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Research Article

Follicular fluid steroid profile in sows: relationship to follicle size and oocyte quality[†]

N. G. J. Costermans^{1,2}, N. M. Soede², F. van Tricht³, M. Blokland³, B. Kemp², J. Keijer¹ and K. J. Teerds^{1,*}

¹Human and Animal Physiology, Wageningen University and Research, Wageningen, The Netherlands, ²Adaptation Physiology, Wageningen University and Research, Wageningen, The Netherlands, and ³Wageningen Food Safety Research (WFSR), Wageningen University and Research, Wageningen, The Netherlands

*Correspondence: Human and Animal Physiology, Wageningen University and Research, P.O. Box 338, 6700 AH Wageningen, The Netherlands. Tel: +31 317484136; E-mail: katja.teerds@wur.nl.

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Abstract

Identification of reliable characteristics of follicle guality and developmental competence has been pursued in numerous studies, but with inconsistent outcomes. Here, we aimed to identify these characteristics by analysis of the follicular fluid (FF) steroid profile in relation to cumulus-oocyte complex (COC) morphology and follicle size, followed by molecular substantiation. Multiparous sows at weaning were used to facilitate analysis at the start of the follicular phase of the oestrus cycle. Sows with a higher average follicle size (>5 mm vs. < 5 mm) had a higher follicular fluid β estradiol concentration, but did not differ in other measured steroids. Sows with high compared to low percentage high-quality COCs (<70% vs. ≥70% high-quality) had follicular fluid with a higher concentration of β -estradiol, 19-norandrostenedione, progesterone, and α -testosterone, while the concentration of cortisol was lower. Transcriptome analysis of granulosa cells of healthy follicles of sows with a high percentage high-guality COCs showed higher abundance of transcripts involved in ovarian steroidogenesis (e.g., CYP19A2 and 3, POR, VEGFA) and growth (IGF1) and differential abundance of transcripts involved in granulosa cell apoptosis (e.g., GADD45A, INHBB). Differences in aromatase transcript abundance (CYP19A1, 2 and 3) were confirmed at the protein level. In addition, sows with a high percentage high-quality COCs lost less weight during lactation and had higher plasma IGF1 concentration at weaning, which may have affected COC quality. To the best of our knowledge, this study is also the first to report the relation between FF steroid profile and COC quality.

Summary sentence

Sows with a higher percentage high-quality COCs have a distinct follicular fluid steroid profile and concomitant granulosa cell transcriptome.

Key words: steroid hormones, granulosa cells, metabolism, insulin-like growth factor, porcine, follicle

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Introduction

During ovarian follicular development, most follicles degenerate via follicular atresia, while only a few follicles reach the pre-ovulatory state from which an oocyte ovulates to produce viable offspring [1]. Hence, from a reproductive point of view, selection of the best quality follicles is of utmost importance. It is therefore not surprising that a vast number of studies have been dedicated to identifying reliable predictors of follicle and oocyte quality, which could be useful for the selection of occytes with high developmental competence in *in vitro* fertilization (IVF) procedures [2, 3].

Some studies have investigated relationships between morphological characteristics of follicles or cumulus-oocyte complexes (COCs) and oocyte developmental competence, with very variable results. A number of these studies reported the presence of relations between antral follicle size and oocyte developmental competence, where oocytes from larger antral follicles show an increased blastocyst formation rate and implantation rate [4, 5]. Other studies did not find such relations [6, 7]. Distinct morphological classifications have also been used to assess oocyte competence, such as the number of cumulus cell layers [8], darkness of the ooplasm [9] and number of oocyte anomalies [10], again with variable outcomes [11]. Follicular fluid (FF) steroid profiling could provide a new tool to identify reliable markers for follicle quality and developmental competence, as FF steroid composition determines the microenvironment in which the oocyte develops. For example, estradiol and progesterone can improve oocyte developmental competence during in vitro maturation (IVM) in swine and cattle, when administered using optimal timing and dosing [12, 13]. Also, in swine, greater cleavage and blastocyst formation rates were found when oocytes were matured in FF with higher testosterone and androstenedione concentrations during IVM [14]. FF steroid concentrations can therefore influence oocyte developmental competence.

The aim of this study was to identify potential markers for follicle quality, by studying relations between FF steroid profile, COC morphology, and follicle size. As FF composition is also dependent on the steroidogenic activity of granulosa cells [15], we additionally analyzed granulosa cell transcript abundance using whole-genome transcriptome analysis to explain underlying mechanisms of sow differences in COC morphology and FF steroid profiles. We therefore hypothesized that the follicular fluid steroid profile and granulosa cell transcriptome differ between sows with a high vs. low percentage high-quality COCs, already at the start of the follicular phase onwards.

