Ovarian development in intrauterine growth-retarded and normally developed piglets originating from the same litter

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Epidemiological studies in humans linking adult disease to growth in utero indicate that prenatal life is a critical period for the appropriate development of the reproductive axis. The aim of this study was to compare ovarian development in intrauterine growth-retarded and normally grown piglets originating from the same litter. Intrauterine growth-retarded piglets (runts) were identified on the basis of statistical analysis of the birth weight distribution within each litter. At birth, ovaries were collected from runt piglets (n = 14) and their respective mean weight (normal, n = 14) littermates. Ovaries were weighed and fixed, and development of ovarian germ cells was quantified in haematoxylin–eosin-stained paraffin wax sections using an image analysis system. Germ cell loss, using an in situ TdT-mediated dUTP nick-end labelling (TUNEL) assay for DNA fragmentation, and follicle cell activity, using immunohistochemistry to demonstrate vimentin, were studied in ovarian sections. At birth, body weight and absolute ovarian mass were significantly lower in runt piglets compared with their respective normally grown littermates (body weight: 733 ± 38.5 versus 1530 ± 39.7 g; ovarian mass: 51 ± 3.0 versus 108 ± 9.6 mg; P < 0.001 for both).

In the ovary, the proportion of nests of oogonia, the number of oocytes and TUNEL-positive cells, and the localization and intensity of vimentin immunoreactivity were not different between runt and normal littermates. However, runt piglets had more primordial follicles (268 ± 18.6 versus 235 ± 20.1 per mm² of cortex; P < 0.05), fewer primary follicles (11 ± 2.0 versus 20 ± 3.0 per mm² of cortex; P < 0.001) and no secondary follicles compared with normal piglets. These findings indicate that intrauterine growth retardation delayed follicular development in pig ovaries at birth.

Introduction

Perturbed prenatal growth has a major impact on neonatal survival and it has been suggested that growth restriction during fetal life is also associated with a range of adult pathologies in humans (Barker et al., 1993), such as increased risk of developing type 2 diabetes, hypertension and cardiovascular disease (Barker, 1995). Other reports indicate that altered growth in utero may be associated with the development of reproductive disorders in later life, including altered timing of onset of puberty (Ibáñez et al., 2000a), reduced size of the uterus and the ovaries (Ibáñez et al., 2000b), and earlier menopause (Cresswell et al., 1997). In these reports it is postulated that these postnatal disorders may derive from altered fetal reproductive organogenesis. Indeed, the development of the hypothalamic–pituitary–gonadal axis occurs largely during prenatal life and, within the ovary, processes such as migration, proliferation, degeneration and meiosis of germ cells, and folliculogenesis determine the follicle population at birth and potentially the number of follicles that can be recruited during postnatal life.

Studies on animals and humans have begun to address more specifically the effect of altered prenatal growth on fetal ovarian development and to date have reported contradictory findings. De Bruin et al. (2001) failed to demonstrate an effect of severe growth restriction on ovarian development in mid-gestation human fetuses that died prenatally. In contrast, a significantly lower number of follicles was found in growth-restricted late gestation sheep fetuses (Da Silva et al., 2002a). Therefore, there is a need to clarify the relationship between prenatal growth retardation and ovarian development using other animal models, keeping in mind that the possible exhaustion of follicle reserves has stronger implications for humans, as, compared with domestic species, their period of reproductive activity is much longer (Faddy et al., 1992).

In pigs, fetal growth retardation is an important determinant of perinatal mortality (van der Lende et al., 2001), contributing to a considerable economic loss in the pig industry worldwide (Varley, 1995). During the early neonatal period, growth-retarded piglets have a lower capacity to compete with heavier littermates for...
Appropriate development of all types of ovarian cell was examined by immunohistochemical demonstration of the intermediate filament protein vimentin. Reorganization of vimentin-containing filaments is associated with changes in cellular shape and with various cellular functions such as intracellular transport, nucleus–cell surface interactions and nuclear functions (Goldman et al., 1986; Aumüller et al., 1992), and is recognized as a marker of granulosa cell viability (van den Hurk et al., 1995). It is proposed that study of the pig ovary could complement the recent efforts to understand the relationship between impairment of prenatal growth and ovarian development.

