Filopodia-like protrusions of adjacent somatic cells shape the developmental potential of oocytes

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The oocyte must grow and mature before fertilization, thanks to a close dialogue with the somatic cells that surround it. Part of this communication is through filopodia-like protrusions, called transzonal projections (TZPs), sent by the somatic cells to the oocyte membrane. To investigate the contribution of TZPs to oocyte quality, we impaired their structure by generating a full knockout mouse of the TZP structural component myosin-X (MYO10). Using spinning disk and super-resolution microscopy combined with a machine-learning approach to phenotype oocyte morphology, we show that the lack of Myo10 decreases TZP density during oocyte growth. Reduction in TZPs does not prevent oocyte growth but impairs oocyte-matrix integrity. Importantly, we reveal by transcriptomic analysis that gene expression is altered in TZP-deprived oocytes and that oocyte maturation and subsequent early embryonic development are partially affected, effectively reducing mouse fertility. We propose that TZPs play a role in the structural integrity of the germline–somatic complex, which is essential for regulating gene expression in the oocyte and thus its developmental potential.

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Introduction

During oogenesis, mammalian oocytes undergo a process of differentiation that determines their quality and thus their developmental potential as early embryos after fertilization. Oocyte differentiation begins with the entry into a growth phase during which the oocyte increases in size and stores large amounts of macromolecules necessary to complete its development and early embryogenesis. During growth, the oocyte remains arrested in prophase and gradually acquires the competence to resume meiosis (1). Once fully grown, it further differentiates by undergoing meiotic maturation. During maturation, the oocyte completes the first meiotic division (meiosis I) and becomes arrested at metaphase of the second meiotic division (meiosis II). Meiosis completion then occurs if fertilization takes place.

The oocyte does not progressively acquire meiotic and developmental competence by itself in an autonomous way, but through a close dialogue with the somatic cells that surround it, called follicular cells, which themselves undergo a parallel and intricate process of differentiation (2, 3, 4, 5). Together with the oocyte, the follicular cells form a germ–somatic complex called the ovarian follicle.

Although the oocyte communicates with its surroundings via secreted factors (2, 4, 6), somatic communication occurs mostly through direct cell–cell contacts mediated by actin-rich filopodia-like (7) protrusions called transzonal projections (TZPs) (6, 8, 9, 10). These TZPs cross the zona pellucida, the extracellular matrix surrounding the oocyte, and contact the oocyte membrane at their tips. They allow follicular cells to metabolically support the oocyte and prevent early meiotic resumption (3, 6). In addition, TZPs may provide structural support in the follicle by maintaining cohesion between the oocyte and follicular cells (9, 11).

Although the dialogue within the ovarian follicle is increasingly emerging as a major player in the emergence of oocyte quality, the role of direct cell–cell contact communication between the oocyte and follicular cells is still poorly understood and may shed additional light on the acquisition of oocyte developmental competence. In this study, we sought to characterize the contribution of TZPs to oocyte development by impairing the structure of TZPs. As a candidate gene, we selected myosin-X (MYO10), a structural component of TZPs (7) known to promote filopodium formation (12, 13, 14) and potentially involved in the formation or maintenance of TZPs (15). We generated a full knockout mouse of Myo10 in all cell types to completely deplete it from the ovarian follicle (Myo10\textsuperscript{−/−}) full. Furthermore, we took advantage of our previously generated mouse strain conditionally deleted for Myo10 specifically in oocytes from early growth onward (Myo10\textsuperscript{−/−} oo) (16) to help distinguishing phenotypes caused by loss of somatic MYO10 or loss of oocyte.

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MYO10. We show that the loss of somatic but not oocyte MYO10 greatly decreases the density of TZPs. Surprisingly, oocytes deprived of TZPs develop to a normal size, but display oocyte-matrix defects. Importantly, gene expression is altered in TZP-deprived oocytes, correlating with lower rates of oocyte maturation and subsequent early embryonic development, inducing lower fertility in mice.

Results

Global deletion of Myo10 decreases the density of TZPs without altering ovarian follicular organization

To deplete TZPs, we generated full knockout mice lacking the TZP component MYO10 in all cell types (Fig S1A–D). Loss of MYO10 expression in mice was previously characterized as semi-lethal, with some animals dying during embryogenesis whereas others being able to develop until adulthood (14). Similar to this study, we obtained adult Myo10−/− mice (named Myo10−/− full thereafter) and most showed phenotypes characteristic of the earlier described Myo10 full knockout, such as the presence of a white spot on the abdomen and webbed fingers (14) (Fig S1B). In addition to this strain, we used our previously generated mouse strain conditioned for the oocyte-secreted factor GDF9 (18, 19) led to ovaries with abnormal morphology, lacking fully grown follicles. We therefore tested whether decreasing TZP density altered the integrity of Myo10−/− full ovaries. To gain access to intra-ovarian organization, whole ovaries from adult mice were transilluminated (20). As in control ovaries, Myo10−/− full ovaries contained late antral follicles (Fig S3A). Myo10−/− full mice also showed a variation in the volume of their oocytes (Video 2 and Video 3). Because of this variation, we assessed global ovarian integrity by measuring the follicular density of the ovary (number of follicles normalized by the ovarian area) rather than the absolute number of follicles per ovary. Follicular density was not significantly different between control and Myo10−/− full ovaries (Fig S3B), indicating that the complete loss of MYO10 and subsequent reduction in TZP density do not alter global follicular organization in the ovary. Thus, we confirmed that MYO10 is a structural component of TZPs, essential for TZP formation or maintenance (7, 15).

Oocytes deprived of TZPs reach a canonical size at the end of growth but are morphologically different from controls

To determine the TZP-mediated contribution of follicular cells to oocyte development, we assessed whether Myo10−/− full oocytes...
deprived of TZPs developed properly. We used our previously developed deep-learning approach to characterize in an automatic manner the morphology of Myo10−/− full oocytes at the end of the growth phase with the software Oocytor (21) to assess their meiotic competence (defined as the ability of oocytes to resume meiosis at the end of growth and mature properly). First, Myo10−/− full oocytes grew to normal sizes (Fig 2A and B) and were arrested in prophase. We then measured the thickness and the perimeter of the zona...
Figure 2. TZP-deprived oocytes have morphological alterations.

(A) Brightfield images of fully grown oocytes from the Myo10 full knockout strain (left panel) and the Myo10 oocyte knockout strain (right panel). For each panel, controls are on the left, and Myo10−/− oocytes, on the right. Scale bar = 10 μm.

(B) Scatter plot of the equatorial plane area of control and Myo10−/− full oocytes (in dark gray and green, respectively) and control and Myo10−/− oo oocytes (in light gray and blue, respectively). (n) is the number of oocytes analyzed. Data are the mean ± s.d. with individual data points plotted. Data are from four to 13 independent experiments. Statistical significance of differences was assessed by an ANOVA or a Kruskal–Wallis test.
Cells, SiR-actin (white) distribution, control (upper panel) and Filopodia from somatic cells shape oocyte quality. (Figs 2D–G and 5A–G), allowing us to detect in an unbiased manner the morphological features that differed the most between control and Myo10−/− full oocytes. One of the most important features used by our algorithm to distinguish control from Myo10−/− full oocytes was the composition of the zona pellucida with the presence of vertical structures resembling TZPs crossing the zona pellucida (Figs 2D, 6A, and 5D and G). Intriguingly, these structures were detected more frequently in Myo10−/− full oocytes than in controls (Fig 2G), despite the fact that total TZP density is decreased in Myo10−/− full oocytes. These structures, visible in transmitted light in Myo10 full knockouts, always contain actin (Fig 2F). This implies that even though there are less TZPs in Myo10 full knockouts (Fig 1E), they are more visible by transmitted light than in controls, which are more abundant (Fig 1E) but less visible by transmitted light (Fig 2F). Our observations suggest that in Myo10 full knockouts, the structure of the remaining TZPs and/or that of the zona pellucida is different from that of control oocytes. Interestingly, we occasionally observe some follicular-like cells located ectopically within the perivitelline space (the space between the oocyte and the zona pellucida) of Myo10 full knockouts (Fig 5C), and the texture of the zona pellucida is different between control and Myo10 full knockouts (Fig 2C). We therefore hypothesize that the structure of the zona pellucida (a viscoelastic extracellular matrix) may change as a function of the cross-linking generated by the number of TZPs passing through it. If so, this could change its mechanical properties. We thus assessed the mechanical properties of the zona pellucida of control and Myo10 full knockouts using atomic force microscopy (AFM) as in Reference 22. Our experiments show decreased zona pellucida elasticity in Myo10 full knockouts compared with controls (Fig 5H). This suggests that the reduced number of TZPs could affect the structure, maybe via reduced cross-linking of the zona pellucida, and thus the mechanical properties of the zona pellucida.

