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Regulation of EDEN-dependent deadenylation of Aurora A/Eg2-derived mRNA via phosphorylation and dephosphorylation in *Xenopus laevis* egg extracts

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Summary

Deadenylation is an intimate part of the post-transcriptional regulation of maternal mRNAs in embryos. EDEN-BP is so far the only known member of a complex regulating the deadenylation of maternal mRNA in *Xenopus laevis* embryos in a manner that is dependent on the 3′-untranslated region called EDEN (embryo deadenylation element). In this report, we show that calcium activation of cell-free extracts triggers EDEN binding protein (EDEN-BP) deadenylation and concomitant deadenylation of a chimeric RNA bearing Aurora A/Eg2 EDEN sequence. Deadenylation of mRNA deprived of EDEN sequence (default deadenylation) does not change with egg activation. Kinase and phosphatase inhibitors downregulate EDEN-dependent deadenylation but they do not substantially influence default deadenylation. Using indestructible ∆90 cyclin B to revert interphase extracts to the M-phase, we show that modulation of EDEN-dependent deadenylation is independent of M-phase promoting factor (MPF) activity. These results suggest that the increase in EDEN-dependent deadenylation following egg activation is achieved, at least partially, via dephosphorylation and/or phosphorylation of regulatory proteins, including EDEN-BP dephosphorylation. This regulation proceeds in a manner independent from MPF inactivation.

Key words: Aurora A, Eg2, EDEN-BP, Protein phosphorylation, MPF, RNA deadenylation

Introduction

Calcium oscillations triggered by fertilization stimulate the release of the oocyte from metaphase II (M II) meiotic arrest (Nuccitelli, 1991). Calcium induces inactivation of cytostatic factor (CSF), thereby enabling inactivation of the M-phase promoting factor (MPF) (Lorca et al., 1993). The molecular mechanism of CSF activity is not well characterized; however, we know that at least the c-mos/.../mitogen-activated protein (MAP) kinase extracellular signal-regulated kinase 2 (ERK2)/p90rsk pathway is involved (Sagata et al., 1989; Haccard et al., 1993; Gross et al., 1999) (reviewed by Masui, 2000). MPF is a complex of p34cdc2 protein kinase and cyclin B (Labbé et al., 1989) (reviewed by Dörée and Hunt, 2002). Inactivation of both CSF and MPF is necessary for the successful transition to the first embryonic interphase and the beginning of the developmental programme.

Early development preceding the midblastula transition in *Xenopus laevis* depends entirely on the maternal information stocked in mRNA and proteins accumulated during oogenesis. Gene expression during early development (as well as during oocyte maturation) depends largely on post-transcriptional modification of maternal mRNAs (reviewed by Richter, 1999). Polyadenylation and deadenylation of mRNA synthesized before oocyte maturation are the most important mechanisms involved in this process (Osborne and Richter, 1997). Different classes of maternal mRNA were described in *Xenopus* oocytes and embryos with respect to their pattern of polyadenylation/deadenylation. The Eg family of mRNA containing Eg1/cdk2 (cyclin-dependent kinase 2), Eg2/Aurora A (kinase), Eg5 (kinesin-like protein) and c-mos are polyadenylated during oocyte maturation due to the presence of a cytoplasmic polyadenylation element (CPE) in the 3′-untranslated region (3′-UTR) (Paris and Richter, 1990; McGrew and Richter, 1990) and deadenylated following fertilization due to the presence of an embryo deadenylation element (EDEN) also in the 3′-UTR (Paillard et al., 1998). Most EDEN-containing mRNAs known to date are involved in cell cycle regulation during early development. For example, c-mos must be degraded and its synthesis/accumulation downregulated to enable the exit from M II-arrest after fertilization, as c-mos is a key component of the CSF activity. Ectopic expression of c-mos following fertilization induces cell cycle alterations and eventually cell cycle arrest (Sagata et al., 1989; Murakami and Van de Woude, 1998; Murakami et al., 1999). Deadenylation of c-mos mRNA (Paillard et al., 1998) after c-mos degradation (Lorca et al., 1993) could play an important role in the elimination of c-mos gene products during early development of *Xenopus laevis* (Ueno and Sagata, 2002).

