Nuclei and Microtubule Asters Stimulate Maturation/M Phase Promoting Factor (MPF) Activation in *Xenopus* Eggs and Egg Cytoplasmic Extracts

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**Abstract.** Although maturation/M phase promoting factor (MPF) can activate autonomously in *Xenopus* egg cytoplasm, indirect evidence suggests that nuclei and centrosomes may focus activation within the cell. We have dissected the contribution of these structures to MPF activation in fertilized eggs and in egg fragments containing different combinations of nuclei, centrosomes, and microtubules by following the behavior of Cdc2 (the kinase component of MPF), the regulatory subunit cyclin B, and the activating phosphatase Cdc25. The absence of the entire nucleus–centrosome complex resulted in a marked delay in MPF activation, whereas the absence of the centrosome alone caused a lesser delay. Nocodazole treatment to depolymerize microtubules through first interphase had an effect equivalent to removing the centrosome. Furthermore, microinjection of isolated centrosomes into anucleate eggs promoted MPF activation and advanced the onset of surface contraction waves, which are close indicators of MPF activation and could be triggered by ectopic MPF injection. Finally, we were able to demonstrate stimulation of MPF activation by the nucleus–centriole complex in vitro, as low concentrations of isolated sperm nuclei advanced MPF activation in cycling cytoplasmic extracts. Together these results indicate that nuclei and microtubule asters can independently stimulate MPF activation and that they cooperate to enhance activation locally.

**Key words:** cell cycle • centrosome • maturation promoting factor • microtubule • nucleus • *Xenopus*

**Introduction**

Entry into mitosis in most, if not all, cells depends on the activation of maturation/M phase promoting factor (MPF),† a complex containing the kinase Cdc2 and a regulatory cyclin B subunit (for reviews see Beckhelling and Ford, 1998; Ohi and Gould, 1999). MPF activation has been particularly well studied in *Xenopus* egg cytoplasm, where it is controlled by both the accumulation and periodic degradation of cyclin B (Murray and Kirschner, 1989; Murray et al., 1989) and by the phosphorylation status of Cdc2. Inhibitory phosphorylations on Cdc2, catalyzed by the kinases Weel/Myt1, are reversed by the phosphatase Cdc25 (Lew and Kornbluth, 1996). MPF activation is probably amplified by a positive feedback loop involving Cdc25, which is itself activated by MPF (Gautier et al., 1991; Hoffmann et al., 1993) and polo-like kinases (Kumagai and Dunphy, 1996; Abrieu et al., 1998; Nigg, 1998). A ciste MPF triggers a cascade of phosphorylations that promote mitotic events including nuclear membrane breakdown, the formation of a bipolar mitotic spindle, and, ultimately, cyclin destruction by the anaphase promoting complex to terminate mitosis (Sudakin et al., 1995; Peters, 1999).

Studies of cells from several different species have shown that Cdc2 kinase and various cyclin B isoforms, as well as other proteins involved in controlling MPF activity, move between different subcellular locations during the cell cycle (Ohi and Gould, 1999; Pines, 1999). In particular, cyclin B and Cdc2, along with certain Cdc25 isoforms, shift from predominantly cytoplasmic to predominantly nuclear locations at the beginning of mitosis (Seki et al., 1992; Lopez-Girona et al., 1999; Pines, 1999). Within the cytoplasm, Cdc2 and cyclin B (B1 in human cells, B2 in *Xenopus*) associate with microtubules and centrosomes, particularly during late interphase and M phase (Bailly et al., 1989;
A. I. A. et al., 1990; Pines and Hunter, 1991; Baily et al., 1992; Gallant and Nigg, 1992; Ackman et al., 1995; Okada et al., 1995). Polo-like kinases also concentrate at the centrosome (Llamazares et al., 1991; Golsteyn et al., 1995). Cyclin B2 and Myt1 colocalize with the ER and Golgi apparatus in mammalian cells (Ackman et al., 1995; Liu et al., 1997), whereas Wee1 is nuclear in human cells and yeast (Hald et al., 1993; Baldin and D'commun, 1995; Aigue et al., 1997). In eggs, distinct cortical and perinuclear localization has been reported for cyclin B in amphibians (Sakamoto et al., 1998) and Drosophila (Raff et al., 1990), and Cdc25 cortical localization has been observed in Caenorhabditis elegans eggs (Ashcroft et al., 1999). The significance of these various localizations is not yet clear. A Citation and/or inactivation of MPF and its regulators, in association with particular subcellular structures, may be important to coordinate changes in intracellular organization, in particular those accompanying mitosis. The controlled compartmentalization of these molecules may also be important for the regulation of cell cycle progression itself (Oh and Gould, 1999; Pines, 1999). There are clear examples of such spatial regulation being important in "checkpoint control" responses to DNA damage (Touyoshima et al., 1998; Lopez-Girona et al., 1999), but as yet there has been no direct evidence for a role during the normal process of MPF activation.

