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24 **Ovine placental explants: a new ex vivo model to study host–pathogen interactions in reproductive**
25 **pathogens.**

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Abstract

Reproductive failure is one of the main performance constraints in ruminant livestock. Transmissible agents such as *Toxoplasma gondii* and *Neospora caninum* are commonly involved in the occurrence of abortion in ruminants, but little is known about the mechanisms involved. While in vivo models are optimal for the study of abortion pathogenesis, they have a high economic cost and come with ethical concerns. Unfortunately, alternative in vitro models fail to replicate the complex in vivo placental structure. To overcome the limitations of currently available models, we developed an ex vivo model based on the cultivation of fresh and cryopreserved sheep placental explants, enabling the biobanking of tissues. Reproducible and simple markers of tissue integrity (histology, RNA concentrations), viability (resazurin reduction), and functionality (synthesis of steroid hormones) were also investigated, allowing a clear quality assessment of the model. This work shows that, similar to fresh explants, tissues cryopreserved in ethylene glycol using slow freezing rates maintain not only their structure and function but also their receptivity to *T. gondii* and *N. caninum* infection. In addition, the findings demonstrate that explant lifespan is mainly limited by the culture method, with protocols requiring improvements to extend it beyond 2 days. These findings suggest that cryopreserved tissues can be exploited to study the initial host–pathogen interactions taking place in the placenta, thus deepening the knowledge of the specific mechanisms that trigger reproductive failure in sheep. Importantly, this work paves the way for the development of similar models in related species and contributes to the reduction of experimental animal use in the future.

Keywords: Ovine, placental explant, ex vivo model, reproductive failure, *Neospora caninum*, *Toxoplasma gondii*

71 Highlights

- 72 • An ex vivo model of the ovine placenta using fresh and cryopreserved tissue was developed.
- 73 • Fresh and thawed explants were found to behave similarly and might be useful for studying host–
74 parasite interactions.
- 75 • Several markers of explant integrity, viability, and functionality are defined and characterised.
- 76 • The model presented has the potential to be applied to several reproductive pathogens and even
77 to other related host species.

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1. Introduction

The performance of sheep flocks is heavily influenced by reproductive failure, which is currently considered one of the main economic burdens of ovine production systems. The aetiology of such failure varies and includes alterations in the oestrous cycle, nutritional deficiencies, environmental toxicity and infections from pathogens, some of which have zoonotic potential. In particular, abortions are one of the major constraints for optimal sheep breeding operations, with a high percentage (~50%) of the abortions being caused by transmissible agents displaying placental tropism. These include the apicomplexan parasites *Toxoplasma gondii* and *Neospora caninum*, the bacteria *Brucella melitensis*, *Chlamydia abortus* and *Coxiella burnetii*, and the Border disease virus, among others [1]. In recent years, extensive efforts have been made to describe the specific mechanisms involved in abortions caused by transmitted pathogens. In most cases, the abortions seem to be a consequence of the placental damage caused by replication of the pathogenic agent, resulting in an alteration of key placental functions (e.g., nutrient and oxygen exchange and hormone synthesis). This is confounded by the ability of some of these agents to modify and evade host immune responses, favouring their replication and allowing the chronification of the infection [2-5]. However, little is known about the specific molecular mechanisms triggering such abortions and, consequently, the means to prevent them.

The placenta is an ephemeral but specialised organ finely tailored to provide continuous support to the foetus during pregnancy in mammals. Although its role is similar for all species, covering the function of the respiratory, digestive, excretory, and endocrine systems, it shows incredible morphological and histological diversity among different species [6]. Notably, sheep have chorioallantoic and cotyledonary placentae, in contrast to the haemotrichorial and discoid placentae of mice [7]. Such differences are also reflected in their physiology and immune responses, limiting to some extent the usefulness of mice as animal models for studying the transmissible causes of

119 abortion in ruminants [8]. Considering the higher costs of working with pregnant sheep and also the
120 longer experimental periods and the ethical issues associated with their use, some research groups
121 have opted to develop simplified cell culture models based on the major population of the foetal
122 placenta: the trophoblast. Several studies have demonstrated the usefulness of the ovine trophoblast
123 AH-1 cell line for the study of abortifacient agents such as *C. abortus*, *Waddlia chondrophila*, *N.*
124 *caninum* and *T. gondii* [9-12]. Nevertheless, similar experiments performed with bovine-derived cells
125 isolated from the foetal and maternal placenta (trophoblasts and caruncular cells, respectively) have
126 failed to mimic foetal–maternal interactions and the local immune modulations described in vivo
127 during experimental *N. caninum* infections. This likely reflects the scarce representation of the
128 placental cell populations in these systems [13-15]. Consequently, the use of placental explants could
129 represent an intermediate approach to overcome the limitations of both in vitro and in vivo models,
130 as these explants retain the placental architecture and functions for a short period of time under
131 artificial culture conditions [16].

132 Previous works have shown that ex vivo systems based on placental explants are useful for the study
133 of the pathogenesis of *Brucella abortus*, *Listeria monocytogenes* and *Listeria ivanovii* in cattle [17-20]
134 but also for describing the effect of *Trypanosoma cruzi* and *T. gondii* infections in sheep [21,22].
135 However, a bottleneck for such approaches is the short lifespan and limited availability of tissues
136 suitable for obtaining explants [23]. Explant cryopreservation could represent a convenient tool to
137 circumvent such drawbacks, and has, in fact, proven successful with human and canine placental
138 explants [24,25]. Consequently, the objective of this work was to develop an ex vivo model based on
139 cryopreserved sheep placental explants. Placental tissues from healthy ewes were collected,
140 subjected to different cryopreservation methods, and characterised before and after thawing over
141 different time points. Then, we (i) assessed in detail different markers related to tissue architecture,
142 viability, and functionality to study the effect of culture and cryopreservation on placental explants

143 and (ii) evaluated their suitability to serve as a quality control for the model. We also investigated the
144 susceptibility of fresh and thawed explants to *N. caninum* and *T. gondii* infection. This work paves the
145 way for the generation of tissue biobanks and their exploitation to deepen our understanding of the
146 pathogenic mechanisms of infectious reproductive failure in sheep and the findings reveal promising
147 prospects for related species, such as cattle and goats.

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2. Materials and methods

2.1. Ethics approval and animals

All protocols involving animals followed current Spanish and European regulations (R.D. 53/2013; Council Directive, 2010/63/EU) and were approved by the Animal Welfare Committee of the Community of Madrid, Spain (PROEX 65.6/20, 2 ewes). To reduce the number of animals used, most tissues were obtained from ewes that were culled for other purposes in parallel experiments also approved by the Animal Welfare Committee of the Community of Madrid, Spain (PROEX 68.0/20, 8 ewes). All animals ($n = 10$) were strictly handled under good clinical practices to minimise potential suffering.

Animal selection was performed as described in previous research [26]. In brief, Rasa Aragonesa breed ewes were purchased from a commercial flock after verifying their seronegativity for *T. gondii*, *N. caninum*, Border disease virus, Schmallenberg virus, *Coxiella burnetii* and *Chlamydia abortus* by commercial or in-house ELISAs. After oestrus synchronisation using progestogens and pregnant mare serum gonadotrophin, the ewes were mated with rams for 2 days. Pregnancy was confirmed by ultrasound scanning on Day 40 after mating. Subsequently, pregnant ewes were transferred and housed at the Clinical Veterinary Hospital facilities (Complutense University of Madrid, Spain) until the day of culling, which took place around Day 100 of gestation. Foetal viability was confirmed by ultrasound scanning before culling.

2.2. Isolation of ovine placental explants

Pregnant ewes were intravenously sedated with xylazine (Calier laboratories, Spain) and later euthanised by an overdose of embutramide and mebezonium iodide (T61, MSD, Spain). Necropsies were performed at the Clinical Veterinary Hospital facilities (Complutense University of Madrid, Spain) following standard protocols. Immediately after euthanasia, placentomes with diameters

191 between 3 and 6 cm were randomly selected, dissected from the placenta using sterile material, and
192 transported to the laboratory in an ice-cold sterile washing solution (phosphate buffered saline -PBS-
193 supplemented with 100 IU/ml of penicillin, 100 µg/ml of streptomycin, 0.25 µg/ml of fungizone and
194 100 µg/ml of L-glutamine, all from Gibco, UK). The placentomes were longitudinally cut into 2-3 mm-
195 thick slices using sterile skin graft blades (Swann Morton, UK) inside a laminar flow cabinet.
196 Afterwards, the slices were transversely cut into 60-80 mg sections containing both the maternal and
197 foetal parts of the placentome using switch-blade shears (Feather, Japan) (**Fig. 1a**). Once obtained,
198 the explants were maintained on ice-cold sterile washing solution for less than 30 minutes and then
199 washed 10 times with ice-cold washing solution before cultivation or cryopreservation.

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201 2.3. Experimental design

202 As a first approach, we tested the performance of four different cryopreservation methods by
203 comparing the behaviour of fresh and thawed explants in terms of tissue damage (histology, see
204 Section 2.5) and viability (LDH test, see Section 2.6.1) after 1 to 3 days in culture (**Fig. 1b**). For this
205 purpose, a group of freshly obtained explants from two ewes (OV1 and OV2) were cultured in 24-
206 well plates (12 explants/sheep/time-point, see Section 2.4). In parallel, another group of fresh
207 explants was subjected to cryopreservation (**Fig. 1b**). For this, explants were incubated for 20 minutes
208 with two different cryoprotectants consisting of foetal bovine serum (FBS, South America origin,
209 Gibco, UK) supplemented with 3 M dimethyl sulfoxide (DMSO, Sigma–Aldrich, MA, USA; OV1) or FBS
210 supplemented with 2 M ethylene glycol (EG, Merck Chemicals, Germany; OV2). In both cases, two
211 different cryopreservation speeds were tested: snap freezing in liquid nitrogen and a slow-controlled
212 freezing rate. For the latter, Mr. Frosty® containers (Thermo Fisher Scientific, MA, USA) were used
213 and maintained at -80 °C for 24 h; subsequently, tubes were transferred to liquid nitrogen. To analyse
214 the effect of the cryopreservant in the tissue, three explants were fixed in 10% buffered formalin

215 immediately after incubation with each of them and before cryopreservation. Four weeks after
216 cryopreservation, the explants were thawed in a 37 °C water bath, washed 10 times with washing
217 solution (see Section 2.2), transferred to 24-well plates and cultured with protocols identical to those
218 used for the fresh explants (12 explants/sheep/time-point, see Section 2.4).

219 Based on the results obtained in the first test, in consecutive experiments, a similar approach was
220 followed to compare fresh and thawed explants using placentomes from 6 ewes (OV3-OV8) but using
221 the best cryopreservation method, culturing the explants from 5 h to 2 days, and expanding the
222 parameters measured in all cultured explants (12 explants per sheep for each time point and for each
223 condition, **Fig. 1c**). In this manner, tissue viability (resazurin and XTT tests), integrity (damage scores
224 based on histology, *KRT8* expression) and functionality (*PLAC-1* expression, synthesis of steroid
225 hormones) were studied as indicated in the sections below.

226 Based on works that have described the regeneration of placental explants after 2-4 days in culture
227 [27,28], in subsequent trials, we extended the explant culture up to 9 days using placentomes from 2
228 ewes (OV9 and OV10) to measure tissue viability (resazurin test), integrity, and functionality, as
229 detailed in the sections below (12 explants per sheep for each time point and for each condition, **Fig.**
230 **1c**). To increase the number of replicates and corroborate the obtained results, cryopreserved
231 explants from OV5 to OV9 were thawed, cultured for up to 9 days, and analysed (8-12 explants per
232 sheep for each time point and for each condition).

233 To ensure the reproducibility of the model, all parameters tested in each experimental design and at
234 each time point were measured in at least two different ewes using triplicate wells whenever
235 possible. The highest number of biological replicates was performed in explants cultured from 5 h to
236 2 days (6 ewes on average), while the lowest number was performed in explants cultured from 3 to
237 9 days (2 ewes on average).

