semen culture is not routinely feasible in field conditions, a control plan of CE should be based, where Rev.1 vaccination is not possible, on both systematic clinical and serological examination of rams, followed by the culling of seropositive and/or genital tract alterations carrier rams.

**Keywords:** *Brucella ovis*, Semen, Sheep, Genital disease

**Background**

Contagious Epididymitis (CE) due to *Brucella ovis* (*B. ovis*) is a contagious disease of worldwide importance that impairs fertility in rams due to epididymis, testicle and accessory sexual gland alterations. More rarely, abortions in ewes and increased perinatal mortality rates may be observed as well \([1, 2]\). Transmission occurs via passive venereal infection or direct contact \([2]\). The disease can result in significant economic losses in infected flocks where no control programs are in place, due to reproductive failure, culling of breeding animals as well as ban on trade. In the European Union, no compulsory surveillance of the disease is currently in place in flocks, while neither eradication programme nor compensation scheme for culling animals in infected flocks is foreseen. Nevertheless, in order to avoid the contamination of non-infected areas or flocks through international trade, rams have to undergo serological pre-movement tests \([2]\). On farms, diagnosis mainly relies on a clinical detection and a serological test when the palpation of testicles reveals lesions or when there is significant infertility in the flock \([1, 3]\).
An increased incidence of CE has been observed in South Eastern France ("PACA" region) since the Rev. 1 vaccination against *B. melitensis* has been stopped in 2008 due to eradication of brucellosis in domestic ruminants 5 years before in the whole country [3]. In 2011, the flock prevalence rate (at least 1 seropositive ram per flock) ranged from 5.5 % to more than 50 % according to the area.

The aim of this study was to examine the relationship between *B. ovis* infection, evaluated through clinical, serological and bacteriological findings, and the sexual function, in rams originated from infected flocks of this region where CE appears enzootic.

**Methods**

**Animals**

Two-hundred eighteen sexually-mature rams from 11 seropositive flocks from the PACA region of South-Eastern France were included in this study in September 2012. Both seropositive and seronegative animals (based on results of tests performed in former spring) were selected by voluntary breeders in a balanced manner, as much as possible. Breeds of rams were as follows: Merinos (n = 69), Préalpes du Sud (n = 30), meat breeds (Ile de France and Texel; n = 91); Merinos cross-breeds (n = 15), others (n = 13).

**Breeding soundness assessment**

For each animal, the following parameters were recorded: age, body condition score (BCS), scrotal circumference (SC, in cm), as well as clinical alterations of the genital tract investigated by palpation and, in suspicious cases, by ultrasonography [4].

The Body Condition Score (BCS) was assessed according to the technique described by Russel [5]. Briefly, the BCS was noted from 0 (extremely emaciated) to 5 (obese) by the palpation of the lumbar region (spinous and transverse processes, muscles and fatty tissues), a BCS of 3 to 4 being considered as optimal during the breeding season [6].

Testicle alterations included asymmetry, indurations, degeneration and/or atrophy, while epididymis alterations included epididymitis, indurations, nodules, hypertrophy and/or cysts.

In order to classify the rams for their breeding potential, a clinical score and a seminal score based upon a point-score system adapted from Ley et al. [7] were calculated for each ram. The clinical score took into account the age, BCS, SC and the presence/absence of clinical signs, as shown in Table 1.

**Animals’ handling and sample collection**

All samples were collected in the frame of the *B. ovis* infection control programme implemented by the Fédération Régionale des Groupements de Défense Sanitaire Provence Alpes Côte d’Azur (FRGDS PACA) as approved by its advisory board and in accordance with the EU (in particular Directive 2010/63/EU) and French regulations regarding ethics and best practices of veterinary care.

Animals’ handling and sample collection were performed by trained technicians of the FRGDS PACA, veterinary surgeons and graduating veterinary students (under the supervision of 2 professors), with the help of the stockbreeders who had brought their own rams and given their informed consent. Blood samples were collected from each ram for serological examinations. Semen was collected by electroejaculation (Electrojac, Ideal Instruments, MI, USA) for further laboratory examinations. To preserve the welfare of the ram, the stimulation was discontinued if signs of stress or physical discomfort were detected.