The second important feature used by our algorithm was the distribution of the perivitelline space around the oocyte, which was detected to be more uniform in Myo10−/− full oocytes than in controls (Fig 2D, quantification in E; Fig 5A and G), as if oocytes lacking TZPs were more homogeneously detached from the zona pellucida. Furthermore, our machine-learning algorithm identified Myo10−/− full oocytes as more circular than control oocytes (Fig 5D, E, and G), as indicated by a lower aspect ratio in TZP-deprived oocytes (Fig 5E). In accordance with this feature, the perimeter of Myo10−/− full oocytes was slightly smaller than control oocytes for a similar area (Fig 5F).

Thus, using automatic approaches, we show that Myo10−/− full oocytes can reach normal sizes but display morphological differences from controls, in particular regarding the zona pellucida texture, a feature known to predict meiosis entry and maturation outcome (21).

Gene expression is modified in oocytes deprived of TZPs

We next wondered whether the differences both in the density of communicating structures between the oocyte and its follicular cells and in oocyte morphology translated into differences in oocyte developmental potential. The oocyte acquires its developmental potential during its growth, where it accumulates a large number of transcripts. Transcription decreases dramatically at the end of oocyte growth and increases after fertilization in the zygote (23); thus, protein synthesis during the meiotic divisions relies predominantly on the translational regulation of transcripts accumulated during growth, making this storage critical for oocyte quality (24, 25, 26). Interestingly, follicular cells have been shown to modulate chromatin configuration and thus global transcriptional activity of the oocyte during its growth (24, 27). Because Myo10−/− full oocytes have reduced ability of the oocyte to resume meiosis (21), and showed that they are similar between Myo10−/− full oocytes and controls (Fig 5A and B). However, the texture of the zona pellucida, one of the two most important features to assess oocyte maturation potential (21), was lower in Myo10−/− full oocytes (Fig 2C). Thus, the morphology of Myo10−/− full oocytes at the end of growth suggests that they have a similar potential for meiosis resumption as controls, but a decreased potential for oocyte maturation.

test depending on whether the data followed a Gaussian distribution, \( P = 0.4459 \) (n.s, Ctrl full versus Myo10−/− full), \( P = 0.7355 \) (n.s, Ctrl full versus Ctrl oo), \( P > 0.9999 \) (n.s, Myo10−/− full versus Ctrl oo), and \( P > 0.9999 \) (n.s, Ctrl oo versus Myo10−/− oo). n.s, not significant. (C) Scatter plot of the zona pellucida texture of control and Myo10−/− full oocytes (in dark gray and green, respectively) and control and Myo10−/− oo oocytes (in light gray and blue, respectively). (n) is the number of oocytes analyzed. Data are the mean ± s.d. with individual data points plotted. Data are from four to 13 independent experiments. Statistical significance of differences was assessed by an ANOVA or a Kruskal–Wallis test depending on whether the data followed a Gaussian distribution, \( P = 0.0000 \) (****, Ctrl full versus Myo10−/− full), \( P = 0.0001 \) (****, Myo10−/− full versus Ctrl oo), \( P > 0.9999 \) (n.s, Ctrl oo versus Myo10−/− oo), and \( P > 0.9999 \) (n.s, Ctrl oo versus Myo10−/− oo). n.s, not significant. (D) Brightfield images of a fully grown control (upper panel) and Myo10−/− full oocyte (bottom panel). For each panel, the right image displays the automatic segmentation of the oocyte perivitelline space (perivitelline space, yellow). Scale bar = 10 μm. (E) Scatter plot of the coefficient of variation of the perivitelline space thickness, as represented by the schemes on the left. Control and Myo10−/− full oocytes are in dark gray and green, respectively. Control and Myo10−/− oo oocytes are in light gray and blue, respectively. (n) is the number of oocytes analyzed. Data are the mean ± s.d. with individual data points plotted. Data are from four to 13 independent experiments. Statistical significance of differences was assessed by an ANOVA or a Kruskal–Wallis test depending on whether the data followed a Gaussian distribution, \( P < 0.0001 \) (****, Ctrl full versus Myo10−/− full), \( P = 0.9999 \) (n.s, Ctrl full versus Ctrl oo), \( P = 0.3139 \) (n.s, Ctrl full versus Myo10−/− oo), \( P = 0.0001 \) (****, Myo10−/− full versus Myo10−/− oo), \( P > 0.0001 \) (****, Myo10−/− full versus Ctrl oo), \( P > 0.9999 \) (n.s, Ctrl oo versus Myo10−/− oo), and \( P > 0.9999 \) (n.s, Ctrl oo versus Myo10−/− oo). n.s, not significant. (F) Images of the flattened zona pellucida obtained from oocyte segmentation. The top panel shows a control zona pellucida, and the lower one, a Myo10−/− full zona pellucida. For each panel, the bottom image is a magnification of the brightfield image of the zona pellucida at the top. Scale bar = 10 μm. Below each flattened zona pellucida are examples of the zona pellucida from control and Myo10−/− full oocytes. Brightfield images are on the left panels, SiR-actin (white)–labeled TZPs are on the middle panels, and merge images (SiR-actin in magenta) are on the right panels. Scale bar = 9 μm. (G) Scatter plot of the presence of vertical TZP-like structures in the zona pellucida (ZP), as represented by the schemes on the left. Control and Myo10−/− full oocytes are in dark gray and green, respectively. Control and Myo10−/− oo oocytes are in light gray and blue, respectively. (n) is the number of oocytes analyzed. Data are the mean ± s.d. with individual data points plotted. Data are from four to 13 independent experiments. Statistical significance of differences was assessed by an ANOVA or a Kruskal–Wallis test depending on whether the data followed a Gaussian distribution, \( P > 0.9999 \) (n.s, full versus Myo10−/− full), \( P = 0.7183 \) (n.s, full versus Ctrl oo), \( P = 0.7291 \) (n.s, full versus Myo10−/− oo), \( P > 0.9999 \) (n.s, full versus Myo10−/− oo), \( P = 0.0001 \) (****, Myo10−/− full versus Ctrl oo), \( P = 0.0001 \) (****, Myo10−/− full versus Myo10−/− oo), and \( P = 0.0001 \) (****, Myo10−/− full versus Myo10−/− oo). n.s, not significant.
Figure 3. Gene expression is modified in TSP-deprived oocytes.

(A) Volcano plot of differential gene expression between fully grown Myo10+/− and Myo10−/− full oocytes. Differential transcriptomic analysis was performed by RNA-Seq. Up-regulated transcripts are shown in red (up, 605 transcripts); down-regulated transcripts, in blue (down, 80 transcripts, including Myo10−/−); and non-deregulated transcripts, in gray (not significant, 15,702 transcripts). Significance was set at P-adj < 0.05. Myo10−/− full oocytes were collected from two Myo10fl/fl; Cre+ mice and Myo10+/− oocytes from two Myo10wt/fl; Cre+ mice. For each condition, three biological replicates (each containing 10 oocytes) and three technical replicates were performed.

(B) Pie chart showing the distribution of deregulated transcripts. The chart highlights the proportion of protein coding, pseudogene, and other transcript types.

(C) Bar graph comparing relative mRNA quantity for different conditions.

(D) Images showing growing and fully grown oocytes with and without EU treatment.

(E) Graphs depicting relative EU intensity with and without EU treatment.
oocytes lack follicular cell contacts and potentially benefit less from exogenous transcriptional regulation, we tested whether their transcriptome differed from that of control oocytes.

By performing differential transcriptomic analysis by RNA-Seq between Myo10+/- and Myo10-/- full oocytes at the end of the growth phase (23), we found 685 genes significantly (P-adj < 0.05) misregulated in Myo10-/- full oocytes (Fig 3A and Table S1), most of which were up-regulated (605 of 685 representing 88.3% of deregulated transcripts). Most of the deregulated genes were protein-coding genes (Fig 3B), 10 of which were validated by RT-qPCR (Fig 3C). Among them, Myo10 mRNA was confirmed to be down-regulated in Myo10-/- full oocytes, validating both our RNA-Seq analysis and our genetic deletion tool (Fig 3C). Because most of the deregulated transcripts in Myo10-/- full oocytes were up-regulated, this could mean that the reduction in TZP density had resulted in either a higher transcription rate or increased stability of these transcripts.

We thus tested whether this deregulation reflected a change in the global transcriptional rate of TZP-deprived oocytes (23). Fully grown and smaller growing Myo10-/- full oocytes were incubated with the uridine analog 5-ethynyl uridine (EU) to label RNA synthesis (Fig 3D and E). Oocytes were also incubated without EU to serve as a negative control. To measure global RNA synthesis, the EU signal intensity was normalized to the DAPI signal intensity. Interestingly, global RNA synthesis was not different in growing and fully grown Myo10-/- full oocytes compared with control oocytes (Fig 3E).

In summary, we demonstrate here that the lack of somatic contacts caused by decreased TZP density leads to a deregulation of oocyte gene expression at the end of its growth.

