Vascular supply as a discriminating factor for pig preantral follicle selection

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Abstract

This research analyses how somatic and vascular compartments change during preantral follicle growth. To address this aim, theca–granulosa (somatic) proliferation indexes (PIs), proportion of proliferating endothelial cells (PE), vascular area (VA) and vascular endothelial growth factor A (VEGFA) expression were simultaneously recorded on single healthy preantral follicles, classified into six different stages on the basis of the diameter and the granulosa layers. An autonomous blood vessel network starts to appear only in class 3. Vascular remodelling requires VEGFA expression, and VEGFA mRNA and VA significantly increase between class 3 and classes 4 and 5 and, further, in class 6. In addition, a positive correlation exists between these parameters in classes 3–5. Despite variation in angiogenesis results from classes 3 to 5, the statistical analysis reveals that the vascular parameters are positively and strictly correlated with somatic PIs. Conversely, class 6, also characterized by higher values of somatic PIs, displays a stable proportion of PEs (≥40%) without showing any correlation among the different parameters analysed. To identify follicular subpopulations within different classes, a multivariate hierarchical cluster analysis was performed. This analysis reveals that the majority of classes 3 and 4 are quiescent follicles or structures that grow very slowly. Class 5 represents a transitory category, where half of the follicles maintain a low activity and the remaining express significantly higher levels of granulosa PI and VA. The follicles with this high activity are probably able to reach class 6 becoming dominant structures where somatic and vascular parameters are constantly on high levels and the VA remains the unique differentiating element.

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Introduction

The regulation of folliculogenesis in the mammalian ovary is a complex process of cellular interactions able to create the local condition to sustain the development of a competent oocyte and the secretion of adequate steroids level in order to drive the reproductive cycle (Knight & Glister 2006). Prior to puberty, as well as into more advanced reproductive age, most of the follicles are primordial, and composed by an immature sized oocyte surrounded by a single layer of flattened granulosa cells. As a consequence of adequate stimuli, the primordial structures are activated. The follicles entering into the growing phase are characterized by morphological and functional modifications (Gougeon 2004, Yang & Fortune 2006).

The first phases of follicle growth can be divided into three distinct stages: activation of primordial follicles, transition from primary to preantral and from preantral to early tertiary/antral follicles (Fortune 2003). During these latter developmental phases, ovarian follicles become an active endocrine structure and sustain the growing phase of the oocyte that acquires the ability to undergo meiotic progression at the time of antral differentiation (Hunter et al. 2004, Knox 2005, Drummond 2006). Even if the process of follicle recruitment occurs cyclically, few of the growing ovarian structures physiologically reach the final stages, while the majority of them undergo a premature regression process known as atresia, which produces an irreversible and progressive loss of germ cells (Gougeon 2004).

The mechanisms involved in ovarian follicle recruitment and regression are only partially known, in particular those that regulate the early stages of folliculogenesis. Nevertheless, interest in this field has increased as the use of the pool of immature follicles is considered as a strategy to improve the efficiency of the assisted reproductive technologies (Fortune 2003).

The development of immature follicles is modulated by a variety of autocrine, paracrine and endocrine signals (Erickson & Danforth 1995). In addition, morphological adjustments of the vascular supply may be crucial to regulate follicle growth and differentiation, although such a cause effect has yet to be established (Redmer & Reynolds 1996, Barboni et al. 2000, Berardinelli et al. 2002). In particular, during the final antral stages, follicle growth is coupled with an increased blood vessel area (Barboni et al. 2004), close to ovulation, changes its organization into a mature vascular network (Martelli et al. 2006). In fact, until the periovulatory phase, the vascular network is
represented by capillaries that, a few hours before ovulation, become covered by pericytes, and new large blood vessels appear in the follicular wall (Zimmermann et al. 2003, Martelli et al. 2006). The acquisition of a specific vascular supply seems to represent a limiting step in the selection of a dominant follicle destined to ovulation (Mattioli et al. 2001, Stouffer et al. 2001), while its disappearance is correlated with the process of early follicular atresia (Moor & Seamark 1986).

The ovarian follicle has an active role in the process of capillary remodelling by producing several angiogenic factors. In particular, the vascular endothelial growth factor A (VEGFA) seems to play a major role in ovarian angiogenesis as demonstrated recently by several pieces of evidence (Wulff et al. 2001, 2002, Zimmermann et al. 2003, Kaczmarek et al. 2005, Shimizu & Sato 2005, Abramovich et al. 2006, Shimizu 2006). Even if relevant progress has been made in understanding the mechanisms involved in the angiogenic events during the dominant final phase of tertiary follicle development (Fraser 2006, Martelli et al. 2006), the role of blood vessel remodelling in early follicular stages is unclear. In regard to preantral follicles, different reports indicate that these are able to synthesize VEGFA and are sensitive to its stimulatory effect (Danforth et al. 2003). Analogously, the VEGFA stimulatory effect is indirectly confirmed by the immunoneutralization experiments where a decrease in theca cell proliferation and a reduction in the endothelial cell area in late preantral follicles were observed (Wulff et al. 2002).

Starting from this limited information on the functional correlation existing between the vascular remodelling and the preantral follicle development, the present research has been designed to determine how follicular and vascular morphological parameters change during preantral follicle growth. To address this aim, several parameters such as granulosa, theca and endothelial cell proliferation, vascular area (VA), and VEGFA intrafollicular distribution have been studied. These analyses were simultaneously performed on single preantral follicles that were previously classified on the basis of the morphological stage of development (Morbeck et al. 1992). The variables were then statistically evaluated to examine the relationship existing between the above-examined parameters and the different categories of preantral follicles. Moreover, the multivariate hierarchical cluster analysis was designed to identify subpopulations within each class of preantral follicle.

