Role of age and length of oestrous cycle in alteration of the oocyte and intrauterine environment in the rat

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Summary. Zygotes were transferred, on the day of fertilization, from young and old rats with 4- or 6-day oestrous cycles into the ovarian bursa of young recipients with 4-day cycles, and zygotes from young rats with 4-day cycles were transferred into young and old recipients with 4- or 6-day cycles. Young rats with 4-day cycles served as controls for both donors and recipients. An increase in length of cycle or maternal age of donor caused an increase in unfertilized and/or abnormal eggs at the pronuclear stage (non-transferred zygotes). Increased age of donor or length of cycle decreased the implantation rate observed on Day 11 of pregnancy. Likewise, increased age of recipient or length of cycle decreased implantation rate observed on Day 11 of pregnancy. The increase in both age and length of cycle of donor or recipient caused the greatest decline in implantation rate and percentage of normal embryos observed on Day 11 of pregnancy.

Introduction

Mature female rats exhibit oestrous cycles of 4 or 5 days in length, until 10–12 months of age. At this age, they begin to show oestrous cycles of abnormal lengths (Lu, Hopper, Vargo & Yen, 1979), including cycles of 6 days (a natural delay of ovulation). Abnormal embryonic development increases in aged rats, with more abnormalities occurring when the length of cycle is 6 days, than when it is 4 or 5 days (Fugo & Butcher, 1971). Decreased fertilization rate, increased embryonic anomalies and increased embryonic resorption are consequences of this natural delay of ovulation. Delay of ovulation in young rats can be induced experimentally with pentobarbitone sodium (Everett & Sawyer, 1950) and results in an increase in chromosomal anomalies (Butcher & Fugo, 1967) as well as morphologically abnormal embryonic and fetal development (Fugo & Butcher, 1966; Butcher, Blue & Fugo, 1969).

It has been proposed that the effect of delayed ovulation on developmental anomalies and pregnancy loss is mediated, at least in part, by an increased exposure of the intrafollicular oocyte to oestradiol (Butcher, Collins & Fugo, 1975; Butcher & Pope, 1979; Butcher & Page, 1981; Page & Butcher, 1982). Effects of intrauterine environment also contributed to pregnancy loss in old animals (Butcher & Page, 1981). Talbert & Krohn (1966) reported that only 14% of zygotes from young mice survived to term in old hosts compared to 48% surviving to term in young hosts. Butcher et al. (1969) performed reciprocal transfers of blastocysts between young rats with 4-day cycles and young rats in which ovulation had been delayed (6-day cycles). Compared to control transfers

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between donors and recipients with 4-day cycles, an induced delay of ovulation in young rats caused detrimental changes in the zygotes and in the intrauterine environment. Because transfers were performed at the blastocyst stage, effects due to the oocyte could not be separated from those of oviducal and uterine environment during the 4 days before transfer. Although the developmental potential of the blastocyst has been reported not to be influenced by the endocrine status of the oviduct (Beyer & Zeilmaker, 1973), contribution of the oviduct due to age or length of oestrous cycle has not been assessed.

The present experiment was designed to separate effects of intrafollicular ageing of the oocyte from effects of intraoviducal/intrauterine environment by transferring zygotes into the ovarian bursa within 3–5 h after fertilization. Moreover, by using young and old rats with normal oestrous cycles and with delayed ovulation, effects of both maternal age and of delayed ovulation can be evaluated.

Materials and Methods

Animals. Young (170–200 g) and old (retired breeders > 1 year of age) female Holtzman rats were housed under controlled temperature, humidity and lighting. All animals received 12 h light per day. However, the lighting schedules in the rooms for donors and recipients were different, for at least 14 days before beginning the experiment; lights were on from 06:00 to 18:00 h for donors and 10:00 to 22:00 h for recipients. In preliminary experiments, an increase in pregnancy rate was obtained when the lighting schedule for the recipients was delayed by 4 h compared to that of the donors. The difference in lighting schedule allowed eggs from donors to be collected 3–5 h after fertilization (6–8 h after ovulation) and to be transferred into recipients that were 2–4 h after ovulation. The time of transfer of zygotes was limited to the period of 09:30–10:30 h.

