Requirement for Atr in phosphorylation of Chk1 and cell cycle regulation in response to DNA replication blocks and UV-damaged DNA in Xenopus egg extracts

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The checkpoint kinase Xchk1 becomes phosphorylated in Xenopus egg extracts in response to DNA replication blocks or UV-damaged DNA. Xchk1 is also required for the cell cycle delay that is induced by unreplicated or UV-damaged DNA. In this report, we have removed the Xenopus homolog of ATR (Xatr) from egg extracts by immunodepletion. In Xatr-depleted extracts, the checkpoint-associated phosphorylation of Xchk1 is abolished, and the cell cycle delay induced by replication blocks is strongly compromised. Xatr from egg extracts phosphorylated recombinant Xchk1 in vitro, but not a mutant form of Xchk1 (Xchk1-4AQ) containing nonphosphorylatable residues in its four conserved SQ/TQ motifs. Recombinant human ATR, but not a kinase-inactive mutant, phosphorylated the same sites in Xchk1. Furthermore, the Xchk1-4AQ mutant was found to be defective in mediating a checkpoint response in egg extracts. These findings suggest that Xchk1 is a functionally important target of Xatr during a checkpoint response to unreplicated or UV-damaged DNA.

[Key Words: Atr; Chk1; phosphorylation; replication checkpoint; mitosis]

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During the cell cycle, the fidelity of DNA synthesis and the occurrence of DNA damage are monitored by checkpoint pathways, which ensure that chromosomal DNA is accurately replicated and transmitted in an undamaged form to daughter cells (Elledge 1996; O’Connell et al. 2000). Many components of such checkpoint pathways have been identified from yeast to vertebrates. In the fission yeast Schizosaccharomyces pombe, a group of eight proteins (Rad1, Rad3, Rad9, Rad17, Rad26, Hus1, Cut5, and Crb2) is thought to participate in detecting structures characteristic of DNA damage and/or incomplete DNA replication. Homologous components exist in the budding yeast Saccharomyces cerevisiae and higher eukaryotes, including Xenopus and humans (Elledge 1996). Among these sensor proteins, Rad3 encodes a protein kinase that plays a pivotal role in checkpoint signaling (Bentley et al. 1996; O’Connell et al. 2000). Its budding yeast and human homologs are Mec1 and ATR, respectively (Cimprich et al. 1996; Keegan et al. 1996). Rad3, Mec1, and ATR belong to a larger subfamily of protein kinases with a phosphoinositide kinase (PIK)-related domain at the C terminus. A member of this family, ATM, is mutated in the disease ataxia telangiectasia (AT; Lavin and Shiloh 1997). Cells derived from AT patients exhibit defective checkpoint responses to DNA damage induced by ionizing radiation (IR). The budding yeast homolog of ATM, Tel1, is also involved in damage responses (Sanchez et al. 1996). Another member, DNA-PKcs, is well characterized as a protein kinase that binds to DNA and is activated by double-stranded DNA ends (Smith and Jackson 1999).

After detection of damaged DNA or specific DNA-replication structures, a signal is transduced to effector molecules, including the protein kinases Chk1 and Cds1 (Elledge 1996). Yeast Rad3 or Mec1 are absolutely required for signaling through the effector kinases Chk1 and Cds1/Rad53, which are phosphorylated because of the presence of DNA damage or replication blocks. In human cells, ATM controls the phosphorylation of the Cds1 homolog Chk2 after exposure to ionizing radiation but not ultraviolet (UV) light or DNA replication blocks (Matsuoka et al. 1998; Blasina et al. 1999a; Brown et al. 1999; Chaturvedi et al. 1999; Tominaga et al. 1999).
In contrast to ATM, ATR is essential for early embryonic development of mice [Brown and Baltimore 2000; de Klein et al. 2000]. Overexpression of a kinase inactive ATR in human fibroblasts causes increased sensitivity to IR, UV, and hydroxyurea [HU] and abrogates the cell cycle arrest after DNA damage [Cliby et al. 1998; Wright et al. 1998]. Recently, evidence has been presented that ATR regulates the phosphorylation of human Chk1 in response to DNA damage [Liu et al. 2000]. Both ATM and ATR phosphorylate p53, and ATM phosphorylates Nbs1 and Brca1 [Banin et al. 1998; Canman et al. 1998; Cortez et al. 1999; Tibbetts et al. 1999; Gatei et al. 2000; Lim et al. 2000]. Nonetheless, the functional effects of these phosphorylations have yet to be elucidated. Chk1 and Chk2/Cds1 phosphorylate Cdc25C at a 14–3–3 binding site, leading to its cytoplasmic sequestration through the binding of 14–3–3 proteins [Kumagai and Dunphy 1999; Lopez-Girona et al. 1999; Yang et al. 1999; Zeng and Piwnica-Worms 1999]. Chk1 and Chk2/Cds1 also phosphorylate p53 on Ser 20, resulting in stabilization of p53 [Chehab et al. 2000; Hirao et al. 2000; Shieh et al. 2000]. The p53 target gene product 14–3–3, induced by DNA damage through the function of p53, interacts with Cdc2-cyclin B1 in the cytoplasm [Chan et al. 1999]. The cytoplasmic sequestration of Cdc25C and Cdc2-cyclin B1 appears to contribute, at least in part, to DNA checkpoint-induced cell cycle arrest mechanisms.