The EDEN-dependent deadenylation of maternal mRNA depends on the presence of a trans-factor, EDEN binding protein (EDEN-BP) (Paillard et al., 1998). EDEN-BP is a homologue of human Nab50, or CUG-BP protein (Timchenko...
et al., 1996). This protein was shown to be involved in maturation of mRNA in human somatic cells (Philips et al., 1998). It is a phosphoprotein with a high potential to be hyper- or hypophosphorylated. Phosphorylation and dephosphorylation of CUG-BP was thought to be one of the key mechanisms involved in regulation of the localization (nuclear versus cytoplasmic) and activity of this protein in mRNA maturation (Roberts et al., 1997).

Because EDEN-dependent deadenylation is active in early Xenopus embryos, whereas the quantities of EDEN-BP protein are similar before and after fertilization (Paillard et al., 1998), we investigated whether post-translational modifications, like phosphorylation or dephosphorylation, regulate this sequence-specific deadenylation activity. In this paper we show that EDEN-dependent deadenylation is sensitive to kinase and phosphatase inhibitors. EDEN-BP is phosphorylated during oocyte maturation and dephosphorylated following egg activation. Dephosphorylation of EDEN-BP is calcium dependent and correlates with the increase in the activity of EDEN-dependent deadenylation. It suggests that these post-translational modifications could regulate the efficiency of EDEN-dependent deadenylation. MPF reactivation following egg extract activation does not influence EDEN-dependent deadenylation, suggesting that the regulation does not depend directly on MPF activity. These results link calcium signals and the cell cycle regulatory machinery to the deadenylation of certain maternal mRNAs at the very beginning of the embryonic development.

Materials and Methods

Oocytes and extracts

Oocytes were collected as described by Blot et al. (Blot et al., 2002). The stage VI oocytes were then incubated overnight in Merriam buffer (Merriam, 1971) [NaCl 88 mM, Ca(NO3)2 0.33 mM, KCl 1 mM, MgCl2 0.41 mM, MgSO4 0.82 mM and HEPES 10 mM, pH 7.5]. Maturation was induced by addition of progesterone to 1 µM final concentration in the buffer. Oocytes were collected at different times, frozen in liquid nitrogen and kept at –70°C.

Unfertilized eggs were dejellied with 2% L-cysteine pH 8.2 in XB buffer (Murray, 1991) (KCl 100 mM, MgCl2 2 mM, HEPES 10 mM, pH 7.65, sucrose 50 mM) and washed in XB. Some of the eggs were collected and the others were activated using 0.5 mg/ml calcium ionophore A23187 for 2 minutes and then extensively washed in XB. Eggs collected at different times after activation were frozen in liquid nitrogen and kept at –70°C.

Extracts were prepared by the addition of homogenization buffer [β-glycerophosphate 60 mM, p-nitrophenyl phosphate 15 mM, MOPS 25 mM pH 7.2, EGTA 15 mM, MgCl2 15 mM, dithiothreitol 2 mM, Na-orthovanadate 1 mM, NaF 1 mM, di-Na-phosphophosphate 1 mM, leupeptin 10 µg/ml, aprotinin 10 µg/ml, chymotatin 10 µg/ml and pepstatin 10 µg/ml; 200 µl for 40 oocytes] to the thawing oocytes and pipetting. After centrifugation at 12,000 g at 4°C the supernatant was used for the experiment.

CSF extracts were prepared using joint protocols previously described (Lohka and Masui, 1984) and Blow (Blow, 1993) with few modifications: additional EGTA up to 7 mM final concentration and sucrose 50 mM in the CSF extraction buffer (Blow et al., 1993) (KCl 50 mM, HEPES 50 mM pH 7.6, MgCl2 5 mM, EGTA 5 mM and DTT 2 mM) and two centrifugations at 15,000 g. Calcium-activated extracts were prepared from CSF extracts supplemented with CaCl2 (0.6 mM final concentration). To check the integrity of our extracts, 10-20 µl of extract were incubated at 21°C for 1 to 1.5 hours with sperm nuclei. The chromatin condensation (CSF extract) and chromatin decondensation (activated extract) was visualized by fluorescence and phase contrast microscopy using Hoechst dye in fixing media. In some cases we also measured histone H1 kinase activity or determined the state of phosphorylation of Cdc25 phosphatase by western blotting.

Calcium ionophore-activated embryo extract was prepared as described by Legagneux et al. (Legagneux et al., 1995).

