Introduction

Oocytes arrest at metaphase of the second meiotic division (MetII) before fertilization because of an activity termed cytosstatic factor (CSF; Masui, 2000; Tunquist and Maller, 2003; Jones, 2005). Sperm break this arrest via a Ca$^{2+}$ signal (Jones, 1998; Stricker, 1999; Runft et al., 2002), and in so doing, oocytes complete the second meiotic division before entering the embryonic cell cycles.

CSF activity, a terminology that was first defined several decades ago (Masui and Market, 1971), is now known to constitute an inhibitor of the anaphase-promoting complex/cyclosome (APC; Tunquist and Maller, 2003). The APC is an E3 ubiquitin ligase whose activity is required for the metaphase–anaphase transition to polyubiquitinate key cell cycle proteins, thereby earmarking them for immediate proteolysis through association with its key coactivator cdc20 (Fang et al., 1998; Kramer et al., 1998; Harper et al., 2002; Eytan et al., 2006). The reduced APCcdc20 activity in MetII oocytes prevents the destruction of both M-phase (maturation)–promoting factor (MPF) activity (CDK1/cyclin B1) and cohesin, which holds sister chromatids together (Nixon et al., 2002; Madgwick et al., 2004). Resumption of meiosis in mammalian oocytes is achieved by a sperm-borne phospholipase C activity (Saunders et al., 2002; Knott et al., 2005), which generates an oscillatory Ca$^{2+}$ signal, switching on APCcdc20 (Nixon et al., 2002; Madgwick et al., 2004) through a signaling pathway involving calmodulin-dependent protein kinase II (CamKII; Markoulaki et al., 2003, 2004; Madgwick et al., 2005). This signaling process is conserved and was first demonstrated in frog eggs (Lorca et al., 1993, 1994). Activation of the APC in MetII oocytes induces the destruction of MPF and sister chromatid cohesion through the polyubiquitination of cyclin B1 and securin, respectively (Morgan, 1999; Zachariae and Nasmyth, 1999; Peters, 2002). Loss of cyclin B1 causes a reduction in MPF, and the loss of securin frees separase to act on the kleisin component of cohesin (Zachariae and Nasmyth, 1999; Peters, 2002; Blow and Tanaka, 2005; Nasmyth and Haering, 2005).

Many proteins have been associated with the establishment and/or maintenance of CSF activity. Factors responsible for setting up a second meiotic spindle after completion of meiosis I do not, a priori, have to be the same as those that are responsible for maintaining arrest. Indeed, proteins have been described that are involved in establishing MetII, but not in maintaining arrest once it has been achieved (Tunquist and Maller, 2003).

Emi2 antisense morpholino knockdown during oocyte maturation did not affect polar body (PB) extrusion. However, in interkinesis the central spindle microtubules from meiosis I persisted for a short time, and a MetII spindle failed to assemble. The chromatin in the oocyte quickly decondensed and a nucleus formed. All of these effects were caused by the essential role of Emi2 in stabilizing cyclin B1 after the first PB extrusion because in Emi2 knockdown oocytes a MetII spindle was recovered by Emi2 rescue or by expression of nondegradable cyclin B1 after meiosis I.

Mouse Emi2 is required to enter meiosis II by reestablishing cyclin B1 during interkinesis

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During interkinesis, a metaphase II (MetII) spindle is built immediately after the completion of meiosis I. Oocytes then remain MetII arrested until fertilization. In mouse, we find that early mitotic inhibitor 2 (Emi2), which is an anaphase-promoting complex inhibitor, is involved in both the establishment and the maintenance of MetII arrest. In MetII oocytes, Emi2 needs to be degraded for oocytes to exit meiosis, and such degradation, as visualized by fluorescent protein tagging, occurred tens of minutes ahead of cyclin B1.

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Abbreviations used in this paper: APC, anaphase-promoting complex/cyclosome; Bub, budding uninhibited by benzimidazole; CSF, cytosstatic factor; Emi, early mitotic inhibitor protein; GV, germinal vesicle; GVBD, GV breakdown; Mad, mitotic arrest deficient; MISS, MAPK-interacting and spindle-stabilizing protein; MO, morpholino; MPF, M-phase (maturation)–promoting factor; PB, polar body.

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The mechanism of CSF is most well characterized in the frog, where various groups have firmly defined the c-Mos–MAPK–90-kD ribosomal protein S6 kinase (p90rsk)–budding uninhibited by benzimidazole 1 (Bub1) pathway in establishing CSF activity (Sagata et al., 1989; Abriou et al., 1996; Bhatt and Ferrell, 1999; Gross et al., 1999, 2000; Tunquist et al., 2002). Other activities that are fundamentally involved in the establishment of CSF in frog include cyclin E/Cdk2 (Tunquist et al., 2002) and mitotic arrest deficient 2 (Mad2; Tunquist et al., 2003). However, once established, p90rsk, Mad2, Bub1, and cyclin E/Cdk2 are all dispensable for the maintenance of CSF activity (Bhatt and Ferrell, 1999; Tunquist et al., 2002, 2003).