Previously, by using Xenopus egg extracts, we have demonstrated that the effector kinases Cds1 and Chk1 respond to quite different signals from the genome. Xenopus Cds1 [Xcds1] becomes phosphorylated and activated in response to DNA molecules with double-stranded ends [Guo and Dunphy 2000], whereas Xenopus Chk1 [Xchk1] responds to DNA replication blocks and UV damage [Kumagai et al. 1998]. Similar, but less restricted, responses of Cds1 and Chk1 to DNA signals have been observed in other vertebrates, including mice and humans [Sanchez et al. 1997; Matsuoka et al. 1998; Blasina et al. 1999a; Brown et al. 1999; Tominaga et al. 1999; Piras et al. 2000; Tsuchiya et al. 2000; Liu et al. 2000]. To explore the role of Atr in these pathways, we have cloned and characterized a Xenopus homolog of ATR (Xatr). Xatr binds to DNA molecules in egg extracts, and this form of Xatr displays greatly increased kinase activity. Immunodepletion of Xatr from egg extracts abolishes the response of Xchk1 to incompletely replicated and UV-damaged DNA, whereas Xcds1 still responds to double-stranded DNA ends in Xatr-depleted extracts. We present evidence that Xchk1 is a direct and functionally important target of Xatr in the unreplicated/UV-damaged DNA checkpoint(s).

**Results**

**Isolation of Xenopus Atr**

To investigate the function of ATR in cell cycle regulation, we have isolated a cDNA encoding the Xenopus homolog of ATR [Xatr] by using a degenerate polymerase chain reaction [PCR] approach. Through multiple rounds of library screening, three overlapping Xenopus oocyte cDNAs were isolated that contain ~90% of the COOH-terminal coding region and the 3´ untranslated region. The NH2-terminal 10% of the coding sequence and the 5´ untranslated region (~1.2 kb) were obtained by reverse transcription–coupled PCR using Xenopus oocyte total RNA as template. The four cDNA sequences were ligated to generate a product that encodes a 301-kD protein containing 2654 amino acids. As the 5´ untranslated region of this cDNA contains three in-frame translation termination codons, its open-reading frame most likely represents the full-length coding region. Xatr is most related to human ATR [70% identical and 80% similar] and is 29%, 28%, and 23% identical to Dro sophila melanogaster Mei-41, S. pombe Rad3, and S. cerevisiae Mec1, respectively. The COOH-terminal kinase domain, shown in Figure 1A, is the most conserved region among the ATR homologs.

**Xatr binds to DNA in egg extracts**

To functionally characterize Xatr, polyclonal antibodies were generated by using either the truncated fusion protein His6-Xatr [2351–2654] or a 14 amino acid peptide [EKTNPKGTRGEPK, residues 1617–1630] as the antigen. Affinity-purified antibodies recognized a ~300-kD protein in Xenopus egg extracts [Fig. 1B, lane 1]. The antibodies also recognized a recombinant GST [glutathione S-transferase]-Xatr fusion protein expressed in budding yeast [Fig. 1B, lane 2]. GST–Xatr migrated about 30 kD larger than endogenous Xatr on SDS-PAGE gels, which corroborates the fact that the Xatr cDNA is full length.

