Successful transfer of day 10 horse embryos: influence of donor–recipient asynchrony on embryo development

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Abstract

A total of 78 day 10 horse embryos were transferred non-surgically to recipient mares that had ovulated 9, 7, 6, 5, 4, 3, 2 or 1 day after (negative asynchrony), on the same day (synchronous), or 2 or 4 days before (positive asynchrony) the donor (n = 6 or 8 mares per group). Pregnancy rates between 100% (6/6) and 63% (5/8) were seen in recipient mares that were between +2 and −6 days asynchronous. Embryo survival to the heartbeat stage declined in recipients that were −7 days asynchronous and no embryos survived in recipients that were −9 days asynchronous. Irrespective of uterine asynchrony, cessation of embryo mobility and fixation at the base of a uterine horn occurred when the conceptus was ~17 days old. Conceptus growth and development was slowed when embryos were placed in negatively asynchronous uteri. At the greatest degree of negative asynchrony at which embryos survived to the heartbeat stage, i.e. −7 and −6 days, development of the embryo proper and allantois was retarded. Luteostasis was achieved in recipient mares when day 10 embryos were transferred to recipient mares at any stage of asynchrony between −9 and +2 days with respect to the donor. These results indicate that in the horse, there is a wide window for establishment of pregnancy following embryo transfer to asynchronous recipients. Although progesterone priming of the uterus to a stage equivalent to that of the transferred embryo does not appear to be a prerequisite for embryo survival, it does nonetheless influence embryonic development.

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Introduction

It is a generally accepted tenet of equine embryo transfer that optimum pregnancy rates are achieved when transferring embryos to recipient mares that have ovulated 1 day before to 2 days after the donor (denoted as +1 to −2 days asynchrony: Allen 1982, Squires et al. 1982, McCue & Troedsson 2003, Stout 2003, 2006). Work in other large animal species has shown that embryos placed in an asynchronous uterine environment may be exposed to inappropriate levels of growth factors, secreted proteins and steroid hormones so that the rate of embryonic growth and development is altered or embryonic death occurs (see Pope 1988, Barnes 2000 for reviews).

Current practice in commercial embryo transfer is to flush the uterus of a donor mare on day 7 or 8 after ovulation; flushing before this stage results in significantly lower embryo recovery rates (Boyle et al. 1989) due to the equine embryo’s 6-day sojourn in the oviduct before it enters the uterine lumen between 156 and 168 h after ovulation (Battut et al. 1997, 2001). Older day 9 or 10 equine embryos have been widely considered too large and fragile to survive routine embryo transfer procedures. For example, Squires et al. (1982) reported that more mares became pregnant following the transfer of day 8 embryos (32%) than day 9 embryos (9%). Similarly, Vogelsang et al. (1985) obtained no pregnancies following the transfer of day 9 or 10 embryos, compared to a 61% pregnancy rate following transfer of day 6 embryos, 55% from day 7 embryos and 25% from day 8 embryos. Sirois et al. (1987) achieved no pregnancies when they transferred 4 day 10 horse embryos to recipients that were +1 to −1 days asynchronous with the donor, whereas Fleury et al. (1989) achieved a 69% pregnancy rate when they transferred 16 day 9 horse embryos. In attempting to explain these variable results, Squires et al. (1985) suggested that the increased fluid volume-to-surface area ratio of days 9 and 10 embryos made them more prone to damage during the collection and transfer procedures. However, it must be borne in mind that all these earlier studies were undertaken before the technique of non-surgical embryo transfer had become widely used and operators had acquired much practical experience.

Transfer of day 10 embryos to asynchronous recipient mares could benefit the management of commercial equine transfer programmes and may provide a useful model for investigation of maternal influences on
early embryogenesis. We reported recently on the successful transfer of day 10 horse embryos to 7 days asynchronous recipient mares (Wilsher & Allen 2009), which clearly demonstrated an overriding influence of uterine environment on embryonic growth and early placentation. This paper extends that report and describes the rates of embryo survival following transfer of day 10 embryos to recipients that had ovulated from 9 days after to 4 days before the donor mares and discusses the implication of such asynchronous transfers on embryonic growth and conceptus development.

Results

Embryo recovery

Prior to the embryo flush carried out on day 10, ultrasound scanning of the uterus revealed the presence of an embryo or embryos in 64/121 (53%) of the donor mares; all these embryos measured ≥ 3 mm in diameter. In 5/121 (4%) instances, the subsequent flushing of the mare’s uterus produced an embryo > 3 mm diameter that had not been visualised ultrasonographically. In 3/121 (2%) mares, an embryo of > 5 mm diameter was seen ultrasonographically but it proved impossible to recover that embryo despite repeated flushing of the uterus.

