The consequences of porcine IVM medium supplementation with follicular fluid become reflected in embryo quality, yield and gene expression patterns

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Oocyte and embryo developmental competence are shaped by multiple extrinsic and intrinsic factors. One of the most extensive research areas in the last decade is the regulation of lipid metabolism in oocytes and embryos of different species. We hypothesized that differences in developmental competence of oocytes and embryos between prepubertal and cyclic gilts may arise due to distinct fatty acid profiles in follicular fluid. We found that supplementation of oocyte maturation media with follicular fluid from prepubertal pigs affected quality and development of embryos from prepubertal pigs while embryos of cyclic pigs were not affected. PLIN2, SCD and ACACA transcripts involved in lipid metabolism were upregulated in embryos originating from oocytes of prepubertal pigs matured with autologous follicular fluid. The surface occupied by lipid droplets tend to increase in oocytes matured with follicular fluid from prepubertal pigs regardless oocyte origin. The change into follicular fluid of cyclic pigs increased the efficiency of embryo culture and improved quality, while gene expression was similar to embryos obtained from cyclic gilts. We assume that the follicular fluids of prepubertal and cyclic pigs influenced the quality of oocytes and embryos obtained from prepubertal pigs which are more susceptible to suboptimal in vitro culture conditions.

Lipid metabolism is considered as one of the major factors responsible for shaping the developmental competence of oocytes and embryos. Porcine oocytes containing exceptionally high levels of intracellular lipids may serve as a model to study lipid metabolism during early embryonic development. Follicular fluid (FF), providing the natural environment for oocyte growth, serves as a supplemental source of fatty acids and growth factors to porcine oocyte in vitro maturation media (IVM)1–3. Qualitative and quantitative composition of FF may be directly translated into oocyte growth and maturation since it is influenced by animal age, nutrition and size of the follicle from which it was collected2,3. Moreover, the essential role of FF for oocyte nuclear maturation, cumulus cells expansion, apoptosis, polyspermy prevention and embryo culture efficiency has been documented2,4,5. A study of Grupen et al. showed the significance of FF origin for IVM of oocytes from prepubertal and adult gilts6. Earlier studies showed that the developmental competence of oocytes obtained from prepubertal gilts increase with the diameter of follicles from which they were collected7–12. This property however, has not been noted in oocytes collected from adult sows which show high competence despite the follicle origin. The search for the causes of reduced developmental competence of oocytes obtained from prepubertal animals focused on different mammalian species, however without satisfactory results. A common observation was the reduced blastocyst yield, but the causes still remain mostly unknown. The efficiency of nuclear maturation in vitro did not provide the unequivocal answer, while the hypothesis on the insufficient cytoplasmic maturation is valid13–15. However, the influence of environmental factors on oocyte and preimplantation embryo quality have...
Higher concentration of estradiol was found in FF of cyclic pigs (21.5 + 0.6). However, no differences in the ratio of P4 to E2 were significant between groups (3.9 + 0.64). The number of lipid droplets in oocytes of cyclic gilts increased significantly irrespective of the supplemented FF (P < 0.05). The FF collected either from P or from C pigs showed no adverse effects on embryos from cyclic (polyunsaturated fatty acids), linoleic and arachidonic fatty acids (P > 0.05). Total fatty acid concentration was noted in FF of prepubertal pigs (P < 0.01). Previous work on prepubertal (P) and cyclic (C) gilts has led to the discovery of differences in the amount of fatty acids (FA) in the follicular fluid. The majority of papers describing the composition of FF comes from studies done on cattle (impaired reproduction under the influence of negative energy balance) and humans (searching for markers of oocyte quality allowing to predict embryonic developmental potential). In both species, however, FF is not used for IVM therefore, the interpretation is limited to the effect of individual follicles on oocytes derived from them. The advantage of pig model, allows more experimental approach in elucidating how FF and its components influence full oocyte maturation and developmental competence. So far it has been shown that some FAs may negatively or positively affect the metabolism and the development of the embryo in vitro. It is also known that the fate of the embryonic development does not solely rely on single molecules, but on their relative proportions, in a way that one FA can reduce or neutralize the inhibiting effects of another. Therefore, our aim was to analyse the influence of FF collected from prepubertal and cyclic pigs (which differ in FA content) on lipid droplet (LD) number and on the area occupied by the droplets in relation to the expression of genes responsible for FAs metabolism in oocytes and embryos. Determining the role of FAs in FF will lead to better understanding of the lipid metabolism in ovarian follicle and potentially reveal new markers for prediction of oocyte developmental competence and embryo quality in IVF programs.

**Results**

**In vitro embryo production (IVP).** The analysis of the effectiveness of embryo culture to the blastocyst stage showed an interesting correlation between applied FF and its effect on embryos. The least likelihood of embryos reaching the blastocysts stage was observed in experiments where oocytes collected from prepubertal (P) females were matured in IVM medium supplemented with the FF collected from the same experimental group (20.6%). However, if the same experimental group was supplemented with FF collected from cyclic (C) females a significant improvement in embryo culture efficiency was noticed (27.8%, P < 0.05; Table 1). Interestingly, no effect of FF was observed on cyclic embryos, and in both cases, the culture efficiency was high (C FF P = 29.6%, C FF C = 29.5%). It seems most likely that the FF collected from prepubertal animals showed no adverse effects on embryos, but FF from cyclic pigs improved conditions for oocyte maturation and improved the blastocyst yield.