A noteworthy characteristic of the PIK-related protein kinases ATM, ATR, and DNA-PKcs is that they bind to DNA and/or associate with chromosomes, although it remains to be determined whether this association represents direct contact. DNA-PKcs binds to DNA through two accessory proteins Ku70 and Ku80, and its kinase activity is greatly increased by double-stranded DNA ends [Smith and Jackson 1999]. It has been reported that both ATR and ATM are associated with meiotic chromosomes [Keegan et al. 1996]. Evidence has also been presented that ATM binds to double-stranded DNA and that this interaction is enhanced by treatment of the DNA with ionizing radiation or restriction enzymes [Smith et al. 1999, Suzuki et al. 1999]. To test whether Xatr is bound to DNA molecules in Xenopus egg extracts, we incubated DNA cellulose [M13 single-stranded DNA or pBluescript plasmid DNA] with interphase egg cytosol. After extensive washing, the Xatr protein was found to associate with both types of DNA cellulose but not with control cellulose [Fig. 2A]. Addition of the DNA polymerase inhibitor aphidicolin or protease inhibitors to cytosol did not significantly enhance the binding. The association of Xatr with DNA cellulose was reduced if the resin was treated with DNase I after incubation with cytosol [Fig. 2B], indicating that DNA digestion had partially released Xatr. As a control,...
RPA70, the largest subunit of the RPA complex, which binds tightly to single- and double-stranded DNA in egg extracts (Adachi and Laemmli 1992), also specifically bound to DNA cellulose, and its binding was reduced by DNaseI treatment (Fig. 2B).

Because DNA interacts with cellulose through physical adsorption, it can be released by a variety of conditions, under which tightly bound proteins may still associate with DNA and might be coimmunoprecipitated because of bridging DNA. Indeed, RPA70 was found to coimmunoprecipitate with Xatr in 0.5% NP-40 eluates (Fig. 2C). In contrast, RPA70 could not be coimmunoprecipitated with Xatr after elution from DNA cellulose by DNaseI treatment (Fig. 2C).

Enhanced kinase activity of Xatr isolated with DNA cellulose

Next, we characterized the kinase activity of Xatr and its potential regulation by DNA. By using affinity purified antibodies, we found that anti-Xatr immunoprecipitates from Xenopus egg extracts contained a kinase activity that phosphorylated the model substrate protein PHAS-I in vitro (Fig. 2D, lane 1). Control immunoprecipitates with nonspecific IgG did not contain this activity (Fig. 2D, lane 6). Furthermore, the Xatr-associated kinase activity was efficiently inhibited by caffeine (half-maximal inhibition at about 0.9 mM caffeine; Fig. 2D), which is consistent with the reports that the PIK-related kinases ATR, ATM, and TOR are sensitive to caffeine, though to different extents (Blasina et al. 1999b; Hall-Jackson et al. 1999; Sarkaria et al. 1999). Significantly, caffeine also inhibits the phosphorylation of Xchk1 in response to unreplicated DNA in aphidicolin-treated Xenopus egg extracts at essentially the same half-maximally effective dose (data not shown), indicating that Xatr and the kinase activity that phosphorylates Xchk1 in these extracts display similar sensitivities to caffeine. To investigate the role of binding to DNA on Xatr-associated kinase activity, we incubated egg cytosol with DNA cellulose to allow the binding of Xatr, digested the DNA with DNaseI, and then immunoprecipitated the released...
Xatr with anti-Xatr antibodies. Anti-Xatr immunoprecipitates prepared in this manner displayed approximately 10- to 20-fold higher kinase activity toward PHAS-I than Xatr that was immunoprecipitated directly from egg cytosol (Fig. 2E). Addition of DNA templates (e.g., M13 and pBluescript) directly to anti-Xatr immunoprecipitates from egg cytosol did not result in elevation of Xatr-associated kinase activity (data not shown). Therefore, the DNA-cellulose procedure either results in activation of Xatr or is useful for the isolation of a population of Xatr that is already activated in the extracts.

Response of Xchk1 to DNA replication blocks or UV-damaged DNA is dependent on Xatr

To study the signaling pathways between Xatr and potential downstream effectors, we immunodepleted Xatr from Xenopus egg extracts (Fig. 3). As shown in Figure 3A, Xatr could be completely removed from these extracts with polyclonal anti-Xatr antibodies. Normally, the Xchk1 protein in the nuclear fraction of egg extracts undergoes extensive phosphorylation in response to the DNA polymerase inhibitor aphidicolin or to the presence of UV-damaged DNA (Kumagai et al. 1998; Fig. 3B). However, in Xatr-depleted extracts, the phosphorylation of Xchk1 in the presence of aphidicolin or UV-damaged DNA was completely abolished in comparison with mock-depleted extracts (Fig. 3B). In parallel, we assessed the role of Xatr in the phosphorylation of the Xenopus Cds1 homolog Xcds1. As described recently, Xcds1 responds to DNA templates distinct from those that elicit phosphorylation of Xchk1 (Guo and Dunphy 2000). Xcds1 is not affected by unreplicated or UV-damaged DNA. Instead, Xcds1, but not Xchk1, becomes highly phosphorylated in response to DNA replication blocks or UV-damaged DNA.
phosphorylated in response to linearized plasmids, double-stranded oligonucleotides, M13 DNA, and poly(dT)$_{40}$. As M13 DNA, which is partially nicked, and poly(dT)$_{40}$ are efficiently replicated to a double-stranded form in egg extracts, it appears that only DNA molecules with double-stranded ends can bring about the phosphorylation of Xcds1 (Guo and Dunphy 2000). As shown in Figure 3C, the phosphorylation of Xcds1 in response to M13 DNA or poly(dT)$_{40}$ was not abolished by removal of Xatr from egg extracts. Therefore, immunodepletion of Xatr causes a selective defect in the phosphorylation of Xchk1.