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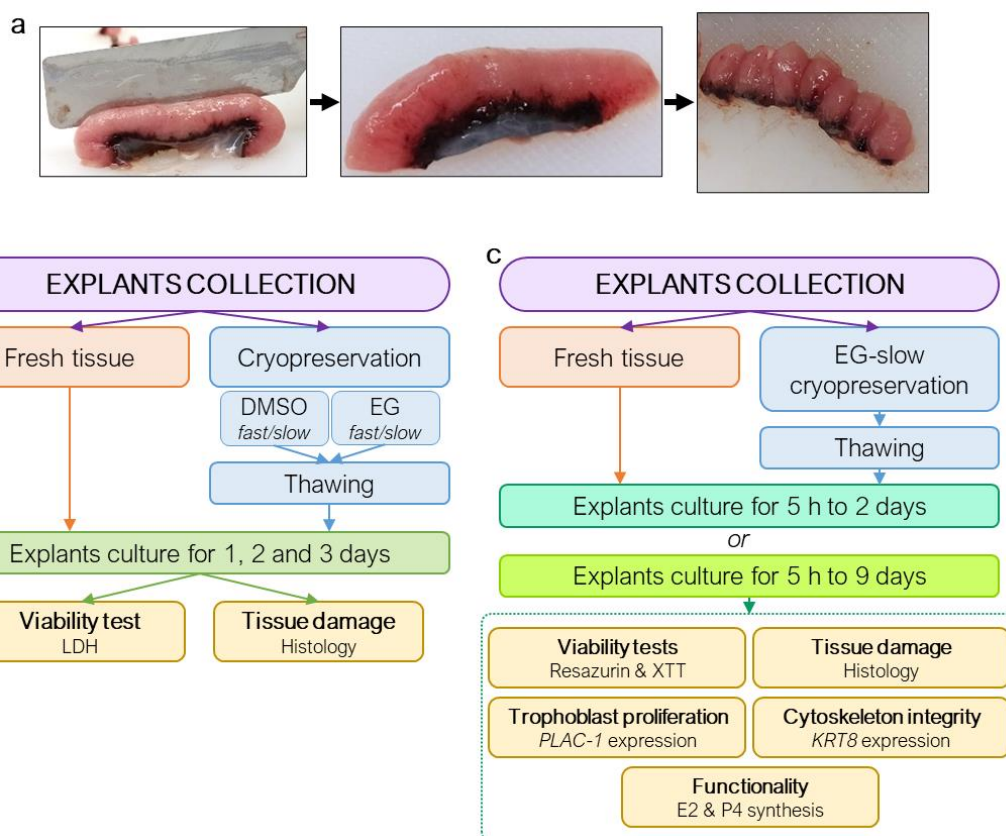


Fig. 1. Collection of ovine placental explants and experimental design. **a:** Placentomes were cut longitudinally and transversely into 60-80 mg sections containing both the maternal and foetal components. **b:** Workflow diagram followed by explants obtained from pregnant ewes to determine the best cryopreservation method. **c:** Workflow diagram followed by explants obtained from pregnant ewes to further characterise the differences between fresh and EG-slow-cryopreserved explants cultured up to 2 days or up to 9 days. DMSO: dimethyl sulfoxide; EG: ethylene glycol; fast/slow: speed at which freezing was performed.

2.4. Culture of fresh and thawed placental explants

Placental explants were individually allocated in 24-well plates previously filled with 2 mL of M-199 culture media without phenol red (Gibco, UK) (phenol red stimulates oestrogenic receptors)

252 supplemented with 20 mM HEPES (HyClone, Cytiva, MA, USA), 20% FBS (South America origin, Gibco,
253 UK), and an antibiotic-antimycotic mixture (100 UI/ml penicillin, 100 µg/ml streptomycin and 0.25
254 µg/ml Fungizone from Gibco, UK). The explants were then incubated from 5 h to 9 days at 37 °C and
255 5% CO₂. The culture media was replaced on a daily basis. In the absence of indicators in the media,
256 the pH was regularly measured at each time-point using pH indicator strips (Fisherbrand, Thermo
257 Fisher Scientific, MA, USA).

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259 2.5. Histological analyses, damage scoring, and immunohistochemical staining

260 Histological studies were considered our gold-standard technique for assessing the extent of tissue
261 damage in all explants. These were conducted in fresh and thawed tissues collected from all ewes at
262 all the time points of the cultures tested (5 h to 9 days). For each placental condition (fresh or
263 thawed), triplicate explants were fixed in buffered formalin for 5 days and subsequently processed
264 for paraffin embedding and haematoxylin-eosin staining for histological evaluation.

265 The tissue sections were examined in a blinded manner, and the damage to the tissue was scored
266 from 0 to 3 according to the following criteria (**Fig. 2**). A **score of 0** indicated the absence of any
267 evident or significant histological changes. A **score of 1** indicated the presence of a layer of sloughed
268 epithelial cells dispersed among cellular debris, as well as retraction of cotyledonary and caruncular
269 villi, isolated apoptosis of epithelial cells, and/or mild autolysis. A **score of 2** indicated evident loss of
270 the epithelial layer in both cotyledonary and caruncular villi, with the latter being shrunken and
271 eosinophilic; frequent apoptosis of epithelial cells was also noted, along with loss of defined borders
272 in the cells that were still attached to the villi. Finally, a **score of 3** indicated generalised autolysis of
273 tissue with evident apoptosis, lack of nuclear definition in most epithelial cells, and/or
274 indistinguishable cell borders. After evaluation of single sections from three different explants, a
275 single agreement score was assigned for each placental condition tested following the

276 aforementioned criteria (**Fig. 2**). In general, the appearance of the tissue was generally homogeneous
277 within the same sample and among the replicates from the same animal, although variations were
278 occasionally observed. In cases where differences were present between replicates or within the
279 same sample, the highest score was assigned.
280 Immunohistochemical labelling of *N. caninum* in tissue sections was also performed as described
281 previously [29].

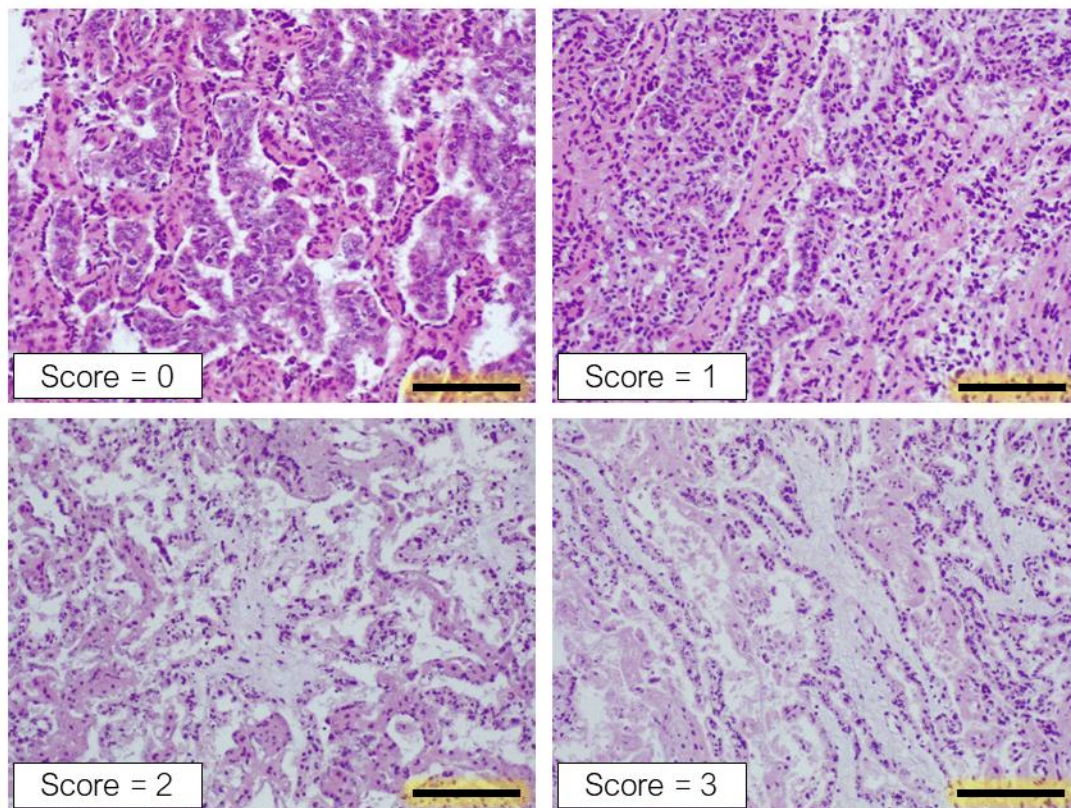


Fig. 2. Representative microscopy images of the tissue damage scores of placental explants cultured ex vivo. Haematoxylin-eosin staining, 10x magnification. Bars: 100 µm.

289 2.6. Viability assays

290 2.6.1. Lactate dehydrogenase (LDH) assay

291 As an abundant intracellular enzyme, extracellular detection of LDH is routinely used as an indirect
292 measure to study the extent of breakage of cellular membranes. To assess the damage to placental
293 explants due to cultivation and/or freezing, we quantified LDH levels in culture supernatants of fresh
294 and thawed explants obtained from two sheep (OV1 and 2). A total of six supernatants per animal
295 and placental condition were collected and kept at -80 °C until further analysis. The LDH levels were
296 quantified using the CyQUANT™ LDH Cytotoxicity Assay Kit (Thermo Fisher Scientific, MA, USA)
297 according to the manufacturer's instructions. All samples were diluted 1:20 in fresh culture medium
298 and measured using a Synergy H1 microplate reader (Biotek, 490 nm).

299 2.6.2. XTT assay

300 XTT assays were used to detect the presence of metabolic activity at different culture time points on
301 fresh and thawed explants in duplicate or triplicate. For this purpose, we used the Cell Proliferation
302 Kit II (Sigma-Aldrich, MA, USA) following the manufacturer's instructions. The explants were
303 incubated with the reagent for 18 h at 37 °C and 5% CO₂, and after incubation, 200 µL from each well
304 was transferred to an ELISA microplate (Maxisorp®, Nunc, Denmark) and measured using a Synergy
305 H1 microplate reader (450 nm, using a wavelength of 603 nm as reference; BioTek, VT, USA). A
306 background control (culture medium) and a negative control (nonviable explants previously fixed in
307 4% paraformaldehyde for 30 min) were included in all the assays.

308 2.6.3. Resazurin assay

309 Resazurin assays were also used to detect the occurrence of metabolic activity at different culture
310 time points on fresh and thawed explants in triplicate. For this purpose, 0.2 µm-filtered resazurin
311 (0.25 mg/mL, Acros Organics, Belgium) was added to each well at the pertinent time points and
312 incubated for 18 h at 37 °C and 5% CO₂. After incubation, 200 µL from each well was transferred to

313 an ELISA microplate (Maxisorp®, Nunc, Denmark), and the optical density was measured using a
314 Synergy H1 microplate reader (570 nm, using a wavelength of 600 nm as a reference; BioTek, VT,
315 USA). Background and negative controls were included in all assays as detailed for the XTT assay
316 (Section 2.6.2).

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318 2.7. Quantification of cell proliferation and cytoskeleton integrity by RT–qPCR

319 Total RNA was extracted using the Maxwell® 16 LEV simplyRNA Purification Kit (Promega, WI, USA)
320 following the manufacturer's recommendations. For this purpose, 20 mg from each explant (two or
321 three per condition tested) was homogenised in the provided buffer using a Politron® PT1600E
322 homogeniser (Kinematica AG, Switzerland). RNA quality was assessed by the visualisation of the 18S
323 and 28S ribosomal fragments after electrophoresis on bleach agarose gels (0.8% agarose, 0.05%
324 sodium hypochlorite, Sigma–Aldrich, MA, USA) and by spectrophotometry (NanoPhotometer®,
325 Implen, Germany). Reverse transcription to complementary DNA (cDNA) was carried out using the
326 SuperScript® VILO™ cDNA Synthesis Kit (Thermo Fisher Scientific, MA, USA) using a maximum of 2.5
327 µg of RNA per reaction.

328 Quantitative reverse transcription PCR (RT–qPCR) was employed to quantify the mRNA expression
329 levels of *PLAC-1*, a marker of trophoblast proliferation, *KRT8*, a keratin protein of the villi cytoskeleton
330 used as a marker of structural integrity, and *β-actin*, the housekeeping gene, using the primers
331 detailed in **Table 1** [30]. For each gene, a seven-point standard curve based on tenfold serial dilutions
332 of PCR products was included in each batch of amplifications using glycogen as a carrier (33 µg/mL,
333 Thermo Fisher Scientific, MA, USA). The primers used for the generation of standard curves are
334 detailed in **Table 1**. All cDNA samples were diluted 1:5 in nuclease-free water prior to analysis. The
335 reaction mix contained 12.5 µL of Power SYBR® Green PCR Master Mix (Applied Biosystems, MA,
336 USA), 10 pmol of each primer and 5 µL of the diluted cDNA. The reaction was performed in an ABI

337 7500 Fast Real-Time PCR System (Applied Biosystems, MA, USA). mRNA expression levels for each
338 target were normalised by the $2^{-\Delta Ct}$ method [31].

339

340 **Table 1. Primers used for the generation of standard curves and RT-qPCR assays to quantify**
341 ***PLAC-1* and *KRT8* expression in cultured explants. F: forward, R: reverse.**

342

Gene	GenBank accession number	Primer sequence for standard curves (5'-3')	Primer sequence for RT-qPCR (5'-3')
<i>β-actin</i>	NM_001009784.3	F: AGACCGCGTCCGCCCGCCAG R: GTACATGGCAGGGGTGTTGA	F: ACACCGCAACCAGTTCGCCAT R: GTCAGGATGCCTCTCTTGCT
<i>PLAC-1</i>	XM_012107306.3	F: GCCTGAACTCCTCAAGAGATAGC R: GCTAAGTTCCTGGAGCAGGAC	F: CCGGACAAAATCCAATGACTGT R: AACCCAGGCCCAAGTGTAATCA
<i>KRT8</i>	XM_012174208.3	F: CTGATGAGATCAACTTCTACAGGC R: CTGCTTGGCATTCTCAGAG	F: ATGGACAACAACCGCAACCT R: CCAGTGTCTGCAGCTCCTCAT

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344 2.8. Functionality assays: oestradiol and progesterone synthesis

345 To assess the functionality of fresh and thawed explants at the different time points studied, the
346 levels of 17-β-oestradiol (E2) and 17-OH-progesterone (P4) released into the medium were
347 quantified in duplicate or triplicate using a commercial competitive ELISA test (E2-ELISA [KAP0621]
348 and 17OH-Progesterone ELISA [KAP1401], DiaSource, Belgium) following the manufacturer's
349 instructions. Blank controls (media supplemented with FBS) were included in all runs to determine
350 the basal levels of both hormones in the media.