**Results**

**Preantral follicle morphological classification**

The present research was performed on healthy preantral follicles identified using haematoxylin and eosin (HE) and caspase-3 *in situ* TUNEL staining. Preantral follicles were definitively judged as healthy when they showed a regular shaped oocyte, surrounded by granulosa cells regularly apposed on an intact basement membrane, as well as granulosa cell nuclei without signs of pycnosis. Moreover, healthy preantral follicles were confirmed when less than 20% of the granulosa cells displayed signs of apoptosis, revealed by the simultaneous positivity for caspase-3 and nuclear fragmentation (data not shown). Follicles not fulfilling these criteria were classified as unsuitable for the analyses. Healthy preantral follicles were divided, as summarized in Table 1, into different classes on the basis of the follicular diameter, number of granulosa layers, oocyte diameter and, when present, theca layer thickness.

Class 1 preantral follicles did not display the theca compartment that started to be evident as an undifferentiated structure in a low percentage of class 2 preantral follicles (<10%). In class 2, the theca compartment was characterized by few connective cells aligned to each other and localized near to the basal lamina.

Theca compartment characterized by a well-distinguishable inner and outer layer started to appear in class 3 (12%). All follicles belonging to classes 4–6 showed a complete follicular morphological organization.

**Granulosa and theca proliferation index (PI)**

Class 1 and 2 preantral follicles rarely displayed Ki-67 immunopositive cells (data not shown). By contrast, Ki-67 positive cells were variably recorded in preantral follicles from classes 3 to 6 (Table 2; Fig. 1A and A’). The statistical analysis of granulosa and theca PI (percentage of Ki-67 positive cells/total cells of each follicular compartment) did not reveal a Gaussian distribution. For this reason, the results were expressed as median ranging between 25 and 75%, as summarized in Table 2. Granulosa and theca PI of follicles from classes 3 to 5

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**Table 1** Pig preantral follicle classification.

<table>
<thead>
<tr>
<th>Class</th>
<th>Follicular diameter (µm)</th>
<th>Granulosa layers</th>
<th>Oocytes diameter (µm)</th>
<th>Number of follicles</th>
<th>Theca thickness (µm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>&lt;65</td>
<td>1–2</td>
<td>35–40</td>
<td>94</td>
<td>&lt;5</td>
</tr>
<tr>
<td>2</td>
<td>&gt;65–110</td>
<td>2–4</td>
<td>41–50</td>
<td>53</td>
<td>5–12</td>
</tr>
<tr>
<td>3</td>
<td>&gt;111–180</td>
<td>5–7</td>
<td>51–55</td>
<td>47</td>
<td>11–13</td>
</tr>
<tr>
<td>4</td>
<td>&gt;181–240</td>
<td>8–10</td>
<td>56–70</td>
<td>42</td>
<td>11–13</td>
</tr>
<tr>
<td>6</td>
<td>&gt;300</td>
<td>&gt;13</td>
<td>81–100</td>
<td>40</td>
<td>16–23</td>
</tr>
</tbody>
</table>
Table 2 Somatic (granulosa and theca) proliferation index (PI), vascular area (VA), vascular endothelial growth factor A (VEGFA) mRNA and proportion of proliferating endothelial cells (PE/PT) recorded in the considered classes of pig preantral follicles.

<table>
<thead>
<tr>
<th>Classes</th>
<th>Somatic PI (%)</th>
<th>VA (μm²/field)</th>
<th>VEGFA mRNA (n cells/field)</th>
<th>PE/PT (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>Granulosa</td>
<td>12.50 (4.71–22.61)</td>
<td>18.11 (3.51–27.52)*</td>
<td>3.41 (1.32–7.21)</td>
</tr>
<tr>
<td></td>
<td>Theca</td>
<td>4.12 (2.67–21.17)</td>
<td>40.52 (9.02–46.75)*</td>
<td>5.41 (0.70–9.61)</td>
</tr>
<tr>
<td>4</td>
<td>Granulosa</td>
<td>12.55 (2.87–17.77)</td>
<td>4.25 (2.53–8.06)§</td>
<td>2.01 (0.51–4.32)</td>
</tr>
<tr>
<td></td>
<td>Theca</td>
<td>4.91 (2.85–9.85)*</td>
<td>4.89 (2.45–11.25)§</td>
<td>18.25 (9.72–23.92)</td>
</tr>
<tr>
<td>5</td>
<td>Granulosa</td>
<td>12.61 (2.72–15.67)</td>
<td>41.52 (8.25–56.03) †</td>
<td>2.01 (0.51–4.32)</td>
</tr>
<tr>
<td></td>
<td>Theca</td>
<td>4.91 (2.85–9.85)*</td>
<td>4.89 (2.45–11.25)§</td>
<td>18.25 (9.72–23.92)</td>
</tr>
<tr>
<td></td>
<td>Theca</td>
<td>5.41 (0.70–9.61)</td>
<td>4.25 (2.53–8.06)§</td>
<td>2.01 (0.51–4.32)</td>
</tr>
</tbody>
</table>

The values are expressed as median between the 25 and 75%, since all values did not display a Gaussian distribution. *,†,‡,§Values with different superscripts within each column of parameters indicate data significantly different at least for P<0.05. Granulosa and theca values were compared separately.

showed a similar median distribution (P>0.05) that became significantly higher (P<0.05) in class 6 preantral follicles.

**von Willebrand factor (vWF) and VA**

A specific follicular blood vessel network, identified with the endothelial marker vWF, started to appear in class 3 when preantral follicles had more than four granulosa layers, class 3 and 4 preantral follicles displayed small capillaries scattered within the follicular wall (Fig. 1A and A’). While a specific blood vessel network localized close to the basal membrane, characterized by small vessels, started to appear in classes 5 and 6 (Fig. 1A and A’). In addition, in the outer theca of class 6 preantral follicles, some large blood vessels were recorded (Fig. 1A’).