Figure 1 Examples of day 10 embryos recovered from donor mares. (a) Capsule material recovered from an embryo that was ruptured during the flushing procedure; (b) collapse of the trophoblast within an intact capsule of a day 10 embryo, damaged during flushing; (c) an abnormal, degenerating embryo, one of twins. The other embryo was normal and was transferred successfully.
An embryo or embryos were recovered from 94/121 (78%) flushes; two embryos were recovered from 16 of the flushes to yield a total of 110 embryos. Of these, 9/110 (8%) were broken and either the capsule had ruptured (Fig. 1a) or the trophoblast had collapsed within the intact capsule (Fig. 1b); a further 6/110 (5%) embryos were of poor quality (Fig. 1c). The 78 embryos that were transferred exhibited a mean diameter of 3.89 ± 0.14 mm (range 1.80–7.00 mm). All embryos, including multiple embryos from the same donor, were allocated randomly to the recipient mares.

**Embryo survival**

All inseminated control mares were diagnosed pregnant 12 days after ovulation (9/9; 100%), and all went on to develop an embryo proper with a heartbeat (9/9; 100%). Pregnancy rates and conceptus viability, judged by an embryonic heartbeat, did not differ between the control mares and the synchronous transfers. The numbers of recipient mares in which a) an embryonic vesicle was present at the first ultrasound examination 2 days after transfer, b) a vesicle was still present at 10 days post transfer and c) a viable embryo with a heartbeat was seen ultrasonographically are detailed in Table 1. None of the embryos transferred to recipient mares with a −9 day asynchrony survived to the heartbeat stage, although anembryonic vesicles persisted until days 23–25 post transfer in two of these recipients. Similarly, three embryonic vesicles failed to develop an embryo proper in recipients with a −7 day asynchrony, and two more anembryonic vesicles were seen in recipients with an asynchrony of −4 and −2 days respectively.

Transfer of day 10 embryos to a uterine environment that was 2 days advanced resulted in survival of all six transferred embryos to the heartbeat stage. In contrast, none of the six embryos transferred to recipient mares with a +4 day asynchrony survived. Nevertheless, three of these six were seen ultrasonographically 2 days after transfer despite obvious oedema in the endometrium and a serum progestagen level of <1.0 ng/ml in these recipients; the vesicles had disappeared 2 days later.

Despite elevated serum progesterone concentrations, transient endometrial oedema was noted between days 16 and 18 after ovulation in three of the six +2 recipient mares. All six of the recipient mares with a +4 day asynchrony in which no embryos survived spontaneously returned to oestrus and, hence, showed significantly lower mean serum progesterone concentrations 2 and 4 days post transfer compared to all the other groups of recipient and control mares (P < 0.001). However, there were no differences in mean serum progesterone concentration profiles between the recipient mares in which the transferred embryos survived versus those in which the embryo failed (P > 0.05 in all cases). Furthermore, mean progesterone concentration profiles did not differ between any group (recipient or control) in which a pregnancy was established (P > 0.05 in all cases). Table 2 shows the mean ± S.E.M. serum progesterone concentrations of the recipient mares on the day of embryo transfer and those of the control mares at day 10 of gestation. Recipient mares with a −9 day asynchrony had a significantly lower mean serum progesterone concentration on the day of transfer compared to all other groups (P < 0.05), with the exception of the −7 and +4 day asynchronous recipient groups.

Figure 2a shows the mean embryonic vesicle diameters of the control (non-transferred embryos derived from artificial insemination) and transferred embryos that survived until conceptus day 22 (recipient days 15–24), in relation to the level of asynchrony of the recipient

### Table 1: Influence of donor–recipient asynchrony on pregnancy rates and early embryonic loss rates following transfer of day 10 embryos.