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352 2.9. Explant susceptibility to *T. gondii* or *N. caninum* infection

353 Fresh and thawed explants were experimentally infected with *T. gondii* (Tg-PRU isolate) or *N.*
354 *caninum* (Nc-Spain7 isolate) to assess tissue receptivity to two relevant ruminant reproductive
355 pathogens. The parasites were propagated, harvested, and dosed following previous protocols [32].

356 Since the culture media used to grow the parasites differed from that used to culture the explants,
357 infected monolayers were washed three times with washing solution and replenished with
358 supplemented M-199 media before harvest. In a first trial, fresh explants were infected in triplicate
359 with 10^5 or 10^6 *T. gondii* or *N. caninum* tachyzoites for 1, 3, and 5 days. In a second trial, EG-thawed
360 explants were infected in triplicate with both parasites at the same time points, but only with the
361 higher tachyzoite dose (10^6). In both trials, media was replaced on a daily basis, and explant viability
362 was recorded at every collection time as detailed in Section 2.6.3. The infected explants were
363 individually collected in microcentrifuge tubes and immediately stored at -80 °C until DNA extraction,
364 which was performed using the DNeasy Blood & Tissue Kit (Qiagen, Germany). For that purpose, the
365 explants were thawed, individually supplemented with 180 µl of ATL buffer (Qiagen, Germany) per
366 25 mg of tissue and homogenised using a Politron® PT1600E homogeniser (Kinematica AG,
367 Switzerland). Only 25 mg (~180 µl) of tissue was subjected to DNA extraction. The parasite burden
368 was quantified by qPCR using previously described methods [33,34] and the Go Taq® qPCR Master
369 Mix (Promega, WI, USA). In both protocols, *β-actin* was used as a housekeeping gene, while the TgRE
370 (529-bp repeat element) and Nc-5 sequences were targeted for quantification of *T. gondii* and *N.*
371 *caninum*, respectively. All primers used for qPCR are listed in **Table 2**.

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Table 2. Primers used for qPCR assays to quantify *N. caninum* and *T. gondii* replication in infected explants. F: forward, R: reverse.

Gene	GenBank accession number	Primer sequence for qPCR (5'-3')
<i>β-actin</i>	NM_001009784.3	F: AGCGCAAGTACTCCGTGTG R: CGGACTCATCGTACTCCTGCTT
<i>Nc-5</i>	X84238.1	F: ACTGGAGGCACGCTGAACAC R: AACAATGCTTCGCAAGAGGAA
<i>TgRE</i>	AF146527.1	F: CACAGAAGGGACAGAAGT R: TCGCCTTCATCTACAGTC

2.10. Statistical analysis

Statistical analysis was performed using GraphPad Prism software, v.8 (Dotmatics, UK). Data normality was checked using the Shapiro–Wilk test prior to performing any further statistical analyses. Data fitting a normal distribution were analysed with one-way ANOVA coupled to Tukey's multiple comparison test. The rest of the data were analysed using the Kruskal–Wallis test coupled to Dunn's multiple comparison. Statistical significance for all analyses was established at $p < 0.05$.

3. Results

3.1. Selection of optimal cryopreservation methods for ovine placental explants

The performance of four cryopreservation protocols based on the use of DMSO and EG at two different freezing speeds was assessed through LDH-based viability assays and histological studies of fresh and thawed explants cultured from 1 to 3 days (**Fig. 1b**).

The LDH analyses were performed on supernatants collected from fresh explants (OV1 and 2) and from thawed explants previously subjected to fast and slow freezing rates with DMSO (OV1) or EG (OV2). Overall, enzyme levels remained high under all conditions tested. No significant differences were observed between fresh explants obtained from different animals or between fresh and thawed explants obtained from the same animal, regardless of the speed at which cryopreservation was performed ($p > 0.05$, one-way ANOVA and Tukey's multiple comparison test) (**Fig. 3**). Under all experimental conditions, we observed a drop in LDH activity at 3 days of culture, but this was only significant for fresh-OV1 (Day 1 vs. Day 3; **Fig. 2a**), DMSO-Fast (Day 1 vs. Day 3; **Fig. 2b**), and EG-Slow (Day 1 vs. Day 3; **Fig. 2c**) ($p < 0.05$, one-way ANOVA and Tukey's multiple comparison test).

Given the absence of obvious differences in the LDH secretion levels between the fresh and thawed explants, the damage score of the tissue was used as a gold-standard criterion to determine the best cryopreservation method to be applied in subsequent trials. The histological analyses revealed that the mere incubation of fresh explants with DMSO induced certain levels of tissue alteration (score = 1), and this was not observed in explants incubated with EG (score = 0) (data not shown). After thawing, DMSO-treated explants showed worse tissue damage scores than those preserved with EG (**Fig. 4**). In addition, the use of slow freezing rates in EG-thawed explants resulted in lower damage scores (**Fig. 4**). Therefore, cryopreservation with EG at slow freezing rates was selected for subsequent experiments.

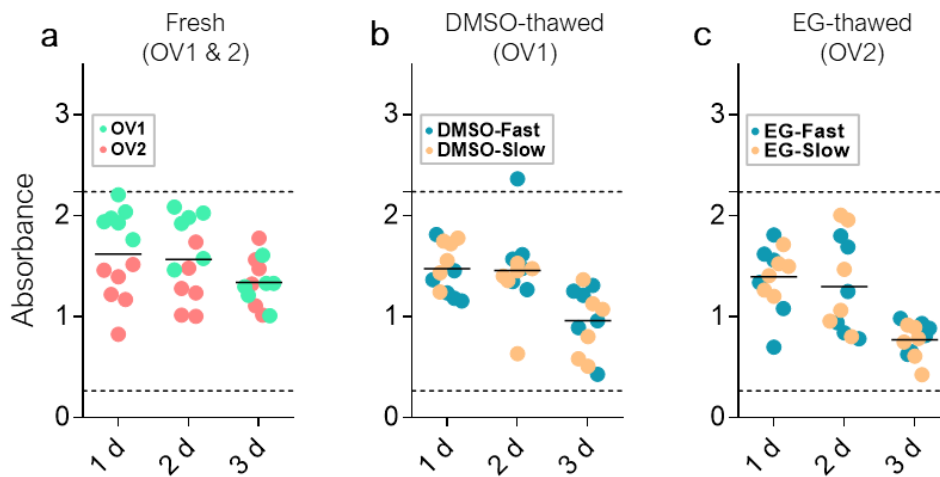
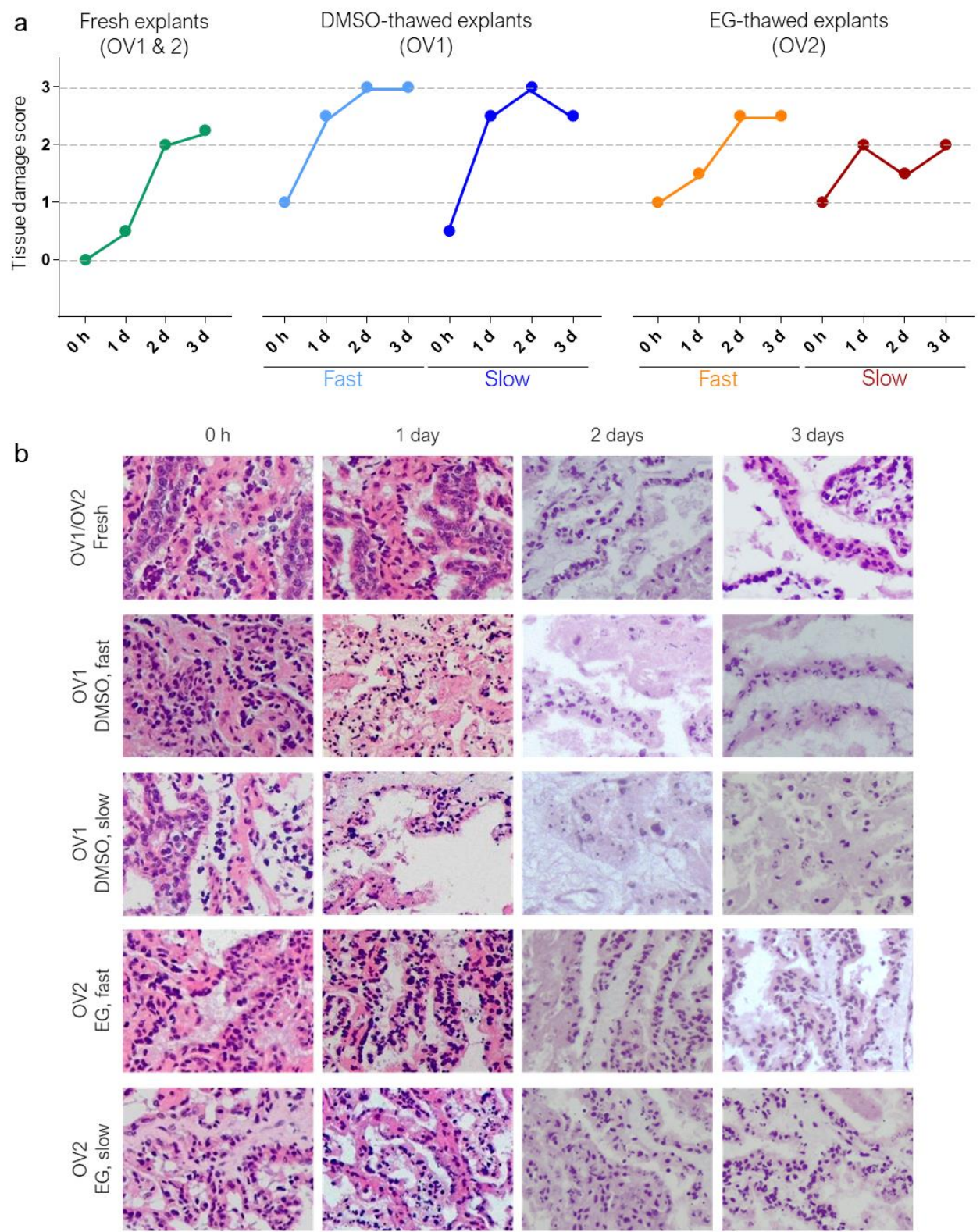


Fig. 3. LDH levels in culture supernatants of fresh and thawed explants. **a:** Fresh explants; **b:** DMSO-thawed explants; **c:** EG-thawed explants. In all graphs, the top dotted line represents the absorbance levels of the kit's positive control, while the bottom dotted line represents the LDH levels present in fresh culture media without explants. Each dot represents individual explants, and bars represent grand mean values.

3.2. Histological analyses and tissue damage scoring

Further histological analyses were performed to assess the effect of cultivation and EG-slow cryopreservation on the explants' tissue integrity in eight additional sheep using the tissue damage score defined in Section 2.5. In fresh explants, the structural integrity was intact during the first day of culture and displayed progressive degradation with time (**Fig. 5a**). In the EG-thawed explants, we observed a pattern of tissue degradation comparable to that observed in the fresh explants (**Fig. 5a**). When comparing the fresh and EG-thawed explants, the latter tended to display earlier signs of tissue degradation, although no significant differences were found between them ($p > 0.05$, Kruskal–Wallis test and Dunn's multiple comparison). These results suggest that both fresh and EG-thawed explants

440 might be suitable for performing ex vivo experiments lasting up to 1 to 2 days. Nevertheless, the lack
441 of signs of severe autolysis and loss of membrane integrity (score 3) until Days 3-7 of culture may
442 suggest that these assays could last for longer periods.



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Fig. 4. Tissue damage of fresh, DMSO-thawed and EG-thawed explants cultured for 1 to 3 days. a: Graphical representation of the tissue damage scores of fresh and thawed explants. Each dot represents the average score of three different explants cultured under the same conditions. **b:** Representative histological sections obtained from fresh, DMSO-thawed and EG-thawed explants. OV1/OV2: sheep identification; Fast/Slow: speed at which freezing was performed.

3.3. RNA integrity

Considering that RNA abundance can be exploited as an indirect marker of the transcriptomic activity of the explants, we took advantage of the extractions performed for RT-qPCR assays to analyse the effect that culture of fresh and EG-thawed explants had on the isolated concentrations of RNA (**Fig. 5b**). Under both conditions, and especially during the first 2 days of culture, we observed a significant progressive decrease in RNA concentrations as the culture time increased ($p < 0.05$, one-way ANOVA and Tukey's multiple comparison test). This decrease was more marked in EG-thawed explants, whose RNA concentrations were lower at 5 h of culture compared to those of fresh explants ($p < 0.05$, one-way ANOVA and Tukey's multiple comparison test). These findings mirrored the behaviour observed in terms of tissue damage, since RNA concentration decreased as tissue degradation increased (**Fig. 5**).

