Background. Ovarian tissue cryopreservation (OTC) is important for fertility preservation and conservation. Delay in OTC can occur for transport or workflow management, yet little is understood about the effect of this on the tissue. Objective. To determine if a delay of 24–48 h to OTC affects primordial follicle (PF) health. Methods. Ovaries (n = 6 sheep) were processed immediately or after storage at 4 °C (24 h, 48 h). Tissue was fixed fresh, after cryopreservation or 10-day xenotransplantation. Morphological assessment of follicle tissue can be stored for up to 48 h prior to cryopreservation with no net impact on PF morphology which is indicative of health. Findings. A total of 1,541 follicles were analysed. A 24 h processing delay did not impact PF health in fresh or cryopreserved tissue. In fresh tissue, a 48 h delay had an adverse effect on PF health (OR = 2.47, 95% CI 1.29–4.71). Interestingly, a 48 h delay resulted in cryopreserved tissue being less likely to be graded as abnormal compared to control (OR = 0.56, 95% CI 0.36–0.87). There was no difference in PF health or development across groups following xenotransplantation. Conclusion. Ovarian tissue can be stored for up to 48 h prior to cryopreservation with no net impact on PF morphology which is indicative of health.

Keywords fertility preservation; delayed processing; ovarian cryopreservation; follicle; human; sheep; conservation

1. Introduction
At present, over 80% of children and adolescents diagnosed with cancer will survive the disease and reach adulthood [1]. Furthermore, as of 2011, there are an estimated 388,000 childhood cancer survivors in the US [2] and over 30,000 in the UK [3]. Childhood cancer treatment can, however, often be associated with a significant negative impact on the growth, development and, commonly, the fertility of young patients [4]. Thus, fertility preservation is of importance to young oncology patients, in order to prevent further emotional distress at the time of cancer diagnosis and long-term negative impact on the recovery and future wellbeing of childhood cancer survivors [5].

Ovarian tissue cryopreservation (OTC) is a method of fertility preservation that is becoming increasingly well-established world-wide. This involves dissection, cryoprotection and freezing of the ovarian cortex with the aim of preserving the primordial pool of follicles contained within. Importantly, although the intervention is still considered experimental in several countries, as of 2017, an estimated 130 pregnancies have been achieved after ovarian tissue transplantation globally [6]. However, OTC remains a specialised procedure, which requires practical experience, specialist equipment and dedicated facilities, which comply with EU and/or national regulations. As a result, although individual experimental cases have been reported from 21 countries [7], the availability of the technique as a modality of fertility preservation is limited [8], while the need for it may continue to rise.

One suitable approach to improve accessibility to OTC involves ovarian tissue being procured at various satellite clinics and transported to a central cryopreservation facility; this has been termed the “Danish model” [8,9]. However, the Danish model may require overnight storage of the tissue to allow for long-distance transportation of the samples and coordination and workload management at the cryopreservation facility. This may create a delay between ovarian tissue retrieval and processing for cryopreservation, during which the ovary is in a state of ischaemia, which can cause nutrient deprivation, hypoxia and, as a result, cell death [10]. The process of tissue cryopreservation and thawing is further associated with great physiological stress to living cells and tissues due to the dramatic changes in osmolarity and temperature experienced by the tissue, as well as the formation of structurally disruptive ice crystals [11]. Hence, it is of concern that pre-exposure of the ovarian tissue to challenging conditions such as prolonged cold ischaemia has the potential to exacerbate the damaging effects of freeze-thaw on tissue health and integrity, even if no direct effects were observed in fresh tissue.
At present, two case reports of cryopreservation delay for up to 20 h at 4 °C with one subsequent live birth have provided proof of principle that follicles can survive freezedelay and that ovarian cortical tissue can retain graft functionality [8,12]. However, the impact of cryopreservation delay on the health of the primordial pool remains controversial, with studies reporting either no effect on follicle density and health [13] or a negative effect on both [14].

