A chromatin-bound kinase, ERK8, protects genomic integrity by inhibiting HDM2-mediated degradation of the DNA clamp PCNA

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Proliferating cell nuclear antigen (PCNA) acts as a scaffold, coordinator, and stimulator of numerous processes required for faithful transmission of genetic information. Maintaining PCNA levels above a critical threshold is essential, but little is known about PCNA protein turnover. We now show that ERK8 (extracellular signal-regulated kinase 8) is required for PCNA protein stability. ERK8 contains a conserved PCNA-interacting protein (PIP) box. Chromatin-bound ERK8 (ERK8CHROMATIN) interacts via this motif with PCNACHROMATIN, which acts as a platform for numerous proteins involved in DNA metabolism. Silencing ERK8 decreases PCNA levels and increases DNA damage. Ectopic expression of PCNA blocks DNA damage induced by ERK8 loss. ERK8 prevents HDM2-mediated PCNA destruction by inhibiting the association of PCNA with HDM2. This regulation is physiologically relevant as ERK8 activity is inhibited in transformed mammary cells. Our results reveal an unanticipated mechanism to control PCNA levels in normal cycling mammary epithelial cells and implicate ERK8 in the regulation of genomic stability.

Introduction

The physiological functions of ERK8 (extracellular signal-regulated kinase 8) are unknown. It is most closely related to ERK1/2 and ERK5 by virtue of a Thr-Glu-Tyr (T-E-Y) activation motif that must be phosphorylated in order for the kinase to be active (Abe et al., 2002). Unlike ERK1/2, ERK8 and ERK5 contain C-terminal extensions. ERK8 appears to be a rapidly evolving kinase (Coulombe and Meloche, 2007). The C-terminal tail of ERK8 is ~51% conserved between primates and other mammals, in contrast to ERK5 in which the tail is ~96% conserved. Given the paucity of information on ERK8, we investigated its function.

Proliferating cell nuclear antigen (PCNA) is a key player in a variety of DNA metabolic pathways in which it coordinates and regulates the functions of numerous proteins that perform enzymatic reactions on DNA (Moldovan et al., 2007). PCNA exists as a homotrimer that encircles DNA and provides a platform for its interacting partners. In addition to facilitating complex assembly, PCNA is able to stimulate or inhibit the activity of some of its binding partners (Gomes and Burgers, 2000; Azam et al., 2001). Intensive efforts have revealed the docking domains that facilitate interaction with PCNA (Moldovan et al., 2007; Gilljam et al., 2009; Havens and Walter, 2009). It is hypothesized that PCNA controls the myriad processes involved in genomic stability by direct physical association with its numerous binding partners (Moldovan et al., 2007). A decrease in PCNA protein levels prevents the formation of complexes necessary for DNA metabolism and results in catastrophic consequences (Jaskulski et al., 1988; Henderson et al., 1994). Thus, it is essential for cell survival to maintain PCNA protein levels at a critical threshold. PCNA is covalently modified by ubiquitin (Andersen et al., 2008; Das-Bradoo et al., 2010) and SUMO (small ubiquitin-like modifier; Hoege et al., 2002), but these modifications are thought to control the association with binding partners and not in the regulation of degradation. Thus, little is known about the regulation of PCNA protein turnover.