**Laboratory examinations**

Each blood serum was subjected to both complement fixation test (CFT) and indirect ELISA (I-ELISA; IDEXX

| Table 1 Clinical score calculation and rams classification |
|-----------------------------------------------------------|
| **Note** | 1 | 4 | 7 |
| Age (years) | ≥5 | | 1 | 2-4 |
| BCS* | <2 | | >3 | 2-3 |
| SC (cm)** | <33 | | ≥33-< 39 | | ≥39 |
| Epididymis/testicle alterations*** | Presence (note = 0) | | N/A**** | Absence (note = 7) |
| Clinical score | | | | |
| Rams classification | 3-12 | 13-21 | 22-28 | |
| Poor | | | | |
| Fair | | | | |
| Good | | | | |

*BSC: body condition score (0–5)

**SC: scrotal circumference

***Testicle alterations: asymmetry, indurations, degeneration and/or atrophy; Epididymis alterations: epididymitis, indurations, nodules, hypertrophy and/or cysts

****N/A: not applicable
**Brucella ovis** Ab Test; Idexx Montpellier, France) as described by Praud et al. [8]. The CF antigen (Anses, Maisons-Alfort, France) was standardised against the International Standard anti-*Brucella ovis* Serum (ISaBoS) and the positivity threshold was 50 international CFT units (ICFTU)/mL according to OIE requirements [2]. I-ELISA was standardized against ISaBoS with the 1/64 pre-dilution of the ISaBoS made up in a negative pool of sera as the minimum detection requirement and the 1/256 dilution at which the standard must be classified as negative. All tests were performed by the same two technicians at the Anses Brucellosis Reference Laboratory (Anses, Maisons-Alfort, France). In a previous work, Praud et al. [8] suggested that depending on the aim of the test performance (screening, diagnosis confirmation, export control) and disease situation (free, enzootic), different cut-offs could be used (30 %, 45 % or 60 %) but found that the 45 % cut-off gave the best concordance with CFT. In this study, I-ELISA results were therefore interpreted either with a unique cut-off of 45 % (negative or positive) or with a double-cut-off of 30 % and 60 % respectively (negative, doubtful or positive).

For the semen examination, the following characteristics were recorded: volume (mL), gross and progressive motility, sperm morphology and abnormalities after eosin-nigrosin staining; sperm concentration (spermatozoa per mL) was measured by spectrophotometry (Accuread, IMV technologies, L’Aigle, France). The seminal score was based on the number of spermatozoa in the ejaculate, the individual motility and the percentage of normal spermatozoa, as shown in Table 2.

Semen cultures for *B. ovis* were performed on samples stored frozen, on non-selective 5 % equine-serum added blood-agar base N°2 (Oxoid, France) in an atmosphere of 5–10 % CO₂, at the ANSES Brucellosis Reference Laboratory [2].

**Statistical methods**
Statistical analysis was performed using the Fisher exact test and a logistic regression model (binomial logit) with the R software (http://www.R-project.org). A threshold of 5 % (p < 0.05) was considered as significant.

Kappa statistics were also used to assess the agreement between different diagnostic methods.

**Results**

**Clinical and semen laboratory examinations**
The clinical and seminal characteristics of the 218 rams are presented in Table 3. In 73 rams, clinical alterations were observed by palpation, mainly on epididymes (n = 60; head and tail hypertrophy, indurations and nodules) and testicles (n = 13; asymmetry, indurations, degeneration and atrophy); 17 alterations were bilateral, 56 unilateral. 145 rams had no macroscopic alteration of the genital tract. Respectively 26 % and 37 % of rams had a good clinical or seminal score.

There was a significant relationship between seminal and clinical score (p = 0.0084), mainly explained by the presence (or not) of clinical alterations but not by the SC or age.

| Table 2 Seminal score calculation and rams classification |
|----------------------------------------------------------|
| Note | 1 | 4 | 7 |
| Total number of spermatozoa/ejaculate (x10⁹) | =< 1 | >1-≤ 2 | >2 |
| Individual motility (%) | =< 30 | >30-≤ 70 | >70 |
| Normal spermatozoa (%) | =< 50 | >50-≤ 70 | >70 |
| Rams classification | 3-9 | 12-15 | 18-21 |
| Poor | | | |
| Fair | | | |
| Good | | | |

| Table 3 Clinical and seminal characteristics of the 218 rams |
|-------------------------------------------------------------|
| Parameter | Mean | SD | Median | Min. | Max. |
| Age (years) | 3.3 | 1.7 | 3 | 1 | 9 |
| BCS* | 2.4 | 0.7 | 2.3 | 1.5 | 3.8 |
| SC (cm)** | 36.4 | 3.6 | 34 | 26 | 46 |
| Number of spermatozoa/ejaculate (x10⁹) | 1.9 | 1.8 | 1.3 | 0.004 | 8.9 |
| Individual motility (%) | 48.9 | 29.9 | 57.5 | 0 | 90 |
| Normal spermatozoa (%) | 70.8 | 22.5 | 78.7 | 2.6 | 98.5 |
| Clinical score | Seminal score |
| Poor | Fair | Good | Total |
| Poor | 23 | 21 | 12 | 56 |
| Fair | 31 | 36 | 38 | 105 |
| Good | 12 | 14 | 31 | 57 |
| Total | 66 | 71 | 81 | 218 |