In this context, the present study set out to examine the effect of delayed processing and cryopreservation on ovarian tissue stored for fertility preservation using tissue processed in a clinically relevant manner. The sheep was used as a model organism for three reasons: (1) its structural similarity with human ovaries [15]; (2) human OTC has been largely optimised using the sheep as a model; and (3) it is the only species, aside from rodents and primates, in which a live birth has been achieved after transplantation of frozen-thawed ovarian tissue [16]. As the clinical value of cryopreserved ovarian tissue is largely determined by its potential to restore fertility following autologous grafting, xenotransplantation into immunodeficient mice was employed to examine the effect of delayed processing on in vivo development after cryopreservation and thawing. The aim of the study was to establish if processing delay of 24 h or 48 h had an effect on primordial follicle morphology in fresh, frozen-thawed and xenotransplanted ovine ovarian cortical tissue.

2. Materials and methods

2.1. Tissue collection

Pairs of ovaries from six female lambs (Ovis aries, breed unknown) were collected from a local abattoir and transported on ice in 100 mL of transport medium (Leibovitz’s L-15 medium [Thermo Fisher, UK] supplemented with 100 U/mL penicillin and 100 µg/mL streptomycin [Sigma], and 3 mg/mL bovine serum albumin (BSA, Fisher Scientific, UK)) for 1 h on ice. Cryopreservation was carried out in a controlled-rate freezer (IceCube 14S, SY-LAB, Sweden) using the following cooling programme: start temperature: 4 °C; cooling rate I: −2 °C/min to −9 °C; cooling rate II: −0.3 °C/min to −40 °C; cooling rate III: −10 °C/min to −140 °C. Manual seeding was performed at −9 °C and after freezing samples were stored in vapour phase liquid nitrogen.

2.2. Ovarian tissue processing and cryopreservation

Ovaries were processed for cryopreservation using procedures aligned with the relevant standard operating procedures (SOPs) applied at the Oxford Cell and Tissue Biobank (Oxford, UK) [17,18], with some modification. Each ovary was bivalved and the ovarian medulla was gently dissected away using curved scissors under aseptic conditions. The outermost 1 mm of the ovarian cortex was cut to strips of approximately 5 × 2 × 1 mm using a surgical scalpel blade. All tissue handling was carried out in transport media on a Medicool ice block. Cortical strips were either fixed directly after processing (fresh samples) or cryopreserved. For cryopreservation, individual cortical strips were transferred to Nunc cryotubes and equilibrated in 1 mL of cryoprotectant medium containing L-15 supplemented with 1.5 M ethylene glycol (Sigma), 0.1 M sucrose (Sigma), and 3 mg/mL bovine serum albumin (BSA, Sigma, UK) for 1 h on ice. Cryopreservation was performed in a controlled-rate freezer (IceCube 14S, SY-LAB, Sweden) using the following cooling programme: start temperature: 4 °C; cooling rate I: −2 °C/min to −9 °C; cooling rate II: −0.3 °C/min to −40 °C; cooling rate III: −10 °C/min to −140 °C. Manual seeding was performed at −9 °C and after freezing samples were stored in vapour phase liquid nitrogen.

2.3. Tissue thawing

Cryotubes were thawed in a water bath at 30 °C for 3 min, after which cortical strips were washed through three thawing solutions, containing a decreasing gradient of ethylene glycol (1 M, 0.5 M and 0 M), 0.1 M sucrose and 3 mg/mL BSA in L-15 medium, for 5 min each at room temperature. For samples that were to be xenotransplanted, 2.5 µg/mL amphotericin B, 100 µg/mL penicillin, and 100 µg/mL streptomycin were added to the thawing solutions. Tissue was transplanted within 1 h of thawing.

2.4. Mice

Immunodeficient female mice (SCID; CB17/Icr-Prkdcscid/IcrIcoCrl) were obtained from Charles River Laboratories (Kent, UK) and housed together in a cage with filtered air supply under a 12:12 h light-dark cycle with ad libitum access to sterile food and water. The animals were acclimatised for two weeks prior to surgery and xenotransplantation was carried out at 8 weeks of age.