**Materials and Methods**

**Animals and experimental design**

The experiment was approved by the Animal Ethics Committee of the Veterinary Faculty of Utrecht University. The weight and sex of live-born piglets from litters produced by cross-bred sows (Dutch Landrace/Yorkshire) in the same experimental farm (Sterksel, The Netherlands) were examined within 12–24 h after birth. Litters were statistically analysed at the farm on the basis of number of live piglets born during one farrowing by the biological mother, duration of gestation and variation in birth weight within each litter. The statistical computerized method used to determine the within-litter birth weight distribution was based on procedures for the identification of multiple ‘outliers’ in small samples and has been described in detail by van der Lende et al. (1990). The use of this method allows identification of low birth weight piglets that are ‘outliers’ in an otherwise normally distributed population and are referred to as intrauterine growth-retarded piglets (colloquially denoted as ‘runts’). In the present study, 14 of 73 litters contained intrauterine growth-retarded female piglets; seven litters had one runt piglet and seven litters had two runt piglets, but only one runt piglet was studied per litter. The mean ± SEM duration of gestation and litter size for these litters were 115 ± 0.3 days and 12 ± 0.6 piglets (n = 14), respectively.

At the farm, each identified intrauterine growth-retarded female piglet was pair-matched with a normally developed littermate, which was selected as the female piglet with the birth weight closest to the mean birth weight of the litter which consisted of both female and male piglets. The aim of the present study was to compare (i) germ cell development, (ii) germ cell loss and (iii) the structural integrity of all types of ovarian cell and viability of the granulosa cell layer between oocytes collected from naturally occurring intrauterine growth-retarded newborn piglets and their respective normally developed littersmates. Germ cell development was determined by classical histological procedures, whereas germ cell loss was examined by detection of DNA fragmentation (using TdT-mediated dUTP nick-end labelling (TUNEL) assay).

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**Fig. 1.** Illustration of representative litters that contained intrauterine growth-retarded piglets analysed on the basis of the intrinsic variation in birth weight. For the present study, an identified runt female piglet in each litter (○: 690 g birth weight; △: 1010 g birth weight) was pair-matched with a normally sized female littermate (arrows), which was selected as the female piglet with the birth weight closest to the mean weight of the litter which consisted of both female and male piglets. △ and ○: each symbol represents a littermate of the runt female piglet.
Tissue fixation and preparation

The right ovary was immersion-fixed in Bouin’s solution at room temperature (20–22°C) for 20 h and was then rinsed and stored in 70% (v/v) ethanol. The ovary was dehydrated in a series of ethanol, cleared in xylene and impregnated in paraffin wax. The ovary was subsequently bisected transversely and both parts were embedded in the same paraffin wax block. The ovaries were cut into serial sections (thickness 5 μm) and each twentieth section was stained routinely with haematoxylin–eosin for estimation of germ cell development. A few sections were also mounted on polylysine (Menzel-Gläser, Braunschweig)-coated slides for immunohistochemical study of vimentin. The left ovary was immersion-fixed in 4% (w/v) phosphate-buffered formaldehyde at 4°C for 20 h, rinsed in 70% (v/v) ethanol and embedded in paraffin wax as described above. The ovaries were cut into serial sections (5 μm thickness) and each twentieth section was mounted on polylysine-coated slides for examination of DNA fragmentation, which is characteristic of apoptosis.