VA (vWF positive area/10 000 μm², VA) did not follow a Gaussian distribution and for this reason has been expressed as median values (Table 2). Two significant increases in the VA were observed during preantral follicle development: first during the transition from class 3 to classes 4 and 5, and then from classes 4 and 5 to class 6 (P<0.05).

**VEGFA mRNA expression and VEGFA protein localization**

A few somatic cells hybridized for VEGFA mRNA were occasionally observed in class 1 and 2 preantral follicles (data not shown). By contrast, VEGFA mRNA hybridization was regularly observed in cells from class 3 follicles (Fig. 1B and B’).

The median of VEGFA mRNA (percentage of hybridized cells) in granulosa and theca cells of preantral follicles significantly increased during the transition from class 3 to classes 4 and 5 (P<0.05). Moreover, a further significant increase in VEGFA mRNA was then recorded in class 6 both in granulosa and theca cells (Table 2; P<0.05).

A limited and irregular VEGFA immunostaining was observed in follicles of classes 1 and 2 (data not shown). VEGFA positivity, even if highly variable, involved several cells of theca and granulosa compartment of class 3–5 preantral follicles (Fig. 1C and C’). By contrast, all class 6 preantral follicles displayed a high immunopositivity for VEGFA protein. Analysing the global VEGFA pattern in each preantral follicle, it has been observed that there is a positive link between VEGFA mRNA and protein. In fact, on the same follicle and on sequential sections of class 3–5 preantral follicles when the transcript was observed, also the protein of the angiogenic factor was expressed, while in the absence of VEGFA mRNA, there was no immunopositivity for the protein. By contrast, all class 6 preantral follicles showed both VEGFA mRNA and protein (Fig. 1B’ and C’). As in preantral follicles, an analogous VEGFA mRNA and protein expression was also recorded in the oocyte (Fig. 1B’ and C’). VEGFA oocyte expression was confirmed by immunoblot and RT-PCR analyses performed on a pool of oocytes collected from preantral follicles of different classes (Fig. 2).

**Proliferating endothelial cells/proliferating thecal cells (PE/PT)**

A double immunostaining for Ki-67 and vWF (Fig. 1A’) was performed to calculate the PE/PT. The quantification of the PE/PT within the different classes of preantral follicles is summarized in Table 2. In classes 3–5, only some preantral follicles (n=10, 11, and 20 respectively) showed no PEs. By contrast, all class 6 preantral follicles showed PEs. The statistical analysis did not show any significant differences in the PE/PT among all follicular classes considered (P>0.05).

**Correlation between follicular developmental parameters**

A correlation test was carried out to assess how the parameters considered above were related to each other within the different classes of preantral follicles. As summarized in Table 3, the statistical analysis revealed that in classes 3–5, there was a positive correlation between all parameters considered, both in granulosa and in theca compartment (P<0.05 or P<0.01). However, these correlations were lost in class 6 (P>0.05).
Cluster analysis of preantral follicles

By evaluating the distribution of the different parameters recorded for each follicle analysed, multivariate hierarchical analysis revealed several subpopulations within each preantral follicle class (Figs 3 and 4). All classes of follicles could be divided into two subpopulations (1 and 2) that differed significantly for somatic PI, VA and granulosa/theca VEGFA mRNA ($P<0.01$) and PE/PT for class 5 ($P<0.01$).

The subpopulations 1 and 2 of classes 3 and 4 (Fig. 3) could be subsequently divided into four subpopulations (1a and 1b, 2a and 2b). The first two subpopulations (1a and 1b) differed significantly for somatic PI ($P<0.01$ for class 3 and $P<0.05$ for class 4), VA ($P<0.01$) and granulosa/theca VEGFA mRNA ($P<0.01$). The other two subsets (2a and 2b) were significantly different ($P<0.01$) only for the VA.

Preantral follicles of class 5 (Fig. 4) showed a complex variety of subpopulations. Subpopulations 1a and 1b were different in terms of theca ($P<0.05$), granulosa PI ($P<0.01$), VA ($P<0.01$) and granulosa/theca VEGFA mRNA ($P<0.05$). Subpopulations 2a and 2b differed in terms of granulosa PI ($P<0.01$) and VA ($P<0.01$).

Moreover, the subpopulation 1a could be further divided into 1a' and 1a'' on the basis of somatic PI ($P<0.05$), VA ($P<0.01$) and VEGFA mRNA ($P<0.05$ for theca and $P<0.01$ for granulosa). The subpopulation 2a generated two other subpopulations 2a' and 2a'' with differing granulosa PI ($P<0.01$) and VA ($P<0.01$).

Finally, it was possible to recognize in class 6 preantral follicles (Fig. 4) four subpopulations (1a and 1b, 2a and 2b) that were significantly different only for VA ($P<0.01$).

The dendrograms allowed the building of a model that describes the different subpopulations within each class of preantral follicles. As shown in Fig. 5, this model ordinates in a one-dimensional space the subpopulations using the distances between the bifurcations inside the dendrograms, which are proportional to the homologies between the subpopulations themselves.