The FF used for IVM influenced also the quality of blastocyst stage embryos revealed by the blastomere number. The lowest number of cells was observed in embryos obtained from prepubertal pigs after supplementation with the FF collected from the same group (29), but the change into FF of cyclic pigs resulted in a significant increase in the observed mean cell number (37, P < 0.05) comparable to embryos of cyclic gilts (37 cells – C FF C; 44 cells - C FF P). The FF collected either from P or from C pigs showed no adverse effects on embryos from cyclic females neither on the quality nor on the efficiency of embryo culture.

**Fatty acid profile and hormones concentration in follicular fluid.** Fatty acid profile in P and C pigs differed in terms of total FA content, as well as palmitic, oleic, linoleic and arachidonic acids (Fig. 1). The highest total FA concentration was noted in FF of prepubertal pigs (P < 0.05) together with higher content of palmitic and oleic acids (P < 0.05). On the other hand, FF from cyclic pigs was characterized with increased content of PUFA (polyunsaturated fatty acids), linoleic and arachidonic fatty acids (P < 0.05, Fig. 2).

Progesteron and prostaglandin 2A (PGE2) concentration did not differ between FF collected from prepubertal and cyclic gilts (35.6 +/- 8.1 vs 39.5 +/- 9.6) and (19.2 +/- 14.4 vs 17.8 +/- 13.9) respectively. A significantly higher concentration of estradiol was found in FF of cyclic pigs (21.5 +/- 13.3 vs 9.1 +/- 5.9, P < 0.01). The differences in ratio of P4 to E2 was not significant between groups (3.9 +/- 1.33 vs 2.9 +/- 0.64).

**Lipid droplets.** Statistical analysis (including a total of 268 oocytes, P - 123, C - 145) showed a significantly greater number of LD in oocytes of prepubertal gilts before IVM (P < 0.05). After in vitro maturation, the number of lipid droplets in oocytes of cyclic gilts increased significantly irrespective of the supplemented FF (P < 0.05). The number of lipid droplets in P oocytes did not change significantly after IVM if matured in P follicular fluid, however a significant decrease was noticed after the use of cyclic FF (Fig. 3A; P < 0.05). After IVM no significant differences has been noticed between experimental groups.

The results describing the area occupied by the LD showed lower variability than mean LD number. The oocyte area shared by lipid droplets was lowest before IVM and no correlation with gilt category was noticed (Table 2). After maturation, the area occupied by LD increased in all groups with noticeable trends (Fig. 3B).

| Group | Number Of Activated Oocytes | % Blastocyst/ Oocytes | Mean Cell Number | Min/Max Cell Number |
|-------|-----------------------------|-----------------------|-----------------|---------------------|
| P FF P | 315                         | 20.6*                 | 29,1*           | 18/51               |
| C FF P | 446                         | 29,6*                 | 44,6*           | 19/86               |
| P FF C | 312                         | 27,8*                 | 37,3*           | 25/50               |
| C FF C | 348                         | 29,7*                 | 37,0*           | 22/53               |
| TOTAL  | 1421                        | 26,9                  | 37,4            | 18/86               |

Table 1. Total number of oocytes activated, the efficiency of blastocyst stage embryo production and quality of blastocysts as revealed by the cell number. Different letters indicate statistically significant differences between groups (P < 0.05).
Maturation of oocytes in FF collected form prepubertal gilts resulted in increased but not statistically significant lipid droplet area in both P and C oocytes compared to cyclic FF (Table 2).

Gene expression (mRNA level). Gene expression on mRNA level was analysed for 26 oocyte samples (5 biological replicates before IVM in each group and 4 replicates after IVM). No significant differences in the
expression level of genes related to developmental competence and quality (BMP15, GDF9, OCT4) were noted between groups before IVM and after IVM (Supplementary Fig. 3). Similarly, the analysis of genes related to fatty acid metabolism (ACACA, SCD, FADS2, FADS1, FASN, PLIN2, ELOVL2, ELOVL6) showed no statistically significant differences in mRNA expression level between oocytes of P and C pigs after IVM (P > 0.05, Fig. 4). No change in the gene expression was noticed independently of the fluid used for IVM.

The differences resulting from distinct IVM conditions were evident at the blastocyst stage of embryonic development. The PLIN2, SCD, ACACA gene expression was significantly increased (P < 0.05; Fig. 4) in blastocysts of prepubertal pigs originating from oocytes matured in FF of the same group compared to the other three groups. Supplementation of IVM with FF of cyclic pigs for maturation of oocytes from prepubertal pigs resulted in gene expression similar to embryos of cyclic pigs, which expressed above genes at comparable levels regardless the FF origin. Supplementation of IVM with FF from prepubertal pigs had no significant effect on ELOVL2, ELOVL6, FASN, FADS1 and FADS2 gene expression in embryos. The OCT4 gene expression level did not differ between blastocysts of different groups (Supplementary Fig. 4).