<table>
<thead>
<tr>
<th>Degree of donor–recipient asynchrony (days)</th>
<th>Days after ovulation of the recipient on day of ET</th>
<th>2 days post ET</th>
<th>10 days post ET</th>
<th>With embryonic heartbeat</th>
</tr>
</thead>
<tbody>
<tr>
<td>+4</td>
<td>14</td>
<td>3/6 (50%)</td>
<td>0/6 (0%)</td>
<td>0/6 (0%)</td>
</tr>
<tr>
<td>+2</td>
<td>12</td>
<td>6/6 (100%)</td>
<td>6/6 (100%)†,‡</td>
<td>6/6 (100%)†,‡</td>
</tr>
<tr>
<td>0</td>
<td>10</td>
<td>6/6 (100%)</td>
<td>6/6 (100%)†,‡</td>
<td>6/6 (100%)†,‡</td>
</tr>
<tr>
<td>−1</td>
<td>9</td>
<td>6/6 (75%)</td>
<td>5/8 (63%)†,‡</td>
<td>5/8 (63%)†,‡</td>
</tr>
<tr>
<td>−2</td>
<td>8</td>
<td>6/6 (75%)</td>
<td>6/6 (75%)†,‡</td>
<td>6/6 (75%)†,‡</td>
</tr>
<tr>
<td>−3</td>
<td>7</td>
<td>6/6 (75%)</td>
<td>6/6 (75%)†,‡</td>
<td>6/6 (75%)†,‡</td>
</tr>
<tr>
<td>−4</td>
<td>6</td>
<td>8/8 (100%)</td>
<td>8/8 (100%)†,‡</td>
<td>7/8 (83%)†,‡</td>
</tr>
<tr>
<td>−5</td>
<td>5</td>
<td>6/6 (75%)</td>
<td>6/6 (75%)†,‡</td>
<td>6/6 (75%)†,‡</td>
</tr>
<tr>
<td>−6</td>
<td>4</td>
<td>5/6 (83%)†</td>
<td>5/6 (83%)†</td>
<td>5/6 (83%)†,‡</td>
</tr>
<tr>
<td>−7†</td>
<td>3</td>
<td>8/8 (100%)</td>
<td>6/6 (75%)†,‡</td>
<td>3/8 (38%)†,‡</td>
</tr>
<tr>
<td>−9</td>
<td>1</td>
<td>4/6 (67%)</td>
<td>2/6 (33%)†</td>
<td>0/6 (0%)†</td>
</tr>
<tr>
<td>Control</td>
<td>Al derived, non-ET pregancies</td>
<td>9/9 (100%)</td>
<td>9/9 (100%)†</td>
<td>9/9 (100%)†</td>
</tr>
</tbody>
</table>

*Values with different superscripts within a column are significantly different at P < 0.05.
†A full description of these transfers and the subsequent embryonic growth was reported by Wilsher & Allen (2009).
mare, while Fig. 2b shows the same data plotted in relation to embryonic age. Most groups showed a typical growth profile for an equine embryo; namely, exponential expansion of the vesicle between days 10 and 16 followed by a plateau in growth after this time. However, the embryos in both the −7 and −6 days asynchronous recipient groups continued to expand beyond conceptus day 18 (recipient day 11 or 12). Although the control embryos showed the lowest mean vesicle diameter between days 18 and 22, vesicle diameter did not differ significantly from the diameters of embryos transferred to recipient mares that had ovulated between 5 days before (−) and 2 days after (+) the donor (P>0.05). The mean diameters of embryos on days 2, 4 and 6 after transfer were significantly lower in recipient mares with a

Table 2 The mean ± S.E.M. serum progesterone concentrations of recipient mares on the day of embryo transfer and those of the control mares at day 10 of gestation.

<table>
<thead>
<tr>
<th>Embryo transfer recipients: days after ovulation when transfer of a day 10 embryo occurred (a)</th>
<th>AI-derived pregnancies on day 10 of gestation (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control (9)</td>
<td>1 (6)</td>
</tr>
<tr>
<td>2 (6)</td>
<td>3 (8)</td>
</tr>
<tr>
<td>4 (6)</td>
<td>5 (8)</td>
</tr>
<tr>
<td>6 (8)</td>
<td>7 (8)</td>
</tr>
<tr>
<td>8 (8)</td>
<td>9 (8)</td>
</tr>
<tr>
<td>10 (6)</td>
<td>12 (6)</td>
</tr>
<tr>
<td>14 (6)</td>
<td></td>
</tr>
<tr>
<td>1.0 ± 0.7</td>
<td></td>
</tr>
<tr>
<td>7.8 ± 0.7</td>
<td></td>
</tr>
<tr>
<td>11.1 ± 0.2</td>
<td></td>
</tr>
<tr>
<td>4.8 ± 0.5</td>
<td></td>
</tr>
<tr>
<td>6.7 ± 0.5</td>
<td></td>
</tr>
<tr>
<td>9.1 ± 0.9</td>
<td></td>
</tr>
<tr>
<td>8.0 ± 0.9</td>
<td></td>
</tr>
<tr>
<td>8.0 ± 0.8</td>
<td></td>
</tr>
<tr>
<td>9.0 ± 1.3</td>
<td></td>
</tr>
<tr>
<td>8.3 ± 1.3</td>
<td></td>
</tr>
<tr>
<td>7.5 ± 1.4</td>
<td></td>
</tr>
<tr>
<td>7.5 ± 1.5</td>
<td></td>
</tr>
<tr>
<td>6.7 ± 1.0</td>
<td></td>
</tr>
<tr>
<td>5.6 ± 1.0</td>
<td></td>
</tr>
</tbody>
</table>

Values with different superscripts within a row are significantly different at P<0.05.