Studies on follicle and oocyte developmental capacity have mainly used widely available pre-pubertal porcine or bovine ovaries from the slaughterhouse, where the stage of the oestrus cycle is not well controlled or absent. Therefore, we used sows at the moment of weaning, as sows have a well-defined start of the follicular phase at the end of lactation [16]. In many animal species, but especially in sows, the metabolic state highly influences follicular development and FF content [17, 18]. Therefore, we additionally analyzed the sows' metabolic state and its relations with follicular development and FF content.

Materials and methods

The experiment was approved by the Animal Care and Use Committee of Wageningen University (DEC2016036) and performed according to national and EU guidelines.

Animals

A total of 29 multiparous Dutch Landrace sows (parity 3 to 5; Topigs Norsvin, Vught, the Netherlands) with an average parity of 3.8 ± 0.2 were used. The sows were weighed approximately 1 week before parturition and immediately after weaning. Sow weight after parturition was estimated to calculate body weight loss during lactation, as described in Costermans *et al.* [19].

Blood and ovary collection

After a lactation of 26.1 ± 0.2 days, sows were killed by stunning and exsanguination within 2 h after weaning. Blood was collected in 9 ml ice-cold EDTA activator tubes (Greiner Bio-One, Monroe, NC, USA) and centrifuged at $3000 \times g$ for 10 min at 4°C to collect plasma. The samples were stored at -20° C until further analysis. Left ovaries were immediately placed in an insulated container and after arrival in the lab placed in PBS pH 7.4 in a water bath at 37° C until follicle aspiration. Right ovaries were immediately snap-frozen in liquid N₂ and stored in -80° C until follicle size, follicle health, and transcriptome analyses were performed.

Measurements

The experimental procedure used to assess ovarian measurements is summarized in Figure 1.

Left ovary

COC morphology + amount of follicular fluid. The 15 largest follicles of the left ovary were used for follicle aspiration within 5 h from slaughter. The contents of the 15 follicles were pooled and collected in a tube and allowed to settle for 5 min. The supernatant was removed and centrifuged at 1900 × g at 4°C for 30 min to separate cells from the follicular fluid. The total volume of follicular fluid was assessed using reverse pipetting and subsequently stored at -20° C until further analysis. The recovered cumulus-oocyte complexes (COCs) were morphologically classified under a dissection microscope as high-quality (intact cumulus and normal-shaped oocyte) or low-quality (degraded cumulus or degenerated oocyte) similar to Alvarez *et al.* [20]. For each sow, the percentage high-quality COCs was calculated.

Steroid profiling. The follicular fluid of the 15 largest follicles of the left ovary was pooled to obtain a sufficient sample volume needed for endogenous steroid hormone profiling. A modified UHPLC-MS/MS method as described in Blokland *et al.* [21] was used to detect endogenous aglycons in FF. In short: 900 µl water was added to 100 µl follicular fluid and followed by solid phase extraction using an Oasis HLB 96-well SPE plate (Waters, Milford, CA, USA), derivatization of aglycons was performed with picolinic acid. Chromatographic separation of the aglycons was achieved on a Waters BEH C₁₈ column (Waters) followed by analysis on a Xevo TQS mass spectrometer (Waters) in positive ESI mode.

Right ovary

Follicle size + *follicle health*. Right frozen ovaries were held against a ruler and photographed from different sides to measure the size of the 15 largest follicles using ImageJ (version 1.51f, National Institutes of Health). These 15 largest follicles were assumed to represent approximately half of the ovulatory follicle pool, as ovulation rates in modern sows are around 25–30 [22]. Subsequently, follicle health was assessed using an immunohistochemical staining against the apoptotic marker cleaved-Caspase 3 (cCASP), as described in