2.5. Ethics approval

Animal work was carried out in accordance with project license number 30/3352 granted to Dr Suzannah Williams under The Animals (Scientific Procedures) Act 1986.

2.6. Xenotransplantation

Thawed cortical strips (1 × 1 × 3 mm) were transplanted subcutaneously to the left flank of immunocompromised mice (Supplementary Figure 1). Four strips were transplanted to each mouse via two incisions each leading to two separate pockets. For each mouse, strips from one ovarian pair (no delay and 24 h delay) were transplanted into two pockets created from the first incision and strips from another ovarian pair (no delay and 48 h delay) were transplanted into two pockets created from the second incision. Each graft’s position was secured in the pocket before the incision was closed. Mice were housed individually for the first 24 h after surgery. Recovery was monitored by observation and measuring body weight post-transplant; mice were weighed twice daily in the first 48 h and once daily thereafter. After
Figure 1: Health grading of primordial follicles based on morphological appearance. Bouin’s-fixed ovine ovarian tissue was analysed for the presence of primordial follicles (oocyte surrounded by a single layer of flattened pregranulosa cells) using haematoxylin and eosin (H&E) staining. Follicles were scored based on the presence of morphological markers (dense chromatin, eosinophilia, shrunken ooplasm [one point each] or pyknosis [two points]) as normal (0), affected (1), degenerating (2) or atretic (≥3). Scale bar: 10 µm.

2.7. Histological analysis
Cortical strips were fixed in Bouin’s solution overnight at room temperature and embedded in paraffin wax. Serial sections were obtained at 5 µm thickness and slides were stained with haematoxylin and eosin (H&E). Samples were imaged at ×400 magnification using a Leica DM2500 microscope. Images were captured using QCapture Pro 7 software. Follicles were graded in a blinded manner by two independent researchers, with a concordance rate of over 90%.

Primordial follicles were identified based on morphology defined by a single layer of flattened granulosa cells [19]. Oocyte health was assessed as described previously [18], with slight modification. Oocytes were graded as normal (grade 0) if they had no morphological markers, affected (grade 1) if they had either dense chromatin, eosinophilia or shrinkage of the ooplasm (one point each), degenerating (grade 2) if they had two of the above or nuclear pyknosis (two points), and atretic (grade ≥3) if they had more than two of the above markers (Figure 1). Up to 50 primordial follicles were analysed within each tissue fragment, except for xenotransplants, where all primordial follicles were included in the analysis.

2.8. Data analysis
All statistical analyses were performed using R statistical software, version 3.5.0. A proportional odds model (clmm2; [20]) was used to determine whether delay before processing affected follicle health, adjusting for individual sheep (random effect). A logistic regression model (glmer; [21]) was used to determine the odds of follicles being classified as...
Effect of delayed processing on primordial follicle morphology in fresh ovarian tissue. Ovine ovarian tissue was either processed immediately (no delay; \(n = 6\)), 24 h (\(n = 3\)) or 48 h (\(n = 3\)) after collection and fixed in Bouin’s solution. (a) Representative images of fresh ovine ovarian tissue. Scale bar: 50 \(\mu\)m. (b) Health distribution of primordial follicles in fresh ovine ovarian tissue. Numbers at the base of columns refer to the total number of follicles analysed. Primordial follicles (oocyte surrounded by a single layer of flattened pregranulosa cells) were analysed using H&E staining and scored based on the presence of morphological markers (dense chromatin, eosinophilia, shrunken ooplasm [one point each] or pyknosis [two points]) as normal (0), affected (1), degenerating (2) or atretic (\(\geq 3\)). (c) Same data as in (b), showing the distribution of health grades between animals (mean ± SD). (d) Proportional ORs and 95% CIs of follicles being graded as \(\geq 1\) when processed after 24 h or 48 h compared to the no delay group.