Figure 2 Embryonic vesicle diameters of transferred and control embryos. (a) The mean embryonic vesicle diameters of the non-transferred control and transferred embryos that survived until day 22 (control, n=9; −4 days asynchrony, n=7; +2, 0, −3 and −5 asynchrony, n=6; −1, −2 and −6 days asynchrony, n=5; −7 days asynchrony, n=3; anembryonic vesicles are not included) in relation to the level of asynchrony of the recipient mare; (b) mean embryonic vesicle diameter in relation to embryonic age. The S.E.M. of mean vesicle diameters ranged from 0.3 to 3.2. The diameters of transferred embryos on day 10 were obtained from in vitro measurement prior to transfer. All other values were derived from ultrasound measurements made in vivo in the recipient or control mares.
—7 day asynchrony compared to all other groups with an asynchrony of —5 days or less (P≤0.01). Despite the initial slowing in growth rate in embryos transferred to uteri with a —7 or —6 day asynchrony, expansion continued so that by 12 days post transfer they showed a significantly greater mean diameter than either the control or the synchronously transferred embryos (P≤0.01, in both cases).

Conceptus development

Figure 3 illustrates conceptus development, as viewed ultrasonographically, of day 10 embryos transferred to recipient mares with varying degrees of asynchrony. Table 3 shows the mean ± S.E.M. day for recognisable events associated with early development in embryos transferred to recipient mares at varying stages of asynchrony, compared to control pregnancies.
Uterine–embryo asynchrony did not affect the age at which the embryo became fixed at the base of one of the uterine horns ($P=0.887$). However, with increasing negative asynchrony, the mean conceptus age at which disproportionate thickening of the endometrial folds in the mesometrial quadrant of the uterus was noted, and the conceptus age when an embryo with a discernable heartbeat first appeared, increased such that both events occurred at a significantly advanced age in embryos transferred to uteri with a $\geq 5$ day asynchrony compared both to recipients with an asynchrony of $-2$ to $+2$ days and control, inseminated mares ($P\leq0.001$, in all cases; Table 3). No enhancement of embryonic growth occurred in embryos transferred to an advanced uterine environment ($+2$ day asynchrony).

Development of the five surviving embryos transferred to recipient mares with a $-6$ day asynchrony is illustrated in Figs 3a–e and 4a–t. As previously reported for the day 10 embryos transferred to $-7$ day asynchronous uteri (Wilsher & Allen 2009), embryos transferred to uteri with an asynchrony of $-6$ days showed significantly delayed development compared to Al-derived or synchronously transferred embryos. For example, the tiny echogenic spot believed to represent the embryo proper was often noted in the ventral quadrant of the vesicle between conceptus days 20 and 24 (recipient days 14 and 18), but this was not seen again until the appearance of a much larger embryo more than 4 days later (Figs 3c–e and 4f–h, j–l and m–p). This was coupled with a delayed appearance, and then with rapid expansion, of the allantois between conceptus days 28 and 32 (recipient days 22 and 26). This phenomenon was exemplified in two of these mares in which, despite careful ultrasonographic examinations, an embryo proper could not be visualised (Fig. 4k–l and o–p).

Discussion

In conventional embryo transfer programmes, horse embryos are usually transferred to recipient mares that have ovulated 1 day before to 3 days after the donor. When the recipient mare has ovulated $\geq 2$ days ahead of the donor, pregnancy rates are reduced (Allen 1982, Squires et al. 1982, McCue & Troedsson 2003, Stout 2003, 2006) and early pregnancy losses are increased (Carnevale et al. 2000). Two major reasons are put forward for this decline in pregnancy rate following transfer to recipients that ovulate ahead of the donor. First, the uterine environment changes under the dominance of progesterone (Zavy et al. 1982, Hinrichs et al. 1989), so that an embryo transferred to a positively asynchronous uterus may be exposed to an inappropriate environment (Barnes 2000). Secondly, it is believed that the equine blastocyst must transmit its antiluteolytic maternal recognition of pregnancy signal before day 10 after ovulation if it is to suppress the upregulation of oxytocin receptors and prevent luteolysis (Goff et al. 1987, Stout et al. 1999, 2000). Evidence from cattle suggests that it is difficult to inhibit upregulation of oxytocin receptors once the process has commenced (Gilbert et al. 1989). Furthermore, recipient mares that ovulate 2 or 3 days ahead of the donor do not maintain the pregnancy even if their return to oestrus is prevented by the administration of exogenous progesterone (Clarke et al. 1987, Pool et al. 1987) with pregnancy failure presumably resulting from the advanced uterine

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**Table 3** Influence of donor–recipient asynchrony on embryo fixation, endometrial thickening and the appearance of a heartbeat in transferred day 10 embryos.