Morphometric analysis

Germ cells were classified histologically as oogonia (germ cells devoid of follicle cells with an intact nuclear membrane), oocytes (germ cells undergoing meiosis and devoid of follicle cells), primordial follicles (oocyte surrounded by a layer of flattened follicle cells), primary follicles (enlarged oocyte completely surrounded by one layer of cuboidal follicle cells) or secondary follicles (enlarged oocyte surrounded by two or more concentric layers of cuboidal cells) as described by Oxender et al. (1979). Oocytes surrounded by a single granulosa layer that consisted of both flattened and cuboidal cells were classified as primordial follicles. Stained ovarian sections were examined under a microscope using a semi-automatic image analysis system. The equipment used consisted of a light microscope (Zeiss Photomicroskop II) with a x1 or x16 objective coupled to a camera (Sony b/w CCD camera type XC-77CE, frame size 752 × 574 pixels; 256 grey levels) and a specific computer program which was designed using the image analysis KS400 version 3.0 software package (Carl Zeiss Vision, Munich). Ovary area was measured in each twentieth stained section, using a x1 objective, to select the sections around the largest diameter. Each section was produced on to the surface epithelium) were projected on to the computer screen and using the computer program, all nests of oogonia, oocytes and primordial, primary and secondary follicles were outlined by the operator using the computer mouse. These microscopic images were measured automatically and converted into mm². Counts were exported to a Microsoft Excel program and the number of germ cells per mm² of cortex was calculated. This procedure was repeated for the two sections around the largest diameter (100 μm apart) and an index of the number of oocytes and follicles was calculated by averaging the results. Only follicles in which the nucleus of the oocyte was visible were counted. The same quantitative method was used uniformly for all ovaries. In this way, at least 35–40% of the ovarian cortex of a section was quantified. The applicability of this procedure was verified in another study (Da Silva et al., 2002b) in which the results obtained using two sections at the largest diameter of a fetal pig ovary (day 100 after mating) were found to represent the mean number of germ cells per mm² counted in approximately every twentieth section of an entire ovary (the correlation was 0.79).

TUNEL assay

TUNEL assay for DNA fragmentation was performed using an in situ cell death detection kit (Roche, Mannheim) according to the manufacturer’s protocol. In brief, sections approximately around the largest diameter were deparaffinized and rehydrated. Endogenous cellular peroxidase was blocked with 3% (v/v) H₂O₂ in methanol for 10 min. After a phosphate-buffered saline (PBS) wash, terminal deoxynucleotidyl transferase (TdT) enzyme solution was added to each section to label single and double DNA strand breaks with fluorescein, and the reaction was incubated for 1 h in a humidified chamber at 37°C. A negative control section without the addition of the TdT enzyme was also included. After the sections were washed in PBS, anti-fluorescein antibody Fab fragments conjugated with horseradish peroxidase were added to each section and incubated for 30 min in a humidified chamber at 37°C. The sections were washed in PBS and bound antibodies were covered with Mayer’s haematoxylin and dehydrated in increasing concentrations of ethanol and mounted. TUNEL-positive cells (brown staining) were counted in the two sections (100 μm apart) using a light microscope at a magnification of x400. As the presence of TUNEL-positive cells was observed in the ovarian cortex only, the positive cell counts were then averaged and expressed per mm² of ovarian cortex.

Immunohistochemical staining

The intermediate filament vimentin was used as a marker for structural integrity of all types of ovarian cell, with special emphasis on the viability of the granulosa cell layer. The immunohistochemical staining technique was similar to that reported by van den Hurk et al.
Table 1. Body weight, body measurements, ovary mass and number of germ cells in normal and intrauterine growth-retarded (runt) piglets at birth