**TZP-deprived oocytes are prone to stop development in meiosis I despite correct spindle morphogenesis and chromosome alignment**

We showed above that reduction in somatic contacts modulates the oocyte transcriptome. Importantly, oocytes depend on the transcripts synthesized during growth to develop until fertilization (25, 26). We thus tested whether TZP-deprived oocytes successfully underwent the developmental step after growth, namely, meiotic divisions. After nuclear envelope breakdown (NEBD), the oocyte experiences an asymmetric division in size, leading to a large cell, the oocyte, and a tiny cell, the polar body. This size difference is essential to keep the nutrients stored during growth in the prospective fertilized oocyte for embryogenesis (28). Importantly, we observed that Myo10-/- full oocytes deprived of TZPs showed an increased frequency of developmental arrest at meiosis I, without extruding a polar body (Video 4 and Fig 4A and B; 38% of TZP-deprived oocytes do not extrude a polar body, compared with 15% in the controls), consistent with our morphological characterization using Oocytor (Fig 2C). We previously showed that MYO10 expressed in oocytes is dispensable for meiotic divisions and female fertility (16), suggesting that the meiotic arrest observed in a subset of Myo10-/- full oocytes may be a consequence of decreased follicular cell contacts before meiosis resumption. We also observed that the subpopulation of Myo10-/- full oocytes that extruded a polar body did so with a timing similar to controls (Video 4 and Fig 4A and C), indicating that the complete loss of MYO10 and subsequent TZP depletion did not delay polar body extrusion in oocytes, which could achieve it. Therefore, oocytes deprived of TZPs have a lower developmental potential than oocytes that retain a canonical density of follicular cell contacts, reinforcing the importance of the TZP-mediated dialogue between the oocyte and its niche for a fully efficient acquisition of oocyte competency.

To gain insight into the origin of the meiotic arrest observed in some TZP-deprived oocytes, we first investigated whether the meiotic spindles were correctly formed in Myo10-/- full oocytes. After NEBD, microtubules organize into a ball near the chromosomes that gradually bipolarizes to form a barrel-shaped spindle (29, 30, 31). To assess spindle morphogenesis, oocytes were stained with SiR-tubulin (SiR-tub) to label microtubules and followed throughout meiotic maturation. We observed that in both Myo10-/- full oocytes that extruded a polar body and those that did not, meiosis I spindles bipolarized properly and within similar timings as in controls (Video 5 and Fig 5A–D). To further evaluate spindle
morphogenesis in Myo10−/− full oocytes, we measured spindle interpolar length and central width 7 h after NEBD once spindle morphogenesis is complete and showed that Myo10−/− full spindles were not significantly different from control spindles (Fig 5B and D).

Importantly, we observed that Myo10−/− full oocytes that did not extrude a polar body arrested development in metaphase I, without experiencing anaphase (Video 5 and Fig 5A).

In addition to assessing spindle formation, we also examined meiosis I spindle positioning in Myo10−/− full oocytes. During meiosis I, the spindle first assembles in the center of the oocyte and then migrates to the cell cortex by a process involving F-actin, resulting in an asymmetric division in size (32, 33, 34, 35). We monitored the positioning of SiR-tub–labeled spindles throughout meiosis I and analyzed their localization 30 min before anaphase for oocytes that extruded a polar body or up to 12 h after NEBD for those that did not. The percentage of Myo10−/− full oocytes with an off-centered spindle was not different from control oocytes (Video 5 and Fig 5A and E). In agreement with unaltered spindle positioning, we did not observe a defect in the organization of the F-actin cytoplasmic network (including the actin cage) mediating spindle migration in fixed Myo10−/− full oocytes labeled with phalloidin (Fig 5A–C). These results refine our understanding of the meiotic arrest of some Myo10−/− full oocytes by showing that they arrest in metaphase I with a normally assembled and positioned spindle.

Because severe chromosome misalignment could prevent anaphase onset by sustaining the spindle assembly checkpoint in mouse oocytes (36), we assessed chromosome alignment in

Figure 4. TZP-deprived oocytes tend to arrest in meiosis I. (A) Brightfield images from spinning disk videos from 3 h to 18 h after NEBD. The top images show a control oocyte; the middle ones, a Myo10−/− full oocyte that extruded a polar body (PBE); and the bottom ones, a Myo10−/− full oocyte that did not extrude a polar body (no PBE). The blue asterisk indicates a lysed polar body. Scale bar = 10 μm. (B) Stacked bars of first polar body extrusion rate as a percentage of control and Myo10−/− full oocytes. The gray bars represent the percentage of oocytes that extruded a polar body (PBE), and the white bars, the percentage of oocytes that did not extrude a polar body (no PBE). (n) is the number of oocytes analyzed. Data are from three independent experiments. Statistical significance of differences was assessed by a two-sided Fisher exact test, P = 0.0113. (C) Scatter plot of the timing in hours after NEBD of first polar body extrusion of control (dark gray) and Myo10−/− (green) full oocytes. (n) is the number of oocytes analyzed. Data are the mean ± s.d. with individual data points plotted. Data are from six independent experiments. Statistical significance of differences was assessed by a two-tailed Mann–Whitney test, P = 0.3194. n.s, not significant.
Figure 5. TZP-deprived oocytes have correctly formed and positioned spindles and aligned chromosomes. 
(A) Maximum-intensity Z projection images of oocytes stained with SiR-tubulin to label microtubules (SiR-tub, yellow) merged with corresponding brightfield images. Images are extracted from spinning disk videos from 30 min to 14 h 30 min after NEBD. The top images show a control oocyte; the middle ones, a Myo10−/− full oocyte that extruded a polar body (PBE); and the bottom ones, a Myo10−/− full oocyte that did not extrude a polar body (no PBE). Scale bar = 10 μm. (B) Maximum-intensity Z projection images of oocytes stained with SiR-tub (yellow) 7 h after NEBD. The top image shows the spindle of a control oocyte; the middle one, the spindle of a Myo10−/− full oocyte.
**Discussion**

In this study, we aimed to gain more insight into the contribution of cellular protrusions established by surrounding somatic cells to postnatal oocyte development. By genetically deleting the TZP component MYO10 in all murine cell types, including follicular cells, we greatly reduced the density of oocyte–follicular cell contacts and assessed the resulting oocyte development.

Surprisingly, although the complete loss of MYO10 decreased by fivefold the density of TZPs in fully grown oocytes, MYO10−/− full oocytes grew to normal sizes (Fig 2A and B) and were arrested in prophase. Hence, and in contrast to previous studies where the impairment of oocyte–follicular cell communication by deletion of connexin 37 or GDF9 led to, respectively, smaller or larger oocytes than controls (17, 18, 19), reducing TZP density did not prevent oocytes from growing to a canonical size. Interestingly, a small fraction of TZPs remained in MYO10−/− full oocytes, and these remaining TZPs were functional allowing exchanges between the oocyte and follicular cells (Fig S2C and D). This implies that a subpopulation of TZPs may be MYO10-independent and potentially explains how oocytes managed to grow to a canonical size and maintained the meiotic arrest in prophase despite global reduced intercellular communication.