To pursue these observations further, we examined whether immunodepletion of Xatr would affect cell cycle progression in egg extracts undergoing a checkpoint delay. For this purpose, we incubated Xatr-depleted extracts or mock-depleted extracts in the absence or presence of aphidicolin [Fig. 3D]. We observed that the Xatr-depleted extracts treated with aphidicolin entered mitosis much sooner than mock-depleted extracts containing this replication inhibitor, indicating that removal of Xatr had significantly compromised the checkpoint response to unreplicated DNA. Interestingly, however, aphidicolin still induced a partial delay in Xatr-depleted extracts in comparison with extracts lacking aphidicolin. As described previously, this characteristic is shared with aphidicolin-treated extracts from which Xatr1 has been immunodepleted, as the replication checkpoint in this system involves both Xatr1-dependent and Xatr1-independent pathways [Kumagai et al. 1998]. Thus, the response of Xatr-depleted extracts to DNA replication blocks is strongly compromised but not completely abolished, as is also the case for Xcds1-depleted extracts.

**Xatr and human ATR phosphorylate Xchk1 in vitro**

As the checkpoint-associated phosphorylation of Xchk1 did not occur in Xatr-depleted extracts, we asked whether Xchk1 could be a substrate of Xatr. To investigate this possibility, we incubated a kinase-inactive GST-Xchk1(306–352) protein with Xatr that had been immunoprecipitated from DNA cellulose eluates. As shown in Fig. 4, the GST-Xchk1(306–352) protein was strongly phosphorylated by anti-Xatr immunoprecipitates, but not by control immunoprecipitates. It is known that DNA-PKcs, ATM, and ATR display a preference to phosphorylate SQ or TQ motifs in their substrates [Kim et al. 1999]. Xchk1 contains one TQ [Thr 314] and three SQ [Ser 344, Ser 356, and Ser 365], which contain Thr 314, Ser 344, and Ser 352 sequences in an amino acid region. To assess whether Xatr could phosphorylate these sites, we prepared a modified version of GST-Xchk1(306–352) in which Thr 314, Ser 344, Ser 356, and Ser 365 sequences in a 52-amino acid region. To determine whether Xatr could phosphorylate these sites, we prepared a modified version of GST-Xchk1(306–352) in which Thr 314, Ser 344, Ser 356, and Ser 365 sequences in a 52-amino acid region. To assess whether Xatr could phosphorylate these sites, we prepared a modified version of GST-Xchk1(306–352) in which Thr 314, Ser 344, Ser 356, and Ser 365 sequences in a 52-amino acid region. To determine whether Xatr could phosphorylate these sites, we prepared a modified version of GST-Xchk1(306–352) in which Thr 314, Ser 344, Ser 356, and Ser 365 sequences in a 52-amino acid region.
Phosphorylation of Xchk1 in egg extracts is required for the DNA replication checkpoint

To evaluate the contribution of the SQ/TQ domain to the function of Xchk1, we carried out the following experiments. First, we examined the phosphorylation of these motifs in *Xenopus* egg extracts undergoing a checkpoint delay. As one approach, $^{35}$S-labeled wild-type

Xchk1 and various SQ/TQ mutant Xchk1 proteins were incubated in egg extracts that either contained or lacked aphidicolin. Subsequently, the radiolabeled Xchk1 proteins were isolated from the nuclear fraction of the extracts and analyzed for phosphorylation-dependent shifts in mobility during gel electrophoresis. As shown in Figure 6A, in the case of the S344A and S356A mutants, the aphidicolin-induced phosphorylation of Xchk1 was strongly reduced. The T314A and S365A mutants still underwent substantial phosphorylation under these conditions. Finally, the checkpoint-dependent upshift in electrophoretic mobility was abolished in the Xchk1-4AQ mutant in which Thr 314, Ser 344, Ser 356, and Ser 365 were all changed to alanine.