<table>
<thead>
<tr>
<th>Mare groups</th>
<th>Number of mares</th>
<th>Embryo fixation</th>
<th>Loss of spherical shape plus endometrial thickening</th>
<th>Appearance of a heartbeat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Degree of asynchrony donor:recipient (days)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$+2$</td>
<td>6</td>
<td>17.7±0.33</td>
<td>18.7±0.42*</td>
<td>21.7±0.42*</td>
</tr>
<tr>
<td>0</td>
<td>6</td>
<td>17.3±0.42</td>
<td>18.3±0.43*</td>
<td>21.3±0.42*</td>
</tr>
<tr>
<td>$-1$</td>
<td>5</td>
<td>17.6±0.40</td>
<td>19.6±0.75*</td>
<td>22.8±0.49*</td>
</tr>
<tr>
<td>$-2$</td>
<td>5</td>
<td>17.6±0.40</td>
<td>19.6±0.52*</td>
<td>23.4±0.89*</td>
</tr>
<tr>
<td>$-3$</td>
<td>6</td>
<td>17.3±0.42</td>
<td>20.0±0.52*</td>
<td>23.0±0.45*</td>
</tr>
<tr>
<td>$-4$</td>
<td>7</td>
<td>17.7±0.29</td>
<td>21.4±0.43*</td>
<td>23.7±0.29*</td>
</tr>
<tr>
<td>$-5$</td>
<td>6</td>
<td>16.7±0.42</td>
<td>22.3±0.33*</td>
<td>25.0±0.45*</td>
</tr>
<tr>
<td>$-6$</td>
<td>5</td>
<td>17.2±0.49</td>
<td>23.6±0.75*</td>
<td>28.8±0.67*</td>
</tr>
<tr>
<td>$-7$</td>
<td>3</td>
<td>17.3±0.67</td>
<td>22.7±0.67*</td>
<td>27.3±0.67*</td>
</tr>
<tr>
<td>Control</td>
<td>9</td>
<td>17.1±0.35</td>
<td>19.1±0.35*</td>
<td>21.6±0.29*</td>
</tr>
</tbody>
</table>

*Values with different superscripts within a column are significantly different at $P<0.05$. ET, embryo transfer. Ov, day of ovulation (day 0). Ultrasound examinations were performed only on alternate days and, hence, observations recorded on any single day could be subject to an error of ±1 day. The number of mares in which the embryo survived until the heartbeat stage; anembryonic vesicles are not included. Embryo fixation was defined as the first day that the vesicle was located at the base of the uterine horn and noted to be in the same location on all subsequent examinations.
Successful transfer of day 10 horse embryos

environment in which the embryo finds itself. However, our findings that all the day 10 embryos transferred to recipient mares that were either 10 or 12 days post ovulation (0 or +2 day asynchrony) survived suggests the former dogma should be re-evaluated since the luteolytic cascade is either stoppable once initiated on day 10 or, more likely, the cascade does not actually begin until after day 12 post ovulation. By day 14 post ovulation, the luteolytic cascade in the mare has clearly commenced and cannot be halted since all six recipients at this stage into which a day 10 embryo was transferred spontaneously returned to oestrus, even in those mares in which an embryonic vesicle was still ultrasonographically visible.

At the negative end of the window of asynchrony investigated in the present experiment, day 10 embryos transferred to recipients with an asynchrony of up to −9 days suppressed luteolysis. This occurred regardless of the developmental competence of the conceptus, both in terms of its complete lack of development (i.e. an anembryonic or trophoblastic vesicle) or delayed development of the embryo and membranes. The ability of trophoblastic vesicles to suppress cyclicity was not unexpected since Ball et al. (1989) had reported luteal maintenance following the transfer of trophoblastic vesicles to day 10 recipients. Although the delayed development of the embryo and membranes in the present experiment could have resulted in the luteostatic signal being delayed, it was nonetheless surprising to find that at the time when maternal recognition of pregnancy is assumed to occur (i.e. days 10–16 in the recipient mare) the transferred conceptus, although retarded was, in some instances, 19–25 days old. Furthermore, although previous work has suggested that uterine migration of the embryo until the time of fixation is essential for prevent luteolysis (McDowell et al. 1988, Griffin & Ginther 1993), all of the transferred embryos became fixed when they were ~17 days old regardless of the degree of donor–recipient asynchrony. Hence, embryo movement ceased in some recipients when the uterus was only 9 days