*BBCS : body condition score; **SC : scrotal circumference*
Serological results

Detailed serological results are given in Table 4. As expected from the inclusion criteria, about half of animals were effectively seropositive but more clearly in I-ELISA (53.7 %) when a unique cut-off of 45 % was chosen according to Praud et al. [8] than in CFT (37.2 %).

One serum presented an anti-complementary activity and was removed from the analysis. The two tests gave concordant results for 181 animals (83.0 %) with a higher number of positive results with I-ELISA than with CFT (117 vs. 81). The agreement between CFT and I-ELISA (45 %), evaluated by the Cohen’s kappa coefficient, was good: $\kappa = 0.676$ (95 % CI [0.584;0.767]).

### Relationship between serological results and seminal score

Most of I-ELISA-positive animals (83 %) presented a poor or fair seminal score and 40 % of I-ELISA-negative rams presented such a poor or fair seminal score as well (Tables 6 and 7). The Fisher exact test showed a significant relationship between the positive serological status of rams and the low seminal score ($p < 0.001$).

### Bacteriological results

Only 198 results were interpretable; for 16 samples, cultures were contaminated by overgrowing microorganisms that made the interpretation impossible; and 4 samples were not tested due to an insufficient quantity of semen. *B. ovis* was isolated in the semen of 89 rams, *i.e.* 44.95 % of 198 sampled animals with culture results.

There was a significant relationship ($p < 0.05$) between culture results and clinical score (Table 6). Among the parameters determining the clinical score, only BCS ($p < 0.01$) and lesions ($p < 0.001$) influenced significantly the bacteria shedding.

There was a significant relationship ($p < 0.001$) between culture results and seminal score (Table 8). A strong relationship between culture and CFT positive results ($p < 0.001$) as well as I-ELISA positive results ($p < 0.01$) was also evidenced: almost all rams shedding *B. ovis* in their semen gave I-ELISA positive results (Table 9). The agreement between culture and I-ELISA (45 %) ($\kappa = 0.710$; 95 % CI [0.615; 0.806]).

### Discussion

Our study clearly demonstrates that the *B. ovis* infection induces genital lesions and alters the semen quality, leading to an alteration of sexual function of the rams. However this study performed in field conditions on 218 rams included a bias associated to the selection of flocks (voluntary instead of randomly selected) and rams (selected by their owners instead of random), leading to a study sample that could be not representative of the ram population in South-Eastern France.

#### Table 4 Distribution of the results (number of rams) and concordance of serological tests

| CFT*          | I-ELISA (30–60 %)** | CFT I-ELISA (45 %)** |
|---------------|---------------------|----------------------|
|               | Negative   | Doubtful | Positive | Total | Negative | Positive | Total |
| Negative      | 95         | 9        | 32       | 136    | 100      | 36       | 136   |
| Positive      | -          | 1        | 80       | 81     | -        | 81       | 81    |
| Total         | 95         | 10       | 112      | 217    | 100      | 117      | 217   |

*One anti-complementary serum not included in the analysis

**Cut-off: 30-60 % or 45 %

$\kappa$: Cohen’s kappa coefficient; 95 % CI = 95 % confidence interval

#### Table 5 Relationship between serological tests and clinical (testicular and/or epididymal) alterations

| Clinical alterations | Serological results | I-ELISA (45 %)** |
|----------------------|---------------------|-------------------|
|                      | CFT* Negative | Positive |  p < 0.001*** | Negative | Positive |  p < 0.001*** |
| Yes                  | 26         | 47       |               | 14       | 59       |               |
| No                   | 110        | 34       |               | 87       | 58       |               |
| Total                | 136        | 81       |               | 101      | 117      |               |

*One anti-complementary serum not included in the analysis

**Cut-off: 45 %

*** Fisher exact test
Another bias is due to the staining kit that probably induced a high percentage of artifactual sperm abnormalities, mainly reflex midpieces, as reported by Kimberling and Parsons [9]. As a consequence, we did not take into account this anomaly for the calculation of the percentage of normal spermatozoa.