Discussion

The present research analyses how important parameters of the somatic and vascular compartment change during the process of preantral follicle growth. To achieve this aim, somatic index (PI), proportion of PEs (PE/PT), VA and VEGFA expression (VEGFA mRNA and protein) were simultaneously recorded on single healthy preantral follicles classified into six different stages of development as proposed by Morbeck et al. (1992). This precise morphological classification was adopted to better interpret the data obtained, distinct from other studies where preantral follicles were considered as early or late structures when the granulosa compartment was

Figure 1 Representative micrographs of single pig healthy preantral follicle sections belonging to the different classes (from 3 to 6) considered. Left panel (A and A'): intrafollicular distribution of vWF (red stain) and Ki-67, a cellular proliferative marker (green stain), were revealed using a double immunohistochemical technique. The cell nuclei were counterstained with DAPI (blue stain) to visualize the tissue morphology and to identify the follicular compartments. Middle panel (B and B'): in situ hybridization is shown to evaluate the intrafollicular distribution of the VEGFA mRNA. Right panel (C and C'): location of the VEGFA protein within the follicular and germinal compartments. For the classes 3–5, two different examples were given: the three panels on the top (A, B and C) and bottom (A', B' and C') display respectively the sequential sections of a representative follicle with low and high activity. Bar = 30 μm.
was occasionally observed in the follicles of classes 1 and 2, on the contrary, it was always detected in preantral follicles during the late stages of development. Furthermore, the analysis of VEGFA expression revealed that the angiogenic stimulus progressively increases from class 3 to classes 4 and 5, reaching its highest levels during the final stage of development (class 6). The theca VEGFA mRNA that appeared low in preantral follicles from classes 3 to 5 becomes significantly higher in class 6, when a parallel increase in the VA was recorded. Stable and high levels of VEGFA expression both in the theca and in granulosa compartments characterizes the final stage of preantral follicle development when the follicular walls are colonized by a widespread capillary network near the basal membrane and high levels of angiogenic factors may be required to maintain and stabilize these immature blood vessels (Carmeliet et al. 1996, Ferrara 1996, Ferrara & Davis-Smyth 1997).

In addition, high levels of VEGFA appeared within the follicular structure just before the process of antrum formation. In this context, microvessel permeability may be imposed by VEGFA to stimulate plasma extravasations allowing the accumulation of fluids within the differentiating follicular cavity (Isobe et al. 2005).

The VEGFA within preantral follicles also seems to be dependent on the oocyte in order to express and synthesize the angiogenic factor. Similar to other molecules, VEGFA could be included among the signals involved in the bidirectional control of oocyte–follicle function (Díaz et al. 2007, Hutt & Albertini 2007), by guaranteeing that follicle development can be synchronized with the growth of the metabolically active germinal cells. Even if it has been demonstrated that there are also post-transcriptional and post-translational mechanisms involved in VEGFA expression in the ovary and other tissues (Hazzard et al. 1999, Neufeld et al. 1999, Martelli et al. 2006), in this research all preantral follicles showed a relationship between VEGFA mRNA and protein production. In fact, the follicles and the oocytes that expressed the transcript were also positive.
Figure 3 Dendrograms derived from cluster analysis of the data obtained on single follicles of classes 3 and 4. The cutting values (arrows) were determined evaluating the statistically differences among the examined parameters. A, granulosa/theca PI; B, VA; C, granulosa/theca VEGFA mRNA; D, proportion of proliferating endothelial cells (PE/PT). *, **Significantly different values for $P < 0.05$ and $P < 0.01$ respectively.
Figure 4 Dendrograms derived from cluster analysis of the data obtained on single follicles of classes 5 and 6. The cutting values (arrows) were determined evaluating the statistically differences among the examined parameters. A, granulosa/theca PI; B, VA; C, granulosa/theca VEGFA mRNA; D, proportion of proliferating endothelial cells (PE/PT). *,**Significantly different values for $P<0.05$ and $P<0.01$ respectively.
to the related protein. By contrast, preantral follicles that were characterized by the absence of VEGFA mRNA were also negative to the protein.

Follicular angiogenesis in classes 3–5 preantral follicles was quite variable. In fact, some structures did not display PEs (21, 26 and 45% in classes 3, 4 and 5 respectively), while, in the remaining follicles belonging to these classes, the incidence of mitosis within the endothelial cells ranged between 30 and 60%. These data suggest that the follicular angiogenesis in classes 3–5 could be in switch-off or switch-on position in response to specific stimuli. On the contrary, when preantral follicles reached the late stage of growth (class 6), probably under the stimulus of high and stable levels of VEGFA, angiogenesis remained active and the proportion of PEs was high.

These results seem to be in agreement with the recent experiments of Yang & Fortune (2007) showing that

**Figure 5** A general model that describes the distribution of the follicular subpopulations identified inside each follicular class by using cluster analysis. The subpopulations were represented within each class and arranged on the y-axis by considering the distances between the bifurcation obtained in the dendograms. The grey scale represents significant differences recovered for somatic and vascular parameters. The thickness of the dotted line represents the statistical difference among classes, tPl, theca proliferation index; gPl, granulosa proliferation index; VA, vascular area; tVEGFA, theca VEGFA mRNA; gVEGFA, granulosa VEGFA mRNA; PE/PT, proportion of proliferating endothelial cells; % value, percentage of preantral follicles belonging to the subpopulation.
**Vasculature during preantral follicle development**

**vitro** VEGFA is able to stimulate the early preantral follicles but is totally ineffective on the progression of the later ones, even after long periods of culture.

Interpreting the results of the correlation analysis between all the vascular parameters recorded (VEGFA mRNA, VA and proportion of PEs) and the somatic PI in classes 3–5, it could be hypothesized that the small (classes 3 and 4) and middle (class 5) preantral structures are divided into two sets of follicles: one able to increase its vascular network, and another one inactive in terms of angiogenesis and is quiescent.

On the basis of these data, it could be concluded that the process of angiogenesis, also during the preantral stage of folliculogenesis, appears crucial to sustain the increasing metabolic requirements during the process of growth. In addition, other new roles for the PEs could be supposed (Lammert et al. 2001, Roberts et al. 2007).