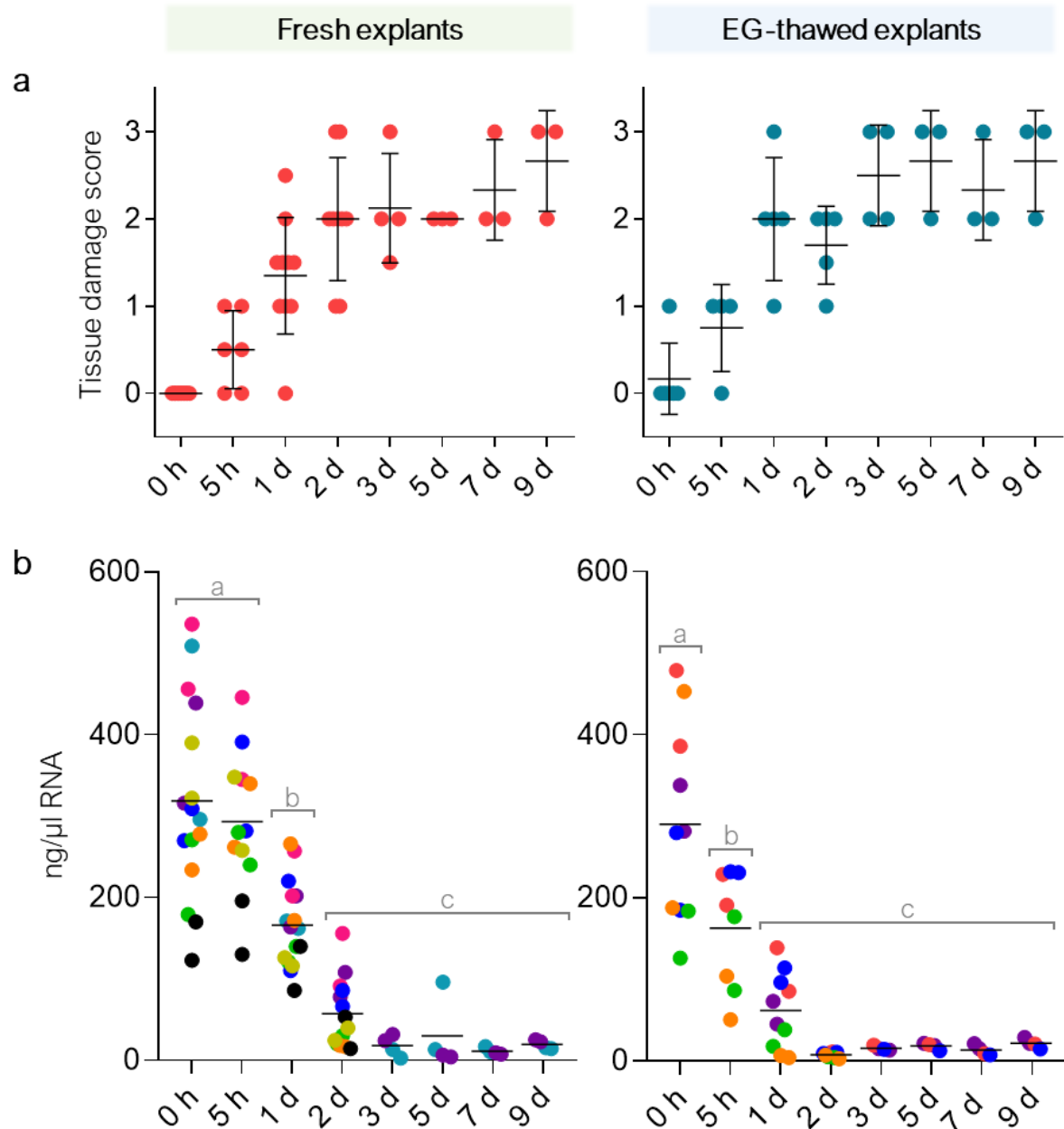


Fig. 5. Evolution of the tissue damage and RNA concentration of fresh and EG-thawed explants throughout the culture time. a: Tissue damage scores recorded from fresh (red dots) and thawed (blue dots) explants obtained from different sheep and cultured in vitro from 0 h to 9 days. The scores were analysed from an average of six (5 h – 2 days) or two (3–9 days) different ewes at each time point. Each dot represents the average score of three different explants obtained from the same sheep and cultured under the same conditions; bars represent the mean value and the standard

deviation. **b**: Evolution of RNA concentrations extracted from fresh (left graph) and EG-thawed (right graph) explants from different sheep and cultured in vitro from 0 hours (h) to 9 days (d). Each dot represents the RNA concentration of an individual explant; each colour represents different ewes; bars represent grand mean values; groups marked with different letters were significantly different ($p < 0.05$, one-way ANOVA and Tukey's multiple comparison test).

3.4. Viability assays

Since histology is a time-consuming approach, we explored alternative tests readily accessible in most labs. Considering the limitations of LDH assays in discerning the impact of cryopreservation on explants, in consecutive trials, we evaluated alternative methods for assessing explant viability. In this sense, we used XTT and resazurin assays to measure the metabolic activity of fresh and thawed explants cultured for different durations.

The XTT assays were run in duplicate or triplicate samples using explants obtained from four sheep. These tests corroborated the occurrence of metabolic activity from 5 h to 2 days of culture (**Fig. 6, top panel**). Experimental reproducibility between biological replicates was confirmed, as no significant differences were found between animals within each time point under all conditions tested ($p > 0.05$, one-way ANOVA and Tukey's multiple comparison test). Grouped analysis showed that the viability of fresh explants was significantly lower after 2 days of culture than after 5 h and 1 day of culture ($p < 0.05$, one-way ANOVA and Tukey's multiple comparison test). These differences were not observed in thawed explants ($p > 0.05$, one-way ANOVA and Tukey's multiple comparison test). The viability of EG-thawed explants at 5 h and 1 day of culture was significantly lower than that observed for fresh explants at the same time points ($p < 0.05$, one-way ANOVA and Tukey's multiple comparison test). However, the assay's threshold between fully viable (fresh explants cultured for up

492 to 5 h) and nonviable explants (paraformaldehyde-fixed explants) was very tight, preventing us from
493 drawing clear conclusions.

494 Considering the low sensitivity of the XTT assays, we focused our efforts on the resazurin tests, which
495 were run in triplicate using explants obtained from seven sheep and cultured for 5 h, 1 day, or 2 days.
496 All fresh and EG-thawed samples showed metabolic activity from 5 h to 2 days of culture (**Fig. 6,**
497 **bottom panel**). We also confirmed the experimental reproducibility between biological replicates, as
498 no significant differences were found between animals when they were compared within every time-
499 point and placental condition (fresh or EG-thawed) ($p > 0.05$, one-way ANOVA and Tukey's multiple
500 comparison test). In addition, no significant differences were observed between the tested time
501 points in fresh or EG-thawed explants ($p > 0.05$, one-way ANOVA and Tukey's multiple comparison
502 test).

503 Given the good results obtained with the resazurin tests, which largely correlated with what we
504 described using histology, in consecutive experiments, we decided to assess explant viability after 9
505 days of ex vivo culture. All fresh and EG-thawed samples showed metabolic activity from 3 to 9 days
506 of culture (**Fig. 6, bottom panel**). In addition, no significant differences were found between animals
507 when they were compared within every time point and placental condition (fresh or EG-thawed) (p
508 > 0.05 , one-way ANOVA and Tukey's multiple comparison test). The grouped analysis of fresh
509 explants cultured for 3 to 9 days did not show differences between the time points tested ($p > 0.05$,
510 one-way ANOVA and Tukey's multiple comparison test). Further analyses performed from 5 h to 9
511 days of culture demonstrated a slight decrease in explant viability after 2 days of culture compared
512 to that observed at 3, 5 and 7 days ($p < 0.05$, one-way ANOVA and Tukey's multiple comparison test)
513 (**Fig. 6, bottom panel**). In EG-thawed explants, a marked drop in viability was observed at 3 days of
514 culture compared to that measured at 5 h and 1, 2, 5, 7, and 9 days of culture ($p < 0.05$, one-way

ANOVA and Tukey's multiple comparison test). Remarkably, explant viability was markedly recovered from 5 days of culture onwards (**Fig. 6, bottom panel**). Apart from the drop in viability of EG-thawed explants on Day 3, no differences were found in the viability of fresh and thawed explants at equivalent culture durations ($p > 0.05$, one-way ANOVA and Tukey's multiple comparison test).

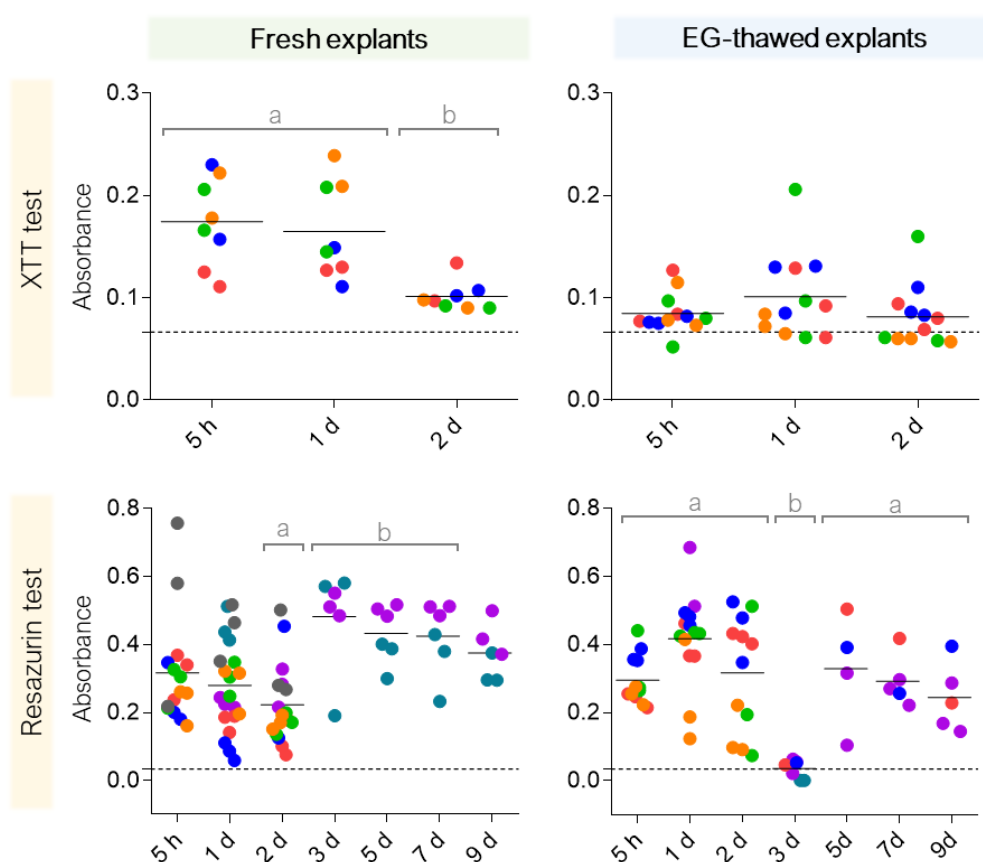


Fig. 6. Quantification of the metabolic activity of fresh and thawed explants using the XTT assay (top panel) and the resazurin assay (bottom panel) over different incubation times. The bottom dotted line represents the average absorbance of explants fixed with 4% paraformaldehyde and used as controls of nonviability. Each dot represents individual explants, each colour represents different ewes, and bars represent grand mean values. Groups marked with different letters were significantly different ($p < 0.05$, one-way ANOVA and Tukey's multiple comparison test). h: hour; d: day(s).

3.5. *PLAC-1* and *KRT8* mRNA expression

To further characterise fresh and thawed explants, we quantified the expression of *PLAC-1*, regarded as a marker of trophoblast proliferation, and *KRT8*, a cytoskeletal protein related to the structural integrity of placental villi. Due to the low amounts of RNA extracted from the explants, we only assessed the expression of both genes from samples obtained from 0 h to 2 days of culture (**Fig. 7**). In fresh explants, *PLAC-1* mRNA expression decreased gradually with the time of culture, being significantly low after 1 and 2 days ($p < 0.05$, Kruskal–Wallis test and Dunn’s multiple comparison). In EG-thawed explants, the expression of *PLAC-1* transcripts was similar from 0 h to 2 days of culture ($p > 0.05$, Kruskal–Wallis test and Dunn’s multiple comparison). When comparing equal time points, no differences in *PLAC-1* expression were found between fresh and EG-thawed explants ($p > 0.05$, Kruskal–Wallis test and Dunn’s multiple comparison).

KRT8 expression in fresh explants increased significantly after 2 days of culture ($p < 0.05$, Kruskal–Wallis test and Dunn’s multiple comparison). This contrasts with what was observed in EG-thawed explants, where a gradual decrease in *KRT8* expression was observed over time, with significantly low expression on Day 2 of culture ($p < 0.05$, Kruskal–Wallis test and Dunn’s multiple comparison). Comparison between fresh and EG-thawed explants at equivalent time points demonstrated that *KRT8* expression was significantly higher in fresh explants at Days 1 and 2 of culture ($p < 0.05$, Kruskal–Wallis test and Dunn’s multiple comparison).