Drugs and recombinant proteins

Extracts were supplemented with recombinant bacterial Δ90 sea urchin cyclin B to 0.2 ng/µl (Murray and Kirschner, 1989). Okadaic acid (OA) (Goris et al., 1989) was used at a concentration of 1 µM, and roscovitine (Meijer et al., 1997) at 100 µM.

Transcripts

Two plasmid constructs pEg2-410 (Legagneux et al., 1992) and pEg2-410a (Bouvet et al., 1994) were used. Both were derived from the 3′-UTR of Eg2/Aurora A mRNA. Eg-410 is a substrate for EDEN-dependent deadenylation. Eg2-410a RNA is a substrate for default deadenylation as it contains neither a CPE nor an EDEN. These plasmids were used to produce capped radiolabelled transcripts as previously described (Legagneux et al., 1992). The radiolabelled transcripts were purified by electrophoresis on a 4% polyacrylamide-urea gel.

In vitro deadenylation and RNA analysis

The deadenylation experiments were realized in vitro as previously described (Legagneux et al., 1995). In brief, 6 µl extracts were incubated with 1 µl capped radiolabelled transcripts Eg2-410 and Eg2-410a at 21°C for 1 to 3 hours. The transcripts were extracted as described by Harland and Mish (Harland and Mish, 1988), then precipitated and analysed by electrophoresis on 4% polyacrylamide-urea gel and phosphorimaging. 0.5 µl of radiolabelled transcripts were loaded directly on the gel without precipitation (lanes T) as a size marker for the input RNA, the starting point of deadenylation.

Immunoprecipitation

Anti-EDEN-BP sepharose beads were prepared with anti-EDEN-BP rabbit serum (Paillard et al., 1998) loaded on protein A-sepharose beads. The IgG were then covalently coupled to the beads using dimethyl pimelimidate coupling agent according to standard procedure.

Oocyte or unfertilized egg extracts were added to anti-EDEN-BP sepharose beads (200 µl of extract for 20 µl of beads) and incubated at 4°C for 40 minutes followed by extensive washing in bead buffer (Tris 50 mM pH 7.4, NaF 5 mM, NaCl 250 mM, EDTA 5 mM, EGTA 5 mM, Nonidet P-40 0.1%, leupeptin 10 µg/ml, aprotinin 10 µg/ml, chymostatin 10 µg/ml and pepstatin 10 µg/ml). The bound material was either used for further experiment or eluted at 90°C in elution buffer (Tris 62.5 mM pH 6.8, 2% sodium dodecyl sulfate) and collected after centrifugation. Concentrated Laemmli denaturating buffer was added to the eluted material. Samples were boiled before analysis.

Phosphatase assays

For these experiments the extracts were made in homogenization buffer without di-Na phenyl phosphate and p-nitrophenyl phosphate. The phosphatase λ (Lambda Protein phosphatase P0735S Biolabs) was incubated according to the manufacturer’s instructions with either nonactivated egg extract or ionophore-activated egg extract (40 units/µl of extract) or with EDEN-BP immunoprecipitated from unfertilized egg extract on sepharose beads (160 Unit/µl of beads).
The incubation was realized at 30°C for 30 minutes to 2 hours. The reaction was stopped by addition of Laemmli denaturing buffer. The samples were boiled before analysis.

Electrophoresis and western blotting

Protein was analysed on a sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS/PAGE) gel (Laemmli, 1970); when required the separating gel consisted of 15% acrylamide and 0.13% bisacrylamide (ratio 29:8:0:2) to allow better separation of different forms of EDEN-BP (Anderson et al., 1973). Otherwise, 10% SDS/PAGE gels were used (Laemmli, 1970). Separated proteins were transferred to nitrocellulose membranes (Hybond C; Amersham) according to standard procedures and probed with either purified rabbit polyclonal anti-EDEN-BP antibodies 1/1000 (nonimmunoprecipitated extracts), polyclonal anti-EDEN-BP guinea pig serum 1/3000 (immunoprecipitation experiments) or anti-Cdc25 (gift from Marcel Dorée, CRBM, Montpellier, France) and followed by alkaline phosphatase-coupled secondary antibody (Sigma) 1/30000.