We have discovered that ERK8 controls PCNA levels in normal cycling cells by preventing its destruction via human
ERK8 regulates proliferation of MCF-10A cells. (A) Analysis of endogenous ERK8 knockdown in MCF-10A cells transduced with shRNA to luciferase (Luc) or one of two different ERK8 targeting sequences for 5 d. The detergent-insoluble fraction (I) of the transduced lysates was normalized for Ran expression and immunoblotted for ERK8. (B) Scatter plot showing the rate of proliferation of transduced MCF-10A cells. Analysis was started 2 d after transduction. Mean is shown (n = 5, quadruplicates), and error bars indicate SEM. (C) A representative flow cytometric analysis of cell cycle progression in MCF-10A cells transduced as in A. Transduced cells were identified by GFP fluorescence, and DNA was stained with DRAQ5. The times indicated are sampling times after the cells were arrested by serum and growth factor depletion, followed by release into the cell cycle by the addition of complete media. The table summarizes the results (mean ± range [n = 2]) generated by ModFit LT (Verity Software House). (D) Analysis of whole cell extracts (WCE) of MCF-10A cells transduced as in A. The lysates were normalized for Ran expression and immunoblotted for regulators of the cell cycle and DNA damage.
homologue of murine double minute (HDM2). Loss of ERK8 resulted in genomic instability by decreasing PCNA levels beyond a tolerated limit. ERK8 was active in a mammary epithelial cell line and in primary mammary cells but was inactivated in breast cancer cell lines. These data argue that ERK8 control of PCNA levels is physiologically relevant, as ERK8 activity is inhibited in transformed cells.

Results

ERK8 regulates the cell cycle by maintaining PCNA levels

We investigated potential physiological functions for ERK8 by silencing its expression in the human breast epithelial line, MCF-10A. Successful silencing of endogenous ERK8, which was localized predominantly to a detergent-insoluble fraction, was observed with two different short hairpin RNAs (shRNAs; Fig. 1 A). Proliferation was measured starting at 2 d after transduction. Based on the doubling time of the control cells (31.9 ± 0.08 h), the transduced cells doubled approximately once before measuring the proliferation rate. Knockdown of ERK8 decreased proliferation approximately twofold, which was prevented by expression of a mutant ERK8 resistant to silencing (rERK8; Fig. 1 B and Fig. S1 A). In agreement with the silencing experiments, ectopic expression of wild-type ERK8 enhanced proliferation (Fig. 1 B). The most effective knockdown of ERK8 occurred 4–5 d after transduction, and therefore, the majority of experiments were performed using cells transduced in this time frame. To investigate the mechanism by which ERK8 controls the cell cycle, we performed flow cytometry using cells synchronized by serum and growth factor deprivation approximately twofold, which was prevented by expression of a mutant ERK8 resistant to silencing (rERK8; Fig. 1 B and Fig. S1 A). In agreement with the silencing experiments, ectopic expression of wild-type ERK8 enhanced proliferation (Fig. 1 B). The most effective knockdown of ERK8 occurred 4–5 d after transduction, and therefore, the majority of experiments were performed using cells transduced in this time frame. To investigate the mechanism by which ERK8 controls the cell cycle, we performed flow cytometry using cells synchronized by serum and growth factor depletion, which resulted in ~70% of the control cells in G1. The addition of serum and growth factors to the control cells resulted in a threefold increase in the number of cells in S phase after 24 h (Fig. 1 C). In contrast to these results, no change in the cell cycle distribution after release from starvation was observed when ERK8 was silenced (Fig. 1 C). Collectively, these results demonstrate that ERK8 is a limiting factor in regulating the proliferation rate of MCF-10A cells and controls entry into S phase.

While analyzing cell cycle markers, we noticed that loss of ERK8 resulted in increases in cyclin D1, cyclin E, and p27 and a decrease in cyclin A levels compared with the control (Fig. 1 D). Furthermore, retinoblastoma (Rb) appeared as a doublet in the immunoblot analysis, and loss of ERK8 reduced the level of the higher molecular weight form (Fig. 1 D). These observations are consistent with a reduction in the phosphorylated form of Rb. Collectively, these changes provide further evidence that loss of ERK8 results in a cell cycle block in the G1/S phase and are in agreement with our flow cytometry results. But, unexpectedly, loss of ERK8 resulted in a decrease in the total level of PCNA (Fig. 1, D and E; and see Fig. 4 B). Silencing ERK8 did not result in higher molecular weight forms of PCNA that could account for the decrease in the unmodified form (Figs. 1 D and 2 E).