<table>
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<th>Characteristic</th>
<th>Normal</th>
<th>Runt</th>
<th>Significance</th>
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<tr>
<td>Number of piglets</td>
<td>14</td>
<td>14</td>
<td>na</td>
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<tr>
<td>Body weight (g) (range)</td>
<td>1530 ± 39.7</td>
<td>733 ± 38.5</td>
<td>( P &lt; 0.001 )</td>
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<tr>
<td>Crown–rump length (cm)</td>
<td>31 ± 0.4</td>
<td>25 ± 0.5</td>
<td>( P &lt; 0.001 )</td>
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<td>Umbilical girth (cm)</td>
<td>25 ± 0.5</td>
<td>19 ± 0.7</td>
<td>( P &lt; 0.001 )</td>
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<td>Biparietal head diameter (cm)</td>
<td>6.0 ± 0.1</td>
<td>5 ± 0.1</td>
<td>( P &lt; 0.001 )</td>
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<tr>
<td>Ovarian mass (mg)</td>
<td>108 ± 9.6</td>
<td>51 ± 3.0</td>
<td>( P &lt; 0.001 )</td>
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<td>Relative ovary mass (g ovary kg body weight(^{-1}))</td>
<td>0.07 ± 0.006</td>
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<td>Nests of oogonia (%)(^{\dagger})</td>
<td>0.8 ± 0.22</td>
<td>1.2 ± 0.22</td>
<td>ns</td>
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<td>Number of oocytes(^{\dagger})</td>
<td>171 ± 17.8</td>
<td>188 ± 19.2</td>
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<td>Number of primordial follicles(^{\dagger})</td>
<td>235 ± 20.1</td>
<td>268 ± 18.6</td>
<td>( P &lt; 0.05 )</td>
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<td>Number of primary follicles(^{\dagger})</td>
<td>20 ± 3.0</td>
<td>11 ± 2.0</td>
<td>( P &lt; 0.001 )</td>
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<td>Number of secondary follicles(^{\dagger})</td>
<td>1.0 ± 0.5</td>
<td>0.0 ± 0.0</td>
<td>( P &lt; 0.05 )</td>
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<td>Number of apoptotic cells(^{\dagger})</td>
<td>10 ± 2.7</td>
<td>10 ± 1.6</td>
<td>ns</td>
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Values are mean ± SEM.  
\(^{\dagger}\)Values are expressed per mm\(^2\) of ovarian cortex.  
n: not applicable; ns: not significant.

(1995) with some modifications. In brief, ovarian sections that had been fixed with Bouin’s solution and paraffin wax were deparaffinized and rehydrated. Endogenous peroxidase activity was blocked with 1% (v/v) \( \text{H}_2\text{O}_2 \) in methanol for 30 min. The intermediate filament protein vimentin was detected with monoclonal mouse antibody (clone V9) purchased from Biogenex (distributed by Klinipath, Duiven) at a dilution of 1:125, was used as the second antibody. The detection of the bound antibody was accomplished by the avidin–biotin immunoperoxidase system (Vectastain Elite ABC kit; Vector Laboratories, Burligame). Negative control slides were prepared by substitution of mouse IgG for the primary antibody. Ovarian sections of 42-day-old pigs, which contain more mature follicles, were used as a positive control. All sections were incorporated in the same immunohistochemical procedure. Staining was evaluated semiquantitatively by light microscopy and estimated on a scale from – to +++ in all types of ovarian cell. A score of –, +, ++ or +++ indicated absent, weak, moderate or strong staining intensity, respectively. The estimated intensity of staining was compared between ovarian sections derived from runt and normally grown piglets.

**Statistical analysis**

All values are presented as mean ± SEM. Body weight and conformation and ovarian data were compared between runt piglets and their respective normally grown littermates using paired Student’s \( t \) test. A Wilcoxon signed rank test was used for the comparison of the number of secondary follicles because number of secondary follicles was the only variable to fail the normality test. Data for numbers of oocytes, follicles and TUNEL-positive cells were transformed (natural logarithm) to normalize variation before statistical analysis (except for number of secondary follicles) and are presented as mean number per mm\(^2\) of ovarian cortex. Correlation analysis was performed using Pearson’s product moment test. Values were considered significant at \( P < 0.05 \).