In amphibian oocytes, eggs and early embryos, MPF activation can clearly occur autonomously in the cytoplasm. It can occur in enucleated oocytes and eggs (Masui, 1972; Newport and Kirschner, 1984; Gautier, 1987; Dabauvalle et al., 1988) as well as in the absence of microtubules (Gerhart et al., 1984; Kimmelman et al., 1987). Despite the independence of MPF activation from nuclei and microtubules, a body of indirect evidence indicates that these structures may modulate the timing and/or the site of MPF activation. First, MPF activation in the intact egg is first detectable in the animal hemisphere (Iwao et al., 1993; Rankin and Kirschner, 1997; Pérez-Mongiovi et al., 1998), where the nuclei and microtubule nucleating centers lie. Second, meiotic MPF activation is compromised in enucleated oocytes (Gautier, 1987; Iwashita et al., 1998). Other evidence comes from analysis of the pairs of periodic cortical reorganizations, or "surface contraction waves" (SCWs), that accompany MPF activation and inactivation during each early cell cycle (Hara et al., 1980; Yoneda et al., 1982; Rankin and Kirschner, 1997; Pérez-Mongiovi et al., 1998). SCW initiation is enhanced in the proximity of the nucleus-centrosome complex, whereas enucleation and disruption of microtubules delay the SCWs (Sakai and Kubota, 1981; Shinagawa, 1983, 1992; Shinagawa et al., 1989). Observations of locally regulated microtubule dynamics in maturing starfish oocytes (Barakat et al., 1994) and in mitotic ctenophore eggs (Houlston et al., 1993) also indicate that regionalized MPF activation occurs in relation to nuclear/spindle position.

The Xenopus egg is large and robust enough to permit physical separation of nuclei and centrosomes into different cytoplasmic fragments, allowing the role of nucleus and centrosome in MPF activation to be addressed directly. We have performed a series of experiments designed to create viable fragments with defined compositions (see Fig. 1), using the microtubule depolymerizing drug nocodazole to disrupt the microtubule network. Microinjection of supernumerary centrosomes into anucleate eggs permitted separation of the effect of the centrosomes from that of the nuclei. We exploited the relationship between MPF activation and the first SCW to aid analysis of the influence of injected centrosomes. Our in vivo experiments were complemented by analysis of cycling cytoplasmic extracts designed to address the role of nuclei in MPF activation in vitro to compare to previous studies, which reported that nuclei inhibited rather than favored MPF activation (Dasso and Newport, 1990).

Materials and Methods

Ovulation and Fertilization

Female X. laevis (Centre National de la Recherche Scientifique Rennes or Blades Biological), preinjected with 50 IU of pregnant mare serum (in some cases), were induced to ovulate by injection of 700 IU human chorionic gonadotropin (Ovagon). Testis mince in 80% Steinberg's solution; 150 mM NaCl, 0.7 mM KCl, 0.8 mM MgSO4, 10 mM Ca(NO3)2, 5 mM Tris-HCl, pH 7.4 was used for insemination. Eggs were dejellied using 1% cysteine, pH 8.0, and cultured in 10% Steinberg's. Variations in ambient temperature affect the length of the first cell cycle (85–110 min), so a normalized time (NT) scale was used to express results. Insemination was 0 NT, and first cleavage (50% of control eggs showing a clear cleavage furrow at the animal pole) was 1.0 NT, with "time units" corresponding to the length of this period (Gerhart et al., 1984). Only batches in which the majority of eggs cleaved within a 5–10 min period were used.

Ligation and Nocodazole Treatment

For "ligation" in vivo, fertilization envelopes were removed from dejellied eggs in 5–10% Ficoll/80% Steinberg's. The eggs were then transferred to 80% Steinberg's in dishes coated with 2% agar, and fragments were separated using 0.3-mm-diameter glass rods (Elinson and Palacek, 1993) placed to produce big and small fragments (~3/4 and 1/4, respectively, of egg volume) with defined compositions (Fig. 1). Fragments were selected for analysis that were completely separated by 0.65 NT, well before the entry into mitosis. The success of the ligation was assessed by examining cleavage patterns in groups of cultured fragments, and only experiments showing a high success (~95%) were considered. Normal cleavage occurred in fragments containing nuclei and centrosomes; abortive "pseudocleavage furrows" occurred in fragments with the female nucleus but no organized centrosome; and no furrow was visible in anucleate fragments. For biochemical analysis, fragments were frozen using liquid nitrogen in the presence of RM (160 mM β-glycerophosphate, 40 mM EGTA, pH 7.5, 30 mM MgCl2, 4 mM DTT) including 0.1 mM NaN3, and protease inhibitors (10 μg/ml each of aprotinin, leupeptin, pepstatin, and 2 mM AEBSF; all from Sigma-Aldrich or ICN Biomedicals).

To depolymerize microtubules, fertilized eggs were transferred to 10 or 30 μM nocodazole (BoIomol) in 20% Steinberg's at 0.2–0.3 NT (early treatments) or at 0.45–0.65 NT (late treatments). Fragments derived from fertilized eggs were ligated in the presence of nocodazole. No nocodazole-treated eggs cleaved or displayed abortive cleavage furrows.

MPF and Centrosome Injections

Fertilized eggs were microinjected in different regions with 4–8 nl of purified recombinant human Cdc2/cyclin B–GST expressed in baculovirus-infected insect cells (gift of J. Gautier, Columbia University, New York, NY). A drummond microinjection apparatus. This MPF was diluted to give histone H1 kinase activity equivalent to that measured in the same volume of mitotic egg cytoplasm. Eggs were immediately mounted and SCWs recorded (see below).