The decrease in PCNA was not correlated with a particular stage in the cell cycle, as total PCNA levels did not change during the cell cycle (Fig. 1 F). In agreement with the literature (Naryzhy and Lee, 2001), the subcellular distribution of PCNA changed during the cell cycle such that during S phase, PCNA preferentially partitioned into a detergent-insoluble fraction (Fig. 1 F and Fig. S1 B). Loss of ERK8 appeared to specifically reduce PCNA levels as we did not observe a decrease in the level of Rad9, a component of the heterotrimeric complex 9-1-1, which, like PCNA, encircles DNA and is involved in DNA damage repair (Fig. 1 D; St Onge et al., 1999). Moreover, the level of RFC-1 (replication factor C-1), a subunit of the clamp loader that loads PCNA onto DNA (Waga and Stillman, 1998), was not changed (Fig. 1 D). We conclude that silencing ERK8 results in a specific decrease in PCNA protein levels.

To determine whether the reduction of PCNA levels caused by ERK8 depletion is physiologically significant, we asked whether silencing ERK8 increases DNA damage, as PCNA is essential for genomic integrity (Umar et al., 1996; Gary et al., 1997; Hoege et al., 2002; Unk et al., 2002; Zhang et al., 2003; Xia et al., 2005). Staining with an antibody to phospho-Ser139 in H2AX (γ-H2AX), a marker of DNA double-stranded breaks (Rogakou et al., 1999; Sedelnikova et al., 2003), resulted in a twofold increase in γ-H2AX staining in ERK8-depleted cells compared with the control, even in the absence of extrinsic genotoxic stress (Fig. 2 A and Fig. S2 A). PCNA acts to repair DNA in response to irradiation with UVC radiation (Essers et al., 2005). Therefore, we determined whether loss of ERK8 would further increase DNA damage in response to UVC irradiation. In response to UVC, both ATR (ataxia telangiectasia and Rad3 related) and Chk2 were activated, although loss of ERK8 reduced the level of Chk2 activation (Fig. S2 B). However, the intensity of γ-H2AX staining increased in parallel in the silenced and control cells up to 2 h after UVC treatment (Fig. 2 A and Fig. S2 A). Strikingly, however, although at 4 h the amount of γ-H2AX staining decreased in the control cells, it remained elevated when ERK8 was absent. We confirmed that the increased γ-H2AX staining was a result of DNA breaks using the comet assay. Silencing ERK8 in the absence of extrinsic genotoxic stress resulted in an approximately threefold increase in comet tail length compared with the control (Fig. 2, B and F). Moreover, in response to UVC treatment, the knockdown of ERK8 resulted in longer tail lengths compared with the control (Fig. 2, B and F). We conclude that ERK8 is not required for the initial response to DNA damage but is required for genomic stability most likely through its ability to maintain PCNA protein levels.