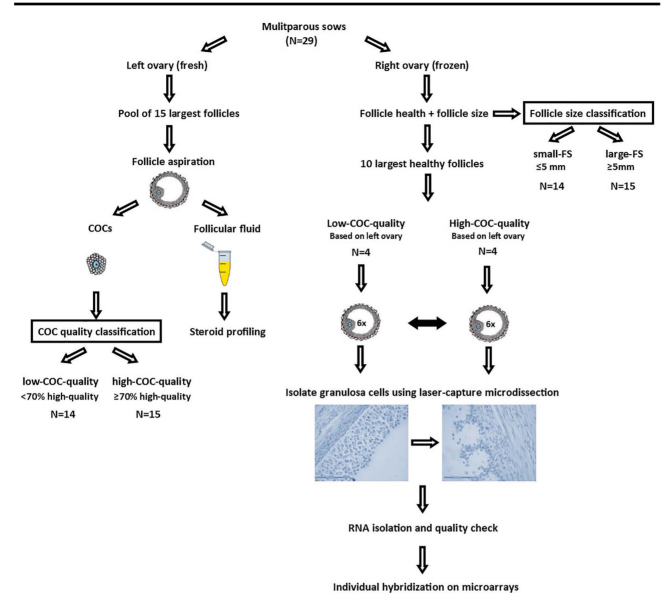


Figure 1. Experimental set-up of a total of 29 multiparous sows; left ovaries were used for follicle aspiration of the 15 largest follicles. Follicular fluid was used for steroid profiling using a modified UHPLC-MS/MS method, and cumulus-oocyte complexes (COCs) were morphologically classified as high-quality or low-quality to determine the percentage of high-quality COCs for each sow. Sows with < 70% high-quality COCs were classified as low-COC-quality and sows with $\ge 70\%$ high-quality COCs as high-COC-quality. Right ovaries were used to determine follicle size and health status of all visible follicles of the right ovary, by using a cleaved-caspase 3 staining. Average follicle size of the 15 largest follicles was used to determine the follicle size class (small-FS < 5 mm vs. large-FS ≥ 5 mm). A subset of 8 out of 29 sows was used for whole-genome transcriptome analysis of granulosa cells, of which 4 were classified as low-COC-quality (54 \pm 12% high-quality COCs) and 4 were classified as high-COC-quality (93 \pm 15% high-quality COCs), as based on the left ovary. For each sow, granulosa cells of 6 healthy follicles of the pool of 10 largest healthy follicles were isolated using laser capture microdissection, after which samples were individually hybridized onto whole-genome microarrays for whole- genome transcriptome analysis.

Costermans *et al.* [19]. Follicles were classified as atretic when they had > 5% positively stained granulosa cells.

Granulosa cell transcriptomics. A subset of 8 out of 29 sows was used for whole-genome transcriptome analysis of granulosa cells, of which 4 were classified as low-COC-quality ($54 \pm 12\%$ high-quality COCs of the largest 15 follicles) and 4 were classified as high-COC-quality ($93 \pm 15\%$ high-quality COCs), based on left ovary classification. The right ovary was used for transcriptome analysis as it is essential for this analysis to obtain high quality intact RNA, for which the ovaries needed to be immediately frozen.

The subset of sows which was used for transcriptome analysis had a similar average follicle size $(5.1 \pm 0.1 \text{ mm vs}, 5.0 \pm 0.1 \text{ mm in})$ low-COC-quality and high-COC-quality, respectively). In short, for each of the 8 sows, granulosa cells of 6 individual healthy follicles of the pool of 10 largest healthy follicles were isolated using laser-capture microdissection and individually hybridized onto 4×44 K Agilent whole-porcine genome arrays (G2519F-026440, Agilent Technologies Inc., Santa Clara, CA, USA) as described in Costermans *et al.* [23]. Microarray data have been deposited in NCBI Gene Expression Omnibus (GEO) under accession number GSE125189. Fold-change is expressed as the ratio of the

normalized expression of sows with low-COC-quality (<70% high-quality COCs) over sows with high-COC-quality ($\geq70\%$ high-quality COCs).

Aromatase immunostaining. The right ovary of the sows was used for quantification of aromatase protein abundance. A subset of 12 out of 29 sows was selected for analysis, 6 sows with low-COCquality $(37 \pm 19\%)$ and 6 sows with high-COC-quality $(98 \pm 5\%)$, based on left ovary classification. For each of the 12 sows, both the largest and smallest follicles of the 10 largest healthy follicles were analyzed. Cryosections were mounted on Superfrost plus glass slides (Menzel- Gläser, Braunschweig, Germany), air-dried for 30 min, and fixed in 4% PAF for 10 min. Slides were subsequently washed in H₂O, microwaved in sub-boiling 0.1 M sodium citrate buffer (pH 6) for 10 min for epitope antigen retrieval, cooled down to room temperature, and next rinsed with PBS pH 7.4. Aldehyde residues were blocked with 0.3% glycine in PBS for 10 min. After rinsing with PBS, sections were pre-incubated with 5% (wt/v) normal goat serum in PBS for 60 min at room temperature. Subsequently, the sections were incubated overnight at 4°C in a humid chamber with anti-aromatase primary antibody (ab18995, lotnr:GR3231482-1, Abcam, Cambridge, UK) diluted 1:50 in PBS-BSA-c (Aurion, Wageningen, The Netherlands). Next, sections were rinsed with PBS and incubated in the dark with a secondary Alexa fluor 488 labeled goat-antirabbit antibody (A-11008, ThermoFisher Scientific, Waltham, MA, USA) diluted 1:200 (v/v) in PBS-BSA-c for 1 h at room temperature. Sections were counterstained with DAPI (1 µg/ml; Sigma-Aldrich, Saint Louis, MO, USA) for 10 min. Sections were imaged at 10 times magnification using a fluorescence microscope (Leica DM6B), a digital camera (DFC365 FX), and imaging software (LasX; all Leica Microsystems, Amsterdam, The Netherlands). Mean staining intensity was determined using ImageI for the complete intact granulosa cell layer. Those parts of the granulosa layer that were visually not intact were excluded from the analysis.