**Immunofluorescent staining.** For the protein products of the several investigated genes we have observed distinct distribution patterns within oocytes and embryos. ACACA was found always on the surface of the lipid droplets while SCD, FADS1 colocalized with whole LDs with different intensity (Fig. 5). FASN foci were found closely localized next to some population of lipid droplets. No differences between oocytes before and after in vitro maturation have been noticed in distribution of analysed proteins. The same patterns like colocalization with lipid droplets or LD coating by these proteins were noticed in blastocyst stage embryos.

**Apoptosis.** In all 47 examined embryos and 1944 blastomeres only few showed apoptotic signals. The frequency of apoptotic cells within parthenogenetic embryos was estimated on as few as 0.007% (Table 3). No correlation between oocyte origin and the frequency of apoptotic blastomeres was noticed.

**Discussion**

The developmental potential of oocytes is shaped in a complex manner and controlled by multiple factors. In recent years, the metabolism of fatty acids and lipids in oocytes has been the subject of meticulous studies carried out in various mammalian species44,55. Our previous studies have indicated the possible link between the fatty acid content of the FF and the quality of oocytes from prepubertal and cyclic gilt. The pig is the only species in which the FF is added to the IVM media therefore, its impact is prolonged and the qualitative and quantitative FF acid content directly translates into the developmental potential of the oocytes.

**Table 2.** Mean lipid droplet number and occupied area within the oocytes of each experimental group. Different letters indicate statistically significant differences between six experimental groups and between pooled PRE-IVM and POST-IVM groups (P < 0.05).

| GROUP | Number Of Analyzed Oocytes | Mean Lipid Droplet Number (Sd) | Min/Max | Area Occupied By Lipid Droplets (%) |
|-------|---------------------------|--------------------------------|---------|------------------------------------|
| C     | 48                        | 450 (159)†                     | 196/1088| 9.3†                               |
| C-FF C| 40                        | 577 (162)†                     | 348/1104| 11.9†                              |
| C-FF P| 35                        | 533 (195)†                     | 246/948 | 13.2†                              |
| P     | 59                        | 572 (253)†                     | 194/1178| 8.1†                               |
| P-FF C| 39                        | 462 (140)†                     | 176/756 | 11.8†                              |
| P-FF P| 47                        | 541 (200)†                     | 224/1074| 12.9†                              |
| TOTAL | 268                       | 522 (198)                      | 176/1178| 10.6                               |
| PRE-IVM| 107                       | 511 (223)                      | 194/1178| 8.6†                               |
| POST-IVM| 161                       | 528 (180)                      | 176/1104| 12.2†                              |

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after IVM which may however arise due to the lack of FF supplementation, and in consequence to the depletion of stored lipids during extensive IVM.\textsuperscript{46} Taken together, the differences in the number of LDs in cyclic oocytes do not correspond to the developmental potential and may not be a marker of their quality.

Unlike Milakovic et al., we noticed the great heterogeneity of LD number in all the groups of oocytes we examined.\textsuperscript{46} Limiting the analysis to the LD number alone thus might not be very informative. Microscopic analysis also revealed a large variation in the LD size in oocytes. We therefore decided to look at this trait and the possible correlation between LD number and size, since they may change in size or shape, grow de novo or fuse and redistribute through the cell.\textsuperscript{27,31,37,47} We assumed that such analysis might more precisely describe the effect of particular FF or maturation conditions on oocyte characteristics. Our results clearly show that the LD area

Figure 4. mRNA expression level of genes responsible for fatty acids metabolism in oocytes (pre-IVM), oocytes matured \textit{in vitro} (post-IVM) and blastocyst stage embryos. Different letters indicate significant differences between groups within each analysed gene (P < 0.05). P – oocytes from prepubertal gilts; C – oocytes from cyclic gilts; P FF P – oocytes collected from prepubertal females matured with autologous, prepubertal follicular fluid; P FF C – oocytes collected from prepubertal females matured with follicular fluid from cyclic gilts; C FF P – oocytes collected from cyclic females matured with follicular fluid from prepubertal gilts; C FF C – oocytes collected from cyclic females matured with autologous, cyclic follicular fluid.
Figure 5. Immunofluorescence staining of oocytes for ACACA, FASN, FADS1, SCD and TUBULIN distribution patterns assessment. NC – negative control (without primary antibodies). BF – brightfield, DAPI – chromatin staining and merged channels. All stacks present (excluding marked GV tubulin staining) oocytes at MII stage of meiosis (visible and stained polar body). Scale bars represent 50 μm.