If the effects of ERK8 are mainly caused by a decrease in PCNA levels, depleting PCNA should phenocopy the knockdown of ERK8. Consistent with our observations with ERK8, we observed that silencing PCNA inhibited proliferation of MCF-10A cells (Fig. 2 C). Moreover, in the absence of extrinsic genotoxic stress, loss of PCNA resulted in an increased level of...
Figure 2.  ERK8 regulates DNA repair by a PCNA-dependent mechanism.  (A) Analysis of DNA damage by γH2AX immunofluorescence in ERK8 knockdown cells.  MCF-10A cells transduced for 5 d were treated with or without (-) 20 J/m² UVC.  The times indicated refer to the length of time after irradiation.  At the indicated time, the cells were treated with detergent, fixed, and immunostained with an anti–γH2AX antibody and an anti–mouse fluorescent secondary antibody.  Nuclei were stained with DRAQ5.  The intensity of γH2AX staining was determined and normalized to the levels obtained in the control cells, which were transduced with luciferase (Luc) shRNA and not irradiated.  Mean is shown (n = 2, duplicates, ≥35 cells/condition), and error bars indicate SEM.  (B) Analysis of DNA damage by comet assay of cells transduced as in A, without irradiation or 2 h after UVC treatment.  DNA was visualized by staining with Sybr green (Trevigen, Inc.).  Bar, 50 µm.  (C) Rate of proliferation of MCF-10A cells transfected with control or PCNA-specific siRNA.  The rate of proliferation was determined over 48 h, starting at 2 d after transfection.  Mean is shown (n = 4, sextuplicate), and error bars indicate SEM.  The right panel shows the extent of PCNA knockdown 4 d after transfection in lysates normalized for Ran.  (D) Analysis of DNA damage by γH2AX immunofluorescence in PCNA knockdown cells.  Cells transfected as in C were treated as in A, and the intensity of γH2AX staining was determined and
γ-H2AX staining compared with the control (Fig. 2 D and Fig. S2 C). Furthermore, at 4 h, although DNA repair started to occur in the control cells, the level of γ-H2AX staining remained elevated in the absence of PCNA. These results are consistent with our hypothesis that ERK8 controls genomic integrity via regulation of PCNA protein levels. To further test our hypothesis that ERK8 acts primarily through PCNA, we asked whether the ectopic expression of PCNA would reverse the decrease in proliferation caused by reduced ERK8 levels. MCF-10A cells were transduced with mRFP-tagged PCNA (mRFP-PCNA) or the mRFP control and then transduced a second time with ERK8-specific or control shRNA. In agreement with our earlier results, silencing ERK8 reduced proliferation; however, importantly, the overexpression of PCNA after silencing ERK8 prevented the decrease in proliferation (Fig. 2 E). Additionally, the forced expression of PCNA alleviated the DNA damage caused by loss of ERK8 (Fig. 2 F and Fig. S2 D). Collectively, these data support the surprising conclusion that ERK8 is essential to maintain PCNA expression at a level that prevents DNA replication stress in normal cycling cells (Halazonetis et al., 2008).

ERK8 interacts and controls PCNA turnover via a PCNA-interacting protein (PIP) box

To determine how ERK8 controls PCNA levels, we asked whether ERK8 and PCNA physically associate. A fraction of wild-type Venus-tagged ERK8 (Venus-ERK8) and endogenous PCNA associated with chromatin after a detergent and high salt wash extraction, which indicates that there is a pool of ERK8 and PCNA that has a high affinity for chromatin (Fig. S3 A). Based on these observations, we used a modified chromatin immunoprecipitation assay and found that, in the presence of DNase I, chromatin-bound HA-tagged ERK8 (HA-ERK8) associated with PCNA (Fig. 3 A). Furthermore, endogenous PCNA associated with endogenous ERK8 (Fig. 3 B). A fraction of PCNA and ERK8 was not associated with chromatin, and HA-ERK8 obtained from this soluble fraction (HA-ERK8) and PCNA did not interact (Fig. 3 A). Most of the interacting partners for PCNA interact through a conserved motif called the PIP box (Warbrick, 1998; Moldovan et al., 2007). We identified a putative PIP box in ERK8, specifically the sequence from 297 to 308 (QALHPYVQRFH), where bold indicates a match with the conserved motif (Fig. S3 B). The spacing between the Q and the V is usually two residues; however, DNA polymerase β has a three-residue spacing, like ERK8 (Kedar et al., 2002). A GST fusion protein containing the putative ERK8 PIP box (GST–ERK8 (PIP)) interacted with PCNA (Fig. 3 C), but a mutant PIP box, in which the residue corresponding to Glu300 was mutated to Ala (GST–ERK8 (PIPm)) did not interact. Further confirmation that ERK8 contains a PIP box was shown by mutation of Gln300 to Ala (GST–ERK8 (Q300A)) mutant, treated with DNase I, and analyzed by immunoblotting. A fraction of the input is shown for comparison to the level of associated PCNA. I, insoluble fraction; IP, immunoprecipitation; S, soluble fraction.