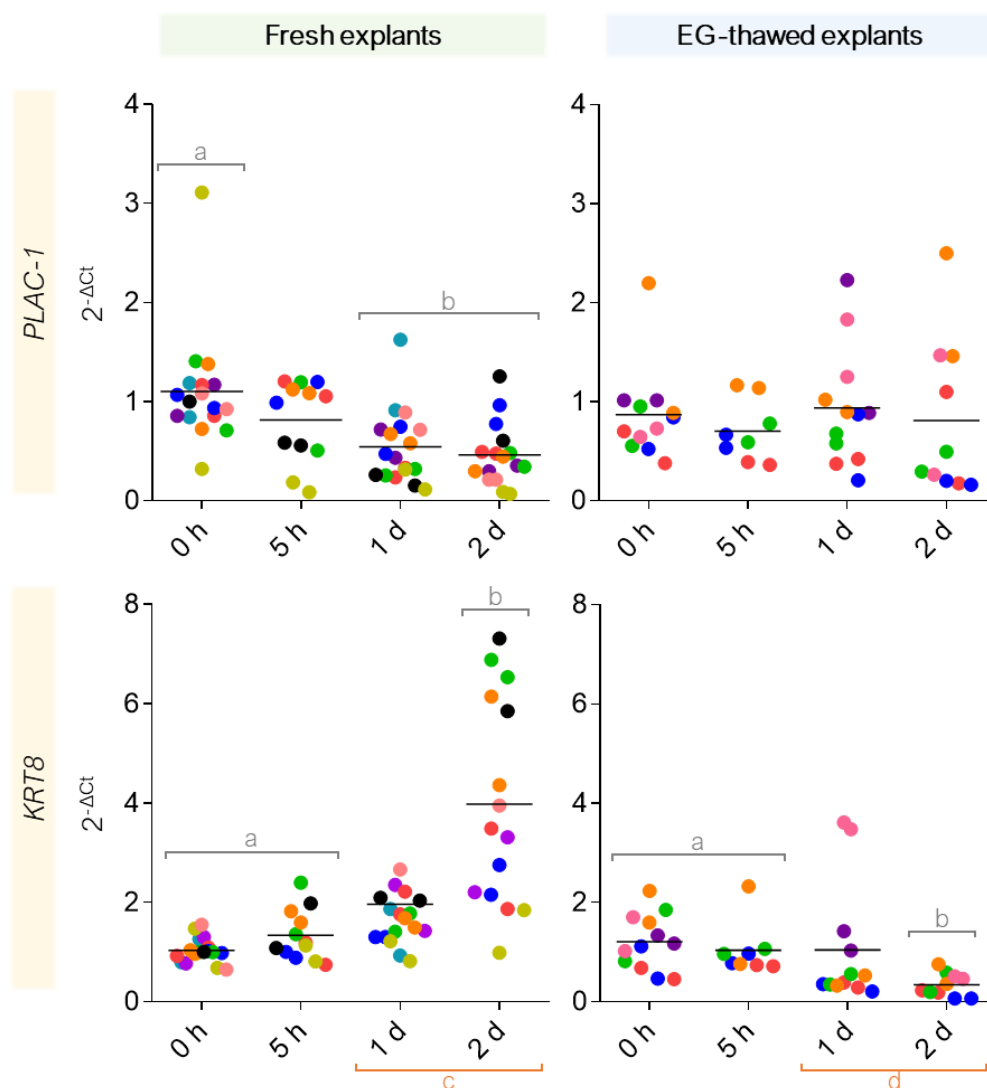


Fig. 7. *PLAC-1* and *KRT8* mRNA expression. Evolution of the abundance of *PLAC-1* and *KRT8* transcripts in fresh (left graph) and EG-thawed (right graph) explants cultured from 0 hours (h) to 2 days (d). Each dot represents the normalised expression value of an individual explant; each colour represents different ewes; bars represent grand mean values; groups marked with different letters were significantly different (*p* < 0.05, Kruskal–Wallis test and Dunn’s multiple comparison) within the same condition (grey letters) or between different conditions (orange letters).

3.6. Steroid hormone biosynthesis

Under physiological conditions, the placenta synthesises steroid hormones such as 17- β -oestradiol (E2) and 17-OH-progesterone (P4). Consequently, we quantified the explants' production of both hormones at different culture time points as an indirect measure of their functionality.

Both fresh and EG-thawed explants actively secreted E2 throughout the culture time (5 h to 9 days) (**Fig. 8, top panel**). Statistical analyses were focused on the experiments with the highest number of replicates (5 h to 2 days) and evidenced no significant differences between biological replicates or between the different time points tested ($p > 0.05$, one-way ANOVA and Tukey's multiple comparison test). Similarly, no differences were observed between fresh and EG-thawed explants when we compared equal time points ($p > 0.05$, one-way ANOVA and Tukey's multiple comparison test).

Regarding P4, variable levels of detection were found in both fresh and EG-thawed explants. Explants obtained from the same individuals and analysed at identical time points displayed uneven P4 levels (**Fig. 8, bottom panel**). As detailed for E2 levels, statistical analyses were focused on the 5-h to 2-day time points. Overall, no significant differences were found between biological replicates or between the time points tested. Similarly, we found no differences between fresh and EG-thawed explants compared at equal time points ($p > 0.05$, one-way ANOVA and Tukey's multiple comparison test). The exception was observed in P4 levels in fresh explants, which were significantly higher at 5 h than at later time points ($p < 0.05$, one-way ANOVA and Tukey's multiple comparison test), although the number of available samples was lower.

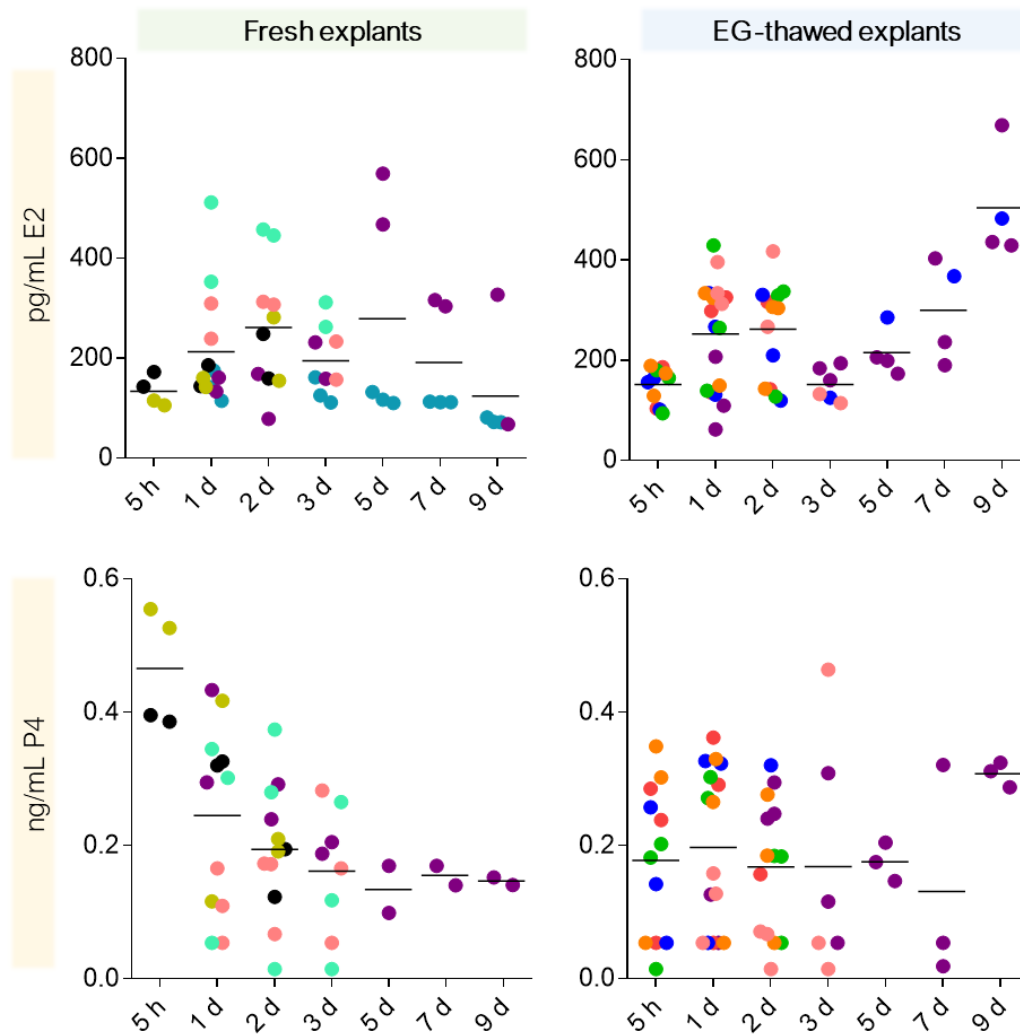


Fig. 8. Secretion of 17-β-oestradiol (E2, top panel) and 17-OH-progesterone (P4, bottom panel) in the supernatants of fresh (left panel) and EG-thawed (right panel) explants throughout the culture time (5 hours to 9 days). Each dot represents the hormone concentration of an individual explant; bars represent grand mean values.

3.7. Receptivity to *T. gondii* or *N. caninum* infection

To assess the suitability of this system for studying the interactions between the sheep and two relevant reproductive pathogens, we experimentally infected fresh and EG-thawed explants (from three ewes and one ewe, respectively) with *T. gondii* or *N. caninum* tachyzoites. These parasites

584 display an extraordinary tropism for placental tissues. In a first assay, we infected fresh explants with
585 5×10^4 tachyzoites of both parasites, but the parasite burden achieved was extremely low (data not
586 shown). Consequently, in successive trials, we opted to increase the challenge dose to 10^5 and 10^6
587 tachyzoites.

588 Infection of fresh explants with *T. gondii* tachyzoites was successful, resulting in active replication of
589 the parasite from 1 to 5 days postinfection (dpi) (**Fig. 9a, top panel**). This increase in the parasite
590 burden was significantly higher from 3 dpi in explants infected with 10^5 and 10^6 tachyzoites ($p < 0.05$,
591 Kruskal–Wallis test and Dunn’s multiple comparison). Since we did not find differences between the
592 two challenge doses used ($p > 0.05$, Kruskal–Wallis test and Dunn’s multiple comparison), EG-thawed
593 explants were only infected with 10^6 *T. gondii* tachyzoites. These explants behaved similarly to the
594 fresh ones, with parasite burdens tending to be higher from 3 dpi onwards. However, no significant
595 differences were found between the time points assayed ($p > 0.05$, Kruskal–Wallis test and Dunn’s
596 multiple comparison). The resazurin tests that were run in parallel with infected and uninfected fresh
597 and EG-thawed explants demonstrated that all tissues remained viable during the experiments, and
598 in fact, comparable viability scores were observed under all conditions (**Fig. 9b**).

599 Fresh explants were also successfully infected by *N. caninum* tachyzoites at 1 dpi, although the
600 replication levels quantified at 3 and 5 dpi were not significantly increased ($p > 0.05$, Kruskal–Wallis
601 test and Dunn’s multiple comparison) (**Fig. 9a, bottom panel**). No differences were observed
602 between the two challenge doses employed ($p > 0.05$, Kruskal–Wallis test and Dunn’s multiple
603 comparison). Since *T. gondii* replication appeared to occur more actively than that of *N. caninum*, we
604 corroborated that the latter was able to form parasitophorous vacuoles in fresh explants through
605 immunohistochemical studies (**Fig. 9c**). Based on these results, we subsequently infected EG-thawed
606 explants only with 10^6 *N. caninum* tachyzoites and observed parasite kinetics similar to those found

607 in fresh explants. No differences were found between fresh and EG-thawed explants ($p > 0.05$,
608 Kruskal–Wallis test and Dunn’s multiple comparison) (**Fig. 9a, bottom panel**). We also carried out
609 parallel resazurin tests under all conditions and these tests demonstrated that all the tissues
610 remained viable during the experiments. In addition, we did not observe differences in viability
611 among all the conditions tested (**Fig. 9b**).