In another method, we used antiphosphopeptide antibodies to monitor the phosphorylation of Ser 344 in the presence and absence of aphidicolin [Fig. 6B,C]. The Ser 344 site was chosen because mutagenesis of this residue caused a severe reduction in the aphidicolin-induced mobility shift of Xchk1. Also, Ser 344 resides in the most highly conserved of the SQ/TQ motifs among the *Xenopus*, human, and *Drosophila* Chk1 proteins. For this experiment, we used recombinant, baculovirus-expressed Xchk1-WT-GST-His6 and Xchk1-S344A-GST-His6 proteins that were double-tagged at the C-terminal end. Xchk1-WT-GST-His6 and Xchk1-S344A-GST-His6 were incubated in egg extracts in the absence or presence of aphidicolin for 90 min, reisolated with glutathione agarose, and then subjected to immunoblotting with either anti-GST antibodies to detect recombinant Xchk1 [Fig. 6C, top panel] or anti-S344-p antibodies to detect phosphorylation of Ser 344 [Fig. 6C, bottom panel]. The results of this analysis indicated that wild-type Xchk1 is phosphorylated on Ser 344 in the presence of but not in the absence of aphidicolin [Fig. 6C, bottom, lanes 1,2]. In contrast, the anti-S344-p antibodies did not react with the S344A mutant of Xchk1 in either condition [Fig. 6C, bottom, lanes 3,4].

To assess the physiological significance of phosphorylation at the SQ/TQ motifs, we examined whether the mutant could function in the DNA replication checkpoint in egg extracts. For this purpose, we removed endogenous Xchk1 completely from egg extracts by immunodepletion with anti-Xchk1 antibodies [Fig. 7A, lane 3]. Next, we divided the Xchk1-depleted extract into three aliquots and added baculovirus-expressed wild-type Xchk1, the Xchk1-4AQ mutant, or no additional protein [Fig. 7A, lanes 3–5]. Finally, we treated the extracts with aphidicolin and monitored the timing of mitosis [Fig. 7B]. Consistent with previous results, the extract containing no Xchk1 entered mitosis inappropriately [Kumagai et al. 1998]. As expected, the depleted extract that was restored with recombinant wild-type Xchk1 remained in interphase for at least 3.5 h [Kumagai et al. 1998]. By contrast, the extract containing the recombinant Xchk1-4AQ mutant underwent mitosis at the same time as the extract lacking Xchk1, indicating that this mutant of Xchk1 cannot respond to the presence of incompletely replicated DNA.
In this report, we have provided evidence that Xchk1 is a direct target of Xatr during a checkpoint response to unreplicated or UV-damaged DNA. Immunodepletion of Xatr from Xenopus egg extracts abolishes the phosphorylation of Xchk1 that occurs when chromosomal DNA replication is blocked by treatment with aphidicolin or when the DNA is damaged by exposure to UV radiation. Furthermore, removal of Xatr from egg extracts greatly compromises the cell cycle delay that is induced by DNA replication blocks. Both Xatr and human ATR phosphorylate Xchk1 in vitro at a number of SQ/TQ sites. Moreover, a mutant of Xchk1 that lacks these sites, and thus cannot serve as substrate for Xatr, is unable to act as an effector of the DNA replication checkpoint in Xenopus egg extracts. Taken together, these findings indicate that Xchk1 is a functionally critical substrate of Xatr.

The cellular response to damaged or incompletely replicated DNA includes a group of proteins that are highly conserved from yeast to humans. The functions of these proteins have been analyzed by treating cells with DNA damaging agents, such as methylmethane sulfonate (MMS, which creates single- and double-stranded DNA breaks), IR [which induces double-stranded breaks effectively], and UV light [which leads to the formation of pyrimidine dimers; Lindahl and Wood 1999]. Agents such as HU and aphidicolin elicit DNA replication blocks initially and may subsequently induce DNA damage. The respective roles of the effector kinases Chk1 and Cds1 in response to the DNA signals generated by such agents or treatments display differences between lower eukaryotes and metazoans. In fission yeast, for example, Chk1 normally responds to DNA damage (induced by MMS, IR, and UV), but not to inhibition of replication with HU, except in cells lacking Cds1 (Walworth and Bernards 1996; Brondello et al. 1999). Conversely, fission yeast Cds1 is activated by blockage of DNA replication (Murakami and Okayama 1995; Lind...
say et al. 1998; Brondello et al. 1999) but is also regulated by DNA damage that is inflicted during S-phase.

The situation appears to be somewhat different in metazoans. For instance, Xchk1 in *Xenopus* egg extracts responds to aphidicolin and UV-treated DNA, which is strongly impaired for replication [Kumagai et al. 1998], but not double-stranded DNA ends (Guo and Dunphy 2000). Likewise, the *Drosophila* Chk1 homolog Grapes is required for the replication checkpoint in early embryos [Fogarty et al. 1997; Sibon et al. 1997]. In embryonic mouse cells lacking Chk1, checkpoint responses to IR, UV, and aphidicolin are compromised [Liu et al. 2000; Takai et al. 2000].