Figure 4 Conceptus development between days 20 and 32 of gestation in a pregnancy achieved by artificial insemination (a–d) compared with that of day 10 embryos transferred to four different recipient mares with a −6 day asynchrony (e–h, i–l, m–p and q–t). Note the tiny echogenic spot denoting the embryo (E) in Mare BB (f), Mare HM (j) and Mare LaC (m) and its slow development compared to the control pregnancy (a–d). At 32 days of gestation, the control conceptus shows enlargement of the allantoic cavity (AC) and regression of the yolk sac (YS) such that the former represents over half of the conceptus (d). By comparison, in the recipient mares carrying the asynchronously transferred embryos the allantoic cavity (AC) occupied less than a third of the conceptus (h, l, p and t). P, embryo age; Ov, days post ovulation in the recipient or control mare. Caliper dots on any line are 0.5 cm apart; scale bar = 1 cm.

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old yet luteostasis still occurred. In a similar vein, Rivera del Alamo et al. (2008) demonstrated that the insertion of a relatively immobile 20 mm water-filled plastic ball into the uterine lumen of cycling mares 2–4 days after ovulation prolonged luteal function in 9 of 12 treated animals, suggesting embryo mobility may not be essential for prolongation of luteal function.

Although uterine tone has also been associated with conceptus fixation (Feo 1980, Ginther 1983a, Griffin & Ginther 1991, Carnevale & Ginther 1992), and no doubt plays a role, intraluminal impediment, with or without increased uterine tone, would appear to be the overriding influence on fixation time since mobility of the transferred embryos ceased at the same conceptus age regardless of the stage of the uterus. This accords with previous work that demonstrated that fixation occurs earlier for larger embryos (Gastal et al. 1996). In recipient mares with the greatest degree of asynchrony in which embryos survived (i.e. –6 and –7 days with respect to the donor), the embryos continued to increase in diameter after fixation around embryo day 17 (recipient day 10 or 11). Turgidity of the normal pregnant uterus increases steadily from day 10 or 11 after ovulation until day 16 (Bonafas et al. 1994), when the now intense uterine tone is believed to be responsible for the plateau in conceptus diameter which occurs between approximately days 16 and 26 (Ginther 1998). Hence, the reduced uterine tone in the day 10 or 11 recipient uteri could have allowed the day 17 conceptuses to continue to remain spherical and expand until uterine tone increased. The delay in uterine tone in relation to conceptus age was further exemplified by the later occurrence of endometrial thickening in the mesometrial quadrant and the loss of a spherical outline seen in the transferred conceptuses with the greatest degree of donor–recipient asynchrony.

Under conditions of embryo–uterus asynchrony in other animal species, a variety of complications can occur, including failure of the embryo to implant, early embryonic mortality or retarded or accelerated growth (see Pope 1988, Barnes 2000 for reviews). For example, sheep embryos transferred to a more advanced uterus increase their rate of cell division (Wilmut et al. 1985) and slow this rate when transferred to a less advanced uterus (Wilmut & Sales 1981, Lawson et al. 1983, Wilmut et al. 1986). In the present experiment, an advanced uterine environment in the mare was not associated with any observable enhancement in embryonic development although, as previously observed in the sheep, embryonic development was modified when embryos were exposed to a retarded uterine environment and was fatally modified in 63% of the embryos transferred to –7 days asynchronous uteri and 100% of embryos transferred to uteri that were –9 days asynchronous.

The observed delay in the development of the embryo proper and the allantois in the pregnancies at the greatest degree of negative asynchrony is difficult to explain. In equids, in the normal course of events, the allantois first appears around days 20–21 after ovulation (van Niekerk & Allen 1975, Latshaw 1987), as an evagination from the hindgut of the embryo. Its emergence is critical for the establishment of the chorioallantoic placenta, and it plays a vital role in vascularising the chorion. Furthermore, failures in the development of the allantois have been cited as a potential cause of embryonic loss in both sheep (Robinson 1951) and cattle (Peterson et al. 2000). In ruminant species, a temporal expression of genes associated with the emergence of the allantois has been identified (Ledgard et al. 2006), and it is likely that a similar stage-dependent regulation of embryonic development occurs in the horse. The role of maternal environment in this regulation is not fully understood, although in mammals that alter their gestation length by employing embryonic diapause, it is uterine factors that induce and maintain the embryo in its developmental arrest. Suppression of cell proliferation at the blastocyst stage occurs in most of the species that undergo diapause but the bat is a notable exception to this general rule in which variations in the rate of later embryonic development have been noted (Rasweiler & Badwaik 1997). Ultrasonographically, this appeared to be the case in the affected mares in the present experiment.