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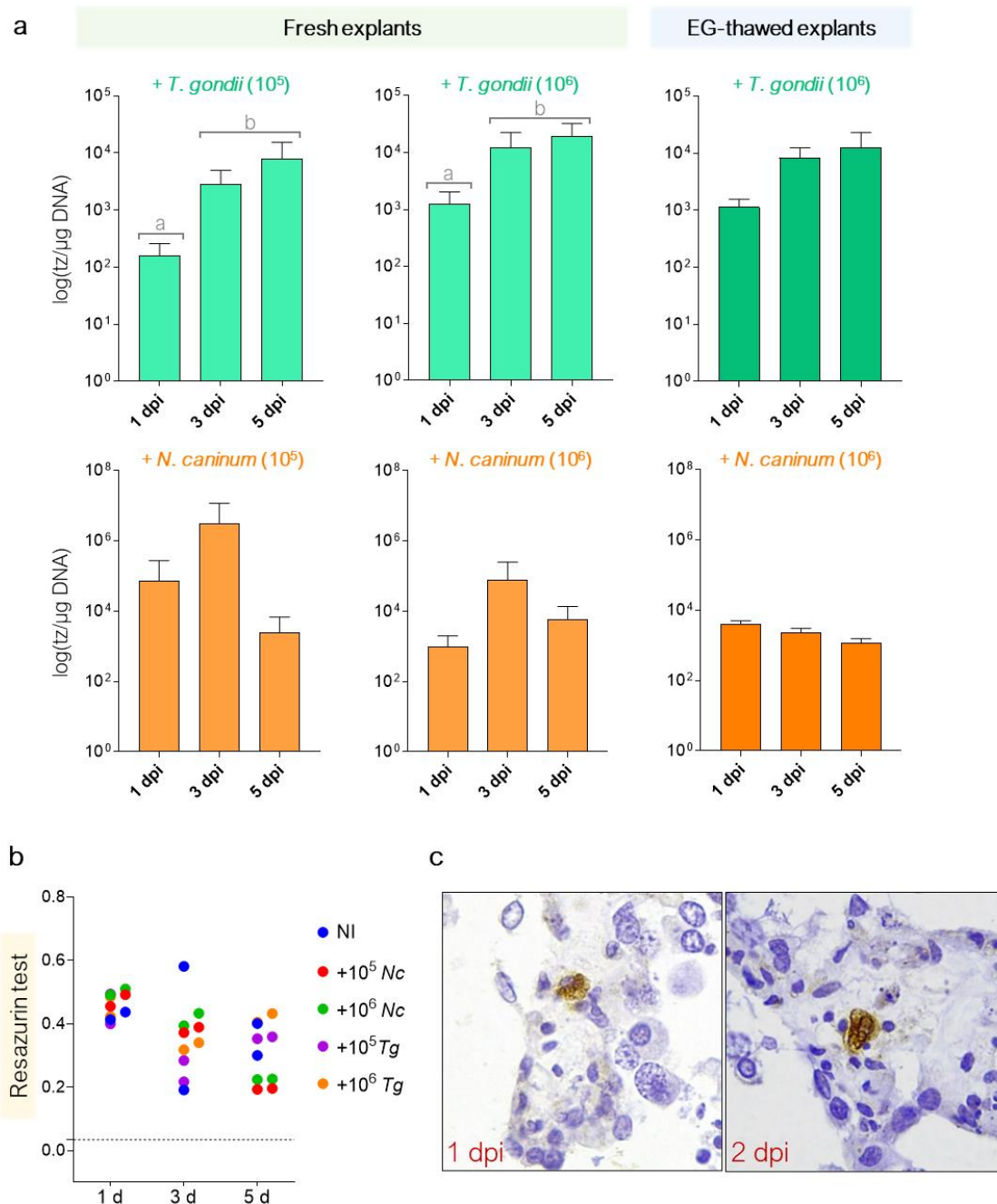


Fig. 9. a: Parasite burdens of fresh (left and centre graphs) and EG-thawed explants (right graphs) experimentally infected with *T. gondii* (top panel) or *N. caninum* (bottom panel). Column bar graph representing the mean and SD values of each condition. The parasite and challenge dose employed for each experiment are indicated on the top of every graph. Groups marked with different letters

619 were significantly different ($p < 0.05$, Kruskal–Wallis test and Dunn’s multiple comparison). **b:**
620 Representative resazurin assay of infected and noninfected fresh explants. The bottom dotted line
621 represents the average absorbance of explants fixed with 4% paraformaldehyde and used as controls
622 of nonviability. Each dot represents individual explants. NI: noninfected; 10^5 and 10^6 : challenge dose;
623 *Nc*: *N. caninum*; *Tg*: *T. gondii*. **c:** Positive IHC labelling of *N. caninum* antigen within the cytoplasm of
624 cells in experimentally infected fresh placental explants at 1 and 2 dpi. 60x.

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4. Discussion

The availability of useful and ethically acceptable models for the study of reproductive failure-causing agents at the foetal–maternal interface is an ongoing effort in veterinary medicine. In the case of apicomplexan parasites, multiple animal models developed in the target ruminant species are suitable to carry out experimental trials that mirror a natural scenario of infection. Nevertheless, their use is constrained by the availability of adequate facilities with trained staff, by the higher costs and longer experimental periods of the animal models, and, increasingly, by ethical concerns. Alternative in vitro systems have brought a cost-effective and animal-free source of experimental material, although the use of these alternative systems cannot replicate the close interaction between the various cell populations locally present in the placenta [23]. Organoids and 'organs-on-chip' are increasingly being proposed as alternatives to the use of animals, but these approaches require technical resources that are still difficult to access for most laboratories and that rely on the use of pluripotent or immortalised cell lines [35]. Therefore, interest in the development of ex vivo systems that are able to reproduce what happens in naturally infected animals is increasing [23]. The use of placental explants for research in human medicine has been extensively reported on thus far. This has been facilitated by the fact that deliveries and voluntary pregnancy interruptions take place under specialised medical care and near research facilities [16,36]. This contrasts with the scenario found in veterinary medicine, where the birth of livestock animals occurs on farms and the slaughter of pregnant females is rare, thus limiting the availability of suitable tissues [23]. Recent publications have reported the collection of placentomes from heifers through caesarean section, but this requires skilled surgeons [37]. Other studies have entailed obtaining explants from sheep full-term placentas and using them to assess the effect of infection with *T. cruzi* and *T. gondii* [21,22,38]. However, these explants lacked the caruncles, which are regarded as the main actors in the local control of parasitic infections [23]. In addition, naturally expelled placentas display extensive

670 disintegration of the chorionic epithelium [39] and possibly modified MHC I expression, which triggers
671 alloimmune rejection [40]. Hence, these tissues would not represent the environment found during
672 gestation. In fact, we decided against the use of full-term placenta explants in experiments
673 performed with three different healthy ewes that delivered lambs in our teaching facilities, as these
674 displayed high tissue damage scores and extremely low RNA concentrations at the time of collection
675 (data not shown).

676 Based on what is described in the literature, we took advantage of ongoing experiments using healthy
677 pregnant ewes to create a tissue biobank. Sheep have been shown to be receptive to infection with
678 both *T. gondii* and *N. caninum* [41] and therefore represent a good starting point to standardise ex
679 vivo models for the study of both agents. To make the most of the collected tissue and basing our
680 approach on previous works that successfully cryopreserved human and canine placental explants,
681 we followed similar methods to develop cryogenic protocols suitable for sheep explants. Using
682 human explants cultured up to 2 days, Huppertz et al. (2011) demonstrated that cryopreservation
683 with DMSO did not impair tissue morphology, viability, proliferation, or functionality after thawing
684 [24]. Following similar protocols, Sahlfeld et al. (2012) isolated proliferative trophoblasts from canine
685 placentas cryopreserved with DMSO, showing the same morphological features observed in fresh
686 cells [25]. In the present study, four different cryopreservation methods were assessed, and these
687 were based on the use of DMSO or EG as cryoprotectants under two different freezing rates. We
688 used the tissue damage scores as a gold-standard criterion to determine the best freezing method.
689 Unfortunately, these results were not supported by our LDH secretion findings, which were
690 inconclusive, as discussed below. In contrast with the findings reported in the aforementioned
691 studies, we observed that pretreatment of fresh explants with DMSO had a negative effect on their
692 tissue integrity but also that DMSO-thawed explants tended to display worse tissue integrity scores.
693 The histological architecture was better preserved in EG-thawed explants, especially in those

694 cryopreserved with slow freezing rates. Hence, we focused on our slow EG cryopreservation protocol
695 to create a tissue biobank for the subsequent experiments.

696 In this study, we also aimed to optimise culture conditions to extend the lifespan of fresh and EG-
697 thawed explants. In parallel experiments, we observed no apparent differences between explants
698 from the same donor incubated with different culture media (M-199, DMEM/F12, RPMI 1640) in
699 terms of media pH, tissue damage scores and viability (data not shown). Similar results have been
700 reported in studies performed with human explants [42]. We opted to use M-199 media based on
701 the report of Wojciechowska et al. (2015), which demonstrated its improved capacity to maintain
702 tissue integrity and hormone secretion in fresh placental explants obtained from cows [30].

703 Once culture conditions were established, we next studied the impact of ex vivo culture (and
704 cryopreservation) on their structural integrity. In general, both types of explants preserved their
705 tissue architecture and displayed progressive degradation with time, which was more evident from
706 Day 2 onwards. EG-thawed explants showed earlier signs of tissue degradation, but compared to
707 fresh explants, no significant differences were found. This indicates that cryopreservation has no
708 effect on tissue degradation and that this is inherent to ex vivo culture. The results also demonstrated
709 that fresh and EG-thawed explants could be suitable for ex vivo studies lasting up to 1 to 2 days.
710 Similar findings have been described for fresh explants obtained from human placentas both at
711 microscopic and ultrastructural levels as early as at 8 h of culture [27,43,44]. Interestingly, (i) culture
712 of human explants under static conditions has been shown to negatively affect tissue structure and
713 biochemical integrity; (ii) nonperfused explants display higher levels of apoptotic cells, especially
714 trophoblasts [42,43]; and (iii) the oxygen tension present in the culture can induce changes in the
715 synthesis of cell adhesion markers, cytokines, and hormones [45]. Altogether, these works indicate
716 that the use of flow culture protocols and oxygen-controlled atmospheres might significantly improve
717 our sheep explant culture model.

718 On the other hand, previous works carried out with human samples have described a marked
719 degeneration of the syncytiotrophoblast after 24 h of culture; this degeneration is followed by an
720 active generation of a new syncytiotrophoblast layer after 2-4 days in culture, which maintains the
721 natural connections with the extracellular matrix and remains morphologically intact for up to 11
722 days [27,28]. Based on that, we extended the explant culture time in some samples up to 9 days and
723 observed that the average tissue damage score of both fresh and EG-thawed explants was close to 2
724 until the seventh day of culture, when the expectation was to find autolytic tissue (score 3). In fact,
725 explants cultured up to 9 days displayed metabolic activity and were able to synthesise steroid
726 hormones. This suggests that our ex vivo model could be used to perform assays lasting beyond 2
727 days of culture. Nevertheless, we did not identify signs of active regeneration of the chorionic
728 epithelium using classic histological approaches. Hence, additional studies focused on the
729 identification of cytoskeleton and apoptosis markers through immunohistochemistry are needed to
730 draw more precise conclusions in this regard [43]. In addition, previous studies have shown that
731 human placental explants can retain their intact tissue structure for up to 10 days when cultured on
732 collagen sponges [46-48]. This indicates that the application of collagen sponges to sheep explants
733 may be advantageous, allowing experiments lasting more than 2 days.

734 To identify whether explant cultures remained metabolically active, we also performed different
735 viability assays. The quantification of LDH levels in culture supernatants is a method broadly
736 employed for human explants [24]. However, its usefulness was limited under the conditions in our
737 study, since enzyme levels remained high under all conditions tested. This was likely a consequence
738 of the placentome slicing, which resulted in a massive rupture of the cells present in the explant
739 edges. A similar effect has been described in experiments carried out with placental explants from
740 different species [49,50]. Based on these results, we opted to use two alternative assays based on XTT
741 and resazurin. Overall, both tests demonstrated that fresh and EG-thawed explants remained

742 metabolically active during all the time points tested, although the XTT assays were less sensitive.
743 Previous studies have shown that the XTT assay itself can be toxic to cells when used for an extended
744 period (> 4 h) and highly inaccurate when culture conditions promote superoxide formation [51,52].
745 For this reason, we focused our efforts on the resazurin assays, which are more sensitive, especially
746 when interpreted through fluorescence emission readings [52]. In the absence of specialised
747 equipment, the reduction of resazurin (dark blue) to resorufin (pink) by metabolically active cells can
748 be qualitatively interpreted with the naked eye, providing an easy and relatively inexpensive indicator
749 of tissue viability. Based on the results from this test, we observed a marked reduction in the viability
750 of EG-thawed explants after 3 days of culture, but this was seemingly recovered from Day 5 onwards.
751 This could reflect the activation of tissue recovery mechanisms, but we did not find evidence of that
752 in any of the experiments included here. For this reason, future efforts should target the
753 identification of specific proliferation markers through immunohistochemistry (e.g., Ki-67 protein)
754 [43]. It is important to note that viability values can be affected by the weight of each explant, which
755 reflects the number of cells present per well. This was not considered in our experimental design to
756 reduce the risk of bacterial contamination due to excessive handling and to prevent a delay in explant
757 plating that could have a negative impact on the explant viability. This drawback could be prevented
758 in future experiments using an automated tissue chopper as described elsewhere [50]. In any case,
759 we have been able to demonstrate, at least qualitatively, the occurrence of metabolic activity in both
760 fresh and EG-thawed explants cultured up to 9 days despite the extent of tissue damage observed
761 through histology. Additionally, we used the amount of RNA isolated from all the cultured explants
762 as a marker of the transcriptomic integrity. Here, we observed that RNA concentrations decreased
763 as tissue degradation increased. Therefore, RNA concentrations could be exploited as an indirect
764 marker of tissue damage when a histological approach is not readily accessible.