Centrosomes isolated from cultured human K 37 lymphocytes (Moudjou and Bornens, 1994) were provided by M. Bornens (Institute Curie, Paris, France). They were injected into anucleate eggs obtained by removal of the metaphase II meiotic spindle of unfertilized eggs by aspiration of a small amount of cytoplasm around the second polar body with a fine glass pipette (Houlston and Elinson, 1993). This "enucleation" usu-
In Vitro Experiments

Low speed egg extracts were prepared according to Hutchison et al. (1987, 1988), with modifications. Washed, dejellied eggs taken 20 min after ionophore activation (Lindsay et al., 1995) were packed for 60 s at 1,000 rpm in a Beckman L-2 ultracentrifuge. 10 μg/ml cytochalasin B added before centrifugation at 10,000 rpm for 10 min at 4°C in a Beckman L-2 ultracentrifuge (SW 50.1 rotor). The supernatant was mixed with 5 μg/ml cytochalasin B and 10 μg/ml aprotinin and then centrifuged a second time at 10,000 rpm for 10 min at 4°C to clarify the extract. Deembranated sperm heads were prepared as described (Hutchison et al., 1987) and resuspended in SuNAsp buffer (0.25 M sucrose, 75 mM NaCl, 0.15 mM spermine, 0.5 mM spermidine, 15 mM Hapes, pH 7.5) at 10^4 nuclei/μl, then added to 200 μl aliquots of egg extracts to give 250–4,000 nuclei/μl of extract. Incubations were started by transfer to 21°C. At appropriate time points, the following samples were removed: for protein analysis, 2 μl of incubated extract was mixed with 2 volumes of 2× SDS sample buffer and 1 volume of water and heated for 3 min at 95°C; for histone kinase assay, 2 μl of extract was mixed with 48 μl of histone kinase buffer (see below) and frozen in liquid nitrogen at each time point.

Histone H1 kinase assays were carried out according to Felix et al. (1989), with the modifications of Lindsay et al. (1995). Extract aliquots were diluted 1:25 in histone kinase buffer (80 mM β-glycerophosphate, 20 mM E GTA, 15 mM MgCl₂, 1 mM DTT, pH 7.3, 10 μg/ml aprotinin, 10 μg/ml leupeptin), and 10-μl duplicates were incubated with 10 μl of reaction cocktail (0.6 mM ATP, 2 mg/ml calf thymus histone H1; Sigma-A Aldrich), 75 μCi/ml [γ-32P]ATP (A mersham Pharmacia Biotech) for 15 min at 21°C. The incubations were stopped by transfer to ice. 15 μl of reaction was spotted onto P81 paper (Whatman), washed three times with phosphoric acid (150 mM), and air dried before scintillation counting.

SCW Recordings

Mitochondrial aggregates associated with germ plasm were visualized by incubating dejellied eggs for 3 min in a 2.5-μg/ml solution of DIOC(3) (Molecular Probes) in water and diluted immediately before use from a 2.5-mg/ml stock in ethanol (Savage and Danielik, 1993; Pérez-Mongiovi et al., 1998). Eggs were transferred to 3–5% Ficoll in 80% Steinberg's solution (Savage and Danilchik, 1993; Pérez-Mongiovi et al., 1998). Washed, dejellied eggs taken 20 min after ionophore activation (Lindsay et al., 1995) were packed for 60 s at 1,000 rpm in a Beckman L-2 ultracentrifuge. 10 μg/ml cytochalasin B added before centrifugation at 10,000 rpm for 10 min at 4°C in a Beckman L-2 ultracentrifuge (SW 50.1 rotor). The supernatant was mixed with 5 μg/ml cytochalasin B and 10 μg/ml aprotinin and then centrifuged a second time at 10,000 rpm for 10 min at 4°C to clarify the extract. Deembranated sperm heads were prepared as described (Hutchison et al., 1987) and resuspended in SuNAsp buffer (0.25 M sucrose, 75 mM NaCl, 0.15 mM spermine, 0.5 mM spermidine, 15 mM Hapes, pH 7.5) at 10^4 nuclei/μl, then added to 200 μl aliquots of egg extracts to give 250–4,000 nuclei/μl of extract. Incubations were started by transfer to 21°C. At appropriate time points, the following samples were removed: for protein analysis, 2 μl of incubated extract was mixed with 2 volumes of 2× SDS sample buffer and 1 volume of water and heated for 3 min at 95°C; for histone kinase assay, 2 μl of extract was mixed with 48 μl of histone kinase buffer (see below) and frozen in liquid nitrogen at each time point.

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Results

To assess the contribution of nuclei, centrosomes, and microtubules to MPF activation at first mitosis, we undertook a series of ligation experiments. Careful positioning of the glass rod used to bisect the eggs, in relation to the animal pole and the pigmented sperm entry point, before the time of pronuclear migration (Fig. 1), yielded defined fragments containing both pronuclei, no nuclei, only the male pronucleus (with associated centrosome-containing centrosomes), or only the female pronucleus (which has some poorly organized microtubule nucleating material but no centriole). The behavior of these fragments at the time of first cleavage was used to assess the success of the ligations. Fragments containing nuclei and centrosomes cleaved normally, whereas those containing only the female pro-
nucleus exhibit abortive pseudocleavage furrows, and anucleate fragments show no visible furrowing (Briggs and King, 1953; Houlston and Elinson, 1991). MPF activation in groups of two or three fragments of different types derived from the same eggs frozen simultaneously at successive times was monitored by Western blotting using antibodies detecting cyclin B2 and Cdc2 as described previously in Pérez-Mongiovi et al. (1998), and/or Cdc25C. The three different antibodies were applied to parts of single blots corresponding to the appropriate molecular weight region, so that slight variations in loading could be assessed by comparing Cdc2 and Cdc25C levels between lanes. A titration of MPF is detected as the loss of slower migrating Cdc2 isoforms, corresponding to the subpopulation recruited to the preactive Cdc2–cyclin B complex (Solomon et al., 1990), accompanied by a marked shift in electrophoretic mobility of Cdc25 due to mitotic phosphorylation on multiple sites.