So, how is CSF activity maintained in the frog? Current evidence suggests it is through early mitotic inhibitor 2 (Emi2)/Emi-related protein 1 (Liu and Maller, 2005; Rauh et al., 2005; Schmidt et al., 2005; Tung et al., 2005; Hansen et al., 2006). Emi2, which acts to inhibit the APC accumulated during oocyte maturation, is present and stable in CSF frog egg extracts, but is rapidly degraded on Ca2+ addition (Schmidt et al., 2005). Degradation of Emi2 is induced by phosphorylation through CamKII (Liu and Maller, 2005; Rauh et al., 2005; Hansen et al., 2006) and, thus, would be predicted to occur ahead of APC activation and cyclin B1 degradation, although this has not been tested.

In mouse oocytes, the mechanism of CSF arrest is less well understood. As the mouse Emi2 homologue appears to have a similar function in maintaining CSF activity (Shoji et al., 2006), it would be logical to predict that the mechanism of CSF establishment is also conserved between frog and mouse. However, this is not so. Oocytes from a triple Rsk (1,2,3) knockout mouse arrest normally at MetII (Dumont et al., 2005), demonstrating that p90rsk is not involved in mouse CSF arrest. Furthermore, using dominant-negative mutants Tsurumi et al., (2004) confirmed that the spindle checkpoint proteins Bub1 and Mad2 are not required for either the establishment or the maintenance of mouse CSF. Although c-Mos is known to be involved in CSF maintenance, it does not appear to be involved in the establishment of CSF arrest because c-Mos+/− oocytes remain at MetII for 2–4 h before extruding a second PB (PB2; Verlhac et al., 1996).

As there are no firm candidates suggested to be involved in the establishment of CSF arrest in mouse, we have investigated the potential of mouse Emi2 in this process. We find that Emi2 has activity consistent with CSF, and its degradation in real time is ahead of any change in cyclin B1. We also demonstrate that a function of Emi2 is in restabilizing cyclin B1 upon exit from anaphase I and that, in this study, it contributes to the formation a bipolar spindle. Oocytes matured without Emi2 do not assemble MetII spindles, and, in the absence of cyclin B1, eventually decondense their chromatin.

**Results**

Stable Emi2 is rapidly degraded by Ca2+

To measure real-time changes in Emi2 levels in oocytes we generated cRNA to mouse Emi2 coupled to Venus fluorescent protein (Emi2-V), which is a yellow variant of GFP. MetII mouse oocytes were microinjected with this construct at a dose of either 0.15 or 0.5 pg, and then cultured for a few hours to allow for Emi2 expression. We Western blotted oocytes expressing Emi2-V with a polyclonal antibody against Emi2 to determine the amount of Emi2-V expression relative to endogenous protein. More than one band was detected on oocyte blots using this antibody; however, one band at ~85 kD migrated at the same molecular mass as in vitro–translated Emi2 (and this band was later knocked down by Emi2 morpholino [MO]). At the 0.15-pg dose, Emi2-V levels were less than endogenous protein after 2 h, whereas the 0.5-pg dose was expressed to about the same level as endogenous protein (Fig. 1 A).

Emi2-V was very stable in MetII-arrested oocytes, but became rapidly unstable when cytosolic Ca2+ increased. With either 0.15 pg (Fig. 1 B) or 0.5 pg (not depicted) injections, we observed no degradation of Emi2-V in MetII oocytes after blocking further synthesis by washing into cycloheximide-containing media. However, when this experiment was repeated, but by washing into Sr2+-containing media to induce spermlike Ca2+ spiking (Bos-Mikich et al., 1997), we observed dose-dependent effects on Emi2-V. At the higher 0.5-pg dose, we observed no loss in Emi2-V signal; instead, Emi2-V levels steadily increased...
Figure 2. *Emi2* is degraded ahead of cyclin *B1*. (A) Western blot (WB) of oocytes for *Emi2* (top; *n* = 100 oocytes per lane) and cyclin *B1* (*n* = 30 oocytes per lane), and corresponding membranes stained with Coomassie brilliant blue (CB) to show equivalent loading. Oocytes are activated by washing into Sr²⁺ media, and samples from a pool of oocytes were removed at various time points and probed for either cyclin *B1* or *Emi2* by Western blotting (Fig. 2 A). In these experiments, it was apparent that loss in *Emi2* protein was rapid and complete by 30 min (in agreement with the rapid loss of *Emi-V* shown in Fig. 1 D). Interestingly, *Emi2* levels increased again in pronucleate embryos (6-h time point), which is in agreement with Shoji et al. (2006) and suggests that it may have a further mitotic function (see Discussion). Similar to *Emi2*, we observed the loss of cyclin *B1* at 30 min, as well as increased levels in pronucleate embryos (Fig. 2 A), which is consistent with our observation that the APC is switched off at this time (Nixon et al., 2002).