The four types of animals used were young rats with 4-day oestrous cycles (Y4), young rats with 4-day cycles in which ovulation was delayed for 2 days by treatment with pentobarbitone sodium (Everett & Sawyer, 1950) to produce 6-day cycles (Y6), old rats with 4-day cycles (O4) and old rats with spontaneous 6-day cycles (O6). Zygotes from rats of each of the 4 groups were transferred into Y4 recipients to study the effects of age and length of oestrous cycle on the oocyte. Also, each type of animal served as recipient for zygotes collected from Y4 donors to examine for effects of age and length of cycle on intrauterine environment. Young rats that did not maintain oestrous cycles of 4 days in length were not used in these experiments. Old rats were required to have exhibited at least 2 previous oestrous cycles of 6 days or longer before use. The contributions of the oocyte and the intrauterine environment were assessed at Day 11 of gestation by implantation rate, post-implantation death and embryonic development.

Transfer of zygotes. Vaginal smears were taken daily between 08:00 and 10:00 h to determine the length and stage of oestrous cycle. Each donor female was housed with a fertile male from the afternoon of pro-oestrous to 08:00 h on the day of oestrous. A vaginal smear containing spermatozoa on the day of oestrous confirmed mating (Day 0). Each recipient female was placed with a vasectomized male on the afternoon of pro-oestrous, and the finding of a vaginal plug under the breeding cage or in the vagina the following day confirmed mating. Vaginal smears were taken daily after the day of transfer to confirm that pseudopregnancy had been induced in the recipient.

Donor females were killed by decapitation. The ovary, oviduct and a small portion of the uterus were removed from each side of the reproductive tract and placed in saline (0·9% (w/v) NaCl) at 37°C. Syringes, glassware and saline were at 37°C at the onset of each transfer, but gradually cooled to room temperature (Dickmann, 1971). The uterine remnant was removed from the oviduct, which was then placed in an embryonic watch glass and observed under a dissecting microscope. Zygotes were collected by one of two methods: (1) the swollen ampullary region of the oviduct was opened with fine forceps and the eggs covered with cumulus cells flowed out with slight pressure on the oviduct, (2) the ovarian bursa was opened, a blunt 27-gauge needle (attached to a 1 ml syringe)
was inserted into the ovarian end of the oviduct (ostium tubae) and the zygotes were flushed from the oviduct with 0.5–1.0 ml saline. Embryos were washed in saline before transfer to reduce the probability of transferring spermatozoa. Also, to ensure absence of cross fertilization, 15 transfers were made between genetically marked donors (Long–Evans hooded rats with dark pigmented eyes mated to albino Holtzman males) and Holtzman recipients. All 44 fetuses resulting from these transfers had dark eyes when recovered at Day 19 of gestation, indicating that fertilization of the recipient's oocytes did not occur.

Transfer of zygotes into recipients was accomplished using a capillary glass tube attached to a 10 µl Hamilton syringe. The washed zygotes to be transferred into one ovarian bursa (3–7 per side, maximum of 10 per recipient) were collected into a glass tube in <3 µl saline. The capillary tube was inserted through the small bursal foramen which opens caudally to the utero-oviducal junction (Alden, 1942), and the zygotes were deposited into the bursal cavity. Halothane anaesthesia was used for the surgery; and transfers were completed within 15 min after death of the donor. After surgery, recipients were housed in stock cages with other females. Because the time for transfers was limited to 09:30 to 10:30 h, no more than 4 transfers were done in 1 day.

Recipient animals were killed by decapitation on Day 11 of gestation and the number and relative size of implantation sites were noted. The uterus was opened on the antimesometrial side with care not to rupture the embryonic membranes, placed in Bouin's fixative overnight, and then stored in 70% ethanol. After removal of the embryonic membranes, the embryos were examined for normality of size and development. Abnormalities were based on development-for-size of previously studied rat embryos from 9–12 days of gestation, with the most frequent anomalies being gross malformations, open neural tube and disproportionate growth (e.g. underdeveloped head). The size of each embryo was evaluated by the use of calibrated lines on the bottom of a Petri dish.