**Nuclei Stimulate MPF Activation at First Mitosis**

Comparison of the timing of detectable Cdc2 dephosphorylation and Cdc25 hyperphosphorylation in nucleate and anucleate fragments derived from the same eggs indicated that whereas activation of the cyclin B–Cdc2 population in nucleate fragments occurred at approximately the same time as in intact eggs, it was delayed in anucleate fragments by \( \sim 0.2 \) time units (Fig. 2; see also Figs. 5 and 6, where experiments were performed seven times with similar results). The observed effect cannot be attributed to the smaller size of anucleate fragments, since the timing of MPF activation was equivalent in large and small nucleate fragments (below). Nor was there any evidence for globally reduced or delayed cyclin B2 accumulation in anucleate fragments; in fact, cyclin B2 went on to accumulate to higher levels than in nucleate fragments (Fig. 2). Furthermore, at the time of MPF activation in nucleate fragments, the phosphorylated inactive form of Cdc2 could be detected clearly in anucleate fragments, indicating that inactive Cdc2–cyclin B complex was available for activation. This suggests that the nucleus acts to trigger or amplify the activation of preformed Cdc2–cyclin B. The delay of \( \sim 0.2 \) time units in MPF activation observed in anucleate fragments corresponds closely with that observed previously for SCWs (Shinagawa, 1983, 1992). Thus, we have confirmed directly in vivo that in Xenopus early embryos, the nucleus acts to stimulate MPF activation.

**Stimulation of MPF Activation by Nuclei In Vitro**

The extensive use of cytoplasmic extracts prepared from activated Xenopus eggs, which can undergo autonomous cycles of MPF activation and inactivation (Hutchison et al., 1988; Murray et al., 1989), has reinforced the idea that MPF activation in early amphibian embryos is controlled entirely by cytoplasmic factors. In such “cytological extracts,” neither nuclei nor microtubules are necessary for MPF activation to occur (Dasso and Newport, 1990; Murray and Hunt, 1993; Minshull et al., 1994), whereas exogenous nuclei can cause a delay in MPF activation. This latter effect results from the activation of a “replication checkpoint” by nuclei blocked in or still undergoing S phase (Dasso and Newport, 1990). Our in vivo demonstration that nuclei can advance MPF activation raised a doubt as to whether cytoplasmic extracts fully reproduce the mechanisms of MPF regulation used in the intact egg. We thus tested the influence of nuclei on the time of MPF activation in cytoplasmic extracts by adding different numbers of exogenous sperm nuclei. The kinetics of MPF activation was measured by histone H1 kinase activity (Fig. 3 a), confirmed by Western blotting (Fig. 3 b), and examination of chromatins condensation and nuclear envelope breakdown microscopically (not shown). Control experiments confirmed that no variation in the timing of histone H1 kinase activity peaks was observed between aliquots containing equivalent numbers of nuclei (data not shown). We found that low concentrations of nuclei (250 and 1,000 nuclei/\( \mu l \) of extract) advanced the time of peak MPF activation by 10–20 min at 21°C compared with that seen in extracts without nuclei (Fig. 3, similar results were obtained in three experiments), thus reproducing the effect seen in vivo. To ensure that MPF activation was not being promoted by active MPF or Cdc25C in the sperm preparation,
we probed sperm nuclei for the presence of these proteins by Western blotting. No Cdc2, cyclin B2, or Cdc25 was detected in samples of 4,000 demembranated nuclei (data not shown). It is also unlikely that the preparation of nuclei stimulated cyclin B synthesis, since no consistent differences were observed between cyclin B levels in relation to the numbers of nuclei added. As reported previously (Dasso and Newport, 1990), extracts containing higher concentrations of nuclei (4,000 nuclei/μl) showed delayed MPF activation. This indicates that with higher concentrations of sperm nuclei present, the replication machinery in the extracts was insufficient to allow complete replication by the time MPF activated in control extracts, triggering the replication checkpoint to delay mitosis.

**Participation of Microtubules in MPF Activation**

The experiments described above allowed us to show clearly both in vivo and in vitro that low concentrations of sperm-derived nuclei advance the time of MPF activation. Since sperm-derived nuclei have tightly associated centrioles, we reasoned that microtubule asters nucleated around this structure may contribute to the stimulatory effect. The sperm-derived centriole provides the dominant microtubule nucleating center in the egg during the first 20–40 min after fertilization, such that the vast majority of egg microtubules at this time form a rapidly expanding "sperm aster" around it (Houliston and Elinson, 1992). During the second half of the first interphase period, more generalized microtubule polymerization occurs throughout the egg. Then, just before mitosis, the duplicated centrosome nucleates small prophase asters, which separate to form the mitotic spindle poles, whereas cytoplasmic microtubules depolymerize elsewhere (Fig. 1). To test the influence of these different microtubule structures on the timing of MPF activation, eggs were treated with 10 or 30 μM nocodazole at different times after fertilization. We verified that 10 μM nocodazole caused depolymerization of the vast majority of egg microtubules within 10–15 min of treatment as assessed by antitubulin immunofluorescence (data not shown). Cleavage was blocked in all nocodazole-treated groups. In seven experiments in which nocodazole treatment began between 0.2 and 0.3 NT, Western blot analysis indicated that Cdc2 dephosphorylation and Cdc25C hyperphosphorylation were delayed by ~0.1 time units compared with untreated controls (Fig. 4 and see Fig. 6). However, early nocodazole treatment consistently delayed Cdc2 dephosphorylation and Cdc25C hyperphosphorylation. Later treatments, beginning at 0.45 or 0.65 NT, did not (Fig. 4). Since it takes 10–15 min for the drug to penetrate the egg and cause microtubule network disassembly, the cutoff time for the stimulatory role of microtubules effectively lies between 0.40 and 0.55 NT. This corresponds to the time at which the sperm aster is replaced by a more widespread microtubule network (Fig. 1; Houliston and Elinson, 1992).