The point-score system that we used to classify the rams was adapted from Ley et al. [7]. The clinical and seminal criteria and/or the thresholds selected were close, but slightly different, from those of Ley et al. [7], Kimberling and Parsons [9] and Van Metre et al. [10].

In rams originated from *B. ovis* infected flocks, only 26 % and 37 % of rams had good clinical or seminal score respectively. This result, lower than the 70 % of rams classified as questionable or unsatisfactory breeders in previous studies [7, 10], can be explained by the high prevalence of *B. ovis* infection in our rams population. Furthermore, the low rate of good clinical and seminal scores in the studied population might be due not only to *B. ovis* infection but also to other infectious or non-infectious causes of orchi-epididymitis, as reported previously [2]. A significant relationship was observed between clinical and seminal scores, but this was primarily explained by clinical alterations, not by differences in SC and age. Indeed, SC may be related to genital abnormalities but also to several factors, such as age, breed, season, testicular tone, nutrition, parasite and other concurrent disease status [11].

The comparison of the serological results obtained by the different methods (CFT and I-ELISA) confirms the higher sensibility of the I-ELISA compared to CFT, already reported by Vigliocco et al. [12], Praud et al. [8] and Ridler et al. [13]; moreover, I-ELISA can detect infected rams earlier than CFT [11].

In a recent study, Ridler et al. [13] considered that a “suspicious” result is a disadvantage of the *B. ovis* I-ELISA and concluded that rams presenting such a result should be isolated and re-tested 2 to 4 weeks later.

Previously, Praud et al. [8] considered that the use of one or two thresholds can depend on the strategy of control of the disease. In an *a priori* healthy area, a high threshold limits the risk of “false positive” results but, at the opposite, in an infected area, a low threshold increases the diagnostic sensitivity; thus, according to the situations and strategic choices, the class of “doubtful” animals can be grouped with that of “healthy” or “infected”. Under the conditions of our study, I-ELISA-45, with a unique threshold, as proposed by Praud et al. [6], was more sensitive than CFT to detect rams carrying lesions and producing semen of poor quality.

Almost all rams shedding *B. ovis* in their semen were seropositive (mainly in I-ELISA) but approximately 25 % of the seropositive rams were culture-negative for *B. ovis*. This can be due to the intermittent excretion of *B. ovis*, as previously reported [3] and, also, to the fact that the rams were sampled only once [1, 13].

According to previous reports [4, 13, 14], we observed a significant relationship between culture-positive results (*i.e.* excretion of *B. ovis* in semen) and genital alterations or poor semen quality. Although significantly linked, seropositivity and genital tract alterations were not systematically associated, as previously described by Blasco [3], Ridler and West [15] and Van Metre et al. [10]. Moreover, approximately one third of *B. ovis*-shedding

| Pathological changes of spermatozoa | Mean ± SD* | Median* | Maximum* |
|-----------------------------------|-----------|---------|----------|
| Detached head                     | 10.8 ± 14.6| 4.1     | 80.7     |
| Elongated head                    | 0.1 ± 0.4  | 0       | 4.0      |
| Proximal droplet                  | 0.4 ± 1.6  | 0       | 19.5     |
| Distal droplet                    | 0.1 ± 0.6  | 0       | 6.1      |
| Isolated tail                     | 5.2 ± 7.3  | 1.6     | 38.5     |
| Bent tail                         | 7.6 ± 7.2  | 5.6     | 51.5     |
| Coiled tail                       | 3.8 ± 4.8  | 2.0     | 28.5     |
| Broken tail                       | 1.2 ± 1.5  | 0.5     | 8.0      |

*% |

Table 7 Relationship between I-ELISA results and seminal score

| Seminal score | Poor | Fair | Good | Total |
|---------------|------|------|------|-------|
| **I-ELISA (30-60 %)** |       |      |      |       |
| Negative      | 6    | 52   | 57   | 95    |
| Doubtful      | 5    | 6    | 11   | 22    |
| Positive      | 60   | 34   | 18   | 112   |
| Total         | 66   | 71   | 81   | 218   |

| **I-ELISA (45 %)** | | | | |
| Negative | 6 | 34 | 61 | 101 |
| Positive | 60 | 37 | 20 | 117 |
| Total | 66 | 71 | 81 | 218 |
rams presented clinical alterations [15], suggesting that B. ovis isolation in semen could be more precocious than the detection of epididymitis, as reported by Ridler et al. [13]. Furthermore, clinical alterations may be undetectable in the presence of active excretion of B. ovis and serological positive results, especially in chronically infected rams, as described previously by Worthington et al. [16]. In this study, 73 rams showed detectable lesions but 89 gave culture positive results while 81 and 117 gave positive results in CFT and I-ELISA respectively.