Results

EDEN-dependent deadenylation is accelerated on calcium addition to the CSF extract

The influence of calcium on EDEN-dependant deadenylation was evaluated using CSF extracts prepared as described in Materials and Methods. The extracts were supplemented with CaCl2 to a final concentration of 0.8 mM, as well as with radiolabelled chimeric mRNAs: Eg2-410 that contains the EDEN sequence and Eg2-410a that does not contain the EDEN sequence. These RNAs are derived from the 3' UTR of Eg2 mRNA (Bouvet et al., 1994). The functional activity of these extracts was verified by measuring the accumulation of the poly(A)+ form of the chimeric RNAs. Samples were collected at 1, 2 and 3 hours after adding the radiolabelled transcripts. Deadenylation of EDEN-containing RNA is processive (Legagneux et al., 1995), which is shown by the accumulation of the deadenylated form, whereas the initial adenylated form persists (see A+ of Eg2-410 in Fig. 1A); default deadenylation is distributive (Legagneux et al., 1995), which is visible as a smear in the gel (see A+ of Eg2-410a in Fig. 1A). The analysis of the transcripts shows a weak EDEN-dependent deadenylation (Eg2-410 RNA) in CSF extract (Fig. 1A, left panel) that is substantially increased in extracts activated with 0.8 mM calcium (Fig. 1A, right panel). Only a slight change in the pattern of accumulation of deadenylated forms of Eg2-410a chimeric RNA was detected on calcium-induced activation of the extracts (Fig. 1A, compare Eg2-410a in left and right panels). Interphase versus M-phase state of extracts was checked by observing the chromatin condensing or decondensing activity; demembranated sperm nuclei were added followed by Hoechst staining and phase contrast observation (Fig. 1B). In CSF extracts we found condensed chromosomes forming metaphase plates (Fig. 1B, left), whereas interphase nuclei were formed in activated extracts (Fig. 1B, right). Therefore, the increase in calcium ion concentrations that induced activation of CSF extracts triggered a substantial acceleration of the EDEN-dependent deadenylation, whereas only a slight modification in the dynamics of default deadenylation was observed.

Calcium triggers changes in the electrophoretic mobility of EDEN-BP in CSF extract and M II oocytes

EDEN-BP migrates as a doublet of 55/53 kDa in SDS/PAGE gels (Legagneux et al., 1992). Therefore, we evaluated the effect of increasing the concentration of calcium ions in the CSF extracts on the electrophoretic mobility of EDEN-BP. Using Anderson’s modified SDS/PAGE gels (Anderson et al., 1973), we detected a more complex pattern of EDEN-BP electrophoretic mobility. Three bands (Fig. 2A-B; named a, b, c) were observed. The uppermost weak band (a) is only visible in CSF extracts collected before calcium addition. The second band (b) corresponds to the major band in the CSF extracts and its intensity decreases after calcium activation. This decrease appears sensitive to the amount of calcium added (compare extracts 3 hours after activation with 0.4, 0.8 and 1.6 mM CaCl2, Fig. 2A). The lower major band (c) is clearly observed in the activated extracts; however, it is also present as a minor band in CSF extracts. The intensity of this band increases in parallel with the increase in calcium concentration (compare extracts 3 hours after activation with 0.4, 0.8 and 1.6 mM CaCl2, Fig. 2A).

To determine whether the calcium-induced changes in the electrophoretic mobility of EDEN-BP occur in vivo, we studied the protein in parthenogenetically activated M II oocytes. The protein was detected by western blotting either directly (data not shown) or after immunoprecipitation (Fig. 2B). In nonactivated eggs (arrested in M II of meiosis), similarly to CSF extracts, three EDEN-BP bands (a, b, c) with a strong upper major band (b) and a very weak lower band (c), were detected (Fig. 2B, NA; compare with untreated CSF extracts, time 0, in Fig. 2A). In activated eggs the uppermost band (a) was absent, the intensity of major band (b)
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Fig. 2. Changes in the electrophoretic mobility of EDEN-BP is triggered by calcium. 
(A) Increasing amounts of CaCl2 were added to activate CSF extract. The extracts were 
incubated for 3 hours, analysed on 15% Anderson SDS/PAGE gels, blotted and reacted 
with anti-EDEN-BP antibody. (B) Proteins extracted from nonactivated eggs (NA) and 
calcium ionophore-activated eggs (A) were immunoprecipitated on proteinA-sepharose 
beads covalently coupled with anti-EDEN-BP antibodies. Extracts from nonactivated 
eggs were also loaded on control proteinA-sepharose beads that were not in contact with 
the antibody (C). The bound material was analysed by electrophoresis and 
immunoblotted with anti EDEN-BP antibodies. a, b, b´ and c indicate the different 
-electrophoretic forms of EDEN-BP.