A variety of observations have indicated that vertebrate Cds1 homologs operate in pathways that are involved in sensing double-stranded DNA breaks. In the human system, Chk2/Cds1 is activated on exposure of cells to IR, UV light, and HU. However, the IR-induced activation of Chk2/Cds1 is the strongest, and this response is abolished in cells deficient for ATM, which is clearly involved in a double-stranded DNA break pathway [Matsuoka et al. 1998; Brown et al. 1999]. Significantly, the weaker phosphorylation of Chk2/Cds1 in response to HU and UV light is independent of ATM. Optimal phosphorylation of other ATM substrates such as p53 and Brca1 also appears to require double-stranded breaks [Banin et al. 1998; Canman et al. 1998; Cortez et al. 1999]. Taken together, these investigations indicate that ATM controls the activation of Chk2/Cds1 in response to double-stranded DNA breaks.

In *Xenopus* egg extracts, the Cds1 homolog Xcds1 undergoes activation in response to DNA templates, including relatively simple oligonucleotides, that contain double-stranded ends (Guo and Dunphy 2000). Xcds1 does not respond to the presence of aphidicolin [DNA replication blocks] or UV-treated DNA, which does not undergo replication [Kumagai et al. 1998]. Because Xatr functions upstream of Xchk1, which does not respond to double-stranded DNA ends, the signal that the Xatr-dependent pathway recognizes is probably distinct from double-stranded breaks. Xatr binds well to either single-stranded or double-stranded DNA cellulose in *Xenopus* egg extracts. M13 phage DNA and the plasmid pBlueScript were used as the single- and double-stranded DNA templates, respectively. Double-stranded DNA ends do not appear to be important for the binding of Xatr to DNA because digestion of the double-stranded plasmid with restriction enzymes did not promote the association of Xatr with the cellulose resin [data not shown]. In contrast, double-stranded breaks enhance the interaction of ATM with DNA [Smith et al. 1999; Suzuki et al. 1999], providing another line of evidence that ATM recognizes this structure.

In *Xenopus* egg extracts, single-stranded DNA is a very good template for replication [Mechali and Harland 1982]. However, double-stranded DNA can undergo strand separation in these extracts [Gaudette and Benbow 1986]. Thus, it remains to be determined what structure Xatr actually recognizes, directly or indirectly, and how this interaction leads to the regulation of Xatr. This structure might resemble a signal that accumulates in aphidicolin-treated nuclei in *Xenopus* egg extracts. Exposure to aphidicolin does not block the firing of replication origins in *Xenopus* sperm chromatin but, rather, arrests DNA synthesis at some point subsequent to the priming stage [Mahbubani et al. 1997]. Thus, some aspects of stalled DNA replication forks may lead to the Xatr-dependent phosphorylation of Xchk1.

There appears to be significant insulation between the Chk1 and Cds1 pathways in this vertebrate system, al-
though it is possible that there is some cross-talk that is beyond our detection. As shown here, Xatr is an upstream regulator of Xchk1 [Fig. 8]. Xatr appears not to be a major regulator of Xcds1. In the human system, ATM controls Chk2/Cds1 and the response to double-stranded DNA breaks (Matsuoka et al. 1998; Brown et al. 1999; Tominaga et al. 1999; Chehab et al. 2000; Hirao et al. 2000; Shieh et al. 2000). It is not known whether Xcds1 is a target of Xenopus Atm (Robertson et al. 1999), but this seems plausible. Both the DNA replication blocks \( \rightarrow \) Atr \( \rightarrow \) Chk1 and double-stranded DNA breaks \( \rightarrow \) Atm \( \rightarrow \) Cds1 pathways appear to possess Cdc25 as one ultimate regulatory target. However, some insulation might be required if these pathways had other separate targets that must respond to certain DNA structures but need to ignore others.

In conclusion, we have provided evidence that Xatr regulates Xchk1 during operation of the checkpoint that detects incompletely replicated or UV-damaged DNA. Xatr phosphorylates Xchk1 at a number of conserved SQ/TQ motifs, especially the Ser 344 site. Immunodepletion of either Xatr or Xchk1 from egg extracts abrogates the DNA replication checkpoint to a similar extent. Similarly, egg extracts containing a mutant of Xchk1 that can not be phosphorylated by Xatr are indistinguishable from Xchk1-depleted extracts in their responsiveness to unreplicated DNA. These observations provide a strong argument that Xatr and Xchk1 are functionally linked in a common regulatory pathway. The involvement of Xatr and Xchk1 in a vital function such as monitoring the success of DNA replication would be consistent with the fact that both ATR and Chk1 are essential for early embryonic viability in the mouse (Brown and Baltimore 2000; de Klein et al. 2000; Liu et al. 2000; Takai et al. 2000).