Evaluation of eggs. When more donors than recipient rats were available, the excess donors were killed, the eggs recovered and the cumulus cells dispersed with a few drops of 2% trypsin. Eggs in 0.9% NaCl were placed on a microscope slide with an interrupted vaseline–paraffin wax ring to support the cover slip (Fugo & Butcher, 1966) and were evaluated under phase-contrast microscopy at × 600 for pronuclei, number of sperm tails within the egg, number of accessory spermatozoa, number and size of polar bodies and other anomalies of the zygote.

Statistical analyses. Data were analysed for differences in fertilization rate and normality of zygotes at the 1-cell stage and for differences in pregnancy rate, implantation rate, embryonic survival and normality of embryonic development at Day 11 of gestation. Overall differences in numbers per group were assessed by χ² tests. Percentages for each variable to be analysed were calculated for each rat, and two-way analysis of variance (ANOVA) with age and length of oestrous cycle as main effects was used to compare results of morphology of non-transferred zygotes and for comparisons within donor and recipient groups. When significance was found with the ANOVA, differences between group means were assessed by least significant difference (LSD) (Snedecor & Cochran, 1967).

Results

Data from morphological evaluation of non-transferred embryos at the 1-cell (pronuclear) stage are presented in Table 1. Compared to zygotes from Y4 animals, the fertilization rate was decreased (P < 0.05) in the other three groups. The percentage of zygotes classified as abnormal was increased in both groups of old rats. ANOVA showed a significant effect of age of rat on normality of embryos. From analysis by LSD, there was a significant increase in abnormal zygotes when comparisons were made between embryos from Y4 or Y6 and O4 or O6 rats. The incidence of polyspermy increased, though not significantly, due to age of rat and length of cycle (0.8, 1.2, 1.7 and 3.7% for Y4, Y6, O4 and O6 rats, respectively).

Data on pregnancy, implantation and embryonic development at Day 11 after transfer of
zygotes are shown in Table 2. There were no differences in number of pregnancies due to type of donor, but the number of rats establishing a pregnancy was decreased ($P < 0.05$) in O6 recipients compared to the other 3 types of recipients. However, compared to control transfers, the number of embryos implanting was decreased ($P < 0.05$) in all other donor and recipient groups. There also was a decrease in implantation rate in O6 recipients ($P < 0.05$) compared to all other groups. Even after eliminating the data from rats without implantation sites, there was still a decreased percentage of embryos implanting (data not shown) in groups with Y6 and O4 donors and O6 recipients. Post-implantation survival ($\%$ implantations as embryos) did not differ ($P < 0.05$) among groups, whether analysed by ANOVA or $\chi^2$ or whether all rats with implantation sites or only rats with one or more surviving embryos were used in comparisons. However, the number of recovered embryos classified as normal was reduced when the donor was an O6 rat ($P < 0.05$) or the recipient was an O4 or O6 rat ($P < 0.05$). By ANOVA, there was a significant effect of age of recipient which, when analysed by LSD, was significant ($P < 0.05$) for O4 or O6 compared to Y4 or Y6 recipients.

### Table 2. Effects of age and length of oestrous cycle of donor and recipient rats on implantation and embryonic development of transferred zygotes, as determined at Day 11 of gestation

<table>
<thead>
<tr>
<th>Donor</th>
<th>Recipient</th>
<th>No. of zygotes transferred</th>
<th>No. of rats pregnant/total no.</th>
<th>No. (%)§ of zygotes implanting</th>
<th>No. (%)§ of embryos as embryos</th>
<th>No. of embryos examined</th>
<th>No. (%)§ of normal embryos</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y4</td>
<td>Y4</td>
<td>193</td>
<td>18/20</td>
<td>103 (53 ± 7)*</td>
<td>72 (65 ± 8)</td>
<td>72</td>
<td>66 (95 ± 2)</td>
</tr>
<tr>
<td>Y6</td>
<td>Y4</td>
<td>184</td>
<td>17/20</td>
<td>71 (38 ± 7)*</td>
<td>58 (81 ± 6)</td>
<td>58</td>
<td>51 (86 ± 7)</td>
</tr>
<tr>
<td>O4</td>
<td>Y4</td>
<td>195</td>
<td>19/21</td>
<td>80 (40 ± 6)*</td>
<td>61 (72 ± 7)</td>
<td>61</td>
<td>46 (85 ± 7)</td>
</tr>
<tr>
<td>O6</td>
<td>Y4</td>
<td>174</td>
<td>15/19</td>
<td>67 (37 ± 7)*</td>
<td>42 (58 ± 10)</td>
<td>42</td>
<td>33 (81 ± 9)*</td>
</tr>
</tbody>
</table>

§ Values are total no./group. Percentages were calculated for each rat and the mean ± s.e.m. % is shown in parentheses.