The electrophoretic mobility changes in EDEN-BP occur during oocyte maturation and following egg activation

To characterize the dynamics of the changes in the electrophoretic mobility of EDEN-BP, we studied the protein 
in detail during oocyte maturation and following egg activation. Samples of maturing oocytes were collected every hour 
following progesterone addition (Fig. 3A) and at short intervals following parthogenetic activation of M II oocytes 
with calcium ionophore (Fig. 3B), and analysed as above. Stage VI prophase I-arrested oocytes contain two forms of 
EDEN-BP: the major lower band (c) and a smear of minor ones 

EDEN-BP is post-translationally modified by phosphorylation/dephosphorylation

To verify whether the electrophoretic changes of EDEN-BP were due to changes in the phosphorylation of the protein, 
extracts were treated with the phage λ phosphatase at 30°C for 
0, 0.5 or 1 hour. The samples were then analysed by SDS/PAGE and anti-EDEN-BP western blotting. The phage λ 
phosphatase treatment caused an increase in the electrophoretic mobility of EDEN-BP, whereas without phage λ 
phosphatase treatment, the major upper band (b) of EDEN-BP was still detectable after 1 hour of incubation at 30°C (Fig. 4A). To 
show the direct effect of the phage λ phosphatase on EDEN-BP, we immunoprecipitated EDEN-BP followed by λ 
phosphatase treatment. In these conditions a similar 
modification of EDEN-BP was observed (Fig. 4B). However, 
the conversion of the immunoprecipitated protein appeared to 
be less complete and residual amounts of bands (a) and (b) 
were sometimes observed. This reduced conversion to the 
dephosphorylated form could be due to a masking effect 
because EDEN-BP is complexed with the antibodies during the 
phosphatase treatment. The data presented show that the phage λ 
phosphatase treatment mimics to a large part the down shift
that OA inhibited the deadenylation process in calcium-EDEN-dependent deadenylation; no poly(A)– form of Eg2-410 (Meijer et al., 1997). We observed a complete inhibition of the concentration roscovitine inhibits a large spectrum of kinases roscovitine at high concentration (100 μM) added to CSF extract containing chimeric RNA was detected even after 3 hours of incubation (Fig. 5). Only slight changes (higher accumulation of intermediate forms of Eg2-410a chimeric RNA) in the dynamics of the default deadenylation were observed.

Kinase and phosphatase inhibitors downregulate EDEN-dependent deadenylation

A large number of kinases are inactivated on egg activation (Karsenti et al., 1987). The temporal correlation between the beginning of EDEN-BP dephosphorylation and the acceleration of EDEN-dependent deadenylation following egg activation suggests that the two processes, which are triggered by free calcium increase, could be due to a change in the equilibrium between kinases and phosphatases activities. Therefore, we determined whether kinase and phosphatase inhibitors could influence the process of EDEN-dependent mRNA deadenylation.

To inhibit a broad range of serine/threonine kinases we used roscovitine at high concentration (100 μM) added to CSF extract activated by the addition of CaCl2 (Fig. 5). At this concentration roscovitine inhibits a large spectrum of kinases (Meijer et al., 1997). We observed a complete inhibition of the EDEN-dependent deadenylation; no poly(A)+ form of Eg2-410 chimeric RNA was detected even after 3 hours of incubation (Fig. 5). Only slight changes (higher accumulation of intermediate forms of Eg2-410a chimeric RNA) in the dynamics of the default deadenylation were observed.

Similar experiments were performed by inhibiting a broad range of phosphatases. A mixture of concentrated phosphatase inhibitors was added to the CSF extract containing chimeric RNAs before CaCl2 addition. EDEN-dependent deadenylation was very efficiently arrested by these inhibitors both in CSF and calcium-activated extracts (data not shown).

Okadaic acid is a potent inhibitor of phosphatases 1 and 2A (Goris et al., 1989). Therefore, using this drug enabled us to identify the spectrum of phosphatases inhibiting EDEN-dependent deadenylation. CSF extracts supplemented with OA were incubated at 21°C for 15 minutes and then activated with CaCl2, and radiolabelled transcripts were added. We observed that OA inhibited the deadenylation process in calcium-activated extracts (Fig. 6A). The inhibition of EDEN-dependent deadenylation was correlated with the inhibition of the dephosphorylation of EDEN-BP (Fig. 6B). The dynamics of default deadenylation was only slightly modified by OA, although the accumulation of several partially deadenylated products became visible.