Although the accumulation of cyclin B2 was similar in treated and nontreated eggs, in some experiments it appeared to be slightly enhanced in the presence of nocodazole (three out of seven early treatments, and one out of four later treatments). A variable inhibitory effect of nocodazole on subsequent cyclin B degradation was also observed in 6 of the 11 treated groups. A generalized role for microtubules in encouraging cyclin B2 degradation has been demonstrated previously in unfertilized eggs (Thibier et al., 1997). Increased cyclin B accumulation through the first cell cycle could explain the slight advance in the onset of MPF activation and the delay of inactivation detected in eggs fertilized in the presence of nocodazole (Gerhart et al., 1984). It may also partially offset the stimulatory effect of the sperm aster to produce the more irregular broad MPF activation peak at first mitosis in such eggs described by Gerhart et al. (1984).
although microtubules are not required for MPF activation in Xenopus eggs and may generally oppose MPF activation by favoring cyclin B degradation, the microtubules of the sperm aster stimulate the activation process at first mitosis. The sperm aster may act by reorganizing components in the animal cytoplasm involved later in the MPF activation process (see Discussion).

Additive Effects of Microtubule Asters and Nuclei to Stimulate MPF Activation

The experiments described above showed that elimination of sperm aster microtubules, like enucleation, can delay Cdc2 dephosphorylation during the first cell cycle. Since the delay provoked by early nocodazole treatment was less than that observed in the absence of a nucleus, the absence of the sperm aster in enucleate treatment was unlikely to fully account for the observed delay, suggesting that the nucleus itself contributes to the activation process. The relative contributions of the nucleus, centrosome, and microtubules were assessed by comparing the timing of MPF activation in various types of egg fragments produced, as shown in Fig. 1, with or without nocodazole treatment. An example of an experiment comparing the effect of different combinations of nuclei and centrioles is shown in Fig. 5, and the results obtained from the various experiments are combined in Fig. 6. Note that as shown previously for SCW’s (Shinagawa, 1992), the time of Cdc2 dephosphorylation was not affected by the size of the fragments. It occurred in parallel in whole eggs, large fragments (~3/4 volume of a whole egg), and small fragments (~1/4 volume of a whole egg) as long as nuclei, centrosomes, and microtubules were present. The same was true for different sized fragments containing nuclei but no microtubules.

Fragments lacking both nuclei and sperm asters showed a consistent delay in Cdc2 dephosphorylation of ~0.2 time units, compared with shorter delays of ~0.1 time units for nocodazole-treated eggs or nocodazole-treated fragments containing nuclei (Fig. 6, compare columns 3 and 6 with 4, 5, 9, and 10). Furthermore, the presence of either pronucleus or the zygote nucleus was sufficient to advance MPF activation in the absence of microtubules (Fig. 6, compare columns 5, 9, and 10 with 6). It is also informative that the time of Cdc2 dephosphorylation in fragments lacking a

\[ \text{Figure 4. Early nocodazole treatment delays MPF activation.} \]

Western blots (details and annotations as in the legend to Fig. 2) showing the behavior of Cdc2, Cdc25C, and cyclin B2 during first mitosis after treatment at different times with 10 mM nocodazole. Treatments were started at 0.26, 0.44, and 0.66 NT as indicated. The times below each lane show the period of collection and freezing of control and treated eggs. Eggs treated at 0.26 NT showed a delay in Cdc2 and Cdc25C activation of ~0.1 time units, whereas treatments beginning at 0.44 and 0.66 NT did not. Degradation of cyclin B2 was delayed in the early nocodazole-treated eggs and appeared to degrade more slowly.

\[ \text{Figure 5. MPF activation depends on nuclei and centrosomes.} \]

Cdc2 and cyclin B2 proteins followed by Western blotting (see Fig. 2) in nucleate (Nuc) and anucleate (Anuc) fragments compared with fragments containing a single male or female pronucleus. The centrosome associated with the male pronucleus organizes a microtubule aster. Cdc2 dephosphorylation was first detectable in fragments containing both nuclei and asters, then in fragments containing the female pronucleus, and finally in anucleate fragments. Low cyclin B2 levels in the last lane of anucleate samples (left blot) partially reflects reduced loading levels (compare with Cdc2 band intensities).
Centrosomes Injected into Anucleate Eggs Advance SCWs

To test whether microtubule asters could affect MPF activation in the absence of a nucleus, we injected purified centrosomes into anucleate eggs obtained by removal of the meiotic spindle at the time of activation. Successful elimination of the female pronucleus was judged by failure to show pseudofurrows at the time of mitosis (Briggs and King, 1953; Houliston and Elinson, 1991), which varied between egg batches but comprised at least 60% in the experiments used (5 experiments, with 10–20 eggs followed in each). Centrosome solutions were diluted sufficiently to produce multiple cortical pigment accumulations in most injected eggs during first interphase, indicating the presence of microtubule asters. Eggs showing these pigment patterns reliably underwent multiple pseudofurrows, indicating the presence of active centrosomes (Heidemann and Kirschner, 1975).