Table 3. The incidence of apoptosis in embryos of different experimental groups. Different letters indicate statistically significant differences between groups (P < 0.05).
within post-IVM oocytes of C and P pigs significantly increased, indicating utilization of the FF components. We also observe higher LD areas in oocytes undergoing maturation with FF of prepubertal pigs, however, despite this marked trend, the difference was not significant. The percentage increase in LD area accounted for 1.1% and 1.3% in P and C oocytes, respectively. This indicates the important role of FF and its composition, which differs considerably depending on the follicles from which it is collected. It would thus be interesting to supplement IVM by FF with extreme amounts of FAs. Such approach would allow observation of the lipid accumulation, changes in LD area or the role of cumulus cells in preventing an the excess of FA supply, as has already been demonstrated in cattle. Our results nevertheless show that the presence of FF in the maturation medium increased LD area which means that they are able to incorporate the lipids present in growth environment. This process is thought to be preferred by porcine oocytes and embryos for energy provision until placenta formation.

Considering the recent data of Grupen et al. it is interesting to ask which of the FF components provides optimal conditions for oocyte maturation, and whether such components act alone or synergistically. These authors found that oocytes from adult pigs were more readily recruited for capacitation than those from prepubertal pigs, possibly due to differences in FF steroid content. Our results show that, although prepubertal FF does not seem to inhibit the competence of oocytes from cyclic pigs, it does not promote development of oocytes from prepubertal pigs which was clearly noted when FF from cyclic pigs was used for maturation. This leads to the conclusion that, once reached, full developmental competence cannot be revoked through the subtle physiological conditions provided by FF of prepubertal origin. However, it is possible to shape the quality of oocytes with reduced competence (e.g. prepubertal ones) using the growth environment supporting their maturation (e.g. cyclic FF). The FF of prepubertal pigs may thus be considered a quiet environment that supports the oocyte to the moment of achieving puberty by the donor and suppresses cytoplasmic maturation, unless matured in FF of cyclic pigs where the inhibiting effect is abolished. These data complement similar observations by measuring only steroid content in the FF of prepubertal and adult gilts. We did not see differences in progesterone and PGE2 concentration, but higher concentrations of estradiol in the FF of cyclic pigs was observed. Grupen et al. showed significant differences in progesterone and estradiol concentration in FF from prepubertal and adult pigs. The hormone contents observed by us and by Grupen are similar while the progesterone concentrations could be affected by the estrous cycle, during which it may change twenty-fold or by differences in animal age (adults vs. pubertal, cyclic). Interestingly, we noticed that the concentration of arachidonic acid, a product of PGE2 metabolism, was higher in cyclic pig FF, whereas prostaglandin concentration was not affected by FF origin. Regarding the results of Grupen et al. it should be noted that the pooling of FF might mask the action of some constituents (e.g., growth factors) that might affect the cytoplasmic maturation.