IGF1 measurement. Plasma insulin-like growth factor 1 (IGF1) content was measured by an immunoradiometric assay according to the manufacturer's protocol (A15729, Beckman Coulter, Woerden, The Netherlands) supplemented with an additional acid-ethanol extraction (87.5% v/v EtOH and 2.9 v/v 12 N HCl). Samples were measured in duplicate and intra-assay coefficient of variation was $\leq 6\%$ for all samples.

Statistical analyses. Distributions of the means and residuals were examined to verify model assumptions of normality and homogeneity of variance. Concentrations of β -Estradiol, progesterone, DHEA, α -testosterone, β -testosterone, 11-deoxycorticosterone, 11-deoxycortisol, cortisone, cortisol, DHT, 17 α -OH-progesterone, and corticosterone were log transformed to obtain normality.

The presence of outliers was tested by calculating the studentized residuals using the REG procedure (SAS 9.4, Cary, NC, USA); identified outliers were removed from further analyses. One outlier was removed for β -estradiol, progesterone, dehydroepiandrostenedion (DHEA), α -testosterone, 11-deoxycortisol, 4-androsten-3,17-dione, and 17α -OH-progesterone. Differences between follicle size (FS) classes (average follicle size of 15 largest follicles: small-FS < 5 mm (N = 14) and large-FS ≥ 5 mm (N = 15)) and COC quality classes (percentage high-quality COCs of the 15 largest follicles: low-COC-quality<70% (N = 14) and high-COC-quality \geq 70% (N = 15)) were analyzed in the same model using the procedure GLM in

models that also contained the factor PAR (PAR3 (parity 3, N = 14) and PAR4 + 5 (parity 4 and 5, N = 15)) and the interactions with PAR. The interactions were excluded from the models when not significant. All values are presented as LS means \pm SE unless otherwise stated.

For microarray analysis, differences in transcript abundance were tested with models which included sow COC quality class (high-COC-quality or low-COC-quality) with sow as repeated subject, using the GENMOD procedure in SAS 9.4 (Cary, NC, USA). A false discovery rate (FDR) adjustment for multiple testing correction was done, according to Benjamini-Hochberg.

Fold-change (FC) is defined as the normalized expression of high-COC-quality over low-COC-quality. All transcripts with a FDR adjusted $P \le 0.2$ and absolute FC ≥ 1.2 were considered significantly different. Analysis of differences in aromatase staining intensity was performed using a 2-way ANOVA (Graphpad Prism version 5.04, GraphPad Software, La Jolla, CA, USA) using sow COC quality class (high-COC-quality (N = 4) and low-COC-quality (N = 4) and follicle size class (large or small) as factors. The interaction between sow COC quality class and follicle size class was not significant.

Results

High-COC-quality sows had lost less body weight on a percentage basis during lactation and had a higher IGF1 plasma concentration at weaning as compared to low-COC-quality sows, but did not differ in follicle size or volume (see Table 1). In follicles of high-COC-quality sows, follicular fluid β -estradiol, progesterone, 19-norandrostenedione, and α -testosterone concentrations were higher as compared to low-COC-quality sows, while cortisol concentration was lower (Figure 2). Concentrations of other steroids did not differ (Supplemental Table S1).

In large-FS sows, average follicle size of the 15 largest follicles were larger as compared to small-FS sows (P < 0.001). No significant effects of follicle size were found in any of the lactational or metabolic measurements (see Table 1). Large-FS sows had a higher concentration of β -estradiol in follicular fluid (5.63 \pm 0.60 vs. 3.51 \pm 0.64 ng/ml, P = 0.03) compared to small-FS sows, but did not differ in any of the other measured steroid concentrations (Figure 3).

Subsequently, whole-genome granulosa cell transcriptomes of healthy antral follicles of the right ovary were assessed in a subset of 8 sows of which 4 were classified as high-COC-quality and 4 were classified as low-COC-quality. The analysis showed a total of 337 unique transcripts, which were differentially expressed. Transcript analysis showed an increased abundance of transcripts involved in ovarian steroidogenesis in sows with high-COC-quality, e.g., CYP19A2 and CYP19A3 (FC = 1.46 and FDR = 0.09 and FC = 1.43 and FDR = 0.03, respectively) and P450 oxidoreductase (POR; FC = 1.22, FDR = 0.03) and vascular endothelial growth factor A (VEGFA; FC = 1.22, FDR = 0.15) (see Table 2 for details). Next to genes involved in steroidogenesis, the top 10 regulated transcripts with a higher abundance in high-COC-quality sows included, e.g., inhibin subunit beta B (INHBB; FC = 1.64 and FDR = 0.05) and metallothionein 1A (MT1A; FC = 1.52 and FDR = 0.07) while the top 10 transcripts with a lower abundance in high-COC-quality sows include, e.g., growth arrest and DNA damage inducible alpha (GADD45A; FC = -1.63 and FDR = 0.08) (for details see Supplemental Table S2). In addition, granulosa cells of antral follicles in high-COC-quality sows had higher IGF1 transcript