765 Despite being sparse, the isolated RNA allowed us to perform RT-qPCR and quantify *PLAC-1* and *KRT8*
766 expression until the second day of ex vivo culture. *PLAC-1* is a cell surface protein whose expression
767 is restricted to trophoblastic cells, determining the proliferative activity of the trophoblast and hence
768 the growth of the placenta [53]. In addition, this gene is involved in the progression of certain types
769 of cancer and in the occurrence of preeclampsia in humans [54]. In our study, only fresh explants
770 displayed a decrease in *PLAC-1* expression after 1-2 days of culture, but this was not observed in EG-
771 thawed explants. This could indicate a decrease in the proliferative activity, but only in fresh explants.
772 In line with this, Wojciechowska et al. (2017) demonstrated in a series of studies that *PLAC-1*
773 expression is downregulated upon treatment of fresh bovine placental explants with pesticides (DDT
774 and DDE) and polychlorinated biphenyls (PCB 153, 126, and 77), and this could be potentially
775 associated with an impairment of the proliferative capacities of the placenta [55,56]. However,
776 considering the low quality of the RNA used for the assays in the present study, the interpretation of
777 our results is difficult. *KRT8* is an intermediate filament protein involved in placental villi formation
778 by chorionic epithelial cells and is considered a marker of normal placental barrier function [57]. In
779 our experiments, a progressive upregulation of *KRT8* expression was observed in fresh explants, with
780 levels peaking on Day 2 of culture. In contrast, EG-thawed explants displayed decreasing *KRT8*
781 transcript levels over the culture duration. This could indicate that only fresh explants retain the
782 ability to repair the placental barrier, as demonstrated in fresh bovine placental explants [30,56].
783 Nevertheless, the interpretation of these findings is challenging considering the low quality of the
784 RNA extracted. Altogether, our results show that the usefulness of *PLAC-1* and *KRT8* as markers of
785 placental proliferative activity and normal placental barrier function, respectively, is limited for this
786 model. Future efforts should aim to detect specific proliferation and cytoskeleton markers through
787 immunohistochemistry to obtain better conclusions [43]. In this sense, we believe that the use of

788 culture systems based on collagen sponges could also improve explant RNA quality. This would
789 enable a comprehensive study of explant modulation through transcriptomic approaches.

790 Once we described the viability, structural integrity, and transcriptomic activity of fresh and EG-
791 thawed explants, we went one step further and tested their functionality in terms of steroid hormone
792 synthesis. As an endocrine organ, the sheep placenta synthesizes E2 and P4 during gestation to
793 regulate blood flow to the pregnant uterus and maintain pregnancy, respectively [58,59]. Overall, the
794 fresh and EG-thawed explants actively secreted E2 and P4 at all the time points tested (0 h–9 days),
795 suggesting that they were still functional. Importantly, the tissue damage observed over the culture
796 time did not seem to impair the explants' ability to synthesise both hormones. In line with this,
797 previous works have described a significant increase in E2 and P4 levels in bovine placental explants
798 stimulated with pesticides and polychlorinated biphenyls [55,56]. Similarly, human villous explants
799 can secrete hCG and hPL for 5-7 days [60]. In our experiments, secretion of P4 was more variable, and
800 we hypothesise that this response might be normal. Under physiological conditions, the corpus
801 luteum supports most of the P4 synthesis needed to maintain pregnancy in ewes, and it is only at
802 late gestation when the placenta overtakes P4 secretion [58]. Thus, considering that all our
803 placentomes were collected at the end of the second term of gestation, this pattern of P4 synthesis
804 could be expected. It is important to highlight that E2 and P4 secretion values were not normalised
805 and thus could be affected by the weight of each explant, as discussed for the resazurin assays. In
806 addition, the commercial tests employed are only validated for human samples and could yield
807 inaccurate results. Nevertheless, recent reports have demonstrated that human-intended kits are
808 suitable for ovine E2 and P4 analyses [61,62]. In any case, and from a qualitative perspective, our
809 results confirm that cultured explants can produce both hormones, confirming their functionality to
810 an extent.

811 Since the explants were demonstrated to be viable and functional, we then infected fresh and EG-
812 thawed explants with *T. gondii* or *N. caninum* tachyzoites and quantified the tissue parasite burdens
813 by qPCR. All the explants were receptive to infection by the two parasites, and no differences were
814 recorded in the parasite burdens present in fresh and EG-thawed explants. Direct evidence of
815 effective *N. caninum* infection was also obtained through immunohistochemistry. Compared to *N.*
816 *caninum*, *T. gondii* tachyzoites showed a higher proliferation capacity, with significant increases in
817 their growth after 3 and 5 days of infection. These findings are aligned with those of previous studies
818 in which explants obtained from ovine term placentas supported *T. gondii* replication [21,22,38]. The
819 results obtained in the explants infected with *N. caninum* were surprising, as pregnant sheep are
820 especially susceptible to experimental infection with this parasite [26]. However, considering that the
821 Nc-Spain7 isolate has a controlled number of passages in cell culture, we hypothesise that the
822 laboratory adaptation of the *T. gondii* strain used for infections (Tg-PRU) might have influenced our
823 results [63]. We also hypothesise that *N. caninum* would need more time than *T. gondii* to achieve its
824 exponential growth. In an attempt to assess whether the receptivity of the explants decreased with
825 their time in culture, a batch of fresh explants was cultured for 6 days in the absence of parasites and
826 then infected with *T. gondii* or *N. caninum* for an additional 1, 3 and 5 days. These explants remained
827 viable at all times in culture and displayed a similar receptivity to freshly obtained/thawed explants
828 (data not shown). Successful infection also supports the occurrence of metabolic activity in the
829 explants, since *T. gondii* and *N. caninum* growth rely on the recruitment of the host organelles in the
830 surroundings of their parasitophorous vacuoles to scavenge or salvage metabolites synthesised in
831 these organelles [64]. It is worth highlighting that the comparison of infected fresh and EG-thawed
832 explants was performed with tissues obtained from a single animal, and this may not be sufficiently
833 representative. However, considering the small variation observed between all animals and
834 conditions in all the parameters measured here (viability, structural integrity, transcriptional activity,

835 and functionality), similar results would be expected in terms of parasite replication. Nevertheless,
836 the susceptibility of EG-thawed tissues to parasitic infection will need to be further characterised in
837 future studies.

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5. Conclusions

In summary, we have developed and characterised an ovine placental ex vivo model suitable for the study of the early interactions between *T. gondii* or *N. caninum* and pregnant sheep. This model is based on the collection of placentomes from healthy ewes, the cryopreservation of placental explants using EG under slow freezing rates, and the use of M-199-based culture media. We have also defined reproducible and simple markers of tissue integrity (damage scores, RNA concentrations), viability (resazurin assays), and functionality (synthesis of E2 and P4) that can easily be exploited to assess the quality of the model. Regarding tissue integrity, RNA quality, and *T. gondii* and *N. caninum* receptivity, explants might be useful for the study of host–parasite interactions for periods of up to 2 days. However, viability and functionality assays suggest that these periods might be longer, up to 9 days. Our findings indicate that explant lifespan is mainly limited by the culture technique employed, and current evidence supports that there is still room for improvement of such methods using flow culture approaches, oxygen-controlled atmospheres, and/or collagen-based culture inserts. Optimisation of explant culture protocols could extend tissue integrity and RNA quality for longer periods, thus boosting their usefulness. This model also has the potential to be applied to other reproductive pathogens affecting sheep (e.g., *Brucella mellitensis*, *Chlamydia abortus*, *Coxiella burnetii*, Border disease virus) and could be easily adapted to other species (e.g., cattle, goats) or even other target tissues (e.g., intestine). While a thorough optimisation and characterisation of this model remains to be achieved, it is expected to have a significant impact on the study of transmissible diseases leading to reproductive failure. This would also contribute to reducing the number of animals used for experimental purposes.

883 CRediT authorship contribution statement

- 884 – **Pilar Horcajo:** conceptualization; data curation; investigation; methodology; validation; writing -
885 review & editing.
- 886 – **Luis Miguel Ortega-Mora:** conceptualization; data curation; funding acquisition; resources;
887 supervision; roles/writing - original draft.
- 888 – **Julio Benavides, Roberto Sánchez-Sánchez and Rafael Amieva:** investigation; methodology;
889 validation; writing - review & editing.
- 890 – **Esther Collantes-Fernández and Iván Pastor-Fernández:** conceptualization; data curation; formal
891 analysis; funding acquisition; investigation; methodology; project administration;
892 resources;supervision; validation; visualization; roles/writing - original draft.

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894 Declaration of competing interest

895 The authors declare that they have no competing interests.

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909 References

910 [1] Menzies PI. Control of important causes of infectious abortion in sheep and goats. Vet Clin North

911 Am Food Anim Pract 2011;27(1):81-93.

912 [2] Innes EA, Mattsson JG. *Neospora caninum* emerges from the shadow of *Toxoplasma gondii*.

913 Trends Parasitol. 2007;23(2):43-4.

914 [3] Głowacka P, Żakowska D, Naylor K, Niemcewicz M, Bielawska-Drózd A. *Brucella* - virulence factors,

915 pathogenesis and treatment. Pol J Microbiol 2018;67(2):151-61.

916 [4] Van den Brom R, van Engelen E, Roest HIJ, van der Hoek W, Vellema P. *Coxiella burnetii* infections

917 in sheep or goats: An opinionated review. Vet Microbiol 2015;181(1-2):119-29.

918 [5] Caspe SG, Livingstone M, Frew D, Aitchison K, Wattegedera SR, Entrican G, et al. The 1B vaccine

919 strain of *Chlamydia abortus* produces placental pathology indistinguishable from a wild type

920 infection. PLoS One 2020;15(11):e0242526.

921 [6] Roberts RM, Green JA, Schulz LC. The evolution of the placenta. Reproduction 2016;152(5):179.

922 [7] Wooding F. The synepitheliochorial placenta of ruminants: Binucleate cell fusions and hormone

923 production. Placenta 1992;13(2):101-13.

924 [8] Entrican G, Wattegedera S, Wheelhouse N, Allan A, Rocchi M. Immunological paradigms and the

925 pathogenesis of ovine chlamydial abortion. Am. J. Reprod. Immunol. 2010;64(4):287-94.

926 [9] Haldorson GJ, Stanton JB, Mathison BA, Suarez CE, Baszler TV. *Neospora caninum*: Antibodies

927 directed against tachyzoite surface protein NcSRS2 inhibit parasite attachment and invasion of

928 placental trophoblasts in vitro. Exp. Parasitol. 2006;112(3):172-8.

929 [10] Wheelhouse N, Wattegedera S, Stanton J, Maley S, Watson D, Jepson C, et al. Ovine trophoblast
930 is a primary source of TNF α during *Chlamydophila abortus* infection. J. Reprod. Immunol. 2009;80(1–
931 2):49-56.

932 [11] Wheelhouse N, Coyle C, Barlow PG, Mitchell S, Greub G, Baszler T, et al. *Waddlia chondrophila*
933 infects and multiplies in ovine trophoblast cells stimulating an inflammatory immune response. PLoS
934 One 2014;9(7):e102386.

935 [12] Fernández-Escobar M, Calero-Bernal R, Regidor-Cerrillo J, Vallejo R, Benavides J, Collantes-
936 Fernández E, et al. In vivo and in vitro models show unexpected degrees of virulence among
937 *Toxoplasma gondii* type II and III isolates from sheep. Vet. Res. 2021;52(1):82-7.

938 [13] Jiménez-Pelayo L, García-Sánchez M, Regidor-Cerrillo J, Horcajo P, Collantes-Fernández E,
939 Gómez-Bautista M, et al. Differential susceptibility of bovine caruncular and trophoblast cell lines to
940 infection with high and low virulence isolates of *Neospora caninum*. Parasit. Vectors 2017;10(1):463-
941 9.

942 [14] Jiménez-Pelayo L, García-Sánchez M, Regidor-Cerrillo J, Horcajo P, Collantes-Fernández E,
943 Gómez-Bautista M, et al. Immune response profile of caruncular and trophoblast cell lines infected
944 by high- (Nc-Spain7) and low-virulence (Nc-Spain1H) isolates of *Neospora caninum*. Parasit Vectors
945 2019;12(1):218.

946 [15] Jiménez-Pelayo L, García-Sánchez M, Collantes-Fernández E, Regidor-Cerrillo J, Horcajo P,
947 Gutiérrez-Expósito D, et al. Crosstalk between *Neospora caninum* and the bovine host at the
948 maternal-foetal interface determines the outcome of infection. Vet Res 2020;51(1):83.

949 [16] Miller RK, Genbacev O, Turner MA, Aplin JD, Caniggia I, Huppertz B. Human placental explants in
950 culture: Approaches and assessments. Placenta 2005;26(6):439-48.

951 [17] Carvalho Neta AV, Styne AP, Paixao TA, Miranda KL, Silva FL, Roux CM, et al. Modulation of the
 952 bovine trophoblastic innate immune response by *Brucella abortus*. Infect. Immun. 2008;76(5):1897-
 953 907.