Even though TZP-deprived oocytes grew to normal sizes, we identified morphological differences, mostly related to the integrity of the zona pellucida. The zona pellucida, which is the extracellular matrix surrounding the oocyte, is composed of proteins secreted by the oocyte throughout growth and assembled in cross-linked filaments (37, 38). Using our machine-learning algorithm to phenotype oocytes, we found that the zona pellucida was more homogeneously detached in TZP-deprived oocytes. It is likely that reduction in TZP density, and consequently the decrease in adherens junctions between the oocyte and follicular cells (9, 11), led to this apparent looser oocyte-matrix adhesion. Furthermore, our machine-learning algorithm detected TZP-like structures more readily in MYO10−/− full oocytes than in controls, despite their decrease in TZP density. These structures always contain actin (Fig 2F). Thus, there are less TZPs in MYO10−/− full knockouts (Fig 1E), but most of these remaining TZPs are visible by transmitted light. This could reflect an alteration in the structure of the remaining TZPs, or alternatively a difference in the structure of the MYO10−/− full zona that extruded a polar body (PBE), and the bottom one, the spindle of a MYO10−/− full oocyte that did not extrude a polar body (no PBE). Scale bar = 5 μm. (C) Graph of the percentage of control (dark gray) and MYO10−/− (green) full oocytes with a bipolarized spindle as a function of time after NEBD. Data are from four independent experiments. For each time point, statistical significance of differences was assessed by a two-sided Fisher exact test, P > 0.9999 for 1 h after NEBD, P = 0.3433 for 2 h after NEBD, P = 0.7765 for 3 h after NEBD, P > 0.9999 for 4 h after NEBD, P = 0.6937 for 5 h after NEBD, and P > 0.9999 for 6 h after NEBD. (D) Scatter plots of spindle interpolar length (left) and central width (right) 7 h after NEBD in control (dark gray) and MYO10−/− (green) full oocytes. (n) is the number of oocytes analyzed. Data are the mean ± s.d. with individual data points plotted. Data are from four independent experiments. Statistical significance of differences was assessed by a two-sided Fisher exact test, P = 0.3832. (n.s., not significant). (I) Maximum-intensity Z projection images from spinning disk videos of oocytes injected with H2B-GFP to label chromosomes (H2B, cyan) in a control oocyte (top images), a MYO10−/− full oocyte that extruded a polar body (PBE, middle images), and a MYO10−/− full oocyte that did not extrude a polar body (no PBE, bottom image). Left images show chromosomes 30 min before anaphase for oocytes extruding a polar body or 12 h after NEBD for the MYO10−/− full oocyte that did not extrude a polar body. Right images show chromosomes after anaphase. Scale bar = 5 μm. (G) Scatter plots of metaphase plate height (left) and width (right), as shown in the lower schemes, 30 min before anaphase for oocytes that extruded a polar body or 12 h after NEBD for those that did not. Control and MYO10−/− full oocytes are in dark gray and green, respectively. (n) is the number of oocytes analyzed. Data are the mean ± s.d. with individual data points plotted. Data are from three independent experiments. Statistical significance of differences was assessed by two-tailed unpaired t tests. P = 0.5331 for the height and P = 0.0395 for the width. n.s., not significant.
pellucida that would make the remaining protrusions crossing it more apparent. All these features taken together may reflect impaired assembly or maintenance of the zona pellucida in oocytes with reduced cell–cell contacts, because the zona pellucida is a viscoelastic extracellular matrix whose properties could change as a function of the cross-linking generated by the number of TZPs passing through it (37, 38). In support of this, we showed using AFM that the reduced number of TZPs passing through it affects the structure, and thus the mechanical properties of the zona pellucida, decreasing its elasticity (Fig S4H). In addition, we occasionally observed some follicular-like cells located ectopically within the perivitelline space of TZP-deprived oocytes, a phenotype not observed in control oocytes (Fig S4C; see red arrow). The same ectopic localization was observed in GDF9-deficient mice, which also displayed oocytes with reduced TZP density, and such abnormal localization of follicular cells was correlated with low oocyte viability (19). Interestingly, the presence of follicular-like cells inside the perivitelline space has been observed in mice lacking ZP1 (39), the secreted zona pellucida protein that mediates the cross-linking of the matrix filaments (37, 38). By modulating matrix filament cross-linking, TZPs may therefore be involved in preserving zona pellucida integrity, which is important for fertilization (37, 38). Interestingly, we recovered more unfertilized oocytes from Myo10−/− full females compared with controls after mating, which suggests a potential effect of the observed alteration of the zona pellucida on fertilization (Fig 6A).

Figure 6. Developmental potential of TZP-deprived oocytes is altered, correlating with subfertility in females. (A) On the left, stacked bars showing the percentage of unfertilized oocytes (black bars), dead zygotes (gray bars), and alive zygotes (white bars) recovered at E0.5. (n) is the number of cells recovered, and (N) is the number of successful matings analyzed. Data are from three independent experiments. Statistical significance of differences was assessed with a chi-squared test, *P < 0.0001. On the right, cells recovered at E0.5 and cultured to E1.5 from a control female (upper panel) and a Myo10−/− full female (lower panel), stained with Hoechst (blue) to visualize DNA. Scale bar = 40 μm. (B) Stacked bars showing the percentage of embryonic arrest (black bars) and blastocysts (white bars). (n) is the number of cells recovered, and (N) is the number of successful matings analyzed. Data are from three independent experiments. Statistical significance of differences was assessed by a two-sided Fisher exact test, *P < 0.0001. (C) Scatter plots of the number of litters per couple, number of pups per litter, and total number of pups per couple from Myo10wt/wt; Cre- female mice (control, gray) and Myo10fl/fl/Cre- female mice (carrying Myo10−/− full oocytes, green). Six independent matings (N) were set for six months each with Myo10wt/wt; Cre- males. Statistical significance of differences was assessed with two-tailed unpaired t tests, *P = 0.04 (for the number of litters per couple), **P = 0.0009 (for the number of pups per litter), and ***P = 0.0052 (for the total number of pups per couple).
Importantly, TZP-deprived oocytes have lower developmental competence than control oocytes, as they more frequently arrest at metaphase of the first meiotic division, and produce fewer embryos with lower viability, impairing female fertility. Consistent with our results, a decrease in oocyte developmental potential correlated with reduced TZP density was also observed in oocytes lacking the focal adhesion kinase/proline-rich tyrosine kinase 2 (40) and oocytes lacking radixin (41). In Myo10lox/lox full oocytes, the basis of the meiotic arrest observed in a subpopulation of oocytes remains to be deciphered. This developmental arrest may be the readout of the altered oocyte transcriptome because we revealed by RNA-Seq analysis that gene expression before meiosis resumption is modified in TZP-deprived oocytes. As oocytes undergo meiotic divisions relying on transcripts synthesized and stored during growth (25, 26), an impairment in this storage could compromise oocyte quality. Interestingly, some deregulated transcripts in TZP-deprived oocytes, including the most deregulated ones, are involved in the oxidative stress response, the metabolism of polyamine, and cell cycle transitions (Osert, Ahk1b3, Oaz1, Psma5, and Cyclin B1; Fig 3C and Table S1), pathways that when deregulated are known to affect subsequent meiotic maturation and early embryogenesis (42, 43, 44, 45, 46, 47).

In conclusion, we have shown that cytoplasmic protrusions of surrounding somatic cells enhance the developmental potential of the oocyte. Consistent with other studies, we suggest that TZPs contribute to zona pellucida integrity and oocyte-matrix adhesion—directly as an intercellular contact structure or indirectly through metabolite exchange optimizing oocyte quality and the establishment of a robust zona pellucida—thus playing a role in the structural integrity of the ovarian follicle. In addition, we propose a novel role of TZPs as modulators of the synthesis/stability of structural integrity and oocyte-matrix adhesion (48). It may notably provide clues as to how cellular development is modulated by exogenous regulation of gene expression through direct intercellular dialogue.

Materials and Methods

Mouse strain generation and genotyping

To generate Myo10wt/lox mice, two loxP sites flanking Myo10 exons 23–25 and an FRT-flanked neomycin selection cassette located between Myo10 exons 22 and 23 were inserted in the genome of C57BL/6 mice by genOway. Mice were then mated with Zp3-Cre mice already housed in the laboratory.

To obtain Myo10 full knockout mice (Myo10lox/lox full), the neomycin selection cassette was retained to allow its promoter to interfere with Myo10 expression independently of Cre expression (49) (Fig S1A). We used the following primers to detect the presence of the neomycin selection cassette at the Myo10 locus: 5′-ACA GCC CAT ATC ACT GTG AGA CCC ATT-3′, 5′-ATC GCC TTT TAC CGT GAC G-3′, and 5′-GAG GAT CCA GAC TTG GAC CGG GT-3′.

Mice conditionally deleted for Myo10 in oocytes (Myo10lox/lox Cre) were previously generated from the above strain by removing the neomycin selection cassette after mating with FLPo mice (16) (Fig S1A). To detect Cre-mediated excision at the Myo10 locus after removing the neomycin selection cassette, the following PCR primers were used: for the Cre, 5′-GGC TGG CAG TAA AAA CTA TC-3′ and 5′-GG AAA CAG CAT TGC TGC TAC TT-3′; and for the Myo10, 5′-ACC CTA GCA GTT GTT CAT ACA TCC TAT AAT CT-3′, 5′-GAC TAC ACC ATT CTG AAT GTG CCT GAT CTC-3′, and 5′-GAG TAT CTG CCA TCT TGT CCC TAA AGG TGG-3′.

For the Myo10 full knockout strain, we used Myo10wt/wt, Cre+, Myo10wt/ox, Cre+, Myo10ox/ox, Cre+, and Myo10ox/ox Cre− as control mice, and Myo10ox/ox, Cre-, Myo10ox/ox, Cre−, and Myo10ox/ox, Cre− as Myo10−/− full mice. For the Myo10 oocyte knockout strain, we used Myo10wt/lox, Cre+, Myo10wt/lox, Cre-, Myo10lox/lox, Cre− as control mice, and Myo10lox/lox, Cre-, and Myo10lox/lox, Cre− as Myo10−/− oo mice.