	Follicle size		COC quality		P-values	
	Small	Large	Low	High	FS	COC
Lactation						
Number of piglets weaned	12.6 ± 0.3	12.7 ± 0.2	12.4 ± 0.3	12.9 ± 0.2	0.99	0.12
Litter growth (kg)	71 ± 3	73 ± 2	72 ± 3	72 ± 2	0.47	0.99
Body condition						
Weight parturition (kg)	254 ± 6	243 ± 5	253 ± 6	242 ± 5	0.12	0.14
Weight weaning (kg) ^a	225 ± 4	220 ± 4	222 ± 4	224 ± 4	0.53	0.82
Weight loss lactation (%)	11 ± 2	10 ± 2	12 ± 2	7 ± 2	0.92	0.03
IGF-1 (ng/ml)	150 ± 17	138 ± 15	120 ± 16	167 ± 15	0.62	0.05
Follicular parameters						
Av. follicle size (mm)	4.4 ± 0.1	5.6 ± 0.1	5.0 ± 0.1	5.1 ± 0.1	< 0.001	0.84
Amount of follicular fluid (µl)	334 ± 40	405 ± 38	363 ± 40	376 ± 38	0.22	0.82
High-quality COCs (%)	67 ± 4	70 ± 4	49 ± 4	89 ± 4	0.67	< 0.001

Table 1. Effects of follicle size (av. follicle size of 15 largest follicles: Small-FS < 5 mm, N = 14 and Large-FS > 5 mm, N = 15) and COC quality class (percentage high-quality COCs: low-COC-quality<70%, N = 14 and high-COC-quality>70%, N = 15) on lactation characteristics, body condition, metabolic parameters, and follicular parameters.

All values are presented as LS means±SE. COC = cumulus-oocyte complex. Follicle size was not related to percentage high-quality COCs.

 $^{a}LS means estimates for the interaction FS*PAR (P < 0.01): Small-FS*PAR3: 228 \pm 8, Small-FS*PAR4 + 5: 259 \pm 6, Large-FS*PAR3: 252 \pm 6, LargeFS*PAR4 + 5: 239 \pm 6.$

abundance (FC = 1.21 and FDR = 0.02). Pigs express multiple homologous (>99%) isoforms of aromatase, CYP19A1, 2 and 3, which are mainly expressed in ovary, placenta, and embryonic tissues, respectively [24]. As *CYP19A2* and *CYP19A3* were regulated, we additionally analyzed aromatase (CYP19A1–3) protein abundance. Sows with high-COC-quality had higher aromatase protein abundance as compared to sows with low-COC-quality (P < 0.01). Aromatase protein abundance did not differ between the large and small healthy antral follicles (P = 0.27) (Figure 4, Supplemental Figure S1).

Discussion

The aim of this study was to identify potential markers for antral follicle quality. We investigated relations between follicular fluid steroid concentrations, follicular development, and COC morphology in multiparous sows at the onset of the follicular phase of the oestrus cycle. To our knowledge, this is the first study in which porcine follicular fluid steroid profiling is reported, let alone in follicles at a specific stage of the estrous cycle.