A nalysis of MPF activation after enucleation and centrosome microinjection is compromised by the variable reliability of these procedures. To circumvent this difficulty, we first chose to monitor the time of MPF activation by videomicroscopy of the SCWs during the mitotic period beyond the time of pseudocleavage in pricked-activated controls. This method has the advantage of allowing the status of each egg to be determined definitively at the time of cleavage. In addition, it does not depend on sample collection at a single time point. A close correlation has been demonstrated between SCW progression and MPF activation propagating across the egg (Rankin and Kirschner, 1997; Pérez-Mongiovi et al., 1998). Both are blocked in isolated vegetal egg fragments and delayed by equivalent amounts in anucleate fragments (Pérez-Mongiovi et al., 1998). To further validate the use of SCWs as reporters of MPF activation, we injected human recombinant MPF into different regions of the egg before mitosis (~0.65 NT). Videomicroscopy of mitochondrial island displacement (Savage and Danilchik, 1993; Pérez-Mongiovi et al., 1998) (Fig. 7) showed that vegetal or equatorial MPF injections provoked ectopic cortical waves, similar to the first SCW, which propagated across the vegetal hemisphere from the injection site (Fig. 7 b). These were 40–50% slower than endogenous SCWs, but still fell within the range of SCW speeds reported previously (Savage and Danilchik, 1993; Pérez-Mongiovi et al., 1998). Subsequent cleavage furrows sometimes deviated around the injected region, perhaps because of impaired degradation of exogenous human Cdc2/cyclinB–GST locally. Injections into the animal hemisphere provoked premature waves indistinguishable from normal SCWs, and premature cleavage furrows, indicating that a premature endogenous MPF activation and inactivation cycle had been triggered. Injections of heat-denatured MPF had no effect.

To assess the effect of centrosomes, simultaneous recordings were made of SCW progression in centrosome-injected anucleate–activated eggs, anucleate eggs without centrosomes, nucleate pricked-activated eggs, and nucleate centrosome-injected eggs (all activated within a 3-min period, Fig. 8 a). The timing of SCWs in the five different experiments performed is illustrated in Fig. 8 b. In anucleate eggs (A), SCWs in the vegetal hemisphere were delayed; typical SCWs coinciding with the end of the cortical rotation were observed in only two of five experiments. In the anucleate eggs in the other three experiments, accelerated aggregation of the mitochondrial islands towards the vegetal pole was observed after the cortical rotation had finished, but before the SCWs arrived. Similar aggregation movements in a prolonged gap between the cortical rotation and the SCWs were observed in some anucleate fragments produced by ligation (Pérez-Mongiovi, D., unpublished observations). However, in all nine “anucleate” eggs injected with centrosomes (A + C, Fig. 8), characteristic SCWs coincided with the end of the cortical rotation and appeared clearly in advance compared with uninjected anucleate eggs. Note that whereas the status of anucleate and activated eggs could be attributed definitively at the end of the recordings by examining cleavage patterns, the status of anucleate eggs injected with centrosomes could not be verified, since either a residual female pronucleus or injected centrosomes can cause furrowing activity. This introduces an error into the data in Fig. 8, A + C, such that 25–40% of these eggs may have possessed a nucleus (and/
or injected centrosome) based on our success at enucleation.

We conclude from these experiments that microtubule asters can restore and/or advance the timing of typical SCWs to anucleate eggs. They thus indicate that centrosomes can stimulate MPF activation in the absence of nuclei.

Centrosome Injection into Anucleate Eggs Stimulates MPF Activation

To obtain biochemical confirmation that the SCWs observed in anucleate eggs injected with centrioles indeed marked precocious MPF activation, we repeated the experiment outlined in Fig. 8 a and harvested eggs for Cdc2, cyclin B, and Cdc25 C blots. The time of sample collection was chosen as 85–90 min after activation. From the timing of SCWs in the vegetal hemisphere (Fig. 8 b), we expected that at this time anucleate eggs would tend to contain inactive pre-MPF, activated eggs to have completed mitosis, and centrosome-injected eggs to be in intermediate states. Note that MPF activation in the animal cytoplasm precedes vegetal SCWs by 15-90 min after activation (Pérez-Mongiovi et al., 1998). Since eggs used for biochemistry cannot be scored for behavior at the time of cleavage, errors of attribution could not be corrected as they could in the SCW experiments. This introduced variation that made initial experiments hard to interpret. We subsequently used only batches of eggs in which pigment patterns favored selection of anucleate eggs before mitosis, and discarded eggs with ambiguous pigment patterns. Although fewer eggs were available for analysis in each experiment, this stringent selection considerably reduced errors in the A and A + C categories.