Figure 3. ERK8 interacts with PCNA via a PIP box. (A and B) Chromatin was immunoprecipitated with antibodies against ectopically expressed HA-tagged ERK8 (A) or endogenous PCNA (B), treated with DNase I, and analyzed by immunoblotting. A fraction of the input is shown for comparison to the level of associated PCNA. I, insoluble fraction; IP, immunoprecipitation; S, soluble fraction.
of PCNA\textsuperscript{CHROMATIN} by $\sim$60\% compared with the control. Consistent with these observations, total PCNA in ERK8-depleted cells was reduced approximately twofold relative to the control after a 12-h treatment with the protein synthesis inhibitor cycloheximide (Fig. S3 E). Thus, loss of ERK8 enhances PCNA\textsuperscript{CHROMATIN} turnover. Mutation of Tyr211 has been reported to destabilize PCNA (Wang et al., 2006). However, we did not detect any Tyr phosphorylation of PCNA (Fig. S3 F). These results are in agreement with orthophosphate labeling analysis of PCNA, which also did not detect phosphorylation of PCNA (Naryzhny and Lee, 2004). We propose that the increase in PCNA\textsuperscript{CHROMATIN} turnover caused by the loss of ERK8 indirectly decreases PCNA\textsuperscript{FREE} levels because of continued recruitment of PCNA\textsuperscript{FREE} to the chromatin in an effort to maintain critical cellular functions (Fig. 5 A). This hypothesis is consistent with the literature in which increasing the extent of DNA damage leads to depletion of PCNA\textsuperscript{FREE} (Mortusewicz and Leonhardt, 2007).

We next asked whether the interaction of ERK8 with PCNA is necessary for stabilization of PCNA. MCF-10A cells were transduced with Venus–ERK8, Venus–ERK8 PIP mutant (Venus–ERK8(Q300A)), or the Venus control and then transduced a second time with ERK8-specific or control shRNA. Expression of resistant ERK8 prevented the decrease in PCNA levels caused by silencing of endogenous ERK8 (Fig. 4 B). However, importantly, the ERK8 PIP mutant was unable to rescue PCNA levels even though the level of rERK8(Q300A) and its ability to associate with the chromatin was similar to that of the wild type (Fig. 4 B and Fig. S3 C). Furthermore, the mutant was as active as the wild type, as shown by the anti-pERK8 antibody (Fig. 4 B). This antibody recognizes the dual phosphorylation of the T-E-Y motif, which must be phosphorylated for members of the ERK1/2 family to be active (Her et al., 1993). Therefore, the pERK8 levels reflect the amount of active kinase. Surprisingly, expression of the ERK8 PIP mutant caused an at least twofold decrease in PCNA levels even in the presence of endogenous ERK8. These results indicate that ERK8(Q300A) acts as a dominant-negative, most likely by competing with endogenous ERK8 for binding to chromatin, thereby blocking association of the endogenous ERK8 with PCNA, which permits PCNA destruction. Collectively, our data support a model in which ERK8\textsuperscript{CHROMATIN} stabilizes PCNA by inhibiting access to PCNA\textsuperscript{CHROMATIN} of a destruction factor. This is the first report of a PCNA-binding partner that regulates PCNA stability during normal cell cycling (Izumi et al., 2001; Wang et al., 2006; Yu et al., 2009).