* $P < 0.05$ compared to Y4–Y4 ($\chi^2$ using total no./group).

† $P < 0.05$ compared to Y4, Y6 and O4 recipients (LSD).

‡ $P < 0.05$ compared to Y4 or Y6 recipients (LSD).
Discussion

The transfer of zygotes was used in the present study to separate the effects of age of rat and length of the oestrous cycle on the oocyte and intrauterine environment, as evaluated by pregnancy rate, implantation and embryonic development. By selection of the time of transfer and use of the bursal foramen for depositing of ova into the ovarian bursa within 3–5 h after fertilization, implantation rates (53%) approached those observed using transfer of blastocysts into the uterus. Preliminary transfers, in which both the donor and recipient were approximately 6–8 h after ovulation, yielded limited success. When the lighting schedule for recipients was delayed 4 h from that of the donors, and the transfers performed between 09:30 and 10:30 h, i.e. recipients were 2–4 h after ovulation, the success rate greatly improved. Casual observations indicated a greater contractile activity of the recipient uterus and oviduct at the time of transfer when the staggered lighting schedules were used.

From the results of morphological evaluation of non-transferred eggs, it was concluded that an increase in length of cycle and increase in age of rat caused a decrease in fertilization rate, while an increase in age led to increases in abnormalities at the pronuclear stage. An increase in length of cycle has previously been shown (Page & Butcher, 1982) to be associated with an early increase in intrafollicular as well as plasma concentration of oestradiol in relation to the time of ovulation. An early increase in oestradiol in animals with induced delay of ovulation has been shown to decrease implantation rate (Butcher & Pope, 1979). The early increase in follicular oestradiol before ovulation may detrimentally affect the intrafollicular oocyte, thus contributing to the decrease in fertilization rate in the Y6 and O6 rats. The mechanism by which age contributes to oocyte abnormalities has not been determined. However, Page & Butcher (1982) proposed that, in old rats with oestrous cycles of normal length, an early development of follicles resulted in altered oestradiol patterns, which could affect the oocytes and the uterus.

There was a reduction in number of pregnant animals on Day 11 only in O6 recipients as compared to the Y4–Y4 group, although more of the transferred zygotes resulted in implantation sites in Y4–Y4 than in any other group. A reduction in implantation rate due to increased age or length of cycle of donor can be accounted for in part by the decreased fertilization rate observed at the 1-cell stage. However, the decline in implantation rate due to age and length of cycle of recipient was independent of fertilization rate, as all embryos transferred to the recipients were from Y4 donors. Therefore, the decreased implantation rate in these recipients was probably due to detrimental effects of the oviducal and uterine environments on embryonic survival before implantation. Although the incidence of post-implantation death of transferred zygotes was much greater than normally found in young unoperated rats, there was no difference due to age or length of oestrous cycle of donor or of recipient. This is in contrast to an increase in post-implantation loss when cycles were 6 days in length in young and old rats in previous studies (Fugo & Butcher, 1966, 1971). Detrimental effects of the transfer procedure, or of the recipient oviduct, on defective zygotes could have produced preimplantation rather than post-implantation loss; whole normal zygotes could have been affected equally in each group to cause similar rates of post-implantation death. The lower incidence of abnormal embryos at Day 11, and the higher embryonic death in all groups compared to earlier studies, lends support to this proposal. The effect on the oviduct and uterus of an early increase in oestradiol in relation to time of ovulation has been proposed (Page & Butcher, 1982) as a possible cause of detrimental effects of the environment within the reproductive tract on embryonic death, both pre- and post-implantation.

In conclusion, increases both in age of rat and length of the oestrous cycle cause changes in the oocyte which decrease fertilization rate. Age and length of cycle of the recipient influence implantation rate and post-implantation embryonic development. These changes in the oocyte and reproductive tract could be caused by an inappropriate sequence of oestrogen in relation to the time of ovulation.

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References


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