Recently, endothelial cells and nascent vessels have been directly associated with the growth and differentiation of tissues/organs providing growth and/or morphogenetic signals (Lammert et al. 2001). Part of this vascular influence seems to be mediated by the VEGFA itself that was hypothesized to have a direct effect on non-vascular elements in a variety of tissues, including the ovary (Roberts et al. 2007). The strict correlation observed between somatic and vascular parameters may represent additional evidence that VEGFA may be capable of direct stimulation of the somatic components, over its indirect influence on follicular development through the control of vascular physiology (Otani et al. 1999, Danforth et al. 2003, Abramovich et al. 2006).

The positive correlation existing between the morphological and vascular parameters disappeared in preantral follicles of class 6, when somatic PI and the angiogenic functions (VEGFA expression, VA and PE/PT) remained at high and stable levels. The high somatic and vascular parameters recorded in class 6 preantral follicles, which differentiates this late stage of growth from all the earlier ones, may be explained by considering previous evidence demonstrating that during this final stage of secondary follicle development sensitivity to the FSH occurs (Sicinski et al. 1996, McGee et al. 1997, McGee & Hsueh 2000).

Furthermore, the data obtained identified the VA as the crucial parameter that is essential to discriminate between growing preantral follicles. In fact, the VA initially increased when follicles passed from class 3 to classes 4 and 5 and then during the late stage of growth (class 6). Thus, the first enhancement in the VA may be necessary to sustain the increased requirements in nutrients and gases in growing structures, while the second increase may guarantee the major delivery of endocrine and paracrine molecules that metabolically change and drive the function of late preantral follicles (Fortune 2003, Gougeon 2004).

The cluster analysis that aggregates the preantral follicles on the basis of reciprocal similarity allowed the building of dendrograms (Figs 3 and 4) representing the data subsets inside each follicular class considered. The statistical analysis stated that this articulation was not a consequence of a mathematical artefact, but that the biological parameters really differed among follicle subpopulations. This statistical technique leads to the establishment of a model representing preantral follicle development (Fig. 5). This model showed that the majority of the small preantral follicles (classes 3 and 4) are present in the ovary as quiescent structures (±70%), or as active follicles that start their growth slowly. In particular, the small active preantral follicles had differing VA, indicating that the extension of the blood vessel network may become the crucial element for the follicles that are recruited to growth. Class 5 represents a transitory category between the earlier (classes 3 and 4) and later (class 6) stage of preantral development. Half of the class 5 follicles maintain a very low activity similar to that recorded in classes 3 and 4, but the remaining follicles express high levels of granulosa PI and VA. These results suggest that in the preantral follicles of class 5 with a higher activity a simultaneous contribution of the somatic and vascular component is required within this class of preantral follicles to guarantee the progression of the follicles towards the final stages of development. In fact, when the follicles enter into class 6 (subpopulation 1a: 35% of follicles) somatic and vascular parameters appear similar to that observed in class 5 preantral follicles showing the highest activity. Class 6 follicles (large follicles) maintained constant high level of somatic and vascular parameters with the proportion of PEs at stable high levels appearing as definitively selected structures. In particular, the class 6 differential element was the VA that is the crucial and often the unique parameter that accompanied the final process of growing in the majority of the stages of preantral follicle development. Hence, the progressive increase in the VA represents the final element of differentiation among the subpopulations in the different stages of preantral follicles, and in those that reach the late stage of development where it is probably required to sustain the process of antrum differentiation.

In conclusion, the classification obtained in the present research permits division of preantral follicles into three different stages (early, mid and late) superimposing the morphological parameters proposed by Fortune (2003) with the developmental parameters analysed in this research. It is known that under physiological conditions, preantral follicle growth occurs within a rich milieu of endocrine, paracrine and autocrine factors (Kotsuji & Tominaga 1994, Gelety & Magoffin 1997). As a consequence, it may be hypothesised that the availability of an adequate vascular supply providing adequate levels of endocrine and paracrine signals may play a key role in follicle selection.

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during preantral follicle growth (Danforth 1995, Hull et al. 2003, Roberts et al. 2007). In conclusion, the data presented suggest that preantral follicles appear as autonomous units engaged in a competition for growth, where the vascular supply represents a crucial prerequisite for the development and selection.

**Materials and Methods**

**Tissue collection and experimental protocols**

Swine ovaries in the luteinic phase of the reproductive cycle were collected at the local slaughterhouse. Once in the laboratory, each ovary was divided into three to five portions. One part was used to mechanically isolate preantral follicles for the biochemical investigations, and the other portions were immediately fixed in 4% paraformaldehyde/PBS (pH 7.4) for 12 h at 4 °C. After dehydration, each tissue sample was embedded in paraffin wax and serially sectioned at 5 μm thickness.

All the follicular sections of each preantral follicle were collected on poly-l-lysine-coated slides (Sigma) and then sequentially processed for HE, immunohistochemical and in situ hybridization analyses. In detail, each preantral follicle was used to perform, at least in double, the following morphological analyses:

- **a)** caspase-3 and in situ TUNEL to distinguish between atretic and healthy follicles (Berardinelli et al. 2004);
- **b)** vWF to identify the endothelial cells and to quantify the extension of the VA (Barboni et al. 2004, Martelli et al. 2006);
- **c)** Ki-67 antigen, a cell proliferation marker, to quantify the degree of mitotic granulosa, theca and endothelial cells (Martelli et al. 2006);
- **d)** VEGFA protein to describe the intrafollicular distribution of the angiogenic factor (Martelli et al. 2006) and
- **e)** VEGFA mRNA detected by in situ hybridization (Barboni et al. 2004, Martelli et al. 2006).