It is unclear whether embryonic diapause is due to the absence of uterine factors necessary for the development of the blastocyst or whether the uterus actively maintains diapause by inhibition of development (see Lopes et al. 2004 for review). Support for the former view can be found in studies that have shown increases in protein synthesis and secretion coinciding with the termination of diapause in roe deer (Lambert et al. 2001) and carnivores (Mead 1989). In the mare, it is known that uterine secretions vary in a stage-dependent manner related to the duration of either endogenous or exogenous progesterone (Zavy et al. 1982, Hinrichs et al. 1989), and it is reasonable to assume that some of these proteins play a role in promoting embryonic development, particularly in view of the increased incidence of early pregnancy failure in older mares suffering fibrotic degenerative changes in the endometrium, which render the secretory glands non-functional (Stewart et al. 1998, Morris & Allen 2002, Allen et al. 2007). The failure of all the day 10 embryos transferred to day 1 uterus to develop an embryo proper, only 3/8 transferred to a day 3 uterus to do so, and the slowed development of 5/6 of the embryos placed in a day 4 uterus strongly suggests the development of stage-dependent factors in the uterine environment, which allow embryonic development to proceed. Furthermore, the mean serum progesterone level in the day 3 or 4 uterus on the day of transfer did not differ significantly, suggesting that the level of progesterone when the day 10 embryo entered the recipient mare’s uterus was not the overriding factor determining embryo survival.
From a practical aspect, the embryo flushing equipment used in the present experiment would need to be modified before being used in a commercial setting. Although overall embryo recovery rate was equivalent to that achieved in embryo transfer programmes using fresh semen from fertile stallions (Losinno et al. 2001), both the failure to recover some ultrasonographically visualised embryos and the damage suffered by 8% of the embryos that were recovered would not be acceptable in a commercial programme. Embryonic damage in all likelihood occurred at the two points in the flushing system that provided the greatest impediment to the passage of the embryo; namely, the entry/exit holes in the flushing catheter and the internal diameter of the entry port into the embryo filter. Although this small percentage of embryos did suffer damage, the fact that so many passed unscathed through the narrow openings is a testament to the protective strength provided by the glycoprotein equine capsule (Betteridge 1989, Oriol et al. 1993a, 1993b). This is not surprising when one considers that the embryo is subjected to strong myometrial contractions during its movement throughout the uterus between days 6 and 17 after ovulation (Ginther 1983b, Allen & Bracher 1992) to the extent that the day 13 or 14 embryo may undergo periodic compressions every 5–14 s with the resilience and elasticity of the capsule allowing temporary distortion of the spherical embryonic yolk sac (Ginther 1985).

One further problematic aspect of the transfer of large day 10 embryos is the use of transfer pipettes with an internal diameter ≥5 mm; the capillary forces required to hold the embryo in its transfer medium in set columns within the pipette do not come into play in these larger bore pipettes. Nevertheless, transfer was successfully accomplished and a high rate of pregnancies resulted despite the perilous movement of the fluid columns within the pipette during the transfer procedure. In conclusion, these results show clearly that a day 10 embryo can survive and develop, to at least the heartbeat stage, in a uterus that is as much as 7 days behind or 2 days ahead in developmental terms. Since conceptus growth and development was delayed when embryos were placed in less advanced uteri, it can be concluded that although progesterone priming of the uterus to a stage equivalent to that of the transferred embryo is not a prerequisite for embryo survival, it does nonetheless influence embryonic development.

Materials and Methods

Management of the mares

The use of the animals was in accordance with the UK Animal (Scientific Procedures) Act (1986). Follicular development and ovulation in experimental donor, recipient and control Thoroughbred mares, aged 3–12 years, was monitored by daily ultrasound scanning of their ovaries during oestrus combined with daily measurement of serum progesterone concentrations (Allen & Sanderson 1987). The donor and control mares were inseminated once with 300–500 × 10⁶ freshly collected, extended, stallion spermatozoa when they exhibited a dominant ovarian follicle of ≥35 mm diameter. Coincidentally, an i.m. injection of 0.75 mg of the GNRH analogue, deslorelin (BET Pharm Laboratories, Kentucky, KY, USA), was administered to induce ovulation of the maturing follicle 40–42 h later (Fleury et al. 2004). Recipient mares underwent the same procedures but were not inseminated. For all mares, the day when ovulation was detected was classified as day 0 of pregnancy or the oestrous cycle.