We next asked whether ERK8 controls PCNA protein stability by pulse-chase labeling with $[^{35}\text{S}]$Met. The degradation rate of PCNA\textsuperscript{FREE}, which does not interact with ERK8, was similar in the presence and absence of ERK8 (Fig. 4 A and Fig. S3 D). However, silencing ERK8 increased the destruction of PCNA\textsuperscript{CHROMATIN} by $\sim$60\% compared with the control. Consistent with these observations, total PCNA in ERK8-depleted cells was reduced approximately twofold relative to the control after a 12-h treatment with the protein synthesis inhibitor cycloheximide (Fig. S3 E). Thus, loss of ERK8 enhances PCNA\textsuperscript{CHROMATIN} turnover. Mutation of Tyr211 has been reported to destabilize PCNA (Wang et al., 2006). However, we did not detect any Tyr phosphorylation of PCNA (Fig. S3 F). These results are in agreement with orthophosphate labeling analysis of PCNA, which also did not detect phosphorylation of PCNA (Naryzhny and Lee, 2004). We propose that the increase in PCNA\textsuperscript{CHROMATIN} turnover caused by the loss of ERK8 indirectly decreases PCNA\textsuperscript{FREE} levels because of continued recruitment of PCNA\textsuperscript{FREE} to the chromatin in an effort to maintain critical cellular functions (Fig. 5 A). This hypothesis is consistent with the literature in which increasing the extent of DNA damage leads to depletion of PCNA\textsuperscript{FREE} (Mortusewicz and Leonhardt, 2007).

ERK8 inhibits the interaction of PCNA with HDM2

PCNA is able to act as a platform to mediate the degradation of some of its binding partners via a specialized PIP box (Havens and Walter, 2009). However, information on how PCNA turnover is regulated in normal cycling cells is lacking. In response to DNA damage to UV, PCNA destruction has been shown to occur via the 26S proteasome pathway (Yu et al., 2009). Therefore, we reasoned that an E3 ligase, which is able to compete for ERK8 binding to PCNA, may also control PCNA turnover in normal cycling cells. HDM2 is the only currently known E3 ligase that interacts with PCNA through a PIP box (Banks et al., 2006),
and therefore, it could possibly compete with ERK8 for binding to PCNA. However, it is not known whether HDM2 targets PCNA for destruction. We observed that silencing HDM2 resulted in an increase in PCNA levels (Fig. 5 A). These results suggest that HDM2 regulates PCNA turnover in normal cycling cells. However, as HDM2 regulates p53 levels, knockdown of HDM2 will result in an increase in p53 levels and disruption of the cell cycle (Martin et al., 1995). To investigate whether elevated p53 levels could increase PCNA levels, we used the inhibitor nutlin-3 (Vassilev et al., 2004). As expected by disrupting the association of HDM2 and p53, the levels of both proteins increased substantially (Fig. S4). We also observed that nutlin-3 altered the electrophoretic mobility of Rb to generate a single immunoreactive band (Fig. S4), which is consistent with the unphosphorylated form of Rb. Unphosphorylated Rb binds and inhibits E2F (Stevens and La Thangue, 2003), which can result in decreased expression of some E2F-regulated genes such as PCNA (Black et al., 2005). However, treatment with nutlin-3 did not alter the levels of PCNA. These results indicate that the increase in PCNA levels in response to the loss of HDM2 is independent of p53 and the phosphorylation status of Rb.

To investigate whether HDM2 facilitates PCNA expression in the absence of ERK8, we silenced both HDM2 and ERK8. Strikingly, the decrease in PCNA levels caused by the loss of ERK8 was reversed by also silencing HDM2 expression (Fig. 5 A). From this data, we would predict that the loss of HDM2 would prevent ubiquitination of PCNA. To test this hypothesis, we silenced ERK8 and then transfected with control or HDM2 siRNA. Additionally, we transfected with a plasmid encoding a six-His tag fused to ubiquitin (His6-Ub). Loss of HDM2 decreased the levels of ubiquitinated PCNA (Fig. 5 A). Knockdown of ERK8 did not alter the levels of HDM2 (Fig. 5 B). However, silencing ERK8 increased the association of PCNA with HDM2 (Fig. 5 B). We conclude that ERK8 inhibits PCNA degradation by inhibiting its association with HDM2, which facilitates PCNA degradation via the 26S proteasome pathway.