The morphological analyses were performed with an Axioskop 2 Plus epifluorescence microscope (Zeiss, Oberkothen, Germany) equipped with a cooled colour charge-coupled device camera (AxioVision Cam, Zeiss) interfaced to a computer workstation and provided with an interactive and automatic image analyser (AxioVision, Zeiss).

To exclude individual differences among animals, the follicles were collected from at least five different pigs.

**Identification and classification of preantral follicles**

Follicular stages were defined according to Wulff et al. (2001): primary follicles (containing only one granulosa cell layer), preantral follicles (more than two granulosa cell layers and no antrum) and early antral follicles (one or more fluid-filled spaces in the granulosa layer).

In this study, only healthy preantral follicles were analysed. Pig healthy follicles were identified combining the HE with the caspase-3/TUNEL staining, according to Wulff et al. (2002) and Berardinelli et al. (2004).

On two or more HE sections, mean follicular diameter was calculated using the KS300 computed image analysis system (Zeiss), set to measure two diameters of the follicle/oocyte section at right angles and only symmetrical follicles (right angle cross sections within 10% of each other) were considered (Morbeck et al. 1992). Since in the early stages of preantral follicles the outer boundary of the theca layers could not be defined precisely, the follicular diameter was always measured from the basement membrane immediately outside the granulosa cells (Lundy et al. 1999, Griffin et al. 2006). When the theca compartment was clearly evident, it was outlined and the mean thecal thickness measured. In all sections of each preantral follicle analysed, the whole cross section was recorded and theca and granulosa compartments were outlined and examined separately. The morphometric parameters (follicle diameters and number of granulosa cell layers) were calculated by two investigators in a blinded fashion, and the correlation coefficient for follicle counts between the two investigators (Zimmermann et al. 2003) was 0.915. On the basis of mean follicular diameter and the number of granulosa layers, preantral follicles were classified into six different categories adopting, with minor modifications, the classification developed by (Morbeck et al. 1992)

- class 1: <65 μm and 1–2 layers of granulosa cells,
- class 2: from 65 to 110 μm with 2–4 layers of granulosa cells,
- class 3: from 111 to 180 μm with 5–7 layers of granulosa cells,
- class 4: from 181 to 240 μm with 8–10 layers of granulosa cells,
- class 5: from 241 to 300 μm with 11–13 layers of granulosa cells and
- class 6: follicular diameter >300 μm with >13 layers of granulosa cells.

At least, 40 healthy follicles were processed and analysed in each preantral follicle category considered.

**Immunohistochemistry**

**Caspase-3 and in situ TUNEL**

At least, two sections of each preantral follicle were simultaneously exposed to immunohistochemical analysis for the detection of the nucleosomal fragmentation and caspase-3 distribution (Berardinelli et al. 2004). According to Berardinelli et al. (2004), on the same section were applied the ApopTag in situ Apoptosis detection kit (Intergen Company, Oxford, UK), and a rabbit polyclonal antibody CPP32 (caspase-3; Novocasta Laboratories Ltd, Newcastle upon Tyne, UK), which recognizes the proenzyme as well as the active form. As a negative control, the sections were incubated with the omission of DNTT and CASP3, and the equilibration buffer was substituted for the volume of the two reagents, as suggested by the manufacturer. As a positive control, normal female rodent mammary gland tissue sections were used (Intergen Company). The specificity of the double immunostaining was verified by localizing each antigen separately (data not shown). Finally, the tissue sections were counterstained with 4',6-
diamidino-2-phenylindole (DAPI, Sigma) at 1:100 in PBS for 10 min (Martelli et al. 2006).

Tissue sections were analysed at a magnification of 400×. The results were expressed as the percentage of apoptotic cells (TUNEL-positive and/or caspase-3-positive cells) on the total number of cells (DAPI-stained nuclei; Wulff et al. 2002, Berardinelli et al. 2004).

Ki-67 and vWF
Double immunostaining for Ki-67 and vWF was performed according to Polak & Van Noorden (1997). Tissue sections, after rehydration, were kept in 2% hydrogen peroxide (Merck) in methanol (Merck) for 5 min to remove endogenous peroxidase activity. The slides were placed in citrate buffer (pH 6), and antigen retrieval was performed by treating the sections twice in a hotbox oven (HO) at 95°C, washed with PBS for 5 min and then in TNT buffer (0.1 M Tris–HCl (pH 7.5), containing 0.15 M NaCl and 0.05% Tween 20) for 5 min twice. After incubation in TNB blocking buffer (0.1 M Tris–HCl (pH 7.5), 0.15 M NaCl and 0.5% of the blocking reagent supplied in the kit) for 1 h to prevent non-specific binding, the sections were incubated with a mouse monoclonal anti-Ki-67 primary antibody (Dako, Glostrup, Denmark) diluted 1:75 in TNT, and with a rabbit anti-vWF polyclonal primary antibody (Dako) diluted 1:400 in TNT, at RT overnight. vWF immunocomplexes were revealed by incubating the tissue sections with CY3-labelled secondary goat anti-rabbit antibody (Chemicon, Billerica, MA, USA) diluted 1:100 in TNT. Ki-67 immunocomplexes were incubated with biotinylated–conjugated anti-mouse antibody (Sigma), and then detected by TSA Fluorescein System (Perkin–Elmer Life Sciences, Boston, MA, USA), according to the manufacturer’s protocol. TNB blocking buffer was used in place of the primary antisera as a negative control.