Immunoblotting

Immunoblotting was performed as in Reference 14 on extracts from Myo10wt/lox, Cre+, Myo10wt/lox, Cre−, and Myo10lox/lox, Cre− mice from the Myo10 full knockout strain. Whole liver, whole spleen, and one kidney per mouse were extracted, weighed, and immediately added to 1× Laemml sample buffer (50 mM Tris, pH 7.4, 2% SDS, 6% glycerol, 0.1 M DTT, and 0.25% bromophenol blue) at 1 ml/20 mg wet tissue weight and boiled for 15 min. Samples were crushed with a glass potter to dissociate the tissue, and the lysates were centrifuged at 20,000g for 30 min at room temperature. The pellets were discarded, and the supernatants were stored at −80°C. Final loading volumes were adjusted for immunoblot based on quantification of the actin band from preliminary tests so that the same protein amount is deposited in all the wells. Samples were boiled for 1 min and subjected to SDS-PAGE using a 4–12% Mini-PROTEAN TGX Stain-Free Protein Gel (Ref. 4568085; Bio-Rad). After SDS-PAGE, lysates were transferred to Immobilon-P PVDF Membrane (Ref. IPV00010; Sigma-Aldrich) at room temperature in transfer buffer (25 mM Tris, 192 mM glycine, 0.3% SDS, and 20% methanol) using Trans-Blot SD Semi-Dry Transfer Cell (Bio-Rad) set at 100 mA constant power with a volt limit of 25 V for 45 min. Before transfer, the PVDF membrane was wetted in methanol, rinsed in distilled water, and equilibrated for 10 min in transfer buffer. After the transfer, the membrane was washed in 50 mM Tris, 150 mM NaCl, and 0.05% Tween-20 (TBST thereafter) and blocked for 1 h in TBST containing 5% dried milk powder. The membrane was washed in TBST and incubated for 1 h in TBST containing 4% dried milk and rabbit anti-MYO10 (1:300, Ref. HPA024223; Sigma-Aldrich) as primary antibodies. The membrane was washed in TBST, blocked for 1 h in TBST containing 5% dried milk powder, rinsed in TBST, and incubated for 45 min in TBST containing 4% dried milk and donkey anti-rabbit IgG HRP-linked (1:10,000, Ref. NA934; Amersham) as secondary antibodies. The membrane was washed in TBST and incubated for 5 min in SuperSignal West Femto Maximum Sensitivity Substrate (Ref. 34095; Thermo Fisher Scientific). Protein expression
was revealed using a chemiluminescence image analyzer (FUSION FX; Vilber, 20-s illumination).

The membrane was then rinsed in TBST and stripped for 10 min in Restore Western Blot Stripping Buffer (Ref. 21059; Life Technologies). The membrane was washed in TBST and incubated for 45 min in TBST containing 4% dried milk and HRP mouse anti-beta-actin AC-15 (1:25,000, Ref. ab49900; Abcam). The membrane was washed in TBST and incubated for 1 min in ECL Western Blotting Substrate (Ref. 32109; Thermo Fisher Scientific). Protein expression was revealed using a chemiluminescence image analyzer (FUSION FX; Vilber, 20-s illumination).

Whole ovary immunostaining and clearing

Ovaries collected from five control mice (Myo10wt/wt, Cre+), between 9 and 12 wk old and five Myo10−/− full mice (Myo10 −/−, Cre+), between 9 and 12 wk old) were fixed in 1X PBS and 4% paraformaldehyde for 4 h with agitation at room temperature, and subsequently washed in 1X PBS where they were stored at 4°C. Bleaching immunostaining, and clearing were performed as in Reference 20. Ovaries were first dehydrated in increasing concentrations of methanol (1X PBS, methanol 50% and 80%, and 100% methanol twice) for 1 h each time with agitation at room temperature. Ovaries were then bleached in 100% methanol + 6% hydrogen peroxide overnight at 4°C and rehydrated in decreasing concentrations of methanol (100% twice, 1X PBS, and methanol 80% and 50%) for 1 h each time with agitation at room temperature. Ovaries were blocked and permeabilized in 1X PBS, 0.2% gelatin, and 0.5% Triton X-100 (PBGST thereafter) for 4 d with rotation at room temperature. Antibody incubations were performed in PBGST + 10X saponin (10 mg/ml) at 37°C with rotation at 70g for 1 wk for primary antibodies and overnight for secondary antibodies. Washes between antibody incubations and after secondary antibody incubation were done in PBGST for 1 d with rotation at room temperature. We used rabbit anti-MYO10 (1:1,000, Ref. HPA024222; Sigma-Aldrich) and rat anti-MECA-32 (1:500, Ref. 355063; BD Pharmingen, data not shown) as primary antibodies and Cy3-conjugated donkey anti-rabbit IgG (1:500, Ref. 711-165-152; Jackson ImmunoResearch) and Alexa Fluor 790-conjugated donkey anti-rat IgG (1:250, Ref. 712-655-153; Jackson ImmunoResearch) as secondary antibodies (data not shown). Incubation with TO-PRO-3 (1:100, Ref. T3605; Thermo Fischer Scientific) was performed along with secondary antibody incubation to stain nucleic acid. Ovaries were embedded in 1X TAE and 1.5% agarose, cleared using an adapted 3DISCO clearing protocol with rotation at 12g, and protected from light at room temperature. Ovaries were dehydrated in H2O and 50% tetrahydrofuran (THF anhydrous with 250 ppm butylated hydroxytoluene inhibitor, Ref.186562; Sigma-Aldrich) overnight, and in H2O, 80% THF, and 100% THF twice for 1 h 30 min each time. Ovaries were then incubated in dichloromethane (DCM, Ref.270997; Sigma-Aldrich) for 30 min for delipidation and cleared overnight in benzyl ether (DBE, Ref.108014; Sigma-Aldrich).

Follicle/oocyte collection and culture

Preantral follicles, cumulus-oocyte complexes, and oocytes isolated from follicular cells were collected from 2- to 5-mo-old mice in M2 + BSA medium supplemented with 1 μM milrinone (50) to maintain oocytes arrested in prophase I. Oocytes were released from follicular cells by repeated pipette aspirations. Meiosis resumption was induced by transferring oocytes to M2 + BSA medium free of milrinone. Culture and live imaging were performed at 37°C under oil. To assess the rate of polar body extrusion, oocytes were cultured overnight in M2 + BSA medium in Heracell 150i CO2 Incubator (Thermo Fisher Scientific) and analyzed the next day for polar body extrusion.

Embryo recovery and culture

Embryos were isolated from superovulated female mice (Myo10fl/fl, Cre− and Myo10−/−, Cre− as Myo10−/− full mice, and Myo10fl/fl, Cre+ and Myo10−/−, Cre+ as control mice) mated with male mice (Myo10−/−, Cre−). Superovulation of female mice was induced by intraperitoneal injection of 5 IU pregnant mare's serum gonadotropin (Ceva, Syncro-part), followed by intraperitoneal injection of 5 IU human chorionic gonadotropin (MSD Animal Health, Chorulon) 44–48 h later. Embryos were recovered at E0.5 from plugged females by opening the ampulla followed by a brief treatment with 37°C 0.3 mg/ml hyaluronidase (H4272-30MG; Sigma-Aldrich) and washing in 37°C FHM. When ampulla was not present, embryos were recovered by flushing oviducts with 37°C FHM (MR-122-D; Millipore) using a modified syringe (1400 LL 23; Acuf orm). Embryos were handled using an aspirator tube (A5177-5EA; Sigma-Aldrich) equipped with a glass pipette pulled from glass micropipettes (Blaubrand intraMark or Warner Instruments). Embryos were placed in KSOM (MR-107-D; Millipore) supplemented with 0.1% BSA (A3311; Sigma-Aldrich) in 10 ml droplets covered in mineral oil (M8410; Sigma-Aldrich). Embryos were cultured in an incubator under a humidified atmosphere supplemented with 5% CO2 at 37°C for 5 d. Embryos were scored for survival and embryonic stage from E0.5 to E4.5.

Fluorescent probes for live imaging

Oocytes were incubated for 1 h in M2 + BSA, milrinone medium supplemented with 100 nM SiR-actin (Ref. SC006; Spirochrome) to label F-actin, 5 μg/ml FM 1–43 (Ref. T33556; Thermo Fisher Scientific) to label membranes, or 100 nM SiR-DNA (Ref. SC007; Spirochrome) to label DNA. Oocytes were incubated in M2 + BSA medium supplemented with 100 nM SiR-tub (Ref. SC006; Spirochrome) at least 1 h before the start of spinning disk videos and throughout video acquisitions to label microtubules. Embryos at E1.5 were incubated for 15 min in M2 + BSA medium supplemented with 5 ng/ml Hoechst (Ref. H3570; Invitrogen) to label DNA.