**ERK8 is activated by chromatin binding**

It has been reported that ERK8 activity is increased in response to various DNA-damaging agents (Klevenic et al., 2009). Using identical conditions as reported by Klevenic et al. (2006, 2009), we observed changes in ERK8 activity only in response to 1 mM H2O2 (Fig. S5 A). We did not observe changes in ERK8 protein levels with any of the DNA-damaging reagents tested. Activation of ERK1/2 was dose dependent on H2O2, but ERK8 activation was observed only at 1 mM H2O2 (Fig. S5 B). This concentration of H2O2 was extremely toxic and resulted in cell death within 3 h, and therefore, it is difficult to determine the physiological relevance of this observation. We conclude that in MCF-10A cells, DNA-damaging agents do not enhance activation of ERK8, and basal levels of ERK8 activation are sufficient to stabilize PCNA levels.

Active ERK8, as shown by pERK8, is mainly present in the insoluble fraction (Fig. S5 C). These results are consistent...
showed defective chromatin association (Fig. 6 D and Fig. S5 E).
We conclude that the PXXXP motif acts as an autoinhibitory regulator of kinase activity. We propose that the mutation mimics the conformational change that occurs upon association of ERK8 with chromatin, which relieves autoinhibition. We conclude that binding to chromatin activates ERK8, independently of PCNA binding.

ERK8 stabilizes PCNA in primary mammary epithelial cells
ERK8 was detected in the detergent-insoluble fraction of primary human mammary epithelial (HME) cells at levels similar to MCF-10A cells (Fig. 7 A). Importantly, silencing ERK8 in HME cells decreased PCNA levels (Fig. 7 B), and this reduction in PCNA was accompanied by a substantial increase in the frequency of DNA breaks for both untreated and UVC-irradiated HME cells (Fig. 7 B). The fold increase in comet length in HME cells ranged from $\leq 2–10$-fold in the absence of UVC, and this range probably reflects the genetic differences between patient samples. Thus, the regulation of PCNA by ERK8 is not an artifact of the MCF-10A immortalized cell line but occurs also in primary mammary cells.

ERK8 is inactive in transformed breast lines
Our results predict that loss of ERK8 will lead to genomic instability. To test this prediction, we silenced ERK8 in MCF-10A cells and maintained the cells in culture for 2 wk. Analysis of the nuclei of

with the immunofluorescence data that show that a kinase-dead mutant of ERK8 (ERK8(K42A)) does not associate with the chromatin (Fig. S3 A). This mutant has the essential Lys residue that is necessary for interaction with ATP mutated to Ala (Abe et al., 2002). It is possible that the interaction of PCNA activates ERK8, like Chk1 (Scorah et al., 2008). However, pERK8 levels are similar for wild type and the ERK8(Q300A) mutant, which does not bind PCNA (Figs. 3 D and 4 B). It is also unlikely that ERK8 is recruited to chromatin by PCNA because, in contrast to PCNA, soluble ERK8 does not redistribute to the insoluble fraction in response to UV (Fig. S5 C). Furthermore, the ERK8(Q300A) mutant associates with chromatin but not with PCNA. Additionally, ERK8 and PCNA only partially co-localize as 42% ± 10.9 of the nuclei positive for Venus-ERK8 costain for PCNA, whereas in nuclei positive for PCNA, 22% ± 5.4 of the cells costain with ERK8. Thus, ERK8 must be able to interact with chromatin in a PCNA-independent manner.

To investigate the mechanism by which ERK8 associates with the chromatin, we analyzed deletion mutants. ERK8 (1–475) was active, associated with chromatin, and stimulated proliferation similar to that of the wild type (Fig. 6, A and B). However, further deletions of the C terminus generated inactive constructs that did not target to chromatin (Fig. 6 A). We identified a repeating PXXXP motif, which is highly conserved among mammals, located between residues 371 and 398 (Fig. S5 D). Unexpectedly, mutations within this motif (ERK8 (P390A/P398A)) substantially increased the level of activation, as indicated by pERK8, compared with the wild type (Fig. 6 C). However, the mutant also
stress. We would speculate that in breast cancer cells, there is a mechanism to prevent HDM2 from associating with PCNA in the absence of active ERK8. Our results also suggest that loss of ERK8 activity is important in the transformation process.