From these experiments, we can conclude that EDEN-dependent deadenylation is regulated by a balance between phosphorylation and dephosphorylation activities and that this process is sensitive to OA.

MPF pathway does not inhibit EDEN-dependent deadenylation

MPF is finely regulated via phosphorylation/dephosphorylation. Cdc25 phosphatase is a major activator of MPF and this phosphatase is itself activated by a series of phosphorylations on the entry into the M-phase (Kumagi and Dunphy, 1991; Gautier et al., 1991). MPF activity depends on the stability of cyclin B (Murray and Kirschner, 1989; Murray et al., 1989). Moreover, OA is known to indirectly activate MPF in Xenopus oocytes and extracts (Goris et al., 1989; Rime et al., 1990; Jessus et al., 1991; Lorca et al., 1991). Therefore, the sensitivity of EDEN-dependent deadenylation to OA could be due to activation of MPF. To determine whether this activity interferes with the process of EDEN-dependent deadenylation we specifically activated MPF via the addition of indestructible Δ90 cyclin B of sea urchin into CSF extracts followed by calcium treatment (Fig. 7A). Addition of exogenous Δ90 cyclin B into CSF extract treated with Ca2+ maintained condensed chromosomes (Fig. 7A, lower panel, compare left and middle images), indicating that the extract remained in M-phase despite calcium addition, whereas interphase nuclei were present in calcium-activated extract (Fig. 7A, lower panel, right). However, the high MPF activity in the extract had no effect on EDEN-dependent deadenylation (Fig. 7A, upper panel).


To check whether MPF activity could affect the deadenylation activity, which had already been accelerated on egg activation, we added Δ90 cyclin B to ionophore-activated embryo extract and followed EDEN-dependent deadenylation (Fig. 7B). In parallel, we checked the inhibitory effect of OA in the same extract, as well as the combined effect of both Δ90 cyclin B and OA (Fig. 7B). As expected, Δ90 cyclin B restored an M-phase state in the extract, which was confirmed in western blot with anti-Cdc25; the disappearance of the fast migrating (dephosphorylated) band is concomitant with the appearance of a new, upshifted band (Fig. 7B, lane 2 in the lower panel). The addition of Δ90 cyclin B did not affect EDEN-dependent deadenylation in this extract when compared with the control, interphase extract (Fig. 7B, upper panel, compare experiments 1 and 2). As already shown for the calcium activated CSF extract, 1 μM OA slowed this process and this reduced rate of deadenylation was not affected when the OA treatment extract was supplemented with Δ90 cyclin B (Fig. 7B, compare experiments 3 and 4). As expected, the combined effect of OA and Δ90 cyclin B treatment induced a more pronounced phosphorylation of Cdc25 (Fig. 7B, lower panel, lane 4). Such a combined treatment seems to have a synergistic effect on the M-phase state of the extract, but does not influence significantly EDEN-dependent deadenylation compared with the OA treatment alone (Fig. 7B, upper panel, compare experiments 3 and 4). From these experiments we conclude that EDEN-dependent deadenylation of RNA is not affected by MPF activity.

Discussion

Calcium signalling upregulates EDEN-dependent deadenylation on egg activation

Calcium plays a major role in egg activation on fertilization or parthenogenetic activation (Stricker, 1999; Runft et al., 2002). Transient rises in intracellular Ca²⁺ are of great importance for numerous biochemical and cell biological processes necessary for further development. Here, we showed that calcium-mediated activation of CSF extracts greatly enhanced EDEN-dependent deadenylation activity, as well as dephosphorylation of EDEN-BP, the only protein known so far to be involved in this process. Moreover, we showed that the same modifications of EDEN-BP occurred when M II oocytes were parthenogenetically activated via calcium ionophore treatment. Because Xenopus laevis EDEN-containing maternal mRNAs are deadenylated following fertilization (Paris et al., 1988; Paris and Philippe, 1990), we believe that the same calcium-dependent mechanism triggers EDEN-dependent deadenylation of maternal mRNAs on fertilization.