Embryo recovery

On day 10 after ovulation, the uterus of each control and donor mare was scanned using a 7.5 MHz transrectal probe attached to a Honda HS-2000VET (BCF Technology, Livingston, Scotland) ultrasound scanner to establish if a conceptus could be observed ultrasonographically. If seen, its diameter was measured using the calliper function within the ultrasound machine. Regardless of the outcome of this ultrasound examination, the uterus of each donor mare was then flushed twice with 500–2000 ml of a commercial embryo transfer flushing medium (Encare Complete; ICPbio Ltd, Auckland, New Zealand) using a standard two-way 100 cm, silicone, 28 French gauge embryo flushing catheter (Stallion Foley Catheter; SurgiVet, Waukesha, WI, USA) and a commercial embryo filter (Sure Flush Filter; Professional Embryo Transfer Supply, Inc., Canton, TX, USA). The flushing medium was recovered by gravity flow alone in the first instance, and recovery was assisted by the i.v. administration of 20 i.u. of oxytocin (Intervet UK Ltd, Milton Keynes, Bucks, UK) to induce myometrial contractions after the second flush was undertaken. Recovered embryos were graded morphologically using the scale proposed by McKinnon & Squires (1988), and their diameters were measured using a stage graticule. They were then placed in a holding and transfer medium (Encare Hold; ICPbio Ltd) and transferred within 1 h of collection.

Embryo transfer

A total of 78 Grade 1 day 10 embryos were transferred nonsurgically to individuals in 11 groups of recipient mares that had ovulated 9, 7, 6, 5, 4, 3, 2 or 1 day after (−) the donor, on the same day (0) as the donor, or 2 or 4 days before (+) the donor (n = 6 or 8 mares per group). The embryos were transferred using a commercial, 63.5 cm, insemination pipette (Reproduction Resources, Walworth, WI, USA) or a rigid plastic tube made from the protective plastic sheath around either an oocyte-pick-up needle or an endometrial swab to which a small piece of silicone tube and a syringe adapter were fitted. For each transfer, the smallest diameter tube into which the embryo would fit comfortably was selected. Non-surgical transfer was performed using the method described by Wilsher & Allen (2004) in which a Polansky speculum is inserted into
the vagina to allow visualisation of the cervix, prior to its retraction using modified forceps. The pipette containing the embryo is then passed through the straightened cervix and the embryo deposited in the uterine lumen.

**Pregnancy diagnosis and monitoring conceptus growth**

Two days after embryo transfer to the recipient mares, or 12 days after ovulation in the control mares, pregnancy diagnosis was performed by transrectal ultrasound of the uterus. These examinations were then repeated on alternate days to measure the size of the vesicle and the degree of development of the embryo. They continued until a viable embryo with a visible heartbeat was seen, or until embryonic death had occurred, when most of the mares received an i.m injection of 250 μg cloprostenol (Estrumate; Schering-Plough, Middlesex, UK) to induce luteolysis and a return to oestrus.

**Hormone measurements**

Jugular vein blood samples were recovered daily from all recipient or control mares from the day of ovulation until day 10 and thereafter on alternate days until termination of the pregnancy or a return to oestrus. The serum was decanted after centrifugation and assayed for progesterone concentrations using the enzyme-linked immunoassay (AELIA) described by Allen & Sanderson (1987), who recorded its sensitivity as 0.135 ng/ml and found its precision to be described by intra- and inter-coefficients of variation of 6 and 8% respectively.

**Statistical analysis**

All statistical analyses were performed using a computer software package (SigmaStat v.2.03; SPSS Inc., Chicago, IL, USA). The χ² test was employed to determine if there was an effect of asynchrony on pregnancy rates in the recipient mares and, if significant, multiple comparisons were made comparing each asynchronous transfer group individually. One-way ANOVA followed by Tukey’s test was used to determine differences in embryonic vesicle growth at different stages of development and to compare serum progesterone concentrations on the day of embryo transfer. Statistical significance was set at $P \leq 0.05$. In addition, one-way RM ANOVA was used to determine if progesterone concentrations in the mares’ blood affected the likelihood of survival of a transferred embryo.

**Declaration of interest**

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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