**Results**

Body size, absolute and relative ovarian masses for normal piglets and their intrauterine growth-retarded littermates at birth are shown (Table 1). Body weight, crown–rump length, umbilical girth and biparietal head diameter were all significantly lower \( (P < 0.001) \) in runt piglets than in their normally sized littermates. Although absolute ovarian mass was lower \( (P < 0.001) \) in runt piglets compared with their normally grown littermates, relative ovary mass (that is, expressed in terms of body weight) was not different between the two groups of piglets (Table 1). Ovarian area and area of the cortical region (around the largest diameter) were significantly lower in runt piglets compared with normal littermates (ovarian area: 5.96 ± 0.49 versus 7.71 ± 0.60 mm\(^2\); area of cortical region: 4.13 ± 0.39 versus 5.04 ± 0.45 mm\(^2\); both \( P < 0.05 \)) and were positively correlated with ovarian mass \( (r = 0.537 \text{ and } 0.648, \text{ respectively; } n = 28; P < 0.01) \).

Neither the proportion of nests of oogonia nor the number of oocytes per unit of ovarian cortex was significantly influenced by naturally occurring growth
Fetal growth retardation and ovarian development

retardation (Table 1), despite the tendency for a higher abundance of oogonia in the cortex of runt piglets (Fig. 2a,b). Runt piglets had higher ($P < 0.05$) numbers of primordial follicles compared with their normally sized littersmates (Table 2); however, there were significantly fewer ($P < 0.001$) primary follicles in runt piglets than in their normal littersmates (Table 1 and Fig. 2c,d). Furthermore, ovaries from runt piglets were devoid of secondary follicles, whereas some secondary follicles could be observed in ovarian sections from their normally sized littersmates (Table 1). Overall, the number of primary follicles was positively correlated with birth weight ($r = 0.399; n = 28; P < 0.05$) and ovary mass ($r = 0.424; n = 28; P < 0.05$).

The number of TUNEL-positive cells present per unit of ovarian cortex did not differ between runt piglets and their normally sized littersmates (Table 1). DNA fragmentation of germ cells predominantly represented isolated oocytes (Fig. 3) and no sign of degeneration was detected in follicular structures in either the oocyte or follicular cells.

Moderate vimentin immunoreactivity was observed in the cell cords surrounding the nests of oogonia and oocytes, but was absent in oogonia, oocytes and in the surface epithelium (Fig. 4). Immunoreactivity was generally weak in pre-granulosa cells of primordial follicles and granulosa cells of primary and secondary follicles, whereas it was strong in follicles of the postnatal ovarian tissue that was used as positive control. However, no differences in either intensity or localization of vimentin staining were observed between runt piglets and normally sized piglets, although no secondary follicles were present in the ovaries of runt piglets.

Discussion

In the present study, the naturally occurring model of growth retardation provided by piglets was used and
Table 2. Estimated proportion of nests of oogonia, number of oocytes and of primordial, primary and secondary follicles per mm² of ovarian cortex in normal and intrauterine growth-retarded runt littermate piglets at birth

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*(Area of nests of oogonia/area of cortex examined) × 100.

Fig. 3. (a) In situ 3’ end-labelling of germ cells in the ovarian cortex of a pig ovary at birth; TdT-mediated dUTP nick end-labelled cells were brown (arrows). (b) Negative control. Scale bar represents 20 μm.

Intrauterine growth-retarded piglets were identified using a well defined statistical method (van der Lende et al., 1990). This statistical approach allowed comparison of ovarian germ cell development in intrauterine growth-retarded (runt) piglets with their respective normally grown (control) littermates, born at normal term, thereby avoiding possible confounding influences of genetic background and housing conditions. The results of the present study indicate that compromised prenatal growth may have delayed the normal pattern of follicular development in pig ovaries as demonstrated, at birth, by the higher number of primordial follicles, the lower number of primary follicles and absence of secondary follicles in the ovaries of runt piglets compared with normally developed littermates.