An example of one experiment is shown in Fig. 9. At the time of sample preparation, all three prick-activated eggs (P) were clearly postmitotic, as indicated by the dephosphorylation of Cdc25 and absence of cyclin B2, whereas one of the two A-category eggs was clearly premitotic, with cyclin B2 and inactive Cdc2 isoforms detectable. Enucleation in the second A -category egg was probably unsuccessful since its protein profile was equivalent to those of prick-activated eggs. In contrast, two of the three A + C eggs had blot profiles clearly distinct from both A and P eggs. They contained fully active MPF as shown by hyperphosphorylation of Cdc25C. Enucleation in the second A +C category egg was probably unsuccessful since its protein profile was equivalent to those of prick-activated eggs. In contrast, two of the three A + C eggs had blot profiles clearly distinct from both A and P eggs. They contained fully active MPF as shown by hyperphosphorylation of Cdc25C. Enucleation was almost certainly successful in all three A + C eggs, since MPF activation was clearly delayed with respect to the prick-activated eggs. Centrosome injection does not appear to interfere with MPF activation, since P + C and P eggs harvested simultaneously had similar blot profiles (not shown). Thus, we can conclude that centrosomes advanced the cell cycle in two of three anucleate eggs injected in this experiment. The third A + C egg was in a premitotic state, expected if the injected centrosomes were not active or were unable to provoke detectable activation at this time.

A second experiment gave very similar results: four of six A + C eggs contained fully active MPF, whereas four of five P eggs were postmitotic, and two of the four A eggs were premitotic (one mitotic and one postmitotic). Thus, whereas MPF activation in anucleate eggs is clearly de-
layed with respect to eggs containing nuclei, albeit to variable degrees, the addition of centrosomes tends to reduce this delay. These data are entirely consistent with the SCW analyses, confirming that SCWs are reliable markers for MPF activation. In addition, they lend biochemical support that centrosomes can stimulate MPF activation in the absence of a nucleus.

Discussion

MPF activation at first mitosis in X. eggs begins at a site within the animal hemisphere (Rankin and Kirschner, 1997; Pérez-Mongiovi et al., 1998). We have been able to demonstrate by a combination of approaches that both nuclei and microtubule asters can stimulate MPF activation and are likely to contribute to this localized effect. To assess the role of centrosomes in anucleate eggs, we used SCWs as an indirect indicator of MPF activity. The close spatiotemporal correspondence between the progress of the SCWs and the waves of MPF activation and inactivation and the identical response of SCWs and MPF activation in ligation and enucleation experiments (Shinagawa, 1985; Pérez-Mongiovi et al., 1998; this study) suggested that MPF activation and SCWs were closely related. We show further evidence of a causal relationship by triggering ectopic SCWs by localized MPF injection. We are confident in using SCW initiation as a marker for the site of MPF activation, and can interpret the comprehensive series of findings of Shinagawa concerning SCWs (Shinagawa, 1983, 1985, 1992; Shinagawa et al., 1989) as revealing stimulatory effects of the nucleus, centrosomes, and microtubules on MPF activation. Taken together, the results from Shinagawa’s group and our current study show that nuclei and microtubule asters can separately enhance...
MPF activation. We propose that these structures act independently, and cooperate to focus MPF activation within the intact egg.

Enhancement of MPF Activation by Structural Components In Vivo and In Vitro

The effect of the nucleus-centrosome complex on MPF activation in vivo was revealed as a clear delay in detectable Cdc2 activation and Cdc25 hyperphosphorylation in anucleate egg fragments compared with nucleate fragments. Surprisingly, we were also able to reproduce the stimulatory effect of the nucleus-centrosome complex in cytoplasmic extracts from activated eggs. Low concentrations of sperm-derived nuclei incubated in these extracts advanced MPF activation by 10–20 min, an interval similar to that seen in egg fragments. Our inability to accelerate MPF activation further by adding more nuclei probably reflects the requirement for cyclin B to accumulate to a sufficient level to allow MPF activation (Murray and Kirschner, 1989; Hartley et al., 1996). It may also partly depend on a limit imposed by the time necessary for S phase (Blow et al., 1989), since a DNA-replication checkpoint ensures that mitosis is delayed until replication is complete. Cycling extracts supplemented with nuclei were originally used to demonstrate this checkpoint: MPF activation was delayed above a high threshold nuclear concentration and the effect was related to the presence of incompletely replicated DNA (Dasso and Newport, 1990). MPF activation was not directly compared in samples with and without low numbers of nuclei in previous studies (Dasso and Newport, 1990; M inshull et al., 1994).

Now that we have clearly established the role of the nucleus and microtubule asters in MPF activation by in vivo experiments, it may be possible to dissect the molecular basis of their action by manipulating egg cytoplasmic extracts.

The Nucleus and MPF Activation

The stimulation of MPF activation by nuclei was detectable in the absence of microtubules, implying that nuclear factors participate directly in the activation process. The known partitioning of MPF regulators between the cytoplasm and the nucleus, for instance the nuclear retention of phosphorylated cyclin B, is likely to favor initiation and/or amplification of MPF activation within the nucleus (see Introduction). In fission yeast, MPF activation in the nucleus is predominant, and mitosis is blocked in response to DNA damage by exclusion of Cdc25 from the nucleus (Lopez-Girona et al., 1999). In Xenopus eggs and other multicellular eukaryotes, where cytoplasmic activation is more significant, it is not clear where MPF first activates. In maturing amphibian oocytes, evidence favors initial cytoplasmic activation whereas nuclear factors aid amplification (Gautier, 1987; Iwashita et al., 1998). The nuclear translocation of cyclin B1 that occurs before meiosis (Li et al., 1997; Yang et al., 1998, 1999; Hagting et al., 1999) is not sufficient to induce MPF activation (Pines, 1999). Likewise, in starfish oocytes MPF activation appears to precede nuclear import of cyclin B (Ookata et al., 1992). In mammalian cells, the cytoplasmically localized B isoform of Cdc25 activates before the predominantly nuclear Cdc25C. Cdc25B is thought to act as an initial MPF activator, and Cdc25C to participate in amplification (Gabrielli et al., 1996, 1997; Nishijima et al., 1997; Lammer et al., 1998; Karlsson et al., 1999). Our data indicate that in the fertilized Xenopus egg, both nuclear and cytoplasmic factors contribute to MPF activation, and that the nucleus is important in initiating and/or amplifying MPF activation locally within the animal half of the egg.