Dephenylation of maternal mRNAs that contain EDEN- sequence (Eg family and c-mos) (Bouvet et al., 1994; Paillard et al., 1998) results in their translational arrest (Ezzeddine et al., 2002; Ueno and Sagata, 2002). Therefore, our data provide a link between calcium signalling and translational repression of maternal genes during early development of Xenopus laevis. We also observed that an increase in the amount of calcium added to CSF extract, induced a more pronounced dephosphorylation of EDEN-BP. Fertilization generates transient intracellular calcium waves. Adding supraphysiological amounts of calcium (1.6 mM) to CSF extract may in one step induce rapid modifications of numerous biochemical processes that in concert enhance EDEN-BP dephosphorylation.

Phosphorylation and dephosphorylation modulate EDEN-dependent deadenylation of mRNA

Calcium waves that occur after fertilization provoke a rapid inactivation of CSF and MPF activity, resulting in metaphase to anaphase transition followed by the first embryonic interphase (Lorca et al., 1993) (reviewed by Maller, 1998). Inactivation of MPF via cyclin B degradation (Murray et al., 1989) triggers changes in the phosphorylation/ dephosphorylation equilibrium in favour of dephosphorylation (Karsenti et al., 1987). Inactivation of multiple kinases participates in this process and correlates with dephosphorylation of EDEN-BP. Using nondegradable sea urchin Δ90 cyclin B, which enables the MPF kinase activity to be artificially maintained in calcium-treated CSF extracts, or to activate MPF in interphase extracts, we show that the EDEN-dependent deadenylation is independent of MPF activity. In contrast, OA reduces the activity of EDEN-dependent deadenylation. OA is known to activate MPF and ERK2 MAP kinase besides inactivating phosphatases PP1 and PP2 (Jessus et al., 1991). Whether the effect of OA on EDEN-dependent deadenylation is via ERK2 activation or phosphatase inactivation is not clear at present. However, in preliminary experiments we observed that inactivation of ERK2 by the addition of recombinant phosphatase CL 100 to OA-treated extract restored the EDEN-dependent deadenylation activity (our unpublished results). In addition,
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the timing of MAP kinase ERK2 activation and inactivation during oocyte maturation and after fertilization correlates with phosphorylation and dephosphorylation of EDEN-BP (Ferrell et al., 1991; Abrieu et al., 1996; Chau and Shibuya, 1999). A positive feedback between MPF and ERK2 activation in CSF extracts through the phosphorylation of c-mos, an upstream regulator of the ERK2, was reported recently by Castro et al. (Castro et al., 2001). In some of our experiments in which Δ90 cyclin B was added to CSF extracts followed by calcium addition, we observed an intermediate rate of EDEN-dependent deadenylation compared with CSF extracts and calcium-activated extracts (data not shown). In these extracts we detected the phosphorylated form of ERK2 MAP kinase. This suggests that the feedback loop described by Castro et al. (Castro et al., 2001) is provoking ERK2 MAP kinase activation in these extracts, supposedly when c-mos was not totally degraded. Further studies are necessary to examine the potential role of ERK2 MAP kinase, and/or a combined role of MPF and MAP kinase pathways, in the regulation of EDEN-dependent deadenylation and, in particular, of EDEN-BP phosphorylation.

The insensitivity of EDEN-dependent deadenylation to MPF contrasts with the recent data by Groisman et al. (Groisman et al., 2002) showing that cyclin B1 mRNA polyadenylation is stimulated via MPF and that its deadenylation is induced by MPF inactivation. There is no proof that cyclin B1 3′-UTR contains an EDEN, suggesting that the deadenylation of this mRNA is independent of EDEN-BP. Therefore, two groups of maternal Xenopus mRNAs can be distinguished on the basis of their behavior regarding deadenylation during early