As a positive control, human breast carcinoma tissue samples (Heffelfinger et al. 2000, Martelli et al. 2006), kindly gifted by Dr De Carolis from the ‘Giuseppe Mazzini’ Hospital of Teramo, were used (Supplementary Fig. 1, which can be viewed online at www.reproduction-online.org/supplemental). The specificity of the double immunostaining was verified by localizing each antigen separately (data not shown). Tissue sections were counterstained with DAPI and analysed at a magnification of 400×. The quantification of the digitized fluorescent signals was accomplished using a semi-automated algorithm written with the image analysis system KS300. A guided program (macro for KS300) was created to count 1) the number of proliferating granulosa or theca cells (green-stained cells), 2) the number of PEs (dual-stained cells for Ki-67 and vWF) and 3) the total number (blue-stained cells) of granulosa or theca cells, inside a fixed area of 10 000 μm² of granulosa/theca compartment. In brief, after background densitometric calibration, the algorithm detected and separated the fluorescent regions of interest from their background and created a new image on the basis of their grey values. Thus, the process created a binary image from true colour image. The analyses were performed on the whole area of the cross section. The somatic (granulosa/theca) PI was calculated as the percentage of Ki-67-positive granulosa or theca cells on the number of total somatic cells in the relative compartment. As a consequence, for each follicle, the PI was expressed either for theca or granulosa layers (Wulff et al. 2001, Watson & Al Zi’abii 2002). Moreover, the proportion of PEs was calculated as the percentage of PEs (number of dual-stained cells for Ki-67 and vWF) on the total number of proliferating theca cells (PE/PT; Wulff et al. 2001, Bauer et al. 2003).

In order to eliminate from the quantitative analyses the background due to the red blood cell autofluorescence, the sections were restained with HE and micrographs of the same fluorescent fields were performed to identify and subtract the red blood cells within each blood vessel analysed.

vWF
In order to identify endothelial cells, after endogenous peroxidase inhibition and HO treatment performed as described previously, the slides were incubated first with normal goat serum (NGS; Sigma) for 30 min and then with a rabbit antibody against vWF (the same antibody used for the double immunostaining; Dako). For this analysis, the secondary antibody used was an anti-rabbit biotinylated–conjugated (1:100 in PBS/1% BSA) applied for 1 h at RT. The samples were finally exposed to the avidin–biotin peroxidase complex ( Vectastain ABC kit; Vector Laboratories, Burlingame, CA, USA) for 1 h at RT. The immunocomplex was then detected using 3,3′-diaminobenzidine (DAB; DBH Laboratory Supplies, Poole, UK) for 7 min. To enhance the DAB reaction products, the ammonium nickel sulphate method was performed (Shu et al. 1988), which turns the brown DAB reaction into a black one. Sections were not counterstained so that quantitative image analysis could be performed (Wulff et al. 2001, Martelli et al. 2006). To quantify the VA (i.e. vWF immunopositivity), the tissue sections were measured at a magnification of 400×. In brief, after geometric and densitometric calibration, the captured grey scale image was thresholded and converted into a binary image. The image analysis system generates one image where the terminal total area appears in black against a white background. Using a semi-automated algorithm, the computer program provides measurements of the area by counting the total number of pixels in the field. The VA was given by the extension of vWF-positive area in μm²/10 000 μm². The analysis was performed on the whole area of the cross section (Barboni et al. 2004, Martelli et al. 2006).

VEGFA
The distribution of VEGFA within the follicular somatic compartments was visualized, after endogenous peroxidase inhibition and HO treatment, with a primary rabbit anti-human VEGFA (Calbiochem, Darmstadt, Germany) diluted 1:10 in PBS/1% BSA, at RT overnight. After a secondary biotinylated–conjugated anti-rabbit antibody (1:100 in PBS/1% BSA; Sigma), the immunocomplex was detected by the DAB reaction products as described above. Human breast carcinoma tissue samples (Heffelfinger et al. 2000, Martelli et al. 2006) were used as a positive control. NGS was used as a negative control in place of primary antibody (Supplementary Fig. 1).

Two or more sections per preantral follicle were qualitatively analysed to evaluate the intrafollicular presence and distribution of the angiogenic factor.

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In situ hybridisation

In situ hybridisation was performed with a commercially available kit (HybriProbe kit; Biognostik, Göttingen, Germany) according to Barboni et al. (2004) and Martelli et al. (2006). A custom synthesized oligonucleotide double-FITC-labelled cDNA probe for porcine VEGFA	extsubscript{164} was obtained from Biognostik. The design process of the oligonucleotide sequence was based on accession no. X81380 (porcine mRNA for VEGF). Probe sequence (5'-3') was as follows: 5'-CAC GTC TGC GGA TCT TGT ACA AAC AAA TGC-3' (Barboni et al. 2004, Martelli et al. 2006).

As negative control, sections were treated with a random double-FITC-labelled oligonucleotide probe containing the same proportion of cytosine and guanine bases as the VEGFA probe supplied by the manufacturer. According to the manufacturer's protocol, as a positive control, the probe for mRNA of housekeeping gene β-actin was used (Supplementary Fig. 1).

To identify the VEGFA mRNA expressing cells, the tissue sections were assessed at 400X magnification. After geometric and densitometric calibration, the image was captured in grey scale, thresholded and converted to a binary image. The results were expressed as the number of positive cells/unit area (10,000 μm²), calculated individually for theca and granulosa compartments (Barboni et al. 2004, Martelli et al. 2006) for each preantral follicle analysed.

Biochemical investigations

Oocytes collection

Preantral follicles were mechanically isolated, with the aid of a stereomicroscope, as described previously (Cecconi et al. 1999, Shuttleworth et al. 2002). After isolation, healthy follicles were selected on the basis of their morphology (Shuttleworth et al. 2002). Follicle diameters were measured with the aid of an inverted phase microscope equipped with an ocular micrometer (40X magnification), and class 3–6 preantral follicles were collected. From each follicle, the cumulus–oocyte complexes were recovered, and classified as healthy if they displayed a compact cumulus and an oocyte with a homogeneous ooplasm (Cecconi et al. 1999). Oocytes were mechanically separated from the surrounding cumulus cells, and zona pellucida, pooled (150 oocytes) and frozen until use for western blot or RT-PCR analyses. To exclude individual differences among animals, the oocytes were collected from at least ten different pigs.