Centrioles, Asters, and MPF Activation

We have been able to demonstrate that isolated centrosomes introduced shortly after egg activation can advance SCW's and stimulate MPF activation in the absence of nuclei. We also observed that nocodazole treatment and/or the absence of a centriole caused a delayed MPF activation in the first cell cycle, albeit less than the delay seen in the absence of the nucleus-centrosome complex. Similarly, microtubule depolymerization has been reported to delay MPF activation in fertilized sea urchin eggs (Walker et al., 1997) and to prevent it in fission yeast (Aifia et al., 1990). We propose that active centrosomes, by nucleating microtubule asters, can organize cytoplasmic components in such a way as to favor local MPF activation in the perinuclear region. The period during which microtubules were required to enhance MPF activation in Xenopus eggs coincides with the presence of the sperm aster, a giant aster that forms around the sperm centriole shortly after fertilization. This implicates the sperm aster in promoting MPF activation. The relatively small prophase asters that form at the beginning of mitosis may also stimulate MPF activation. A local effect of prophase asters on overall MPF activation may be insufficient to be detected, or may be masked, by a general antagonizing effect of microtubules on MPF activation (see below).

Previous studies showed that MPF-activation cycles in Xenopus eggs are microtubule independent (Gerhart et al., 1984; Kimelman et al., 1987). In contrast to our findings and to those concerning SCW's (Shinagawa, 1983), a slight advance in the onset of MPF activation was reported in eggs fertilized and cultured in nocodazole (Gerhart et al., 1984). A stimulation of MPF, assayed by its ability to promote maturation when microinjected into oocytes, then proceeded with relatively slow and uneven kinetics. This may partly explain the apparent discrepancy. Cdc2 dephosphorylation and Cdc25 hyperphosphorylation can only be detected once MPF amplification is well underway, whereas the “rounding-up” monitored by Shinagawa (1983) coincides with MPF inactivation (Rankin and Kirschner, 1997). A weak premature rise in MPF levels in nocodazole-treated eggs would be undetectable by these latter methods. Such kinetics may arise because polymerized microtubules generally favor MPF inactivation by encouraging cyclin B degradation. Such an effect has been demonstrated clearly in unfertilized Xenopus (Thibier et al., 1997) and mouse (Kubiak et al., 1993) eggs. Consistent with this, we observed retarded cyclin B degradation in some nocodazole experiments (see Fig. 4). Mitosis in sea urchin eggs is likewise prolonged when microtubules are depolymerized (Sluder, 1979; Hunt et al., 1992). We propose that in Xenopus eggs, cyclin B accumulates more quickly in the presence of nocodazole due to decreased cyclin B turnover, resulting in generally elevated MPF activ-
ity. However, amplification steps are complicated because regulatory factors are not previously concentrated in a central region by the sperm aster. The opposing actions of microtubules may cancel each other out, depending on the timing of the treatment. Gerhart et al. (1984) preincubated and fertilized eggs in the presence of nocodazole rather than adding it during first interphase, which likely resulted in elevated cyclin B levels during the first cell cycle (Thibier et al., 1997).

The sperm aster might favor precocious MPF activation in the central region of the animal hemisphere by concentrating regulatory molecules via minus end-directed transport or association with astral microtubules, or possibly by stimulating cyclin B synthesis. Candidate regulatory factors include Cdc2, cyclin B, Cdc25, and polo-like kinases, all of which associate with microtubules and/or the centrosome or have been observed in the perinuclear region in prophase (see Introduction). Perinuclear accumulation of cyclin B is known to be microtubule dependent in Drosophila (Raff et al., 1990). Raising cyclin B levels locally could promote MPF activation by accelerating complex formation. Polo-like kinases and Cdc25 would then actively participate in MPF amplification, the former by antagonizing the phosphatase PP2A in the two-step activation of Cdc25 (Karaïskou et al., 1998, 1999). We favor the possibility that a regulator of MPF activity, rather than locally increased levels of cyclin B, is responsible for the sperm aster’s effect because cyclin B2 accumulates to elevated levels before activation in certain types of egg fragments and in nocodazole-treated eggs (Pérez-Mongiovi et al., 1998; this study).

Cooperation between Asters and Nuclei to Stimulate MPF Activation

Taken together, our results indicate that nuclei and microtubule asters have independent but additive effects on MPF activation, and suggest that these structures cooperate to trigger MPF activation within the egg. If MPF activation is initiated within the nuclei, the centrosome-nucleated aster could act to relay the state of the cytoplasm, concentrating molecules such as cyclin B in the perinuclear region and so favoring their transfer into the nucleus for activation. Since we could demonstrate stimulation of cytoplasmic activation by centrosomes in anucleate eggs, we favor the possibility that MPF activates first in the centrosomal region, before translocation to the nucleus.

Regulatory molecules have been the focus of attention in cell cycle research over the past years, and it is becoming increasingly clear that their spatial organization is of great importance (Pines, 1999). Thus, whereas MPF can activate in the absence of nuclei, centrosomes, and microtubules in Xenopus eggs, we have shown that these structural components are not merely effectors, but active protagonists in controlling cell cycle progression.

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