Because of the numbers of oocytes needed for Western blots and the tens of minutes of asynchrony in timing at which Ca²⁺ spiking starts with Sr²⁺ media, (Madgwick et al., 2004), we could not reproducibly resolve *Emi2* degradation ahead of cyclin *B1* by Western blotting groups of oocytes. Therefore, to examine with more accuracy the immediate degradation profiles of cyclin *B1* and *Emi2*, we decided to measure their simultaneous degradation in the same oocyte. Cyclin *B1* was coupled to Cerulean fluorescent protein (cyclin *B1*-C), which is a cyan variant of GFP. There was no overlap in Venus and Cerulean signals, showing that both *Emi2*-V and cyclin *B1*-C, with appropriate filters, could be imaged simultaneously in the same oocyte. Cyclin *B1* was coupled to Cerulean fluorescent protein (cyclin *B1*-C), which is a cyan variant of GFP. There was no overlap in Venus and Cerulean signals, showing that both *Emi2*-V and cyclin *B1*-C, with appropriate filters, could be imaged simultaneously in the same oocyte. Cyclin *B1* was coupled to Cerulean fluorescent protein (cyclin *B1*-C), which is a cyan variant of GFP. There was no overlap in Venus and Cerulean signals, showing that both *Emi2*-V and cyclin *B1*-C, with appropriate filters, could be imaged simultaneously in the same oocyte.

![Figure 2](image_url)
activity. However, these coexpression studies revealed that Emi2 degradation began tens of minutes before that of cyclin B1 (Fig. 2 C). Emi2-V levels were degraded by at least 50% before the start of cyclin B1-C degradation ($n = 7$).

Emi2 degradation is independent of a spindle checkpoint

Degradation of cyclin B1 is dependent on APC$^{cdk20}$ activity, and in mouse oocytes it can be blocked by the induction of a spindle checkpoint (Nixon et al., 2002; Madgwick et al., 2005). In contrast, Emi2 degradation should be checkpoint-independent because its degradation is independent of APC$^{cdk20}$ involvement. Incubation of Emi2- and cyclin B1-expressing oocytes with the spindle poison nocodazole blocks mouse oocytes from exiting MetII arrest when washed into Sr$^{2+}$ media. As expected, the addition of nocodazole completely stabilized cyclin B1 levels ($n = 15/15$; Fig. 3 A). However, nocodazole had no effect on the rate of Emi2-V degradation ($n = 12/12$; Fig. 3 B). Such an observation is consistent with cyclin B1, but not Emi2 degradation, being dependent on the APC.

Therefore, in summary, we have obtained data that are entirely consistent with a model of MetII arrest achieved by Emi2-mediated inhibition of cyclin B1 degradation. Also, the Emi2-V construct generated is a physiologically active, useful tool for both establishing CSF activity and measuring its loss in real time after a Ca$^{2+}$ signal.

Table I. Emi2 MOs used in this study

| MO     | Sequence                                      |
|--------|-----------------------------------------------|
| Mouse Emi2 | A GCC AGC CAC AGA GCA GGA AGC AAT            |
| Emi2 MO | ATT GCT TCC TGC TCT GTG GCT GCC T             |
| 5mp MO | ATT CCT TGC TGC TGT GTG CCT GCC T             |
| Inv MO | TCG GTG GTG TGC TCC TCC TTC GTT A             |

Emi2 MO was designed to target the 5’UTR of mouse Emi2 (underlined). A 5mp MO and an Inv MO were used as control MOs.

Oocytes with Emi2 knockdown extrude a PB, but do not MetII arrest

Emi2 levels are low in both Xenopus laevis and mouse oocytes before they are matured. This would be predicted, as high Emi2 levels during maturation may be deleterious and arrest oocytes at MetI. The increased Emi2 expression during oocyte maturation makes it highly likely that Emi2 expression can be knocked out by an antisense approach. Therefore, we designed an antisense MO to the 5’UTR immediately adjacent to the start codon of mouse Emi2 (Emi2 MO) and used an additional two MO’s as controls (Table I); a 5-base mispair MO (5mp-MO), in which five bases have been altered from the complementary sequence, and an inverted MO (Inv-MO).

To explore the role of Emi2 in the establishment of MetII arrest, we injected Emi2 MO into germinal vesicle (GV) oocytes, which were then matured in vitro. Oocytes were held at the GV stage in milrinone-containing media for 2 h after MO injection; they were then released from GV arrest and allowed to mature for 16 h. Blotting of GV-stage oocytes, Emi2 MO–matured oocytes, and uninjected control matured oocytes demonstrated that Emi2 protein levels increase between GV and MetII stage, and confirmed the Emi2 knockdown in Emi2 MO–injected oocytes (Fig. 4 A).

During maturation, oocytes were scored for the morphological events of oocyte maturation, which are GV breakdown (GVBD) and PB1 extrusion (Fig. 4 B). Both GVBD (Fig. 4 C) and PB1 extrusion (Fig. 4 D) occurred with normal timings in Emi2 MO–injected oocytes. However, after maturation, we observed marked differences in the morphology of control oocytes and those injected with Emi2 MO. When control oocytes (uninjected, 5mp-MO, and Inv-MO–injected) were stained for chromatin, oocytes were morphologically normal. They had a PB1 containing chromatin, which was produced on completion of the first meiotic division, and a fully formed MetII spindle (100%; uninjected oocytes, $n = 60$; 5mp-MO, $n = 32$; Inv-MO, $n = 25$; Fig. 5 A). However, although oocytes injected with Emi2 MO did have a PB1 containing chromatin, the chromatin in the oocyte was decondensed inside a nucleus (93%; $n = 110$; Fig. 5 A).