Western blot analysis

Total proteins were extracted from denuded oocytes according to Cecconi et al. (2008). Briefly, the cells were transferred into 30 μl extraction buffer (Cecconi et al. 2008) and an aliquot was used to evaluate the amount of protein by Lowry’s method (Lowry et al. 1951). The proteins extracted from the granulosa layer (Martelli et al. 2006) were used as positive control of the procedure.

Fifty micrograms of protein were separated by 12% SDS-PAGE and then electrophoretically transferred to a nitrocellulose membrane (Hybond-C Extra; Amersham Pharmacia) for immunoblot analysis, according to standard protocols (Towbin et al. 1979). According to Martelli et al. (2006), the membranes were incubated with the polyclonal anti-human VEGFA-Ab2 (dilution 1:80; Oncogene, San Diego, CA, USA) that recognizes the swine VEGFA protein, and then with the peroxidase-conjugated anti-rabbit antibody (dilution 1:4000; Santa Cruz Biotechnology, Inc., Santa Cruz, CA, USA). After the stripping procedures (Martelli et al. 2006), the filter was reprobed with a monoclonal anti-α-tubulin (dilution 1:5000; Sigma) and then with a peroxidase-conjugated anti-mouse antibody (dilution 1:4000; Santa Cruz Biotechnology, Inc.). The signals were detected using the ECL western blot analysis system (Amersham Pharmacia).

Quantitative data were determined as the mean ratio of the optical density of the specific bands normalized to that of α-tubulin. The densitometric analysis was carried out with the Advanced Image Data Analyzer (Rai Test, GmbH, Straubing, Germany; Martelli et al. 2006). The experiments were repeated three times independently with similar results.

Total RNA extraction and reverse transcription

Total RNA from denuded oocytes or granulosa cells (as positive control) was extracted by TRIzol reagent (Invitrogen Life Technologies), according to the manufacturer’s instructions. Purified RNA was resuspended in 15 μl RNase-free water and was spectrophotometrically quantified (A260 nm). One microgram of RNA was electrophoretically separated in 1% agarose gel in order to determine total RNA quality. Two micrograms of total RNA were reverse transcribed (RT) to cDNA using Omniscript Reverse Transcriptase kit (Qiagen SpA) in a final volume of 20 μl, according to the manufacturer’s instruction. Transcription reactions without RT were performed as a control for DNA contamination.

PCR analyses

Based on swine gene sequences present in the GenBank, the primer for VEGFA	extsubscript{164} splice variant was designed using Primers Express Software (Applied Biosystems, Warrington, UK). The following primers were used:

- VEGFA	extsubscript{164} (accession no. X81380): 5'-GAAGTGGTG-GAAGTTGATGGA-3' (forward) and 5'-GCCTGCAACC-GAGTCTGT-3' (reverse).
- β-Actin (accession no. U07786, (Shimizu et al. 2002)): 5'-ATCGTGCGGGACATCAAGGA-3' (forward) and 5'-AGGAGGAGGTCGTAAGAG-3' (reverse).

They amplify at 507 and 178 bp fragments size respectively. Semi-quantitative PCR was performed in the ThermoCycler (Biometra GmbH, Goettingen) using ReadyMix Taq without MgCl₂ (Sigma). A master mix for the following reaction components was prepared to the following final concentrations: 1.25 μl forward primer (0.25 μM), 1.25 μl reverse primer (0.25 μM), 12.5 μl PCR master mix, 1.5 μl MgCl₂ (1.5 μM), 5 μl cDNA and 3.5 μl RNase/DNase-free water to final volume of 25 μl. All samples were performed in duplicate for all genes. The PCR protocol employed was as follows: start denaturation
for 5 min at 95 °C, 40 cycles at 95 °C for 1 min, 56 °C for 1 min, 72 °C for 1 min and long extension for 4 min at 72 °C. Each sample was run in duplicate, and the results were normalized to the level of housekeeping β-actin gene mRNA levels.

**Statistical analysis**

The data obtained from each preantral follicle were analysed within the different classes of follicles expressed as median and range (first and third quartiles). The values of somatic (granulosa/theca) PI, VA, VEGFA mRNA and ratio of PEs to PTs (PE/PT) variables obtained within the different classes of preantral follicles were then compared using the non-parametric Kruskal–Wallis test followed by the Conover–Inman post hoc test. The correlation existing between each couple of variables was measured by a correlation test. The statistical difference was set at P<0.05 or P<0.01.

Finally, for each class of preantral follicle, a multivariate hierarchical cluster analysis was performed to build a dendrogram aggregating the data of somatic PI, VA, VEGFA mRNA and PE/PT obtained from each single follicle analysed to determine their similarity. In particular, this statistical technique performs the partitioning of a dataset (preantral follicles) into data subsets (clusters) defined on the basis of the considered parameters obtained from each single follicle and analysed altogether. Each subset shares a common trait (the proximity) according to a defined distance measure, which will determine how the similarity of elements is calculated. Hierarchical clustering builds a hierarchy of clusters represented by a tree (called a dendrogram), with individual elements at one end and a single cluster containing every element at the other. Since the dendrogram ordnates all the follicles into data subsets, a non-parametric statistical analysis (as described above) was performed to assess the significance of each partition. Where the difference was not significant (P>0.05), the dendrogram was cut (cutting value).

For all calculations, the StatistiXL 1.7 beta program was used.

**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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