954 [18] Mol JP, Costa EA, Carvalho AF, Sun YH, Tsois RM, Paixao TA, et al. Early transcriptional responses
 955 of bovine chorioallantoic membrane explants to wild type, Δ virB2 or Δ btbB *Brucella abortus* infection.
 956 PLoS One 2014;9(9):e108606.

957 [19] Dos Santos LS, da Silva Mol JP, de Macedo AA, Silva AP, Dos Santos Ribeiro DL, Santos RL, et al.
 958 Transcription of non-classic major histocompatibility complex (MHC) class I in the bovine placenta
 959 throughout gestation and after *Brucella abortus* infection. Vet. Immunol. Immunopathol.
 960 2015;167(3-4):166-70.

961 [20] Rocha CE, Mol JPS, Garcia LNN, Costa LF, Santos RL, Paixao TA. Comparative experimental
 962 infection of *Listeria monocytogenes* and *Listeria ivanovii* in bovine trophoblasts. PLoS One
 963 2017;12(5):e0176911.

964 [21] Liempi A, Castillo C, Medina L, Galanti N, Maya JD, Parraguez VH, et al. Comparative ex vivo
 965 infection with *Trypanosoma cruzi* and *Toxoplasma gondii* of human, canine and ovine placenta:
 966 Analysis of tissue damage and infection efficiency. Parasitol. Int. 2020;76:102065.

967 [22] Medina L, Guerrero-Muñoz J, Castillo C, Liempi A, Fernández-Moya A, Araneda S, et al.
 968 Differential microRNAs expression during ex vivo infection of canine and ovine placental explants
 969 with *Trypanosoma cruzi* and *Toxoplasma gondii*. Acta Trop 2022;235:106651.

970 [23] Pastor-Fernández I, Collantes-Fernández E, Jiménez-Pelayo L, Ortega-Mora LM, Horcajo P.
 971 Modeling the ruminant placenta-pathogen interactions in apicomplexan parasites: Current and
 972 future perspectives. Front. Vet. Sci. 2021;7:634458.

973 [24] Huppertz B, Kivity V, Sammar M, Grimpel Y, Leepaz N, Orendi K, et al. Cryogenic and low
 974 temperature preservation of human placental villous explants - a new way to explore drugs in
 975 pregnancy disorders. *Placenta* 2011;32 Suppl 1:65-76.

976 [25] Sahlfeld L, Hazzard T, Kutzler M. Cellular characteristics of cultured canine trophoblasts. *Reprod.*
 977 *Domest. Anim.* 2012;47 Suppl 6:161-4.

978 [26] Sánchez-Sánchez R, Ferre I, Re M, Regidor-Cerrillo J, Blanco-Murcia J, Ferrer LM, et al. Influence
 979 of dose and route of administration on the outcome of infection with the virulent *Neospora caninum*
 980 isolate Nc-Spain7 in pregnant sheep at mid-gestation. *Vet. Res.* 2018;49(1):42.

981 [27] Palmer ME, Watson AL, Burton GJ. Morphological analysis of degeneration and regeneration of
 982 syncytiotrophoblast in first trimester placental villi during organ culture. *Hum Reprod*
 983 1997;12(2):379-82.

984 [28] Simán CM, Sibley CP, Jones CJ, Turner MA, Greenwood SL. The functional regeneration of
 985 syncytiotrophoblast in cultured explants of term placenta. *Am J Physiol Regul Integr Comp Physiol*
 986 2001;280(4):1116.

987 [29] Arranz-Solís D, Benavides J, Regidor-Cerrillo J, Horcajo P, Castaño P, del Carmen Ferreras M, et
 988 al. Systemic and local immune responses in sheep after *Neospora caninum* experimental infection at
 989 early, mid and late gestation. *Vet. Res.* 2016;47(1):1-13.

990 [30] Wojciechowska A, Mlynarczuk J, Kotwica J. Short-term incubation of bovine placentome sections
 991 as a tool to study xenobiotic mechanism of action. *Reprod. Biol.* 2015;15(4):238-46.

992 [31] Schmittgen TD, Livak KJ. Analyzing real-time PCR data by the comparative C(T) method. *Nat.*
 993 *Protoc.* 2008;3(6):1101-8.

994 [32] Rico-San Román L, Amieva R, Regidor-Cerrillo J, García-Sánchez M, Collantes-Fernández E,
 995 Pastor-Fernández I, et al. NcGRA7 and NcROP40 play a role in the virulence of *Neospora caninum* in
 996 a pregnant mouse model. *Pathogens* 2022;11(9):998. doi: 10.3390/pathogens11090998.

997 [33] Sánchez-Sánchez R, Ferre I, Re M, Vázquez P, Ferrer LM, Blanco-Murcia J, et al. Safety and
998 efficacy of the bumped kinase inhibitor BKI-1553 in pregnant sheep experimentally infected with
999 *Neospora caninum* tachyzoites. Int J Parasitol Drugs Drug Resist 2018;8(1):112-24.

1000 [34] Castaño P, Fuertes M, Regidor-Cerrillo J, Ferre I, Fernández M, Ferreras MC, et al. Experimental
1001 ovine toxoplasmosis: Influence of the gestational stage on the clinical course, lesion development
1002 and parasite distribution. Vet. Res. 2016;47:43-z.

1003 [35] Ramírez-Flores CJ, Tibabuzo Perdomo AM, Gallego-López GM, Knoll LJ. Transcending dimensions
1004 in apicomplexan research: From two-dimensional to three-dimensional in vitro cultures. Microbiol.
1005 Mol. Biol. Rev. 2022;86(2):e0002522,22.

1006 [36] Huckle WR. Cell-and tissue-based models for study of placental development. In: Anonymous
1007 Progress in molecular biology and translational science. Elsevier; 2017, p. 29-37.

1008 [37] Contreras-Correa ZE, Lemire RL, Burnett DD, Lemley CO. Temporal transcript abundance of clock
1009 genes, angiogenic factors and nutrient sensing genes in bovine placental explants. Theriogenology
1010 2020;151:74-80.

1011 [38] Liempi A, Castillo C, Medina L, Rojas-Pirela M, Araneda S, Maya JD, et al. Ex vivo infection of
1012 canine and ovine placental explants with *Trypanosoma cruzi* and *Toxoplasma gondii*: Differential
1013 activation of NF kappa B signaling pathways. Acta Trop 2021;214:105766.

1014 [39] Steven DH. Separation of the placenta in the ewe: An ultrastructural study. Q J Exp Physiol Cogn
1015 Med Sci 1975;60(1):37-44.

1016 [40] Benedictus L, Koets AP, Rutten VPMG. The role of placental MHC class I expression in immune-
1017 assisted separation of the fetal membranes in cattle. J Reprod Immunol 2015;112:11-9.

1018 [41] Benavides J, González-Warleta M, Arteche-Villasol N, Pérez V, Mezo M, Gutiérrez-Expósito D.
1019 Ovine neosporosis: The current global situation. Animals (Basel) 2022;12(16):2074.

1020 [42] Di Santo S, Malek A, Sager R, Andres A-, Schneider H. Trophoblast viability in perfused term
 1021 placental tissue and explant cultures limited to 7-24 hours. *Placenta* 2003;24(8-9):882-94.

1022 [43] Kupper N, Pritz E, Siwetz M, Guettler J, Huppertz B. Placental villous explant culture 2.0: Flow
 1023 culture allows studies closer to the in vivo situation. *Int J Mol Sci* 2021;22(14):7464.

1024 [44] Sooranna SR, Oteng-Ntim E, Meah R, Ryder TA, Bajoria R. Characterization of human placental
 1025 explants: Morphological, biochemical and physiological studies using first and third trimester
 1026 placenta. *Hum Reprod* 1999;14(2):536-41.

1027 [45] Peltier MR, Gurzenda EM, Murthy A, Chawala K, Lerner V, Kharode I, et al. Can oxygen tension
 1028 contribute to an abnormal placental cytokine milieu? *Am J Reprod Immunol* 2011;66(4):279-85.

1029 [46] Lopez H, Benard M, Saint-Aubert E, Baron M, Martin H, Al Saati T, et al. Novel model of placental
 1030 tissue explants infected by cytomegalovirus reveals different permissiveness in early and term
 1031 placentae and inhibition of indoleamine 2,3-dioxygenase activity. *Placenta* 2011;32(7):522-30.

1032 [47] Fitzgerald W, Gomez-Lopez N, Erez O, Romero R, Margolis L. Extracellular vesicles generated by
 1033 placental tissues ex vivo: A transport system for immune mediators and growth factors. *Am J Reprod*
 1034 *Immunol* 2018;80(1):e12860.

1035 [48] Faye A, Pornprasert S, Dolcini G, Ave P, Taïeb J, Taupin JL, et al. Evaluation of the placental
 1036 environment with a new in vitro model of histocultures of early and term placentae: Determination
 1037 of cytokine and chemokine expression profiles. *Placenta* 2005;26(2-3):262-7.

1038 [49] Sato BL, Ward MA, Astern JM, Kendal-Wright CE, Collier AC. Validation of murine and human
 1039 placental explant cultures for use in sex steroid and phase II conjugation toxicology studies. *Toxicol*
 1040 *in Vitro* 2015;29(1):103-12.

1041 [50] Gilligan J, Tong M, Longato L, de la Monte SM, Gundogan F. Precision-cut slice culture method
 1042 for rat placenta. *Placenta* 2012;33(1):67-72.

1043 [51] Wang S, Yu H, Wickliffe JK. Limitation of the MTT and XTT assays for measuring cell viability due
 1044 to superoxide formation induced by nano-scale TiO₂. *Toxicol in Vitro* 2011;25(8):2147-51.

1045 [52] Präbst K, Engelhardt H, Ringgeler S, Hübner H. Basic colorimetric proliferation assays: MTT, WST,
 1046 and resazurin. *Methods Mol Biol* 2017;1601:1-17.

1047 [53] Cocchia M, Huber R, Pantano S, Chen EY, Ma P, Forabosco A, et al. PLAC1, an Xq26 gene with
 1048 placenta-specific expression. *Genomics* 2000;68(3):305-12.

1049 [54] Conese M, Napolitano O, Laselva O, Di Gioia S, Nappi L, Trabace L, et al. The oncogenic theory of
 1050 preeclampsia: Is amniotic mesenchymal stem cells-derived PLAC1 involved? *Int J Mol Sci*
 1051 2023;24(4):3612.

1052 [55] Wojciechowska A, Mlynarczyk J, Kotwica J. Changes in the mRNA expression of structural
 1053 proteins, hormone synthesis and secretion from bovine placentome sections after DDT and DDE
 1054 treatment. *Toxicology* 2017;375:1-9.

1055 [56] Wojciechowska A, Mlynarczyk J, Kotwica J. Disorders in barrier protein mRNA expression and
 1056 placenta secretory activity under the influence of polychlorinated biphenyls in vitro. *Theriogenology*
 1057 2017;89:9-19.

1058 [57] Jaquemar D, Kupriyanov S, Wankell M, Avis J, Benirschke K, Baribault H, et al. Keratin 8 protection
 1059 of placental barrier function. *J Cell Biol* 2003;161(4):749-56.

1060 [58] Reynolds LP, Legacki EL, Corbin CJ, Caton JS, Vonnahme KA, Stanley S, et al. Ovine placental
 1061 steroid synthesis and metabolism in late gestation. *Biol Reprod* 2018;99(3):662-70.

1062 [59] Bairagi S, Grazul-Bilska AT, Borowicz PP, Reyaz A, Valkov V, Reynolds LP. Placental development
 1063 during early pregnancy in sheep: Progesterone and estrogen receptor protein expression.
 1064 *Theriogenology* 2018;114:273-84.

1065 [60] Polliotti BM, Abramowsky C, Schwartz DA, Keesling SS, Lee GR, Caba J, et al. Culture of first-
1066 trimester and full-term human chorionic villus explants: Role of human chorionic gonadotropin and
1067 human placental lactogen as a viability index. Early Pregnancy 1995;1(4):270-80.

1068 [61] Casao A, Pérez-Pé R, Abecia JA, Forcada F, Muiño-Blanco T, Cebrián-Pérez JÁ. The effect of
1069 exogenous melatonin during the non-reproductive season on the seminal plasma hormonal profile
1070 and the antioxidant defence system of rasa aragonesa rams. Anim Reprod Sci 2013;138(3-4):168-74.

1071 [62] Pasciu V, Nieddu M, Baralla E, Porcu C, Sotgiu F, Berlinguer F. Measurement of progesterone in
1072 sheep using a commercial ELISA kit for human plasma. J Vet Diagn Invest 2022;34(1):90-3.

1073 [63] Colos-Arango A, Largo-de la Torre A, Calero-Bernal R, Ortega-Mora L, Regidor-Cerrillo J. Short-
1074 term culture adaptation of *Toxoplasma gondii* archetypal II and III field isolates affects cystogenic
1075 capabilities and modifies virulence in mice. Int J Parasitol 2023:S0020-7.

1076 [64] Nolan SJ, Romano JD, Luechtefeld T, Coppens I. *Neospora caninum* recruits host cell structures
1077 to its parasitophorous vacuole and salvages lipids from organelles. Eukaryot. Cell. 2015;14(5):454-73.

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