3. Results

3.1. The effect of processing delay on fresh ovarian tissue

The vast majority of follicles in all fresh tissues were morphologically normal; 89.0 ± 8.7% for the immediately processed (no delay) group, 90.6 ± 3.0% for the 24 h-delay group, and 79.2 ± 14.2% for the 48 h-delay group (Figures 2(a)–2(c), Table 1). Compared to immediately processed samples, a 24 h delay did not result in an increased likelihood of primordial follicles being graded as morphologically abnormal (OR = 0.78, 95% CI 0.38–1.60, \(P > .05\)). In contrast, primordial follicles from samples processed 48 h after procurement were 2.5 times more likely to be graded as abnormal compared to no delay samples (OR = 2.47, 95% CI 1.29–4.71, \(P < .01\); Figure 2(d)).

3.2. The effect of processing delay on cryopreserved and thawed ovarian tissue

As expected, the process of cryopreservation caused damage to primordial follicles, with follicles in cryopreserved-thawed samples being over 28 times more likely to be graded as abnormal compared to fresh tissue (OR = 28.60, 95% CI 17.91–45.65, \(P < .0001\), Figure 3). Thus, the proportion of normal primordial follicles was lower in cryopreserved-thawed tissue compared to fresh tissue, irrespective of processing time (Figures 4(a)–4(c), Table 1). The mean proportion of normal follicles in cryopreserved-thawed tissue was 26.6 ± 17.0% for the no-delay group and
considerable deviation between individuals (Figure 4(c)).

normal primordial follicles, 45.7 ± 31.3% for the 24 h-delay group. Interestingly, tissue processed 48 h after collection had a higher proportion of normal primordial follicles, 45.7 ± 31.3%, though with considerable deviation between individuals (Figure 4(c)).

This translated to a 44% decrease in the odds of a follicle being graded as abnormal compared to immediately processed tissue (OR = 0.56, 95% CI 0.36–0.87, P < .01; Figure 4(d)). There was no difference in the health of primordial follicles after a 24 h delay before processing compared to immediately processed tissue (OR = 0.93, 95% CI 0.61–1.41, P > .05).

3.3. The effect of processing delay on ovarian tissue xenotransplanted after cryopreservation

Cryopreserved-thawed ovine ovarian tissue from 0 h, 24 h, and 48 h groups was xenografted subcutaneously into SCID mice for 10 days. The number of follicles found in each graft differed substantially across conditions; out of the six control grafts (immediately processed samples) only three contained follicles, with the remaining grafts containing potential degenerating follicle remnants (follicle-like structures with no oocyte; Table 2). Conversely, follicles were observed in two out of three grafts for the 24 h-delay group and all three grafts for the 48 h-delay group. Many primordial follicles were normal after 10-day xenografting, however, considerable variation was observed between grafts, particularly for the 48-h delay group. The proportion of normal primordial follicles was 55.7 ± 9.8% for the immediately processed group, 81.5 ± 3.5% for the 24-h delay group, and 43.0 ± 28.8% for the 48-h delay group (Figures 5(a)–5(c), Table 1). There was no difference in the likelihood of a primordial follicle being graded as abnormal for the 24 h-delay (OR = 0.36, 95% CI 0.081–1.59, P > .05) or 48 h-delay groups (OR = 5.0, 95% CI 0.80–30.83, P > .05) compared to the no delay group (Figure 5(d)).

Of the eight grafts containing follicles, six contained both primordial and growing follicles indicating that follicle activation occurred during the xenografting period (Table 2; Figure 6). The other two grafts that contained follicles

### Table 1: Summary of proportion of follicles per health grade (% ± SD) across conditions (no delay, 24 h delay, 48 h delay) in fresh, cryopreserved-thawed, and xenotransplanted ovarian tissue.