**Conclusion**

This study performed in B. ovis infected flocks of the “PACA” region, confirmed the findings of various previous field or experimental studies. As regards eradication of B. ovis infection, the following recommendations could be proposed: in areas with a low-to-medium prevalence, the eradication of CE is possible by using a test-and-slaughter approach, incorporating compounding tests to increase the overall testing sensitivity and reduce the likelihood of false negatives, as proposed by Blasco [3], Ridler and West [15] Praud et al. [8] and Ridler et al. [13]. Such an approach might include a combination of serological (I-ELISA and CFT) and auxiliary tests (genital palpation and, when possible, semen culture). This approach, which must be applied before each breeding season as well as before any ram introduction in free flocks, might allow for more rapid identification and removal of chronic shedders. In areas with a high prevalence, this strategy might be economically unsustainable. Despite recently developed B. ovis mutant attenuated strains gave promising protecting results in mice [17, 18], the B. melitensis Rev.1 vaccine remains up to now the only vaccine with proven efficacy against B. ovis infection in rams [2]; therefore, Rev.1 vaccination would be certainly the most economical and practicable tool for midterm control in such situations [3], in particular in areas where export of rams or ram semen is not an economical priority and where the official brucellosis-free status is not a short-term goal.

**Abbreviations**

B. ovis: Brucella ovis; PACA: Provence-Alpes-Côte d’Azur; BCS: Body condition score; SC: Scrotal circumference; CE: Contagious epididymitis; CFT: Complement fixation test; I-ELISA: Indirect enzyme-linked immunosorbent assay.

**Competing interests**

The authors declare that they have no competing interests.

**Authors’ contributions**

NPH, JLC, LE, MM, MP, AS and CT performed clinical examination of rams, semen collection and blood sampling for serology. XB, AS and CT performed semen evaluation. BGB directed the laboratory test performance. XB, NPH, FL and MP performed data analyses. NPH, XB and BGB drafted the manuscript. NPH, XB, JLC, MM, MP and BGB contributed to the conception and design of the whole study and revised the manuscript. All the authors approved the final manuscript.

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**Table 8** Relationship between bacteriology and clinical or seminal score

| B. ovis culture results (n = 198) | Clinical score (n = 218) | Seminal score (n = 218) |
|----------------------------------|------------------------|------------------------|
|                                  | Poor | Fair | Good | Poor | Fair | Good |
|----------------------------------|-----|-----|-----|-----|-----|-----|
| Negative                         | No. | 22  | 54  | 33  | 17  | 33  |
|                                  | (%) | (10.1) | (24.8) | (15.1) | (7.8) | (15.1) |
| Positive                         | No. | 29  | 42  | 18  | 43  | 34  |
|                                  | (%) | (13.3) | (19.3) | (8.3) | (19.7) | (15.6) |
| Contaminated or not tested*      | No. | 5   | 9   | 6   | 6   | 4   |
|                                  | (%) | (2.3) | (4.1) | (2.8) | (2.8) | (1.8) |

*16 semen culture contaminated, 4 samples not tested
**Fisher exact test

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**Table 9** Relationship between semen bacteriology and serology

| CFT Serological results | B. ovis culture results |
|-------------------------|-------------------------|
|                         | Negative | Positive |
|                         | No. (%)  | No. (%)  |
| (n = 197)               |          |          |
|Negative                 | 96       | (48.7)   | 25       | (12.7)   |
|Positive                 | 13       | (6.6)    | 63       | (32.0)   |
|K = 0.597**              |          |          |
|I-ELISA (45 %)           | Negative | 84       | (42.4)   | 4        | (2.0)   |
|                         |          |          |          |          |
|                         | Positive | 25       | (12.6)   | 85       | (42.9)   |

*Fisher exact test
**K: Cohen’s kappa coefficient
Acknowledgements
We thank the stock-breeders, PROVALP, Nicolas CORBOZ and the staff of the FRGDS-PACA for their active participation to this study. Moulay-Alla CHEFRA, Yannick CORDE, Antoine DRAPEAU and Gilles LE CARROU (Anses) are also duly acknowledged for their expert technical assistance.

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Received: 8 January 2015 Accepted: 18 May 2015
Published online: 30 May 2015

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