Materials and methods

Cloning of a cDNA encoding Xatr

An internal fragment [140 bp] of a cDNA encoding Xatr was obtained by PCR using the degenerate oligonucleotides CCG GAATTGAT/\( C/GCl[\( A/G\)]/C/G\)T/TIAATGG and CGCGGATCCGCC[A/G]CA/C/T/TCTC[\( A/G\)]T, which were designed according to conserved areas in ATR homologs (Fig. 1). The PCR reactions contained Xenopus oocyte cDNAs as template, 50 pmol of the degenerate oligonucleotides, 200 µM of dNTPs, and 0.5 units of Taq polymerase in the buffer supplied by the manufacturer (GIBCO BRL). PCR reactions were heated at 94°C for 2 min, followed by 30 cycles of amplification. Each cycle consisted of segments of 94°C for 1 min, 45°C for 1 min, and 72°C for 1 min. An extra 10 min were added to the 72°C extension step for the last cycle. The 140-bp probe was used to screen a Xenopus oocyte cDNA library (Mueller et al. 1995). A 3-kb clone was isolated that includes the COOH-terminal sequence of Xatr and 3' untranslated region. By using the 5' end sequence [150 bp] of the 3-kb clone as the probe, a 1.1-kb overlapping clone was isolated from the same library. The 5' end sequence [150 bp] of the 1.1-kb fragment was labeled with \( ^{32}P \) and used to screen another Xenopus oocyte cDNA library (Kinoshita et al. 2000).
1995). A 4-kb clone was obtained that overlaps with the 1.1-kb fragment but not with the 3-kb clone. The NH$_2$-terminal sequence of Xatr and 5’ untranslated region were subsequently cloned by using the 5’ RACE system (GIBCO BRL).

**Antibody production**

An NdeI-EcoRI fragment encoding amino acids 2351–2654 of Xatr was produced by PCR and cloned into pET3 [Novagen]. The His$_6$-Xatr (2351–2654) protein encoded by this plasmid was expressed in *E. coli*, isolated with nickel agarose, purified further by SDS-PAGE, and used for production of antibodies at a commercial facility (Covance Research Products). Antibodies against an internal peptide of Xatr (EKTNPKPGRTEPK, residues 1617–1630) were generated at another commercial facility (Zymed Laboratories).

**Preparation of DNA cellulose and egg extracts**

M13 single-stranded DNA was prepared according to a protocol from the mutagenesis kit from Amersham (oligonucleotide-directed in vitro mutagenesis system version 2). The pBS plasmids were prepared by an alkaline lysis protocol [Sambrook et al. 1989]. To prepare DNA cellulose, 1 mg of M13 single-stranded DNA or pBluescript plasmid DNA dissolved in 1 mL of TE buffer (10 mM Tris-Cl, 1 mM EDTA at pH 8.0) was incubated with 0.3 g of dry, clean cellulose for 5 min at 23°C before lyophilization for 18 h. For naked control cellulose, 1 mL of TE buffer containing no DNA was incubated with cellulose under the same conditions. The thoroughly dried powder was resuspended in 20 volumes of TE buffer and incubated at 4°C for 24 h. DNA cellulose was washed twice with 10 volumes of TE and frozen at −70°C.

*Xenopus* cytostatic factor (CSF)-arrested egg extracts were prepared from unactivated eggs in M-phase as described [Murray 1991]. CaCl$_2$ (0.4 mM) was added to drive these extracts into interphase. In some cases, extracts were arrested in interphase with cycloheximide (100 µg/mL). Interphase cytosol was prepared by centrifugation at 260,000g for 1.5 h at 4°C.

**Binding of Xatr to DNA cellulose**

To allow the binding of Xatr to DNA cellulose, 25–100 µL of DNA cellulose was incubated with 50–500 µL of interphase egg cytosol for 40 min at 23°C with constant rocking. After centrifugation, the cytosol supernatant was removed and the cellulose beads were washed for four times with 1 mL of wash buffer (10 mM HEPES at pH 7.5, 150 mM NaCl, 0.05% NP-40, 30 mM β-glycerophosphate, 0.1 mM Na$_2$VO$_4$, 0.1 mM phenylmethylsulfonyl fluoride, and 10 µg/mL each of pepstatin, chymostatin, and leupeptin). Washed cellulose beads were digested with DNasel (4 U/mL) at 23°C for 10 min. After centrifugation, the supernatant was saved for immunoprecipitation of Xatr.