Considering the standard requirement of the porcine oocytes to be matured with FF supplementation it is almost impossible to identify individual FAs, or a combination with a distinct, stimulating or inhibiting effect on the developmental competence of oocytes and embryos. This approach is routinely used in the bovine model and allows further interpretation of the physiological and pathological status of FF. It has been shown that palmitic and stearic acids negatively affect bovine oocytes and embryos. Conversely, oleic and linoleic acids stimulate the developmental competence of oocytes and embryos. This approach is routinely used in the bovine model and almost impossible to identify individual FAs, or a combination with a distinct, stimulating or inhibiting effect on the developing preimplantation embryo. To date numerous articles have addressed the effect of modified IVM conditions on embryo yield and physiology or cell count most often in cattle. In our study, the embryos of prepubertal pigs were developing at lowest rate and exhibited the lowest number of cells when the autologous FF was used for IVM. However, the use of cyclic pig FF significantly increased the yield and quality of embryos to the levels of embryos from cyclic pigs, in which the quality and embryo culture efficiency was not affected by IVM conditions. Additionally, we have examined the occurrence of apoptosis in porcine parthenogenetic blastocysts, which is almost absent in comparison to IVF embryos or PA embryos of other species. The impact of maturation conditions has been found not only in physiological features of developing embryos, but also on the molecular level describing mostly expression of the genes responsible for cell division, developmental competence, transcription, apoptosis and methylation. Only a few concerns genes related to lipid metabolism. Our data show that oocytes from prepubertal and cyclic pigs that underwent IVM with FF of various origins reached the MII stage at the same rate and expressed genes involved in lipid metabolism at similar levels. However, the differences arising from different maturation conditions were significantly manifested at the blastocyst stage. Interestingly, the selected genes controlling lipid metabolism were upregulated in embryos from the prepubertal pigs. Again, the use of FF from prepubertal pigs, with distinct FA profiles altered the expression pattern of the genes. Two genes (ACACA, the fatty acid synthase) are involved in FA synthesis. The starting substrate for ACACA is acetyl-CoA which is transformed into malonyl-CoA subsequently committing to the first step of FA synthesis. The formation of new lipid droplets which stores lipids mostly in the form of triacylglycerols is controlled by ADRP (also known as PLIN2). Palmitic acid, which is the most abundant in FF and oocytes, can be further converted into other FAs by a number of enzymes encoded by several genes (e.g. those encoded by FADS1, FADS2, SCd, ELOVL2 and ELOVL6). The PLIN2, ACACA and SCd genes showed significantly higher expression in PA embryos of prepubertal pigs derived from oocytes matured in autologous FF. Replacing the FF in the same oocyte group by cyclic FF led to the downregulation of gene expression to the level significantly higher expression in PA embryos of prepubertal pigs derived from oocytes matured in autologous FF. The impact of maturation conditions has been found not only in physiological features of developing embryos, but also on the molecular level describing mostly expression of the genes responsible for cell division, developmental competence, transcription, apoptosis and methylation. Only a few concerns genes related to lipid metabolism. 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Immature COCs were aspirated from categorized ovaries with needle and syringe. COCs were placed in HEPES-NEFA (mixtures of palmitic, stearic and oleic acids) were used. Interestingly, microarrays indicate changes of new LD and elevated lipid storage. The majority of data published on the effect of FA supplementation during PLIN2 to the development of the early mammalian embryo. However, increased expression of this gene in PA showing lipid droplet coating distribution. It may thus be the major functionally conserved component of LDs and thus serve as a marker of lipid metabolism in maturing oocytes. Little is known about the contribution of PLIN2 to the development of the early mammalian embryo. However, increased expression of this gene in PA blastocysts of prepubertal pigs may reflect the growth environment during IVM and contribute to the formation of new LD and elevated lipid storage. The majority of data published on the effect of FA supplementation during IVM on gene expression comes from bovine research, where saturated fatty acid (SA) or high concentrations of NEFA (mixtures of palmitic, stearic and oleic acids) were used. Interestingly, microarrays indicate changes in the lipid gene expression pathway, but also changes in cell metabolism involving changes in glucose consumption, resembling the mechanism of insulin resistance in somatic cells. Van Hoeck et al. described increased expression of genes involved in lipid synthesis, FA uptake or cholesterol biosynthesis in bovine IVP blastocysts due to maturation media supplemented with high concentration of NEFA. Our findings agree with the bovine data on the expression of ACACA which is upregulated in embryos of prepubertal pigs originating from oocytes matured with FF of higher FA content. Van Hoeck et al. found increased expression of ACACA and ACSL1 in bovine blastocysts originating from oocytes matured with high NEFA concentrations. ACSL is involved in the activation of long chain fatty acids for B-oxidation, and also the synthesis of triglycerides. Parallel upregulation of these two genes thus suggests stimulated lipogenesis and higher lipid storage in LDs. Considering the metabolic pathway, one may further explain the increased expression of the SCD gene in embryos of prepubertal pigs. This gene plays an important role in regulating the expression of genes involved in lipogenesis, biosynthesis of membrane phospholipids and triglycerides and in oxidation. The question however remains whether the observed alterations in gene expression are related to embryo development under suboptimal IVM conditions dissociate with oocyte competence or to a metabolic consequence during embryo growth.

In conclusion, the consequences of IVM medium supplementation with FF of distinct origin (prepubertal vs cyclic) became apparent at the blastocyst stage and concerned selected traits of embryo quality and blastocyst yield. Oocytes of prepubertal pigs matured in autologous FF developed to blastocyst at lower rate and displayed reduced cell counts and increased expression of SCD, ACACA and PLIN2 genes (involved in fatty acid metabolism). Blastocyst yield and quality were improved by maturation of these oocytes in the FF collected from cyclic pigs. We assume that the distinct profiles of FAs in FF of prepubertal and cyclic gilts influenced the quality of oocytes of prepubertal piglets which are more susceptible to suboptimal in vitro culture conditions. With regard to oocytes of cyclic pigs, their quality was probably not affected by the FF origin due to full-bodied quality reflected by advanced cytoplasmic maturation. Moreover, the differences in FA content without modifying conditions such as diet or chemical intervention during IVM or IVC, are subtle, and do not seem to elicit the mechanism of lipotoxicity prevention. However, further study of the metabolome, secretome and defined media should reveal detailed influence of particular metabolites in the growth environment on mammalian oocytes and embryos.

**Material and Methods**

**Collection of cumulus-oocyte complexes.** The COCs used in experiment were collected post-mortem from porcine ovarian follicles of 3–6 mm diameter. Ovaries were transported in thermostabilized flask 2 hours after slaughter to the laboratory. Animals were 5–6 months old and weighted 100–110 kg. Ovaries were divided into two groups: P (prepubertal) and C (cyclic). P group was characterized by smaller size follicles and absence of corpus luteum, whereas C group COCs have more follicles >3 mm and presence of multiple corpus luteum. Immature COCs were aspirated from categorized ovaries with needle and syringe. COCs were placed in HEPESTM and scored morphologically. COCs had evenly granulated cytoplasm and 4–5 layers of cumulus cells.