Despite the lack of effect of a 5mp-MO, it remained possible that we had been extremely unlucky in the MO design, such that the observed effects of Emi2 MO were caused by its ability to block the expression of an unrelated protein involved in MetII arrest. We thought this unlikely, given that a similar morphology of Emi2 knockdown oocytes has been reported recently using a double-stranded RNAi approach (Shoji et al., 2006).
However, we decided it was important to confirm the specificity by a rescue to the control phenotype in Emi2-MO–injected oocytes by expression of exogenous Emi2. To recover Emi2, Emi2-V cRNA was microinjected into oocytes 2 h after microinjection of Emi2 MO and immediately before release from GV arrest. This rescue is made possible because Emi2-V lacks the 5′ UTR recognized by the MO.

Injection of Emi2-V cRNA alone into GV oocytes that were matured induced a MetI arrest (n = 40; Fig. 5 A), which is consistent with Emi2-V having CSF activity and the need to keep Emi2 levels low until completion of the first meiotic division. Importantly, Emi2-V cRNA expression could rescue the effects of MO knockdown. Rescue oocytes could progress through meiosis I and arrest as controls with a fully formed MetII spindle. A high dose of Emi2-V (0.5 pg; n = 45) rescued the Emi2 MO phenotype, with an equal mix of either MetI or MetII arrest, whereas oocytes microinjected with the lower Emi2 dose showed a rescue with a predominantly MetII arrest (n = 45; Fig. 5 A).

Lack of Emi2 results in a failure to assemble a metaphase II spindle

By scoring oocytes for decondensed chromatin at only one time point (16 h), we were unable to pinpoint at which stage of meiosis oocytes undergo chromatin decondensation. However, we noted that the vast majority of oocytes with decondensed chromatin extruded only a single PB and contained only one nucleus (88%; n = 102). This observation suggests either decondensation of chromatin occurred after anaphase I, such that oocytes did not form a MetII spindle (Fig. 5 B, i), or, alternatively, that after establishment of a MetII spindle there was no sister chromatid disjunction before decondensation (Fig. 5 B, ii).

We determined whether Emi2 MO knockdown oocytes were able to build a MetII bipolar spindle by staining oocytes for both chromatin and tubulin at various times after PB1 extrusion. Oocytes were fixed at 0.5, 1, and 2 h after PB1 extrusion. In control uninjected oocytes, central spindle microtubules were observed at 0.5 h after PB1 extrusion (Fig. 6 A), and over the next 1.5 h a MetII spindle formed, such that by 2 h after PB1 a fully formed MetII spindle was found in all oocytes (Fig. 6 B). In Emi2-MO–injected oocytes, at 0.5 h after PB1 there was no difference from controls, with central spindle microtubules evident between the chromatin in the ooplasm and the PB1 (Fig. 6 A). However, in contrast to control oocytes, in Emi2-MO–injected oocytes at both the 1 and the 2 h time points we observed no MetII spindle; instead, the chromatin had remained essentially unaltered from the time of PB1 extrusion. The residual central spindle microtubules remained between the chromatin in the oocyte and the PB, and by 2 h the chromatin appeared to be beginning to decondense (Fig. 6 B). As the chromatin started to decondense, spindle microtubule structure was lost, which is consistent with entry into interphase. By 6 h, all Emi2-MO–injected oocytes had a single nucleus containing fully decondensed chromatin. We did not assess if these oocytes underwent S-phase; however, they showed no obvious signs of degeneration over the 6-h time period from PB1 extrusion. Thus, we failed to observe the formation of a bipolar spindle in any Emi2-MO–treated oocytes; instead, oocytes underwent full chromatin decondensation.

Emi2 stabilizes cyclin B1 after PB1 extrusion

Because the formation of a MetII spindle requires an increase in the levels of CDK1 activity after PB1 extrusion, we reasoned that in the absence of Emi2, levels of CDK1’s regulatory partner...
cyclin B1 may not be reestablished. To test this directly, Emi2 MO or control GV oocytes were microinjected with cyclin B1-V cRNA and matured in vitro. Cyclin B1-V levels were monitored in real time during maturation, along with brightfield microscopy to assess progression through meiosis I. As previously reported (Hyslop et al., 2004), in control oocytes there was a period of cyclin B1 degradation that lasted a few hours and was terminated on PB1 extrusion (Fig. 7 A). Immediately after PB1 extrusion in these oocytes, cyclin B1 levels rose to 24.0 ± 1.1% (n = 14) of the peak level before PB1 extrusion (Fig. 7 B). In Emi2-MO–injected oocytes, there was no difference from controls in the timing of initiation or in the rate of cyclin B1-V degradation during meiosis I, which is in agreement with the observed lack of effect of the MO on the timing of PB1 (Fig. 4 D). However, in these Emi2 MO oocytes there was no reelevation of cyclin B1 after PB1 extrusion; cyclin B1-V levels increased to just 1.6 ± 0.3% (n = 15) of the peak level before PB1 extrusion (Fig. 7 B).