Sows at the start of the follicular phase of the oestrus cycle with a large vs. small average follicle size had higher follicular fluid β estradiol concentrations. In pigs, granulosa cell CYP19 abundance and subsequent β -estradiol production increases from the start of the follicular phase until the pre-ovulatory stage approximately 5 days later [25], and consequently as follicles develop, follicular fluid β estradiol concentration increases with follicular diameter. In most studies, effects of follicle size on steroid concentrations were either not investigated in ovaries which were in the same stage of the oestrus cycle [26-28] or were only investigated at the final stages of follicular development, from pro-oestrus to late oestrus [29]. The results of our study are the first to indicate that from the pool of antral follicles, the larger subset of follicles has an increased β -estradiol concentration in follicular fluid already at the start of the follicular phase. Having said this, follicle size is not related to any of the other measured follicular fluid steroids. Follicular progesterone production is highest in the early luteal stage and only starts to increase in the latest stages of the follicular phase [27, 30], which can explain why we did not find a relation between follicular size and follicular fluid progesterone concentration. Although high levels of testosterone and androstenedione are present during the entire follicular phase [26, 30], these and rogens can also be quickly aromatized into β estradiol, which might explain why we did not find a relation between follicle size and testosterone and androstenedione steroid concentrations. Next, we compared the follicular fluid steroid profile of sows with high vs. low percentage of high-quality COCs. COC morphology was used as a marker for oocyte quality as COCs with an intact cumulus layer and normal shaped oocytes have the highest developmental capacity in vitro [20]. Sows with a high percentage high-quality COCs had a higher concentration of FF β estradiol, progesterone, 19-norandrostenedione, and α -testosterone as compared to sows with a low percentage of high-quality COCs at the start of the follicular phase. Similarly, studies in pigs have shown that in early atretic antral follicles, transcript abundance of folliclestimulating hormone receptor (FSHR), luteinizing hormone receptor (LHCGR), and CYP19 is lower [31, 32], resulting in lower FSHinduced estradiol production [33, 34] and lower in vitro cumulus cell progesterone production [35]. It therefore seems reasonable to assume that the follicle pool of 15 largest follicles from which the COCs were isolated and where we did not discriminate between healthy and atretic follicles contained a higher fraction of atretic follicles in sows with a low percentage high-quality COCs. Besides a higher concentration of follicular fluid β -estradiol, progesterone, 19-norandrostenedione, and α -testosterone, we found a lower concentration of cortisol in sows with a higher percentage of highquality COCs. As cortisol is an important regulator of stress-related apoptosis of granulosa cells and oocytes via the FAS system [36], differences in FF cortisol concentration can at least partly explain the observed differences in COC quality.

To better understand differences in COC quality and FF fluid steroid content in sows, we performed whole-genome transcriptome analysis in granulosa cells. We used a subset of eight sows to analyze mural granulosa cell transcriptomes in healthy follicles only. This transcriptome analysis revealed an increased abundance of two transcripts (*CYP19A2* and *CYP19A3*) encoding for cytochrome P450 aromatase, the granulosa cell terminal enzyme in the estrogen biosynthetic pathway responsible for the conversion of androgens into estradiol [29]. Immunostaining against aromatase (CYP19A1–

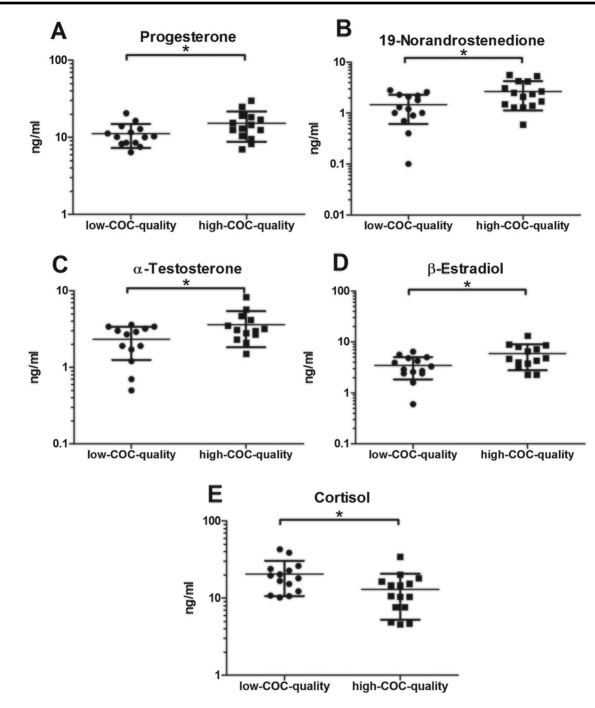


Figure 2. Follicular fluid concentrations of progesterone (A), 19-norandrostenedione (B), α -testosterone (C), β -estradiol (D), and cortisol (E) (ng/ml) of the 15 largest follicles of the left ovary of sows with high-COC-quality (\geq 70% high-quality COCs, N = 14) vs. sows with low-COC-quality (<70% high-quality COCs, N = 15). **P*-value < 0.05.

3) confirms the increased aromatase abundance in granulosa cells of sows with a high percentage of high-quality COCs at the protein level. This increased granulosa cell aromatase transcript and protein abundance may explain the higher FF β -estradiol concentration in sows with high COC quality. In addition, although not differentially regulated, our granulosa cell transcriptome analysis showed that granulosa cells also express other steroidogenic enzymes already at the start of the follicular phase. These include steroidogenic acute regulatory protein (*STAR*), regulating cholesterol transfer within the mitochondria [37], cytochrome P450 family 11 subfamily A member 1 (*CYP11A1*), and hydroxysteroid 17- β dehydrogenase 1 (*HSD17B1*), involved in the conversion of cholesterol into androgens [38] (data not shown). Similarly, in a study by Garmey *et al.* [39], granulosa cells of cyclic gilts had a low transcript abundance of CYP11A and STAR mRNA and protein from approximately day 3 or 5, respectively, of the follicular phase. Apparently, porcine follicular development differs from that of rats and mice, as granulosa cells in the latter species only start to express CYP11A and