Fig. 7. MPF activation through Δ90 cyclin B addition to Ca2+, or ionophore-activated extracts does not influence EDEN-dependent deadenylation. A. Nondegradable sea urchin Δ90 cyclin B was added (+) or not (−) to CSF extract at final concentration of 0.2 ng/μl and incubated for 15 minutes at 21°C to allow its association with cdk1 and the formation of a pool of stable MPF. Both extracts were then supplemented with CaCl2 (to final concentration 0.8 mM) and incubated at 21°C for 1, 2 or 3 hours with radiolabelled transcripts Eg2-410 and Eg2-410a. RNA. Upper panel: autoradiography of the extracted RNAs separated on a urea/acrylamide 4% gel. The lane (T) corresponds to the fully adenylated radiolabelled transcripts without previous incubation with extract and precipitation. The two extracts deadenylate chimeric RNAs with similar dynamics independently from the presence or absence of Δ90 cyclin B. Lower panel: sperm nuclei were incubated with extracts and stained with Hoechst dye following observation in fluorescence and phase contrast. Condensed sperm chromosomes in the control CSF extract (left) and in the calcium activated CSF extract with Δ90 cyclin B (middle) were found at the end of the experiment, whereas decondensed nuclei were present in calcium activated CSF extract (right). This confirms that Δ90 cyclin B-supplemented extract was in M-phase similarly to the control, untreated CSF extract. B. Nondegradable sea urchin Δ90 cyclin B (final concentration 0.2 ng/μl), OA (final concentration 1 μM), or both, were added to ionophore-activated eggs extract prepared 45 minutes after egg activation. The extracts were incubated for 15 minutes at 21°C to allow association of exogenous cyclin with cdk1 and formation of stable MPF or to inactivate protein phosphatases in the case of OA. Then they were supplemented with radiolabelled Eg2-410 transcript and further incubated at 21°C for 3 hours. RNA and proteins were analysed as indicated in Materials and Methods. Upper panel: autoradiography of the extracted RNAs separated on urea/acrylamide 4% gel. The lane (T) corresponds to the fully adenylated radiolabelled transcripts without previous incubation with extract and precipitation. Untreated, as well as Δ90 cyclin B-supplemented extracts dedenylate Eg2-410 transcript rapidly (experiments 1 and 2), whereas OA- and OA+Δ90 cyclin B-supplemented extracts deadenylate only slightly Eg2-410 transcript (experiments 3 and 4). Lower panel: western blot with an anti-Cdc25 antibody. The fast migrating, lowest band represents dephosphorylated (inactive) form of Cdc25. The upshifted forms of Cdc25 produced by different treatments represent phosphorylated (active) forms of the Cdc25 phosphatase. Untreated extract contains inactive Cdc25 (lane 1), whereas Δ90 cyclin B-supplemented extract contains activated form of the phosphatase (lane 2) indicating induction of the M-phase. OA-treated and OA+Δ90 cyclin B-treated extracts contain phosphorylated forms of Cdc25 (lanes 3 and 4, respectively). In OA+Δ90 cyclin B containing extract (lane 4) Cdc25 is phosphorylated to higher degree than in OA-treated extract (lane 3), indicating a synergistic effect of Δ90 cyclin B and OA on Cdc25 phosphorylation.
development: namely, MPF- or cell-cycle-dependent deadenylation, like cyclin B1 mRNA (Groisman et al., 2002), and MPF-independent, but activation-dependent deadenylation, like Eg2 (this paper). Because calcium is as an important regulator of the cell cycle, we cannot exclude that mRNA deadenylation is controlled by cytoplasmic calcium in both cases.

EDEN-BP phosphorylation and dephosphorylation seems a very complex process. We could distinguish four different forms of this protein on western blots. The complex pattern of EDEN-BP phosphorylation suggests that multiple phosphorylation sites are modified during oocyte maturation and following egg activation. The inhibitory effects of both kinase and phosphatase inhibitors on EDEN-dependent deadenylation supports the hypothesis of a network of kinases and phosphatases controlling this process.

Could EDEN-BP phosphorylation regulate EDEN-dependent deadenylation?

EDEN-BP is a good candidate for the regulatory molecule involved in the modulation of EDEN-dependent deadenylation on M II oocyte fertilization or activation. After calcium ionophore-induced activation this protein undergoes a series of dephosphorylations. The hyperphosphorylated band (a) disappears rapidly, whereas the intensity of the dephosphorylated band (c) increases to reach a maximum 2.5 to 3 hours after egg activation. The intensity of the hyperphosphorylated band (b), which is the major band in CSF extracts, decreases gradually and disappears 2 hours after activation. This correlation suggests that phosphorylation of EDEN-BP could be involved in the inhibition of EDEN-dependent deadenylation. However, EDEN-BP is part of a multimeric complex of high molecular weight (Legagneux et al., 1995) that probably contains several other factors required for deadenylation. Therefore, at present we can not exclude that changes in the phosphorylation of one or several of these factors are not involved in the regulation of EDEN-dependent deadenylation. Identification of the amino acids of EDEN-BP that are phosphorylated on oocyte maturation and studies of mutant proteins will permit us to evaluate the exact role of EDEN-BP phosphorylation and dephosphorylation during oocyte maturation and early embryonic development.

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