In keeping with the real-time effects on exogenous cyclin B1-V expression, we also observed a loss of endogenous cyclin B1 after Emi2 knockdown. In vitro–matured oocytes injected with Emi2 MO at the GV stage showed much lower cyclin B1 levels than control in vitro–matured oocytes (Fig. 7 C). Furthermore, cyclin B1 levels in individual oocytes were measured by immunofluorescence at 16 h after release from GV arrest in control and Emi2-MO–injected oocytes (Fig. 7, D and E). We found that cyclin B1 immunofluorescence levels in Emi2 MO–treated oocytes was about one third that of control oocytes (Fig. 7 E), suggesting a relatively uniform level of knockdown.

Nondegradable cyclin B1 and Mad2 rescue a bipolar spindle in Emi2 knockdown oocytes

Because the formation of a MetII spindle requires an elevation in the levels of cyclin B1, we reasoned that the Emi2 MO spindle defect may be recovered by the addition of cRNA to nondegradable cyclin B1. Furthermore, it should also be rescued by inhibiting the APC (Madgwick et al., 2004). Cyclin B1 with deletion of 90 N-terminal amino acids (Δ90 cyclin B1) lacks a D-box and is therefore not a substrate of the APC (Glotzer et al., 1991; Madgwick et al., 2004). Emi2-MO–injected oocytes...
either received no further treatment or were microinjected with 1.5 pg ∆90 cyclin B1 or Mad2 cRNA within 15 min of PB1 extrusion. This injection had to be done immediately after PB1 extrusion because nondegradable cyclin B1 addition to oocytes before PB1, as predicted by the sustained MPF activity, blocks PB extrusion (Herbert et al., 2003). A few hours of ∆90 cyclin B1 cRNA expression generates cyclin B1 at a similar level to endogenous cyclin B1 in a MetII oocyte (Madgwick et al., 2004). This dose of Mad2 causes a metaphase I arrest in maturing oocytes (Homer et al., 2005) and prevents completion of meiosis in MetII oocytes (Madgwick et al., 2005). At 2 h after PB1 extrusion, all oocytes were fixed, stained, and scored for the presence of a bipolar spindle (Fig. 8 A). As observed in Fig. 6 B, oocytes microinjected with Emi2 MO gave decondensing chromatin with a lack of any bipolar spindle structure. However, as with uninjected oocytes, we observed a fully formed bipolar MetII spindle in oocytes microinjected with Emi2 MO and either ∆90 cyclin B1 cRNA or Mad2 (Fig. 8 B).

Discussion

We have demonstrated in live oocytes and in real time that upon a Ca^{2+} signal, Emi2 degradation occurs ahead of cyclin B1, independent of APC inhibition. The current study confirmed

![Figure 7. Cyclin B1 levels remain low after PB extrusion in Emi2 MO—matured oocytes. (A) Venus fluorescence levels in maturing oocytes microinjected with cyclin B1-V cRNA (control) or cyclin B1-V cRNA and Emi2 MO. PB1 extrusion was observed at the times indicated. Note the similar profile of cyclin B1-V degradation in both oocytes, but the failure of cyclin B1-V levels to rise in the oocyte injected with Emi2 MO after PB1. (B) Quantification of the elevation in cyclin B1-V levels after PB1 extrusion in the control oocytes (n = 14) and Emi2 MO oocytes (n = 13) used in A. Endogenous cyclin B1 levels in matured oocytes detected by Western blotting (C) or immunofluorescence (D and E). (C) A Western blot (WB) for cyclin B1 in control matured and oocytes matured after microinjection of Emi2 MO at the GV stage (n = 40), and the corresponding membrane stained with Coomassie brilliant blue (CB) to demonstrate equivalent loading. In D, representative images of cyclin B1 levels in matured oocytes (control) and in matured oocytes microinjected at the GV stage with Emi2 MO detected using a TRITC-conjugated secondary antibody. Mean fluorescence readings are marked for both oocytes on the 8-bit scale bar. (E) Quantification of cyclin B1 immunofluorescence levels in control (n = 25) and Emi2 MO—microinjected (n = 32) oocytes from D, relative to uninjected controls. A is representative of four independent experiments whose data is pooled in B. (C) The Western blot was repeated once. D is representative of two independent experiments whose data is pooled in E.

![Figure 8. Emi2 MO phenotype recovery using ∆90 cyclin B1 or Mad2. (A) Percentage maturation rates in in vitro—matured oocytes after no treatment (n = 12), or microinjection of Emi2 MO at the GV stage with or without further microinjection of ∆90 cyclin B1 or Mad2 after PB1 extrusion (Emi2 MO only, n = 20; with ∆90 cyclin B1, n = 25; or with Mad2, n = 35). Oocytes were fixed and stained 2 h after PB1 extrusion and scored for a bipolar spindle. (B) Tubulin (red) and chromatin (blue) staining in an oocyte microinjected with Emi2 MO at the GV stage and then ∆90 cyclin B1 cRNA 15 min after PB1 extrusion. Image is representative of 20/25 oocytes. Bar, 20 μm. Data is pooled from two independent experiments.}
that Emi2 constitutes CSF activity in mouse MetII oocytes and is responsible for the maintenance of MetII arrest (Shoji et al., 2006). More importantly, we find that Emi2 is necessary and sufficient in stabilizing cyclin B1 levels at the completion of meiosis I, and such stabilization is essential in establishing a bipolar spindle.