In vitro cRNA synthesis and oocyte microinjection

The pRN3–Histone (H2B)–GFP plasmid (51) was linearized with the Sfi restriction enzyme and in vitro–transcribed using mMESSAGE mMACHINE T3 Transcription Kit (Ref. AM1348; Ambion). cRNAs were purified using RNeasy Mini Kit (Ref. 74104; QIAGEN), centrifuged at 4°C, and microinjected into prophase I–arrested oocytes with a Femtojet micropinector (Eppendorf). Oocytes were maintained in prophase I for 2 h to express cRNAs before meiosis resumption.
Analysis of cumulus–oocyte coupling

The oocyte of each cumulus–oocyte complex was microinjected with 10% Lucifer Yellow (Ref. L453; Thermo Fisher Scientific) with a Femtotjet microinjector (Eppendorf). The complexes were incubated for 30 min to allow dye transfer to follicular cells and then examined using spinning disk microscopy. To block gap junction transmission, some complexes were incubated in 150 μM CBX (Ref. C4790; Sigma-Aldrich) for 30 min before, during, and after microinjection. Culture before, during, and after microinjection was performed in M2 + BSA medium supplemented with 1 μM milrinone and 100 nM Sir-actin.

AFM measurements

After removing the cumulus cells, oocytes were immobilized on an electronic microscopy hexagonal nickel grid (Ref. DT300H-Ni, Ø 3.05 mm, thickness 18 μm, hole 73 μm; Gilder) attached to the surface of a petri dish in M2 + BSA medium supplemented with 1 μM milrinone. AFM-indentation measurements were performed using a Nanowizard IV AFM (JPK Instruments) coupled to a widefield microscope (Zeiss Axio Observer with Hamamatsu sCMOS Flash 4.0 Camera). The contact mode was used. MLCT-C tips (Bruker, with silicon nitride cantilevers) with a 17° side angle, a 5-μm average height pyramidal tip, and a 20-μm tip radius were used. For all experiments, before measurements, the spring constant and the sensitivity were calibrated using the contact-based method in the petri dish in Milli-Q water. The spring constant was in a range of 0.05–0.015 N/m. Measurements were performed on a squared grid of 5 μm × 5 μm at the center of the oocyte. For each oocyte, 16 force–displacement curves were acquired, with a 1 μm/s approach velocity and a 0.5 nN set point. The measurement takes no longer than 5 min for each oocyte. With JPK Data Processing software, the force–displacement curves were processed by correcting the piezo displacement curves to obtain apparent Young’s modulus (52).

Immunostaining and global transcription detection

Preantral follicles, cumulus–oocyte complexes, and oocytes isolated from follicular cells were immunostained as in Reference 16. Incorporation and detection of 5-ethyluridine (EU) into nascent oocyte RNAs were performed similar to Reference 23. For all protocols, follicles, complexes, and oocytes were washed in M2 + PVP medium and adhered to gelatin and polylysine-coated coverslips before fixation (53).

Phalloidin labeling of T2Ps

Preantral follicles and cumulus–oocyte complexes were fixed and permeabilized in 1X PBS, Hapes (100 mM, pH 7), EGTA (50 mM, pH 7), and MgSO4 (10 mM) buffer with 2% Triton X-100 and 0.3% formaldehyde for 30 min at 37°C. Follicles and complexes were then washed in 1X PBS where they were stored overnight or 2 d at 4°C. Follicles and complexes were further permeabilized in 1X PBS and 0.5% Triton X-100 for 10 min and washed in 1X PBS, and 1X PBS and 0.1% Tween-20 (PBSTw thereafter). Blocking and antibody incubations were performed in PBSTw and 3% BSA at room temperature for 30 min for blocking, 1 h 30 min for primary antibody incubation, and 1 h for secondary antibody incubation. Washes between antibody incubations and after secondary antibody incubation were done in PBSTw at room temperature, and an additional wash in 1X PBS was performed before mounting. For spinning disk and OMX microscopy, follicles and complexes were mounted on slide wells filled with VECTASHIELD Antifade Mounting Medium with or without DAPI (Ref. H-1200 and H-1000; Vector Laboratories). We used rabbit anti-MYO10 (1,200, Ref. HPA024223; Sigma-Aldrich) as primary antibodies and Cy3-conjugated donkey anti-rabbit IgG (1:150, Ref. 711-165-152; Jackson Immunoresearch) as secondary antibodies. Incubation with Alexa Fluor 488–conjugated phalloidin (10 U/ml, Ref. A12379; Thermo Fisher Scientific) was performed concurrently with secondary antibody incubation to stain F-actin. For STED microscopy, cumulus–oocyte complexes and oocytes freed from follicular cells were mounted on slide wells filled with Abberior Liquid Antifade Mounting Medium (Ref. MM-2009). We used rabbit anti-MYO10 (1,200, Ref. HPA024223; Sigma-Aldrich) as primary antibodies and Abberior STAR 580–conjugated anti-rabbit (1,200, Ref. ST580-1002) as secondary antibodies. Incubation of Abberior STAR RED–conjugated phalloidin (1,200, Ref. STRED-0100) was performed with secondary antibody to stain F-actin.

Phalloidin labeling of oocyte cytoplasmic F-actin

Oocytes isolated from antral follicles were cleared of the zona pellucida in acid Tyrode’s solution 1 h–1 h 30 min before fixation. Immunostaining was performed using a modified protocol from Reference 33. Oocytes were fixed and permeabilized 6 h after NEBD in 1X PBS, Hapes (100 mM, pH 7), ETGA (50 mM, pH 7), and MgSO4 (10 mM) buffer with 0.2% Triton X-100 and 2% formaldehyde for 30 min at 37°C. Oocytes were then washed in 1X PBS and 0.1% Triton X-100 (PBSTx thereafter) where they were stored overnight at 4°C. The remaining immunostaining protocol is similar to that for phalloidin T2P labeling for spinning disk and OMX microscopy, except that oocytes were directly blocked the day after fixation and blocking and antibody incubations were performed in PBSTx and 3% BSA. Oocytes were mounted on slide wells filled with VECTASHIELD Antifade Mounting Medium with DAPI (Ref. H-1200; Vector Laboratories). We used mouse anti-α-tubulin (1,200, Ref. T8203; Sigma-Aldrich) as primary antibodies and Cy3-conjugated donkey anti-mouse IgG (1:150, Ref. 715-165-151; Jackson Immunoresearch) as secondary antibodies. Incubation with Alexa Fluor 488–conjugated phalloidin (10 U/ml, Ref. A12379; Thermo Fisher Scientific) was performed along with secondary antibody incubation.

EU labeling of nascent oocyte RNAs

As in Reference 23, oocytes arrested in prophase I and isolated from follicular cells were cleaved of the zona pellucida in M2 + BSA, and milrinone medium with 0.4% pronase. EU incorporation and detection were performed using Click-IT RNA Alexa Fluor 488 Imaging Kit (Ref. C10329; Thermo Fisher Scientific). Oocytes were incubated in M2 + BSA, and milrinone medium with or without 0.5 mM EU for 3 h at 37°C and fixed in 1X PBS and 4% pafafomaldehyde for 30 min at 37°C. Oocytes were subsequently washed in 1X PBS where they were stored overnight at 4°C. Oocyte permeabilization and washes were...
done at room temperature in 1X PBS and 0.5% Triton X-100 for 10 min and in 1X PBS, respectively. ELU detection was performed as indicated by the manufacturer, and oocytes were then washed twice in Click-IT reaction rinse buffer and mounted on slide wells filled with VECTASHIELD Antifade Mounting Medium with DAPI (Ref. H-1200; Vector Laboratories).

**Imaging**

**Spinning disk microscopy**

Images were acquired with a Planapo 40×/1.25 NA objective on a Leica DMi6000B microscope enclosed in a thermostatic chamber (Life Imaging Service) equipped with a Retiga 3 CCD camera (QImaging) coupled to a Sutter filter wheel (Roper Scientific) and a Yokogawa CSU-X1-M1 spinning disk. Data were collected with MetaMorph software (Universal Imaging).

**STED microscopy**

STED imaging was performed with a STEDYCON module from Abberior mounted on a Zeiss Axio Observer 7 inverted video microscope coupled to a sCMOS camera (Hamamatsu Flash 4) and a Planapo 100× oil-immersion objective (NA = 1.46) for an effective pixel size of 50 nm (Zeiss).

**OMX microscopy**

Structured illumination (3D-SIM) was performed using a DeltaVision OMX Blaze microscope (GE Healthcare) coupled to a Photometrics Evolve 512 EMCCD camera (Photometrics) and a Planapo 60× oil-immersion objective (NA = 1.4) for an effective pixel size of 80 and 40 nm after reconstruction (Olympus).

**Light-sheet fluorescence microscopy**

The cleared samples are acquired with the ultramicroscope I (LaVision BioTec-Miltenyi) assisted by ImSpector Pro software (LaVision BioTec-Miltenyi). The light sheet is created by two cylindrical lenses using the laser with three specific wavelengths: 561, 640, and 785 nm (Coherent Sapphire Laser, LaVision BioTec-Miltenyi). For all the acquisitions, we used a pinocular stereomicroscope (MVX10, Olympus) with a 2× objective (MVPLAPO, Olympus) adding a protective dipping cap with corrective lens. For detection, we used a Zyla sCMOS camera (2,048 × 2,048 pixels) from Andor Oxford Instruments. The images were in 16-bit. We used for zoom magnification 3.2X (pixel sizes x,y = 1.02 μm). Samples were dived in a quartz tank (LaVision BioTec-Miltenyi) filled with DBE to maintain the clearing. The z-step size between each image was 1 μm with 150-ms time exposure.