**Experimental groups.** All procedures were performed in accordance with the “Act on the protection of animals used for scientific purpose” of the Republic of Poland, which complies with the European Union Legislation for the protection of animals used for scientific purposes. According to these regulations ethics approval was not required, as the biological material (ovaries) was collected upon animal slaughter in abattoir. Oocytes of cyclic pigs, their quality was probably not affected by the FF origin due to full-bodied quality reflected by advanced cytoplasmic maturation. Moreover, the differences in FA content without modifying conditions such as diet or chemical intervention during IVM or IVC, are subtle, and do not seem to elicit the mechanism of lipotoxicity prevention. However, further study of the metabolome, secretome and defined media should reveal detailed influence of particular metabolites in the growth environment on mammalian oocytes and embryos.

**In vitro maturation.** IVM media were supplemented with 10% v/v of the FF obtained either from cyclic or prepubertal pigs and of known fatty acids concentration. FF was aspirated from 3–5 mm ovarian follicles and collected to 1.5 ml tubes (Eppendorf, Germany) and centrifuged at 12000 rpm for 1 min. Supernatant was transferred to a new 1.5 ml tubes, frozen in liquid nitrogen and stored at −80°C.

In vitro maturation of COC was performed in NCSU-23 (North Carolina State University Medium-23). Every COC group was incubated in 500 μl of IVM medium in four-well plates (Nunc, New York, USA) in HeraCell 150 incubator (Thermo Scientific) under conditions: 5% CO2 in atmosphere, 39 °C and maximum humidity. The first 20 h of IVM included IVM medium supplemented with hormones: 10U PMSG (pregnant mare serum gonadotropin, Chorulon, MSD Animal Health, Netherlands) and 10U hCG (Folligon, MSD Animal Health,
Next step included transfer of COCs to fresh, equilibrated medium without hormones and incubation for 24 hours.

After IVM all oocytes with first polar body and no degenerative changes were denuded and transferred to PBS with 0.2% PVP (polyvinylpyrrolidone), frozen or fixed for further analyses. Oocytes subjected for embryo production were denuded in fresh equilibrated NCSU23 medium.

**Embryo production.** *In vitro* matured oocytes (44 h) were denuded by pipetting and washed twice in fresh equilibrated NCSU23 medium (North Carolina State University Medium-23). Parthenogenetic activation (PA) was done by incubation of oocytes in TALP supplemented with 5 μM ionomycin for 5 min followed by incubation in 2 mM 6-DMAP (6-Dimethylaminopurine) for 4 h in 5% CO2, 5% O2 and 90% N2 (New Brunswick Galaxy 170R, Eppendorf). Activated oocytes were transferred to 50 μl droplets of NCSU23 with 4 mg/ml BSA in groups of 20 and cultured to day 5 post activation (pa) when half of the medium was replaced with fresh NCSU23 supplemented with PBS (20% v/v). Embryos were cultured for 7 days and resulting blastocysts were fixed in 4% PFA for immunofluorescent staining and TUNEL analysis or frozen for gene expression analysis. Embryos subjected for IF or TUNEL served also for assessing the embryo quality by calculating number of cells.

**Lipid droplets staining.** Oocytes were fixed in 4% PFA for 30 min at 37°C in four-well plates (Nunc, New York, USA). PFA was removed by washing the oocytes twice in PBS with 0.2% PVP. Cells were stored at 4°C no longer than two weeks. Oocytes were permeabilized in 0.2% Triton X-100 solution for 30 min. at RT and washed 2x times in 0.2% PVP/PBS. Fluorescent dye used to stain lipid droplets was 20 μg/ml BODIPY 493/503 (Life Technologies). Incubation was performed in 500 μl of dye solution in PBS at room temperature for one hour. The nucleus and the polar body were visualized by staining the oocytes with 0.5 μg/ml DAPI (4’,6-diamidino-2-phenylindole; Vector Laboratories, Burlingame, CA, USA). Stained and washed oocytes were mounted on glass slide with single concave (Comex, Poland), coverslipped and stored at 4°C. Oocytes were analysed using confocal microscope Zeiss LSM 510 using 488 nm filter with band pass 500–550 nm for Bodipy mounted on glass slide with single concave (Comex, Poland), coverslipped and stored at 4°C no longer than two weeks. Oocytes were permeabilized in 0.2% Triton X-100 solution for 30 min. at RT and washed 2x times in 0.2% PVP/PBS. Fluorescent dye used to stain lipid droplets was 20 μg/ml BODIPY 493/503 (Life Technologies). Incubation was performed in 500 μl of dye solution in PBS at room temperature for one hour. The nucleus and the polar body were visualized by staining the oocytes with 0.5 μg/ml DAPI (4’,6-diamidino-2-phenylindole; Vector Laboratories, Burlingame, CA, USA). Stained and washed oocytes were mounted on glass slide with single concave (Comex, Poland), coverslipped and stored at 4°C. Oocytes were analysed using confocal microscope Zeiss LSM 510 using 488 nm filter with band pass 500–550 nm for Bodipy 493/503 (Laser Argon2) and 420–480 nm for DAPI (Laser Diode 405). Lipid droplets were assessed through several optical sections of 3 μm thickness each (Z-stack) captured every 10 μm were created. After setting the threshold for fluorescent signals the “Watershed” tool was used to separate the same structures on two stacks (Supplementary Fig. 1). Every oocyte was captured from equatorial section to the top of the cell. Objective (Plan Neofluar 40x/1.3 Oil DIC; Zeiss, Germany), pinhole, filters, offset settings were kept constant throughout the experiments. Lipid droplets were counted using AxioVision software and “Events” tool followed by the creation of Excel file with final results for all oocytes. For the estimation of the area occupied by the LD in oocytes the ImageJ (NIH, USA) software was used. Firstly, the binary, 8-bit (black and white) photo were created. After setting the threshold for fluorescent signals the “Watershed” tool was used to separate the overlapping lipid droplets (Supplementary Fig. 5). The lipid droplet area (μm2) from all analysed stacks was dived by the oocyte area (from the corresponding stacks) giving final result in the percentage share of the lipid droplet occupied area within the oocyte.