By definition, CSF activity must accumulate during oocyte maturation, inhibit APCcdc20 activity at MetII, and be inactivated on a Ca2+ signal at fertilization. Although only recently identified, the F-box protein Emi2 is very likely to constitute CSF in frog eggs (Liu and Maller, 2005; Rauh et al., 2005; Schmidt et al., 2005; Tung et al., 2005; Hansen et al., 2006). Emi2 inhibits the ubiquitin ligase activity of the APCcdc20 in vitro, and in keeping with the nature of CSF, it accumulates during oocyte maturation, is present in CSF extracts, and is rapidly degraded by Ca2+ (Schmidt et al., 2005).

**Maintenance of MetII arrest**

We tested the characteristics of the mouse Emi2 homologue in MetII mouse oocytes. This achieved two goals. First, it allowed us to demonstrate that Venus-tagged Emi2 is physiologically active, and second, it provided us with data relating to the timing of Emi2 degradation on activation. Excess *X. laevis* Emi2 protein prevents Ca2+-induced CSF release in egg extracts (Schmidt et al., 2005); similarly, our MetII-arrested mouse oocytes were refractory to Ca2+ through modest rises in Emi2-V. A lower Emi2-V dose, less than endogenous protein, did not block escape from MetII arrest and allowed us to demonstrate that Emi2 is degraded ahead of cyclin B1. If Emi2 inhibits the APC in MetII arrest, it follows that Emi2 destruction must occur before the APC’s accelerated degradation of cyclin B1. Indeed, when both were simultaneously imaged in an oocyte after a Ca2+ rise, Emi2 was degraded by at least 50% before cyclin B1. An anti-Emi2 antibody also revealed Emi2 protein in pronucleate embryos, which is in agreement with the observations of Shoji et al. (2006), demonstrating that Emi2 reaccumulates in one-cell embryos and, thus, interestingly, may have a mitotic function. We have previously reported on Ca2+ spiking during the first mitotic division of mouse embryos (Kono et al., 1996), suggesting that in this study Emi2 may be degraded by a similar Ca2+-dependent process.

In MetII-arrested mouse oocytes, the APC is already active, but the addition of a Ca2+ signal speeds up cyclin B1 degradation sixfold through an increase in APC activity (Nixon et al., 2002). In mouse, the high levels of cyclin B1 required for MetII arrest are therefore achieved through equilibrium between synthesis and degradation (Kubiak et al., 1993). Induction of a spindle-assembly checkpoint by nocodazole prevents cyclin B1 degradation, with or without the presence of a Ca2+ signal (Nixon et al., 2002), showing that in mouse oocytes APC activity can be held in check by both CSF and a spindle-assembly checkpoint. In mouse, the spindle-assembly checkpoint is distinct from CSF (Tsurumi et al., 2004). In contrast to cyclin B1, we found little evidence of Emi2 protein turnover in MetII-arrested oocytes, suggesting that Emi2 itself is not a substrate of the APC. Indeed, upon a Ca2+ signal, destruction of *X. laevis* Emi2 has been shown to occur via the phosphorylation of a known SCFβ-TRCP (Skp1-Cullin-F-boxβ-TRCP) motif (DSGX,S) and not the APC (Rauh et al., 2005; Tung et al., 2005).

**Establishment of MetII arrest**

We used a MO antisense oligonucleotide to ablate the Emi2 rise during mouse oocyte maturation and confirmed a knockdown by Western blotting. By blocking the accumulation of Emi2, we have demonstrated that it is essential for stabilizing cyclin B1 after MetII during interkinesis, and therefore essential for the formation of a MetII spindle. Because of the known inhibitory action of Emi2 on APC activity, we interpret this observation to mean that in the absence of Emi2, the APC remains active, and thus cyclin B1 is continuously degraded. In agreement with this is our observation that nondegradable cyclin B1, but not full-length cyclin B1, rescues a MetII arrest after Emi2 knockdown (Fig. 7 vs. Fig. 8), as does the APCcdc20 inhibitor Mad2. The current data have not ruled out some effect of Emi2 on cyclin B1 synthesis. However, if so, Emi2 would also be able to negatively regulate expression of the microinjected cyclin B1 cRNA (Fig. 7). This lacks the cyclin B1 gene 3′ UTR, which is important for regulating translation through polyadenylation (Tay et al., 2000). Therefore, we would argue that Emi2 is most likely affecting the rate of cyclin B1 degradation, rather than synthesis.