**Oocyte RNA extraction**

Total RNA extraction was performed as in Reference 16 from freshly grown oocytes with the zona pellucida. For the RNA-Seq analysis, three biological replicates were performed each containing 10 oocytes from two Myo10lox/flx, Cre” mice (Myo10−/− full oocytes) and two Myo10flx/lox, Cre” mice (Myo10+/− oocytes). For the RT-qPCR analysis, two biological replicates were made containing 30 and 24 oocytes from two Myo10flx/lox, Cre” mice (Myo10−/− full oocytes) and two Myo10flx/lox, Cre” mice (Myo10+/− oocytes). Total RNA extraction was performed using RNAqueous-Micro Total RNA Isolation Kit (Ref. AM1931; Thermo Fisher Scientific). Briefly, oocytes were washed three times in 1X PBS, transferred to lysis buffer (supplemented with 1 μl β-mercaptoethanol for the RNA-Seq analysis), and frozen in liquid nitrogen where they were stored at −80°C for 1 h. After thawing, RNA extraction was performed as described by the manufacturer. RNAs were eluted twice in 10 μl elution buffer and treated with DNasel.

**RNA-seq and data analysis**

**cDNA libraries and RNA-seq**

Library preparation and Illumina sequencing were performed at the Ecole normale supérieure Genomics Core Facility. 5 ng of total RNA was amplified and reverse-transcribed to cDNA using SMART-Seq v4 Ultra Low Input RNA Kit (Ref. 634889; Clontech). Afterward, an average of 500 pg of amplified cDNA was used to prepare the libraries following the Nextera XT DNA Kit (Ref. FC-131-1096; Illumina). Libraries were multiplexed by 6 on high-output flow cells. A 75-bp read sequencing was performed on a NextSeq 500 device (Illumina). A mean of 26 ± 4 million passing Illumina quality filter reads was obtained for each of the six samples. The sequences can be found on the European Nucleotide Archive website, accession number PRJEB56821.

**Bioinformatics analysis**

The analysis was performed using the Galaxy server of the ARTbio Bioinformatics Platform. Sequencing quality control was carried out with FastQC (Babraham Bioinformatics) (Galaxy version 0.72+galaxy1) and MultiQc tools (v1.6). Alignment of reads to the genome of Mus musculus mm10 reference genome was done with Bowtie 2 (v2.3.4.2) (S4) with fast local option for soft clipping of the Nextera transposable adapter sequence. We used the mm10 GTF file (Ensembl GTF Mus_musculus.GRCm38.94.chr) as annotation files. Counting of reads per gene and differential gene expression analysis were performed with featureCounts (v1.63+galaxy2) and DESeq2 (v2.11.40.4), respectively. Transcripts were determined as differentially expressed based on a P-ad (adjusted P-value with false discovery rate correction) of less than 0.05. Of note, 22 deregulated transcripts were annotated as predicted (32.4% of the deregulated transcripts; Table S1) and were predominantly pseudogenes (84.2% of deregulated predicted transcripts). The Galaxy workflows can be found on our GitHub repository (https://github.com/Terretme/Myosin10-full-KO-RNASeq).

**Gene ontology (GO) enrichment analysis**

The analysis was performed with the PANTHER Classification System Overrepresentation Test (Released 20210224), with the annotation dataset GO biological process complete, using Fisher’s exact test with a P-value with a false discovery rate correction threshold of 0.05. For the GO biological process complete (Table S1), the deregulated transcript list from the RNA-Seq analysis was used as the input list and the Mus musculus genome as the reference list. For the graph in Table S1, to provide an overview of the different enriched pathways, the enriched GO terms were selected as follows: for each term with a fold enrichment >10, we browsed their ancestor chart using QuickGO, and selected the highest ancestor
term (only related by the type "A is a B") that is among the enriched terms with a fold enrichment >10, if any. If no ancestor was found with these criteria, the initial enriched term was mounted in a graph. The selected GO terms are marked in blue in Table S1.

**RT–qPCR**

Reverse transcription of the extracted RNAs was performed using iScript Reverse Transcription Supermix (Ref. 1708840; Bio-Rad). We used SsoAdvanced Universal SYBR Green Supermix (Ref. 1725270; Bio-Rad) and CFX-96 (Bio-Rad) for quantitative PCR of cDNAs using the primers shown below. Two technical replicates were made for each condition. mRNA quantity was normalized to Gapdh, and relative levels of mRNA expression were calculated with the 2-ΔΔCt method. The mean and the SEM were normalized to Myo10+/−. The selected genes (see list below) are marked in green in Table S1.

**Selected genes to validate the RNA-Seq by RT-qPCR.**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gapdh</td>
<td>TGGAGAACCTGCCAAGTATG</td>
<td>GGTCTCACTGTAAGCCAAG</td>
</tr>
<tr>
<td>Ahrtb3</td>
<td>AGAGCATGTTGAAAGGACC</td>
<td>CCGATGCAACAAAGTCCC</td>
</tr>
<tr>
<td>Ccnb1</td>
<td>GAGAGTTGCTCTGAGCAAGC</td>
<td>GAGTTGTGCTCATTCAAC</td>
</tr>
<tr>
<td>Eif4e</td>
<td>GAGGTTGCTAACCCAGAGCA</td>
<td>CATAGGCTCAATCCGGTCTT</td>
</tr>
<tr>
<td>Hsp90a1</td>
<td>CGTAGCTGCTGCTATTCCT</td>
<td>CCAGAGTGTCGGATGAATTC</td>
</tr>
<tr>
<td>Myo10</td>
<td>GGAGCGAAAGCATAATAGAAGG</td>
<td>CTTCGGAACAGATGCTG</td>
</tr>
<tr>
<td>Oaz1</td>
<td>CGAGACGCGAGTAATCAAGGC</td>
<td>CTCCGGAAGAGTACAGG</td>
</tr>
<tr>
<td>Oser1</td>
<td>CCACTGCGAGGATTAATAC</td>
<td>GGCCCTCTCACCAGACAG</td>
</tr>
<tr>
<td>Paox</td>
<td>GTGTCAGCGCTTCTGGAGCTCT</td>
<td>GTAGGAACTCGGGTACG</td>
</tr>
<tr>
<td>Pmsa5</td>
<td>CAGGTGCTATGCTGCAGTCCT</td>
<td>ATGAGCAGGACTGTAGGC</td>
</tr>
<tr>
<td>Rpi19</td>
<td>CGCTCGGGAAGAGTAGG</td>
<td>CCTCTCGGTATGCCCATAGC</td>
</tr>
<tr>
<td>Rps4x</td>
<td>GGGCGGTCCCTCCGTGC</td>
<td>GGTAGCAGGAGGACCAAGAC</td>
</tr>
<tr>
<td>Sxbp5f1</td>
<td>GATTTCGAAAGCAGTTCGCC</td>
<td>TGGCAGTAAACGATCACC</td>
</tr>
</tbody>
</table>

**Machine-learning approach for oocyte morphological characterization**

We used our machine-learning pipeline presented in Reference 21 to automatically measure and compare the morphology of Myo10+/− and control oocytes from brightfield images. Oocyte and zona pellucida contours were automatically determined with Oocytor neural networks. This segmentation allowed us to extract the value of 89 features for each oocyte–zona pellucida complex. These features characterize the size, shape, global intensity repartition, local texture, and spatial organization of the oocyte, perivitelline space, and zona pellucida. The complete list and description of the defined features are given in our GitHub repository (https://github.com/gletort/Oocytor). As the features are not independent between themselves, we automatically selected a subset of uncorrelated features (with a threshold of 0.75) as described in Reference 21. We then trained a Random Forest classifier (55) with these features to distinguish between fully grown Myo10+/− and control oocytes cleared of follicular cells. Importantly for our study, the Random Forest algorithm scored each feature according to its importance for classification, based on the Gini index. We checked that the algorithm was able to recognize the oocyte type with a 10-fold cross-validation on our dataset and obtained a classification accuracy of 82% (with a precision rate of 80.9% and a recall rate of 83%). The most important features for classification concerned the composition of the zona pellucida with the presence of vertical linear structures in it (TZP-like structure), the texture of the cytoplasm, the heterogeneity in the thickness of the perivitelline space, and the circularity of both the oocyte and the internal limit of the zona pellucida. Manual inspection of these features suggested that the difference in the texture of the cytoplasm was mainly due to differences in nucleolus focus rather than actual morphological differences, and this feature was not statistically different between Myo10+/− full and control oocytes during meiotic maturation. We therefore decided not to explore this characteristic further.