In the present study, runt piglets showed a reduction in body weight which was associated with a significant reduction in absolute ovarian mass compared with their normally sized littermates at birth. In contrast, relative ovarian mass was not influenced by growth retardation and the values for the normally developed piglets are in agreement with data reported by Colenbrander et al. (1983). These findings indicate that the decrease in ovary mass was proportional to that in body weight in the present study. The reports regarding the effects of intrauterine growth retardation on organ masses
Fig. 4. Vimentin immunoreactivity in pig ovaries. (a) Vimentin staining present in cords surrounding the nests of oogonia at birth. (b) Vimentin staining in pre-granulosa cells of primordial follicles at birth. (c) Negative control at birth. (d) Vimentin reaction intensified in the granulosa layer of primordial and primary follicles at day 42 of age (positive control). Scale bar represents 20 μm.

generally only include vital organs and rarely include the masses of reproductive organs (Widdowson, 1971; Flecknell et al., 1981; Bauer et al., 1998). Such reports indicate that in growth-retarded piglets, the masses of the liver, heart, kidneys and lungs decrease considerably with decreased body weight, being adequate for their smaller size; whereas the masses of the brain, pituitary and adrenal glands do not change significantly with body weight. Despite this, the effect of growth retardation on ovarian mass reported in the present study does not appear to have been mediated by inadequate development of the pituitary gland, because deprivation of pituitary hormones as a result of decapitation of pig fetuses at day 42 of gestation did not alter ovarian mass at day 112 of gestation (Colenbrander et al., 1983).

Within the ovary, the proportion of nests of oogonia in the ovarian cortex was not significantly different between runt and normal littermate piglets born at normal term. This observation appears to indicate that proliferation and degeneration of ovarian germ cells were not compromised by prenatal growth retardation. However, it does not exclude a possible adverse effect on these two aspects of germ cell development (and subsequent recovery) at an earlier stage. Reduced growth of pig fetuses can already be evident by day 35 (van der Lende et al., 1990) and day 44 (Cooper et al., 1978) of gestation. Finch et al. (1999) reported that poor fetal growth was detected as early as day 30 of gestation, probably related to inadequate placental ability to deliver nutrients into the fetal compartment. By this stage, germ cells are proliferating in the pig ovary, reaching a peak number by about day 50 of gestation at the time when germ cell degeneration becomes markedly accentuated (Black and Erickson, 1968). The slight tendency for more oogonial nests in ovaries from runt piglets at birth may be indicative of the importance of examining germ cell proliferation and
degeneration at mid-gestation to provide better insight into whether compromised fetal growth exerts differential mitotic or degenerative effects or both on development of ovarian germ cells.