The observed Emi2 MO phenotype, PB1 extrusion and decondensed chromatin within a single pronucleus, was not caused by a nonspecific effect of the MO for two reasons. First, we used two control MOs that were without effect. Second, we rescued the Emi2 knockdown by readdition of Emi2 cRNA, an approach that was made possible by the fact that the rescue cRNA did not contain the 5′ UTR recognized by the MO. In some instances, especially where rescue was done with the higher dose of Emi2, oocytes went on to arrest at MetII. This would be consistent with Emi2 inhibiting the APC at this time and the requirement for APC activity in order for mouse oocytes to complete the first meiotic division (Herbert et al., 2003; Terret et al., 2003; Hyslop et al., 2004; Homer et al., 2005). Interestingly, the effect of Emi2 knockdown on oocyte maturation is the same as that obtained when expression of cyclin B1 is prevented by antisense during oocyte maturation (Ledian et al., 2001). In oocytes matured with cyclin B1 antisense, MPF activity is not reestablished after PB1 extrusion, and a single pronucleus forms.

The Emi2 phenotype observed in vitro–matured oocytes to some extent resembles the siRNA Emi2 knockdown phenotype obtained previously (Shoji et al., 2006), with decondensed chromatin in the parthenote. In that study, however, there was no analysis made of the spindle during the normal time course of maturation; instead, observations were made some 30 h after microinjection. Therefore, in such circumstances it was not possible to judge how the oocyte was progressing through meiosis I, for example if a MetII spindle formed and remained stable for some hours before extruding a PB and exiting meiosis II. Also, the difficulty in interpretation at this single time point was exacerbated in this strain of mouse by the degeneration of the PB1 during the 30-h period.

The present experiments were designed to examine in detail the role of Emi2 during the normal timings of meiosis I.
It was found that Emi2 was essential to restabilize cyclin B1 during interkinesis, and this rise in cyclin B1 was necessary to build a bipolar spindle. This is a previously unreported role for Emi2 in either frog or mouse and may add clarity to the function of CSF in mouse. Given that species-specific differences appear to exist, it is unclear whether this role in establishing MetII arrest would be conserved in frog. In frog, several studies have elegantly identified the c-Mos–MAPK–p90rsk–Bub1 pathway (Sagata et al., 1989; Kosako et al., 1994; Bhatt and Ferrell, 1999; Gross et al., 1999, 2000; Tunquist and Maller, 2003) and the spindle checkpoint proteins Mad1 and Mad2 as essential for the establishment of MetII arrest (Tunquist et al., 2003). Once established, Emi2 contributes to the maintenance of MetII arrest (Liu and Maller, 2005; Rauh et al., 2005; Schmidt et al., 2005; Tung et al., 2005; Hansen et al., 2006), and it is perhaps also required for the establishment of MetII arrest; however, this requires further investigation.

When compared with reports in frog, the mechanism of the establishment of MetII arrest in mouse is poorly understood. In contrast to frog, there have been no firm candidates suggested to be involved in establishment. Oocytes from c-Mos−/− mice initially arrest at MetII for 2–4 h, but then fail to maintain this arrest and exhibit spontaneous parthenogenetic activation (Colledge et al., 1994; Hashimoto et al., 1994; Verlhac et al., 1996). Verlhac et al. (1996) demonstrated convincingly that although c-Mos is required for the stabilization of MPF once MetII is achieved, neither c-Mos nor MAPK are involved in MPF reactivation during the MetI–MetII transition. These data suggest that c-Mos is part of the maintenance of MetII arrest, but not its establishment. In agreement with this, the addition of MAPK inhibitors to MetII oocytes induces parthenogenetic activation (Phillips et al., 2002). Therefore, it is clear that the c-Mos–MAPK pathway contributes to MetII arrest once it is established. In any event, MAPK activity declines after cyclin B1 degradation, so MAPK is not targeted immediately by the Ca2+ signal at fertilization (Verlhac et al., 1994). Similarly, the main downstream target of MAPK activity in frog appears to have little role in MetII arrest in mouse because oocytes from p90rsk knockout mice (Dumont et al., 2005) still arrest at MetII. Furthermore, oocytes injected with dominant-negative forms of Bub1 and Mad2 still arrest at MetII (Liu and Maller, 2005; Rauh et al., 2005; Schmidt et al., 2005; Tung et al., 2005; Hansen et al., 2006), and it is perhaps also required for the establishment of MetII arrest; however, this requires further investigation.

Materials and methods

All chemicals were obtained from Sigma-Aldrich, unless otherwise stated, and were of tissue culture or embryo-tested grade where appropriate.

Gamete collection and culture

4–6-wk-old MF1 mice (Harlan) were used. GV oocytes and MetII oocytes were collected from hormonally primed animals, as previously described (Nixon et al., 2002; Hyslop et al., 2004). For bench handling, microinjections, and imaging experiments, GV oocytes and MetII oocytes were cultured in medium M2. When necessary, oocytes were arrested at the GV stage in medium M2 containing 1 μM vinblastine. For longer term incubation, GV oocytes and maturing oocytes were held in a 5% CO2-humidified incubator at 37°C and cultured in MEM (Invitrogen) with 20% fetal calf serum. Parthenogenetic activation of MetII oocytes was achieved by washing oocytes into Ca2+-free M2 media containing 10 mM SrCl2 (Bos-Mikh et al., 1997; Madgwick et al., 2004). Cycloheximide was added to media at a concentration of 10 μg/ml, and nocodazole was added at a concentration of 100 nM. In the case of nocodazole, oocytes were incubated for 15 mins before parthenogenetic activation.