**Description of the features shown in this study**

**Vertical linear structure in the zona pellucida** To quantify the presence of vertical linear structures resembling TZPs, the area in the segmented zona pellucida was first flattened with the “Flatten” option in Fiji (NIH) to obtain a horizontal rectangle. Linear structures were then detected by measuring and thresholding the “tubeness” of each pixel in the image (with “Tubeness” plugin in Fiji, which is based on the eigenvalues of the Hessian matrix of the image). Finally, only fairly vertical structures were kept by applying Sobel filters on the thresholded images.

**Perivitelline space thickness** Thickness was calculated as the distance between the oocyte shape and the internal limit of the zona pellucida at each angle around the oocyte center. The mean, minimum, maximum, SD (data not shown), and coefficient of variation were then calculated from the distribution of distances.

**Oocyte aspect ratio** Aspect ratio was calculated as the ratio between the maximum Feret diameter (longest distance) of the oocyte shape and the minimum Feret diameter (distance in the perpendicular direction to the maximum Feret diameter). A circle has an aspect ratio of 1, whereas an elongated shape has a higher aspect ratio.

**Image analysis**

For all signal intensity measurements, the image background was subtracted using MetaMorph software.

**Signal intensity of SiR-actin–labeled TZPs**

Measurements were performed on growing and fully grown oocytes cleared of follicular cells and labeled with SiR-actin. For each oocyte, the fifth image among eight Z acquisitions spaced 4 μm apart was selected for measurement, which corresponded to the oocyte equatorial plane. After background subtraction, we drew an ellipsoidal ROI of three pixels (0.34 μm) width along the pericenter of the zona pellucida using Fiji. The mean gray value of SiR-actin signal in the ROI was measured as a readout of TZP density. For
each oocyte, the mean gray value was normalized to that of the control oocyte with the highest mean gray value. The diameter of each oocyte was measured in parallel.

**Signal intensity of EU-labeled RNAs**
Measurements were performed using MetaMorph similar to Reference 23 on growing and fully grown oocytes. After background subtraction and sum of Z projection of the DAPI and EU signal images (images were spaced 1 μm apart and selected for Z projection based on the DAPI signal), the integrated intensity of the DAPI and EU signals was measured in a ROI of 77,577 pixels (999 μm²). The same ROI was used for all measurements and was drawn to encompass the DAPI signal of each oocyte. The integrated intensity of the EU signal was then normalized to that of the DAPI signal. For each oocyte, the relative EU intensity value was normalized to that of the control oocyte with the highest relative EU intensity value.

**Signal intensity of phalloidin-labeled cytoplasmic F-actin**
Measurements were done on oocytes fixed 6 h after NEBD using MetaMorph as in Reference 16. After background subtraction, the integrated intensity of the phalloidin signal was measured in a ROI containing the cytoplasm of the oocyte equatorial plane. For each oocyte, we used the same ROI of 216,290 pixels (2,786 μm²) and the integrated intensity of the phalloidin signal was normalized to that of the control oocyte with the highest value of phalloidin signal intensity.

**Ovarian follicular density**
Cleared ovaries were imaged by light-sheet fluorescence microscopy, and ovarian slices, one every 50 slices, were selected for measurement (slices were spaced 1 μm apart). For each image, the area of the ovarian slice was measured by thresholding the TO-PRO-3 signal using Fiji. Briefly, images were preprocessed with a gamma and a median filter and thresholded with the Otsu dark method. When image thresholding led to more than one ovarian area because of discontinuous signal, only the largest thresholded area was selected. For each image, we counted the number of follicles in the thresholded ovarian slice. Follicles were determined based on a defined outline, with an apparent oocyte surrounded by visible follicular cell layers (this includes preantral and antral follicles). The number of follicles in the thresholded ovarian slice was then normalized to the slice area. The follicular density of the ovary corresponds to the mean of the follicular density of the slices analyzed.

**Timing of polar body extrusion**
The timing of the first polar body extrusion was determined in oocytes monitored by live spinning disk microscopy. Only videos with control oocytes extruding a polar body before 12 h after NEBD were selected.

**Spindle bipolarization and positioning**
Spindle morphogenesis and positioning were assessed in SiR-tub-labeled oocytes monitored during meiotic maturation by live spinning disk microscopy. The timing of spindle bipolarization was assessed by scoring the percentage of oocytes with a bipolarized spindle every hour from 1 to 6 h after NEBD. Spindle interpolar length and central width were measured 7 h after NEBD using MetaMorph. Only spindles parallel to the acquisition plane were measured. Spindle localization was assessed by scoring the percentage of oocytes with a central or off-centered spindle 30 min before anaphase for oocytes that extruded a polar body or up to 12 h after NEBD for those that did not.

**Chromosome alignment**
Metaphase plate height and width were measured as in Reference 16 using MetaMorph in H2B-GFP-expressing oocytes monitored during meiotic maturation by live spinning disk microscopy. We measured the height and width of a ROI rectangle enclosing chromosomes 30 min before anaphase for oocytes that extruded a polar body or 12 h after NEBD for those that did not. Only metaphase plates oriented parallel to the acquisition plane were measured.

**Cumulus–oocyte coupling**
Measurements were performed on cumulus–oocyte complexes when single follicular cells were visible, still connected to the oocyte by TZPs (visible with SiR-actin). The coupling was quantified for each follicular cell by measuring the ratio of Lucifer Yellow fluorescence intensity between the follicular cell and the oocyte 30 min after Lucifer Yellow microinjection into the oocyte. For that, after acquiring the whole cumulus–oocyte complex (25 Z acquisitions spaced 5 μm apart), we selected the plane where a single follicular cell connected to the oocyte via TZPs was visible. We drew an ellipsoidal ROI of 19 pixels width and 16 pixels height (4.3 μm × 3.6 μm) centered on the follicular cell using Fiji. The mean gray value of Lucifer Yellow signal in the ROI was measured. We displaced the ROI to put it in the oocyte and measured the mean gray value of Lucifer Yellow signal in the ROI. The mean gray value of Lucifer Yellow signal in the follicular cell was then divided by the one in the oocyte. This was repeated for all single follicular cells.

**Statistical analysis**
We used GraphPad Prism 7.0a software for statistical analysis of RT–qPCR, oocyte polar body extrusion rate, and image data. The Gaussian distribution of values was tested using a D'Agostino & Pearson normality test. Comparison of means was performed by a two-tailed unpaired t test (with Welch’s correction for unequal variances) or by a two-tailed Mann–Whitney test (for a non-Gaussian distribution of values). For comparison of more than two conditions, an ANOVA test comparing the mean of each column with the mean of every other column when the data followed a Gaussian distribution, and a Kruskal–Wallis test comparing the mean rank of each column with the mean rank of every other column when the data did not were performed. We used two-sided Fisher’s exact tests and a chi-squared test for contingency table analysis. All tests were performed with a confidence interval of 95%, n.s (not significant), P ≥ 0.05, *P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001. For each graph, the error bars indicate the SD or the SEM.

**Ethical statement**
All experimental procedures used for the project have been approved by the Ministry of Agriculture to be conducted in our CIRB
Animal Facility (authorization no. 75-1170). The use of all the genetically modified organisms described in this project has been granted by the DGRI (Direction Générale de la Recherche et de l’Innovation: Agrément OGM; DUO-5291).

Data Availability

For the RNA-Seq, the sequences can be found on the European Nucleotide Archive website, accession number PRJEB56821. The Galaxy workflows of the bioinformatics analysis of the RNA-Seq can be found on our GitHub repository (https://github.com/Terretme/Myosin10-full-KO-RNAseq).

Supplementary Information

Supplementary Information is available at https://doi.org/10.26508/lsa.202301963

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F Crozet: conceptualization, formal analysis, investigation, methodology, and writing—original draft, review, and editing.
G Letort: data curation, software, formal analysis, investigation, methodology, and writing—review and editing.
R Bulteau: formal analysis, investigation, methodology, and writing—review and editing.
C Da Silva: formal analysis, investigation, methodology, and writing—review and editing.
A Eichmuller: formal analysis and investigation.
A-F Tortorelli: formal analysis, investigation, and writing—review and editing.
J Blévinial: formal analysis, investigation, and methodology.
M Belle: formal analysis, investigation, methodology, and writing—review and editing.
J Dumont: investigation, methodology, and writing—review and editing.
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A Dauphin: investigation, methodology, and writing—review and editing.
F Coulpier: investigation, methodology, and writing—review and editing.
A Chédotal: methodology.
J-L Maître: methodology and writing—review and editing.
M-H Verlhac: conceptualization, funding acquisition, and writing—review and editing.
HJ Clarke: funding acquisition and writing—review and editing.
M-E Terret: conceptualization, data curation, formal analysis, supervision, funding acquisition, investigation, methodology, and writing—review and editing.

Conflict of Interest Statement

The authors declare that they have no conflict of interest.

References

Filopodia from somatic cells shape oocyte quality  

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