<table>
<thead>
<tr>
<th>Tissue type</th>
<th>Follicle health (grade)</th>
<th>No delay</th>
<th>24 h delay</th>
<th>48 h delay</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Fresh</strong></td>
<td>Normal (0)</td>
<td>89.0 ± 8.7</td>
<td>90.6 ± 3.0</td>
<td>79.2 ± 14.2</td>
</tr>
<tr>
<td></td>
<td>Affected (1)</td>
<td>10.1 ± 7.7</td>
<td>6.7 ± 3.0</td>
<td>18.1 ± 11.1</td>
</tr>
<tr>
<td></td>
<td>Grades 0 and 1</td>
<td>99.0 ± 1.1</td>
<td>97.3 ± 1.2</td>
<td>97.3 ± 3.1</td>
</tr>
<tr>
<td></td>
<td>Degenerating (2)</td>
<td>0.3 ± 0.8</td>
<td>1.3 ± 1.2</td>
<td>2.0 ± 2.0</td>
</tr>
<tr>
<td></td>
<td>Atretic (≥ 3)</td>
<td>0.7 ± 1.0</td>
<td>1.3 ± 1.2</td>
<td>0.7 ± 1.2</td>
</tr>
<tr>
<td><strong>Cryopreserved-thawed</strong></td>
<td>Normal (0)</td>
<td>26.6 ± 17.0</td>
<td>20.4 ± 2.1</td>
<td>45.7 ± 31.3</td>
</tr>
<tr>
<td></td>
<td>Affected (1)</td>
<td>41.6 ± 10.5</td>
<td>50.0 ± 8.5</td>
<td>27.5 ± 4.8</td>
</tr>
<tr>
<td></td>
<td>Grades 0 and 1</td>
<td>68.2 ± 15.3</td>
<td>70.5 ± 10.2</td>
<td>73.2 ± 28.0</td>
</tr>
<tr>
<td></td>
<td>Degenerating (2)</td>
<td>19.0 ± 9.4</td>
<td>23.5 ± 10.5</td>
<td>18.8 ± 19.4</td>
</tr>
<tr>
<td></td>
<td>Atretic (≥ 3)</td>
<td>12.8 ± 9.1</td>
<td>6.0 ± 3.4</td>
<td>8.0 ± 8.7</td>
</tr>
<tr>
<td><strong>Xenotransplanted</strong></td>
<td>Normal (0)</td>
<td>55.7 ± 9.8</td>
<td>81.5 ± 3.5</td>
<td>43.0 ± 28.8</td>
</tr>
<tr>
<td></td>
<td>Affected (1)</td>
<td>16.7 ± 16.5</td>
<td>8.5 ± 2.1</td>
<td>29.3 ± 13.5</td>
</tr>
<tr>
<td></td>
<td>Grades 0 and 1</td>
<td>71.0 ± 17.7</td>
<td>91.1 ± 7.6</td>
<td>73.2 ± 39.6</td>
</tr>
<tr>
<td></td>
<td>Degenerating (2)</td>
<td>28.0 ± 19.1</td>
<td>8.5 ± 7.8</td>
<td>4.7 ± 5.0</td>
</tr>
<tr>
<td></td>
<td>Atretic (≥ 3)</td>
<td>0.0 ± 0.0</td>
<td>1.0 ± 1.4</td>
<td>23.0 ± 39.8</td>
</tr>
</tbody>
</table>

Figure 3: Effect of cryopreservation of ovarian tissue on primordial follicle morphology. Proportional ORs and 95% CIs of follicles being graded as ≥ 1 in fresh tissue compared to cryopreserved-thawed tissue. Ovine ovarian tissue was processed immediately and either fixed or cryopreserved-thawed, and fixed (n = 6) in Bouin’s solution. Primordial follicles (oocyte surrounded by a single layer of flattened pregranulosa cells) were analysed using H&E staining and scored based on the presence of molecular markers (dense chromatin, eosinophilia, shrunken ooplasm [one point each] or pyknosis [two points]) as normal (0), affected (1), degenerating (2) or atretic (≥ 3). Black CI bars indicate P < .05.