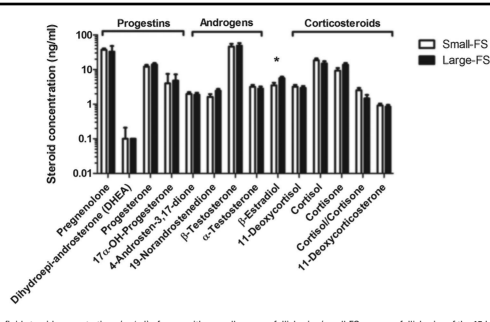


Figure 3. Follicular fluid steroid concentrations (ng/ml) of sows with a small average follicle size (small-FS: average follicle size of the 15 largest follicles of the right ovary \leq 5 mm, N = 14) vs. sows with a large follicle size (large-FS: average follicle size \geq 5 mm, N = 15). **P*-value < 0.05.

Table 2. Significantly different genes (absolute $FC \ge 1.2$, $FDR \le 0.2$) involved in ovarian steroidogenesis (ref: Ovarian Kaleidoscope Database)
between sows with high-COC-quality (93 \pm 15% high-quality COCs (N = 4)) vs. sows with low-COC-quality (54 \pm 12% high-quality COCs
(N = 4))

Gene symbol	Gene description	Accession no	FC 1.46	FDR 0.09
CYP19A2	Cytochrome P450 19A2	NM_214430		
CYP19A3	Cytochrome P450 19A3	NM_214431	1.43	0.03
CYP51	Cytochrome P450, family 51, subfamily A, polypeptide 1	XM_005667583	1.28	0.05
ENSSSCT00000010858	TP53 regulated inhibitor of apoptosis 1	-	1.25	< 0.01
HSD17B7	Hydroxysteroid 17-beta dehydrogenase 7	NM_001185137	1.25	0.05
HSD17B4	Hydroxysteroid 17-beta dehydrogenase 4	XM_021081514	1.24	0.13
POR	P450 oxidoreductase	NM_001129959	1.22	0.03
VEGFA	Vascular endothelial growth factor A	NM_214084	1.22	0.15
EPHX1	Epoxide hydrolase 1	NM_214355	1.21	< 0.01
IHH	Indian hedgehog signaling molecule	NM_001244470	-1.99	0.10
PTCH1	Patched 1	XM_013980137	-1.46	0.09
PPARG	Peroxisome proliferator activated receptor gamma	NM_214084	-1.41	< 0.01
LGALS1	Galectin 1	NM_001001867	-1.24	0.08
GHR	Growth hormone receptor	NM_214254	-1.24	0.20
KLF13	Kruppel-like factor 13	NM_001011505	-1.21	0.04

For each gene, the fold-change (FC) of high-COC-quality sows vs. low-COC-quality sows and significance (FDR) are shown.

STAR in pre-ovulatory follicles, so at a much later stage of follicular development [38, 40]. As *CYP11A1*, *STAR*, and *HSD17B1* were expressed in granulosa cells, they might contribute to follicular androgen production in sows, already at the start of the follicular phase. However, these steroidogenic enzyme transcripts were not differentially regulated in granulosa cells of sows with high vs. low percentage high-quality COCs. Therefore, it cannot explain the higher progesterone, 19-norandrostenedione, and α -testosterone concentrations in FF of sows with high COC quality. Possibly, these steroidogenic enzymes might be differentially regulated at the protein level or have a different enzymatic activity. Thus, although granulosa cells contribute to follicular androgen production in sows, they are at least at this stage of follicular development not responsible for dif-

ferentially regulating androgen concentrations in FF. This confirms the existence of the classical two-cell, two-gonadotropin concept for estrogen biosynthesis in the porcine ovary, in which theca cells play the major role in androgen production and its regulation [41].

Besides genes encoding enzymes participating in steroid metabolism, sows with a high vs. low percentage high-quality COCs had a differential abundance of transcripts, which are involved in regulation of enzymes involved in ovarian steroidogenesis. Examples of these differentially expressed transcripts are *POR* and *VEGFA*, which had a higher abundance in sows with a high percentage high-quality COCS. POR supplies electrons to all CYP enzymes for catalytic activity and is essential for steroid biosynthesis [42]. VEGFA is a growth factor which regulates angiogenesis in the ovary

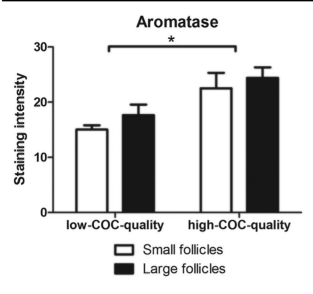


Figure 4. Aromatase protein abundance in granulosa cells of healthy follicles of sows with high-COC-quality (98 \pm 5% high-quality COCs (*N* = 6)) vs. low-COC-quality (37 \pm 19% high-quality COCs (*N* = 6)) in the smallest and largest follicle of the 10 largest healthy follicles. Data are presented as average \pm SD. **P*-value < 0.05.