**Fatty acid profile and hormone concentration in follicular fluid.** Follicular fluid for FA composition analysis was collected by follicle aspiration from ovary pairs collected in abattoir. Each slaughtered gilt was subjected to removal of the reproductive tract, from which the ovaries have been excised and placed in a separate plastic container for transportation. FA profile and hormone concentration were measured in the same FF samples. Altogether 79 ovary pairs (48 prepubertal and 31 cyclic) were used for the experiment. Fatty acid composition of the FF was analyzed by gas chromatography according to the procedure described previously21. Estradiol (High Sensitivity; ADI-900–174), progesterone (ADI-900-011) and prostaglandin 2A (PGE2 High Sensitivity; ADI-930-001) concentration was measured using ELISA kits (Enzo Life Science) and protocol provided by the manufacturer. 50 μl of follicular fluid was used for analysis on Biotek Synergy2 plate reader. For each reaction seven standards have been prepared to build the standard curve, the blank and the NSB and TA wells with conjugate and antibody respectively. All samples were analysed in single assay in order to avoid variations.

**Apoptosis analysis.** TUNEL was carried out on parthenogenetic day 7 blastocysts (7pa). The terminal TUNEL assay kit was used to detect apoptotic cells (DeadEndTM Fluorometric TUNEL system, Promega Biosciences Inc., Madison, WI, USA). Briefly, embryos were fixed in 4% paraformaldehyde solution in PBS for 30 min at 4°C and washed twice in PBS + 0.025% PVP. Afterwards they were permeabilized by incubation in 0.2% Triton® X-100 solution in PBS for 5 min, washed twice in PBS + 0.025% PVP and covered with equilibration buffer provided by the kit. After 15 min, embryos were transferred to 50 μl equilibration buffer drop containing 1 μl of TdT (terminal deoxynucleotidyl transferase) and 5 μl of fluorescein-conjugated dUTP mix and covered with filtered paraffin oil. One-hour incubation at 37°C in a dark chamber was terminated by 15 min washing in 2x SSC. The embryos were washed three times in PBS + 0.025% PVP and placed on slides in groups of 5~10. Slides were mounted with 40 μl of antifade medium containing DAPI (Vectastain, Vector Labs, Burlinghame, CA, USA), covered with glass coverslip and stored at 4°C for maximum 2 weeks, before evaluated. Embryos pre-treated with DNase I (3 U/50 μl; Promega Biosciences Inc.) served as positive controls, whereas those not subjected to terminal TdT transferase, served as negative controls. Slides were evaluated with fluorescence microscope (Zeiss Axiosvert 2.0) equipped with two filters for fluorescence detection: one to view the green fluorescence of fluorescein at 520 ± 20 nm and another for blue DAPI at 460 nm. A blastomere was considered as TUNEL positive when a strong, green fluorescence signal was observed (Supplementary Fig. 2).