20.4 ± 2.1% for the 24 h-delay group. Interestingly, tissue processed 48 h after collection had a higher proportion of normal primordial follicles, 45.7 ± 31.3%, though with considerable deviation between individuals (Figure 4(c)).
Figure 4: Effect of delayed processing on primordial follicle morphology in cryopreserved-thawed ovarian tissue. Ovine ovarian tissue was either processed immediately (no delay; \( n = 6 \)), 24 h (\( n = 3 \)) or 48 h (\( n = 3 \)) after collection, cryopreserved by slow freezing, thawed and fixed in Bouin’s solution. (a) Representative images of cryopreserved-thawed ovine ovarian tissue. Scale bar: 50 \( \mu m \). (b) Health distribution of primordial follicles in cryopreserved-thawed ovine ovarian tissue. Numbers at the base of columns refer to the total number of follicles analysed. Primordial follicles (oocyte surrounded by a single layer of flattened pregranulosa cells) were analysed using H&E staining and scored based on the presence of morphological markers (dense chromatin, eosinophilia, shrunken ooplasm [one point each] or pyknosis [two points]) as normal (0), affected (1), degenerating (2) or atretic (\( \geq 3 \)). (c) Same data as in (b), showing the distribution of health grades between animals (mean \( \pm SD \)). (d) Proportional ORs and 95% CIs of follicles being graded as \( \geq 1 \) when processed after 24 h or 48 h compared to the no delay group. Black CI bars indicate \( P < .05 \).

Table 2: Number of follicles per graft following 10-day xenotransplantation of cryopreserved-thawed ovine ovarian tissue with delayed processing.

<table>
<thead>
<tr>
<th>Animal</th>
<th>No delay</th>
<th>24h delay</th>
<th>48h delay</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sheep 1</td>
<td>25 follicles (14 primordial, 56%)</td>
<td>27 follicles (14 primordial, 52%)</td>
<td>N/A</td>
</tr>
<tr>
<td>Sheep 2</td>
<td>No follicles</td>
<td>154 follicles (86 primordial, 56%)</td>
<td>N/A</td>
</tr>
<tr>
<td>Sheep 3</td>
<td>6 follicles (6 primordial, 100%)</td>
<td>No follicles</td>
<td>N/A</td>
</tr>
<tr>
<td>Sheep 4</td>
<td>No follicles</td>
<td>N/A</td>
<td>75 follicles (50 primordial, 67%)</td>
</tr>
<tr>
<td>Sheep 5</td>
<td>No follicles</td>
<td>N/A</td>
<td>32 follicles (16 primordial, 50%)</td>
</tr>
<tr>
<td>Sheep 6</td>
<td>2 follicles (2 primordial, 100%)</td>
<td>N/A</td>
<td>55 follicles (26 primordial, 47%)</td>
</tr>
<tr>
<td>Total</td>
<td>33 (22 primordial)</td>
<td>181 (100 primordial)</td>
<td>162 (92 primordial)</td>
</tr>
</tbody>
</table>
Figure 5: Effect of delayed processing on primordial follicle morphology in cryopreserved-thawed xenotransplanted ovarian tissue. Ovine ovarian tissue was either processed immediately (no delay; n = 6), 24 h (n = 3) or 48 h (n = 3) after collection and cryopreserved. Samples were thawed and xenotransplanted subcutaneously for 10 days, after which they were fixed in Bouin’s solution. (a) Representative images of cryopreserved-thawed ovine ovarian tissue. Scale bar: 50 μm. (b) Health distribution of primordial follicles in cryopreserved-thawed ovine ovarian tissue. Numbers at the base of columns refer to the total number of follicles analysed. Primordial follicles (oocyte surrounded by a single layer of flattened pregranulosa cells) were analysed using H&E staining and scored based on the presence of morphological markers (dense chromatin, eosinophilia, shrunken ooplasm [one point each] or pyknosis [two points]) as normal (0), affected (1), degenerating (2) or atretic (≥3). (c) Same data as in (b), showing the distribution of health grades between animals (mean ± SD). (d) Proportional ORs and 95% CIs of follicles being graded as ≥1 when processed after 24 h or 48 h compared to the no delay group.