**Immunodepletion of Xatr and Xchk1 from egg extracts**

M-phase extracts (100 µL) were incubated with 20 µg of affinity-purified anti-Xatr antibodies or 10 µg of anti-Xchk1 antibodies bound to 10 µL of AffiPrep protein A beads [Bio-Rad Laboratories] at 4°C for 50 min. The same amount of control rabbit IgG (Zymed Laboratories) was used for mock depletion. After the incubation, the beads were removed by centrifugation. The supernatants were treated again under the same conditions to ensure that Xatr or Xchk1 was quantitatively removed from the extracts.

**Immunoprecipitation and kinase assays**

Polyclonal antibodies against the peptide EKTNPKPGRTEPK were used for immunoprecipitating Xatr from either egg cytosol or DNA cellulose eluates. Immunoprecipitations and kinase assays were performed as described [Guo and Dunphy 2000].

**Preparation of various recombinant Xchk1 proteins**

Individual and combination mutants in which Thr 314, Ser 344, Ser 356, and Ser 365 were changed to alanine were prepared by one or more rounds of mutagenesis of the pBS-Xchk1 plasmid [Kumagai et al. 1998] using the QuikChange kit [Stratagene] and the appropriate oligonucleotides. $\beta$-S-labeled versions of wild-type and mutant Xchk1 proteins were prepared using the TNT in vitro transcription/translation system [Promega]. DNA fragments encoding Xchk1-N135A, Xchk1-N135A-4AQ, and various wild-type and mutated peptides encompassing the SQ/TQ domain of Xchk1 were subcloned into the BamHI and EcoRI sites of pGEX-2T [Amersham Pharmacia Biotech]. GST-Xchk1-N135A, GST-Xchk1-N135A-4AQ, and the various GST fusion proteins containing fragments of Xchk1 were isolated from *E. coli* according to a published protocol [Frangioni and Neel 1993]. The production of Xchk1-WT-GST-His$_6$, Xchk1-S344A-GST-His$_6$, and Xchk1-4AQ-GST-His$_6$, which are double tagged at the C-terminal end, in baculovirus-infected S9 cells will be described elsewhere.

**Detection of Xchk1 with anti-phosphopeptide antibodies**

Interphase extracts (200 µL) containing 100 µg/mL cycloheximide and 3 µM tautomycin were incubated with 2 µg of Xchk1-WT-GST-His$_6$ or Xchk1-S344A-GST-His$_6$ in the presence or absence of 100 µg/mL aphidicolin and 3000 sperm nuclei per µL of extract, as indicated. After an incubation of 90 min, 400 µL of dilution buffer (10 mM HEPES at pH 7.5, 150 mM NaCl, 20 mM β-glycerophosphate, 2.5 mM EGTA, and 0.1% CHAPS) was added. The recombinant Xchk1 proteins were isolated with glutathione agarose, washed four times with dilution buffer, eluted by boiling for 5 min with SDS gel sample buffer, subjected to SDS-PAGE, and immunoblotted either with anti-GST antibodies (Santa Cruz Biotechnology) to detect recombinant Xchk1 or with anti-S344-p antibodies to detect phosphorylation of Ser 344. Anti-S344-p antibodies were produced against a peptide [CGKISFS(p)QPAAPDNM], which is phosphorylated on Ser 344, on S344-p antibodies and used to detect recombinant Xchk1. The antibodies were affinity purified as described [Lim et al. 2000]. A nonphosphorylated version of the same peptide was used to remove antibodies that do not require phosphorylation of Ser 344 for binding.

**Miscellaneous**

To block chromosomal DNA replication, aphidicolin (dissolved in DMSO at 10 µg/mL) was added to egg extracts to a final concentration of 100 µg/mL. To produce the GST–Xatr fusion protein, a BamHI-Sall fragment containing the full-length Xatr coding sequence was subcloned into the yeast expression vector pEG(KT) [Mitchell et al. 1993]. The resulting plasmid pEG(KT)-Xatr was transformed into the host strain JD51 [MATa/mbt-a ara3–52/ura3–52 leu2–3,12/leu2–3,11 his3–200/His3–200 trp1Δ63/Trp1Δ63 lys2–801/Lys2–801] [Dohmen et al. 1995]. Expression and isolation of the GST–Xatr protein were performed according to established procedures [Mitchell et al. 1993]. The recombinant GST–Xatr protein is not catalytically active. Transfection of 293T cells with constructs encoding wild-type.
or kinase-inactive Flag-tagged ATR, immunoprecipitation of Flag-tagged ATR proteins, and kinase assays were performed as described (Canman et al. 1998).

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