**mRNA gene expression.** Gene expression on mRNA level was performed on oocytes before and after IVM and embryos at blastocyst stage. Denuded oocytes (25 per sample) and embryos (3 per sample) were placed in 1.5 ml DNA LoBind tubes (Eppendorf) in PBS and frozen in liquid nitrogen. All samples were stored at −80°C. The total RNA was extracted with mirVANA Paris Kit (Ambion, ThermoFisher Scientific) according to...
the manufacturer’s protocol. Briefly, the samples were incubated in Cell Disruption Buffer and 2x Denaturing Solution. Afterwards the acid phenol chloroform was added to cell lysate and centrifuged for 15 min at 14000 rpm. The upper, clear phase was mixed with isopropanol (1:2.5 v/v) and placed in a filter column and centrifuged at 8000 g for 1 min. The next steps involved 2x washing to remove contaminants. Finally, the total RNA was eluted into a fresh 1.5 ml LoBind tube with 100 μl of prewarmed Elution Buffer. The RNA samples were further precipitated with NF Pellet Paint Co-Precipitant. 1 μl of Pellet Paint, 10 μl of 3 M sodium acetate and 200 μl of freshly prepared 96% EtOH were added to RNA sample. After 5 min incubation, the samples were centrifuged at top speed for 10 min. The RNA pellet was washed followed by centrifugation by 75% and 96% EtOH and dried in 40°C. RNA was resuspended in Molecular Biology Grade water (Sigma-Aldrich) in 10 μl. Next RNA was reverse transcribed using Transcriptor First Strand cDNA synthesis Kit (Roche) following manufactures protocol and using total isolated RNA. The protocol included denaturation of RNA and primers at 65°C for 10 min followed by reverse transcription at 25°C for 5 min and 42°C for one hour and inactivation at 80°C for 10 min. The cDNA samples were stored at −20°C until further analysis. Genes responsible for fatty acid metabolism (ACACA, ELOVL2, ELOVL6, SCD, FASN, FADS1, FADS2, PLIN2) and developmental competence (OCT4, GDF9, BMP15) were analysed. Each cDNA sample was analysed in two independent PCR runs, and the mean value was used for the calculations of relative transcript abundance to the most stable reference genes ACTB and GAPDH (data not shown). List of analysed genes, primer and probe sequences designed by Tib Molbiol (Germany) are shown in Supplementary Table 1. qPCR was conducted using the standard curve method. For this purpose, the desired sequences for all analysed genes were amplified by PCR and visualised on 1.5% agarose gel with the Gene Ruler™ 100 bp DNA Ladder (Fermentas, Canada). The PCR product was excised from the gel, isolated and purified using the Gel Extraction Kit (Fermentas). Based on the DNA concentration measured with a Nanodrop c2000 system (Thermo Scientific, USA), a serial 10-fold dilutions of DNA with a known concentration (standards) were generated. Each standard was used as a separate template for a real-time PCR reaction to produce the appropriate standard curve with the LightCycler 480 II software (Roche, Switzerland). The reaction conditions and efficiency of the reactions for all genes were analysed separately. All reactions were performed using the Light Cycler 480 II system with a set of supplied reagents. The 10 μl reaction mixture consisted of 5 μl of the LightCycler Probe Master, 0.5 μM primers, 0.3 μM probes and 1 μl each of the cDNA and. The reaction conditions were as follows: denaturation: 95°C, 10 min; amplification: 40 cycles of 95°C for 10 s, 60°C for 10 s, 72°C for 10 s; and final cooling at 40°C. The temperature slope was set at 20°C/s during amplification.

**Immunofluorescent staining.** The oocytes and embryos for immunofluorescent staining were fixed in 4% PFA for 30 min at RT and stored in 4°C. The applied staining protocol was previously described by Madeja et al. (Animal) Briefly the staining was initiated by permeabilization of oocytes and embryos for 20 min in 0.55% Triton X-100 in PBS followed by 10 min in NH₄Cl and blocking for 60 min (10% FCS in PBS). Oocytes have been incubated with both primary (1:100) and secondary antibodies (1:200) at 4°C for 24 h. Stained and washed oocytes or embryos were mounted on glass slide with single concave (Comex, Poland), coversons and stored at 4°C. Slides were analysed using confocal microscope Zeiss LSM 880 using 488 nm filter with band pass 500–530 nm (Laser Diode 405) and 543 nm with band pass 550 nm (HeNe1). Objective (Plan Neofluar 40x/1.3 Oil DIC; Zeiss, Germany), pinhole, filters, offset settings were kept constant throughout the experiments. Signal threshold (detector gain) for oocytes and embryos was set after examination of negative controls performed during every staining procedure. Negative control slides were prepared with exactly same protocol without incubation with primary antibody. A-Tubulin was used as a positive control indicating pene-tration of antibodies through zona pellucida and oolemma. ACACA, FASN, SCD and FADS1 localization was performed using Abcam and Santa Cruz antibodies (Supplementary Table 2).

**Statistical analysis.** A comparison of the experimental groups was performed using IBM SPSS Statistics 23.0. All data (before computing) were subjected to testing for normal distribution using the Kolmogorov-Smirnov and Shapiro-Wilk tests. The differences in lipid droplet number and occupied area in oocytes of different groups were calculated using the Kruskal-Wallis and two-tailed Mann-Whitney U tests. mRNA gene expression differences were analyzed using nonparametric two-tailed Mann-Whitney U test. The frequency of apoptosis in oocytes of different groups were calculated using the chi-square and Fisher’s exact tests. Fatty acid and hormones concentration were analysed with t-Student and Mann-Whitney U tests respectively. All data with P < 0.05 were considered statistically significant.

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**Author Contributions**
P.P. designed and performed the experiments (oocyte maturation, embryo culture, gene expression, lipid droplet staining and confocal analysis, hormone concentration), analysed data and wrote the manuscript. A.C. performed analysis of fatty acid profile in follicular fluid, E.W., N.M., E.M. performed experiments (apoptosis, lipid droplet staining and confocal analysis, hormone concentration), analysed data and wrote the manuscript. Z.M. collected samples, Z.M. and D.L. edited the manuscript.

**Additional Information**
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