4. Discussion
This study demonstrates that a processing delay of 24 h at 4 °C did not impact the morphology of the primordial follicles which is indicative of health in fresh or cryopreserved ovine ovarian tissue. In contrast, a processing delay of 48 h resulted in primordial follicles being more likely to be graded as abnormal compared to immediately-processed tissue, while tissue cryopreserved after 48 h was more likely to contain morphologically normal follicles compared to the immediately-processed control. Xenografting of cryopreserved ovarian strips indicated the tissue can support follicle health and development in all experimental groups, with the delay in processing resulting in no difference in the health of primordial follicles, nor in the proportion of growing follicles.

Here we reported that cryopreservation had a significant negative impact on primordial follicle health, with only 20–40% of follicles being classified as normal. Health
Figure 6: Follicle development after xenotransplantation of ovarian tissue with or without delayed processing. Ovine ovarian tissue was either processed immediately (no delay; \( n = 6 \)), 24 h \( (n = 3) \) or 48 h \( (n = 3) \) after collection and cryopreserved. Samples were thawed and xenotransplanted subcutaneously for 10 days, after which they were fixed in Bouin’s solution. (a) Proportion of primordial and growing follicles in cryopreserved-thawed ovine ovarian tissue following 10-day xenotransplantation. Follicles were classified as primordial (P0; oocyte surrounded by a single layer of flattened pregranulosa cells) or growing (Grow; oocyte surrounded by one or more layers of either a mixture of flattened and cuboidal granulosa cells or a complete layer of cuboidal cells. (b) Same data as in (a), showing the distribution of follicle developmental stages between animals (mean ± SD). (c) Proportional ORs and 95% CIs of follicles being classified as growing when processed after 24 h or 48 h compared to the no delay group.

classification criteria vary considerably between studies, and the criteria used in the present study were deliberately strict, in order to detect any discrete morphological changes within the follicles. Others have reported 50–80% normal follicles following cryopreservation, which is consistent with our report if the classification criteria to judge the health of the follicles in this study are adjusted to match [22, 23]. We acknowledge that the results of the present study are somewhat limited by the use of morphology alone to assess primordial follicle health. However, the criteria used were well-defined and, in studies using a combination of morphological and molecular analyses, the two methods are generally in agreement [24]. While the use of molecular markers may have complemented the results presented here, they present their own limitations including challenges in selecting an appropriate marker for the damage mechanism in question, in addition to the transience of some marker expression [25].

A 24 h processing delay had no significant effect on primordial follicle health post-cryopreservation and thawing. This finding is consistent with the presence of morphologically normal primordial follicles after a 20 h processing delay at 4 °C of human ovarian cortical tissue as reported by Rosendahl et al. [12]. Isachenko et al. also provided evidence that a 24 h cryopreservation delay at 4 °C does not affect the proportion of morphologically normal follicles or follicle density in frozen-thawed ovarian cortical tissue [13]. Conversely, another study described a decrease in the proportion of morphologically normal follicles induced by a 24 h freeze-delay prior to thawing and in vitro culture of human ovarian tissue [14], however, this study used vitrification, which relies on very high concentrations of cryoprotectants, while the present study employed slow freezing which is the currently accepted gold-standard method of OTC [26].

Remarkably, a processing delay of 48 h resulted in a 44% decrease in the probability of follicles being graded as affected, degenerating or atretic after cryopreservation, despite showing the opposite effect in fresh tissue. This is consistent with findings from Isachenko et al. who explored a 24 h delay at 4 °C suggesting a role for pre-cooling of human ovarian tissue before cryopreservation [13]. It is possible that acclimatisation of the ovarian tissue at 4 °C for 48 h provides a more gradual temperature transition, which primes the tissue to better respond to the challenges of cryopreservation and, thus, experience a reduced insult on the health of the primordial follicle pool. However, as there was significant variability in morphological follicle health distribution between the three biological replicates in response to a 48 h delay, confirming this result with human ovarian tissue is required before such delays can be used for fertility preservation.