Microinjection and imaging

Microinjection of MOs and cRNA constructs were made as described previously on the heated stage of an inverted microscope (TE300; Nikon) fitted for epifluorescence (Nixon et al., 2002). In brief, fabricated micropipettes were inserted into cells using the negative capacitance overcompensation facility on an electrophysiological amplifier (World Precision Instruments); this procedure ensures a high rate of survival (>90%). Images were recorded using a 20× objective, NA 0.75, and a charge-coupled device camera (Micromax 1300Y; Sony). Filter sets were 430 ± 10 nm and 500 ± 10 nm excitation filters for Cerulean and Venus, respectively (Chroma Technology Corp.). Hoechst staining was examined using a 360 ± 10.9 nm excitation filter (Chroma Technology Corp.). All live cell imaging was performed in medium M2 at 37°C. Confocal sections of Hoechst and Texas red staining were performed on a confocal microscope (SP2, Leica) to capture images of the spindle microtubules and chromatin. MetaMorph and MetaFluor imaging software (Universal Imaging Corp.) were used for image capture and data analysis.

cRNA constructs and MOs

Using primers specifically for the mouse homologue of the Emi2 gene (AK0330012; National Center for Biotechnology Information), we amplified full-length mouse Emi2 by PCR from a mouse MetII-arrested oocyte cDNA library. In brief, the cDNA library was prepared by collecting 750 MetII oocytes in sterile PBS. Oocytes were lysed immediately, and mRNA was isolated directly using a commercial kit according to the manufacturer's instruction (Dynabeads mRNA Direct kit; Dynal). Double-stranded cDNA was synthesized from RNA and amplified by PCR using a cDNA synthesis kit (SMART PCR cDNA Synthesis kit; CLONTECH Laboratories, Inc.). Mouse Emi2 was then cloned from the cDNA library into a modified pRN3 vector, designed to produce mRNA transcript C-terminally coupled to Venus. Full-length cyclin B1 and Δ90 cyclin B1 were amplified by PCR as previously described (Madgwick et al., 2004). Full-length cyclin B1 was cloned into two modified pRN3 vectors designed to produce mRNA transcripts C-terminally coupled to Venus or Cerulean. Δ90 cyclin B1 was cloned into a nonfluorescent pRN3 vector (Madgwick et al., 2004). Mad2 was manufactured as described in Madgwick et al. (2005). cRNA maximal
stability was conferred in all constructs by the presence of a 5’ globin UTR upstream and both 3’ UTR and poly (A)–encoding tracts downstream of the gene. cRNA was synthesized using T3 mMESSAGE mMACHINE (Ambion) and dissolved in nuclease-free water to the required microprecipitate concentration. An antisense MO (Genetools) was designed to recognize the 5’UTR of Emi2. Two further controls were also designed, the invert of this region and a 5’m (for sequences see Table I). All MOs were used at a microprecipitate concentration of 2 mM.

Anti-Emi2 antibody purification

An MBP fusion of the human Emi2 N terminus (residues 1–400; hEmi2N) and a GST fusion of the mouse Emi2 N terminus (residues 1–398; mEmi2N) were bacterially expressed and purified using standard protocols, with the exception that all procedures involving GST-mEmi2N were conducted in the presence of 0.01% Triton X-100 to maintain protein solubility. Sera from rabbits immunized against MBP-hEmi2N were screened for immunoblot cross-reactivity toward in vitro-translated mouse Emi2. GST-mEmi2N, immobilized on a column of CNBr-activated Sepharose 4B (17–0430-01; GE Healthcare), was used to affinity purify anti-mEmi2 antibodies from cross-reactive sera. Antibodies were eluted from the column using 100 mM glycine, pH 2.5.

Immunoblotting

For Emi2, lysates from oocytes were incubated overnight with anti-mEmi2 cross-reactive sera. Antibodies were eluted from the column using 100 mM glycine, pH 2.5.

Immunostaining of oocytes

Immunostaining of cyclin B1 was performed as previously described (Reis et al., 2006). In brief, oocytes were fixed and permeabilized by an incubation in 3.7% paraformaldehyde in PBS (30 min), followed by incubation in 3.7% paraformaldehyde, 2% Triton X-100 in PBS (30 min). Fixed oocytes were then washed extensively in 1% PVP and 1% BSA in PBS. For spindle staining, oocytes were incubated with rat anti-tubulin antibody (Y1/12, 1:40, Abcam). 5 μg/ml Texas red conjugated anti-rat goat IgG (Abcam) was used as a secondary antibody. Anti–mouse IgG (P0447; DakoCytomation) and ECL Plus (RPN2132; GE Healthcare; Fig. 4) and ECL Plus (RPN2132; GE Healthcare) were used as secondary detection reagents.

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