[43] and was shown to increase CYP19A1 granulosa cell transcript abundance after addition to granulosa cell culture medium [44]. Therefore, higher POR and VEGFA might be involved in increasing ovarian steroid biosynthesis at the start of the follicular phase in sows with a high percentage high-quality COCs. Next to genes involved in ovarian steroidogenesis, we also found differential abundance of transcripts regulating ovarian follicular atresia, e.g., INHBB and GADD45A. GADD45A abundance was lower in sows with a high percentage high-quality COCs. GADD45A is involved in the DNA damage response and has a higher abundance in porcine atretic follicles [45]. GADD45A might provide a valuable marker for early atretic follicles, as it was already expressed in the antral follicles of our study which have been classified as healthy, as they did not (yet) express cleaved-caspase 3. Granulosa cell INHBB transcript abundance was higher in sows with a high percentage high-quality COCs. INHBB encodes for the B subunit of activin and inhibin complexes. Follicular atresia has been previously associated with less INHBB transcript abundance in porcine granulosa cells [45, 46].

Other genes that had a higher transcript abundance in granulosa cells of sows with a high percentage high-quality COCs encode for proteins that are protective against oxidative stress e.g., MT1A, GPX3, PRDX3, PRDX6, and SOD2. These proteins might play a role in protecting the ovarian tissues from oxidative stress [47–50]. High levels of reactive oxygen species are produced during steroid biosynthesis, which results in a higher abundance of superoxide dismutase, catalase, and glutathione peroxidase in steroid secreting areas of the ovary [51]. In our study, follicles of sows with a high percentage high-quality COCs produce more steroids, as seen by the higher concentrations of FF steroids and abundance of transcripts involved in steroidogenesis, which might explain the increased abundance of transcripts involved in the oxidative stress response.

Follicle size and health status and COC quality can be affected by environmental factors, such as the nutritional status. In the current study, ovaries were obtained within 2 h after weaning. Therefore, we analyzed the lactational metabolic state of the sows as well, as this might underlie differences in follicular development

[52]. Sows with a higher percentage high-quality COCs at the day of weaning lost a lower percentage of their body weight during lactation and had a higher plasma IGF1 concentration at weaning as compared to sows with a low percentage high-quality COCs. In addition, granulosa cells of sows with a high percentage high-quality COCs express more IGF1. Peripheral IGF1 indirectly influences follicular development by binding to insulin and IGF1 receptors in the hypothalamus and pituitary and therefore stimulates the production of the gonadotropins FSH and LH [53]. In addition, both peripheral IGF1 and locally produced IGF1 are essential for follicular development, as IGF1 can activate IGF1 signaling in granulosa cells and synergize with FSH to stimulate follicular growth and steroidogenesis [54]. This effect is mediated by activation of phosphatidylinositol-dependent kinase/AKT, as shown for bovine granulosa cells in vitro [55]. In human cumulus cells, effects of FSH on cumulus cell transcript abundance of genes involved in cholesterol and steroid metabolism are minimal when IGF1R is blocked [56]. This provides further evidence for the essential role of IGF1 in regulation of steroid production. Next to this, both FSH and IGF1 have been identified as anti-apoptotic factors [57]. Therefore, the relatively low IGF1 concentration in plasma of sows with a low percentage of high-quality COCs might be insufficient to support follicular development and steroidogenesis resulting in an increased risk of follicular atresia. Indeed, the metabolic state of the sows during lactation seems to influence follicular development and FF content.

In conclusion, we found that a positive relation between follicle size and follicular fluid β -estradiol concentration already exists at the start of the follicular phase, as sows with larger follicles had a higher FF β -estradiol concentration. Follicles of sows with a high percentage high-quality COCs contained more β -estradiol, progesterone, 19-noradrenostenedione, and α -testosterone and less cortisol, which could therefore serve as potential markers to identify early atretic follicles with decreased follicular developmental competence.

By comparing transcriptomes of healthy follicles of sows with a high vs. low percentage high-quality COCs, we identified several genes involved in granulosa cell steroidogenesis (e.g., POR and VEGFA) and genes involved in regulating granulosa cell apoptosis (e.g., *GADD45A* and *INHBB*), which could serve as markers for follicle quality.

Supplementary data

Supplementary data is available at BIOLRE online.

Conflict of interest

The authors declare that there are no conflicts of interest that could be perceived as precluding the impartiality of the results reported.

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