At present, the only way to utilise cryopreserved ovarian cortical strips for fertility restoration is autologous transplantation of the tissue either orthotopically or heterotopically. Although morphological analysis after
cryopreservation and thawing granted important insight into the effects of processing delay on primordial follicle health, the role of the tissue in fertility restoration is ultimately defined by its potential to sustain follicle health and growth after transplantation. Therefore, in this study we used xenotransplantation to determine if the delay in OTC had modified functional potential. Importantly, all treatment groups contained grafts that supported follicle development. Nevertheless, there was significant variation in follicle numbers between grafts, with some grafts containing no follicles at all. The variable number of follicles in the grafts was accounted for in the statistical model used. Total absence of follicles was observed in 3 out of 6 grafts that were processed immediately and 1 out of 3 24-h-delay grafts, but all contained degenerating follicle remnants. Most surprisingly, statistical analysis revealed that delayed processing of up to 48 h did not adversely affect follicle development in cryopreserved-transplanted tissue, as primordial and growing follicles were observed in all three of the 48-h-delay grafts. Thus, the presence of follicles in 5 out of 6 grafts confirms that such a delay does not render the tissue deficient in follicles capable of growth, nor does it hinder revascularisation of the tissue, a key component to the success of transplantation. By extension, this data also suggests that a processing delay of 24 h or 48 h did not cause complete loss of function of the vasculature and the supporting stromal cells. This finding is consistent with insight from other studies that also identified follicles in ovarian tissue xenotransplanted after a 24 h delay in cryopreservation [12,13]. Moreover, these results demonstrate that a processing delay of either 24 h or 48 h had no significant impact on primordial follicle health, nor on the proportion of growing follicles after xenotransplantation indicating that delayed processing does not affect the potential for follicle development following transplantation. To the best knowledge of the authors, the work presented here is the first report of normal morphology of the tissue. This, taken together with studies using human tissue, which demonstrated that follicles can survive a processing delay of up to 20 h at 4 °C and lead to a live birth [8, 12, 13], indicates that ovarian tissue responds to delayed cryopreservation favourably without a statistically significant negative impact on the primordial pool.

The results we present here have provided strong evidence to support the idea that ovarian tissue can be chilled for 24 or even 48 h prior to cryopreservation with no statistically significant net negative impact on the health of the primordial pool of follicles after transplantation and in vivo development of the tissue. This lends strength to an emerging body of evidence suggesting that the delayed processing is safe, or even beneficial, for cryopreserved ovarian tissue. Validation of these findings using human ovarian tissue or other endangered species of interest would thereby present a new, feasible approach to increasing access to ovarian tissue freezing for fertility preservation. It should be noted that in the present study ovaries were left intact until the time of processing, therefore, it cannot be concluded that ovarian biopsies would have a similar response. Nevertheless, these findings are of high relevance to fertility preservation and, in particular, the way OTC is organised on a local and national level for humans and endangered species. An update of the current guidelines to reflect an extended window for tissue processing will ensure that the modality is more widely available to patients without the need to create new facilities. Fundamentally, such an advancement is a great step towards ensuring that every cancer patient who has a need, has an option.

Authors’ contributions M. S. Zemyarska and B. D. Bjarkadottir contributed equally to this work.

Acknowledgments The authors wish to thank all staff of the Oxford Cell and Tissue Biobank (OCTB) for their assistance with training in tissue processing and cryopreservation, Prof. David Bennett for use of the surgery facilities, and all staff of Biomedical Sciences John Radcliffe for assistance with the mice used in this study.

Conflict of interest The authors declare that they have no conflict of interest.

References