Naturally occurring growth retardation in piglets in the present study was associated with a higher number of ovarian primordial follicles per mm² of ovarian cortex at birth. This finding was intriguing in view of the fact that the number of oocytes was identical between runt and normally sized littermates. In contrast, there was a significant reduction in the number of primary and secondary follicles in runt piglets and, collectively, these findings point to the occurrence of a delay in follicular development in these piglets, probably in the activation of primordial follicles. Indeed, the gross morphological structure of an ovary from a runt piglet at term resembled that of an ovary collected from a normally developed piglet several weeks before birth (Da Silva et al., 2002b). Fetal growth restriction did not influence the number of follicles in the ovaries of human fetuses at weeks 26–31 of gestation (de Bruin et al., 2001). In contrast, placentally induced fetal growth restriction in sheep reduces the number of follicles in the ovaries of late gestation fetuses (Da Silva et al., 2002a). Nevertheless, comparisons among species need to be addressed with care because the ovary of the pig (polytocous animal), in contrast to sheep and humans, contains proliferating oogonia and numerous oocytes until approximately day 20 of postnatal life (pig: Garrett and Guthrie, 1999; sheep: Sawyer et al., 2002; human: Kurilo, 1981), which could potentially be recruited to form follicles or to simply degenerate. Study of runt and normally sized piglets during early postnatal life is required to address whether a ‘catch up’ process will contribute to the primary follicle pool in the ovaries of runt piglets. Primordial follicles express various factors such as growth differentiation factor 9 (GDF-9) and c-kit–kit ligand complex that help to initiate follicle growth (Picton, 2001; Richards, 2001), and it is possible that their different timing of expression may have been involved in the formation of fewer primary follicles in runt piglets. The significant ovarian differences in the number of primary and secondary follicles, coupled with the major reduction in ovarian mass, between runts and normally grown littermate piglets, offer a unique opportunity to study the factors associated with primordial follicle activation and growth that have the ultimate potential to influence age at puberty, ovulation rate and subsequent litter size in sows. Hence, altered growth in utero, associated with poor early postnatal growth, when ovarian development is still proceeding, may impair fertility in sows and should be the subject of further research. Indeed, it has been reported that slow-growing gilts are older at first mating than are fast-growing gilts (Tummaruk et al., 2000). However, owing to the present interest in relating low birth weight with premature menopause and prospects of fertility in humans, one hypothesis was that perturbed fetal growth could compromise the primordial follicle pool contained in the ovaries at birth. The number of primordial follicles per unit area of pig ovary found in the present study did not confirm this contention. Further methods (for example, collagenase digestion of an entire ovary to assess total number of germ cells) are needed to confirm this observation, as ovarian mass in naturally growth-retarded piglets was significantly lower than that of normal sized littermates.

In the present study, the number of degenerative TUNEL-positive cells was not different between runts and normal piglets. On the basis of their location, size and shape, these cells were predominantly isolated oocytes and occurred in very small numbers (±5%). This finding is supported by the study of Vaskivuo et al. (2001), in which a few apoptotic oocytes were also determined in human fetuses close to term. One possible explanation for the findings of the present study could be that at birth there is little germ cell loss. It has been reported that high cell death occurs just after the number of germ cells reaches a peak (at approximately day 50 of gestation) in pig ovaries (Black and Erickson, 1968; Garrett and Guthrie, 1999). This is also true for other species such as cows and humans (Reynaud and Driancourt, 2000). In women, it has been reported that 9% of the germ cell nuclei are apoptotic in mid-gestation ovaries (De Pol et al., 1997). Thus, as previously suggested, germ cell death should be examined at an earlier stage of gestation. Furthermore, necrosis is another mode of cell death that, although not examined in the present study, could potentially occur in follicles of the pig ovary (van den Hurk et al., 1999).

The findings of the present study regarding the localization of vimentin immunoreactivity are in agreement with previous studies in human (Santini et al., 1993) and cow (van den Hurk et al., 1995) ovaries. Reorganization of vimentin-containing filaments occurs in mitotic cells (Lamb et al., 1989), whereas their increased number is correlated with an alteration of cellular structure (Aumüller et al., 1992). Vimentin-containing filaments are absent in germ cells but they appear to be an important feature of the differentiation events leading to the final phenotype of granulosa cells and, therefore, vimentin is considered as a marker of granulosa cell viability (van den Hurk et al., 1995). The absence of differences in either localization or intensity of vimentin staining between runt and normally grown piglets indicates that the follicles contained healthy active granulosa cells in both groups of piglets.

In conclusion, the results of the present study indicate that, in the pig model, naturally compromised prenatal growth in utero had an effect on ovarian organogenesis. The reduced number of primary follicles and absence of secondary follicles in the ovaries of runts compared with normally grown littermate piglets may be attributable to a delay in activation of primordial follicles. No differences were observed with respect to either normal...
development of follicular cells or their degeneration. It would be interesting to determine whether these ovarian changes that were initiated during fetal life are maintained during postnatal life and to assess their impact on adult reproductive performance.

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