We have identified two important functional domains in ERK8, the PIP box and the PXXXP motif. The PIP box is located within the kinase domain, and the PXXXP motif is in the C-terminal extension. To identify the region within the kinase domain that contains the PIP box, we modeled the ERK8 kinase domain to that of ERK2 and found that the PIP box is adjacent to the conserved docking (CD) domain. The CD domain contributes but is not solely responsible for the interactions of the MAPK superfamily with their docking partners (Tanoue et al., 2000, 2001; Zhang et al., 2003). A corresponding PIP box was not found in other members of the ERK1/2 or p38MAPK families but was present in JNK1/3 (Fig. S5 F). It is not known whether JNK1/3 can interact with PCNA. The CD domain in ERK8 is unusual in that it contains a Cys residue, whereas all other members of the MAPK superfamily have an acidic residue in this position. It is possible that the Cys residue regulates the ability of ERK8 to interact with PCNA. The PXXXP motif functions to regulate the association of ERK8 with chromatin and thereby indirectly regulates association with PCNA as only ERK8CHROMATIN interacts with PCNA. PCNA is known to interact with a plethora of proteins that contain PIP boxes and the mechanism by which these interactions are regulated is unknown.

It is intriguing that a rapidly evolving kinase would control genomic stability via PCNA, which is a highly conserved protein. Curiously, mammary epithelial cells are susceptible to

Discussion

We conclude that ERK8 is as a key controller of genomic stability in mammary epithelial cells via its ability to regulate the formation of complexes containing PCNA. In untransformed mammary cells, ERK8 prevents the destruction of PCNA that is mediated via HDM2 (Fig. 8 D). Remarkably, a twofold reduction in PCNA levels is sufficient to cause extensive DNA damage and mitotic failure even in the absence of extrinsic genotoxic stress. We would speculate that in breast cancer cells, there is a mechanism to prevent HDM2 from associating with PCNA in the absence of active ERK8. Our results also suggest that loss of ERK8 activity is important in the transformation process.

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Figure 7. ERK8 is critical for genomic integrity in mammary epithelial cells. (A) Analysis of endogenous ERK8 knockdown in HME cells (HMEC) transduced with control or ERK8 shRNA for 5 d. The detergent-insoluble fraction (I) of the transduced lysates was normalized for Ran expression. (B) Analysis of DNA damage by comet assay of transduced HME cells treated without irradiation or 2 h after UVC treatment. The left panel shows lysates from transduced HME cells that were normalized for Ran expression and immunoblotted. (C) Analysis of DNA morphology by Hoechst staining of transduced MCF-10A cells kept in long-term culture (~2 wk). Arrows indicate micronuclei. The percentage of the total cell population that contained micronuclei was determined. Mean is shown (≥500 cells), and error bars indicate SEM. Luc, luciferase; LV, lentivirus. Bars: (B) 50 µm; (C) 10 µm.
The short hairpin sequence including the ERK8-targeting shRNAs (in bold) are (ERK8-1) 5′-GATCCCCACATTTACCTGGTGTTTGA and (ERK8-2) 5′-GA\TCCCGACAGATGCCCAGAGAACA TTCAAGAGATGTTCTCTGGGCATCTGTC TTTTGGAAA-3′. The short hairpin sequence including the luciferase-targeting shRNAs (in bold) is 5′-GATCCCCCGTACGCGGAATACTTCGA TTCAAGAGATCGAAGTATTCCGCGTACG TTTTGGAA-3′ (Malliri et al., 2004). GST-PIP and GST-PIPm were generated by subcloning into pGEX2T (GE Healthcare) the annealed oligonucleotides with the upper strand sequences 5′-GATC CCGAGGCACTGCAGCACCCCTACGTGCAGAGGTTCCACTGCCCTGAG-3′ and 5′-GATCCCAGGCACTGGCGCACCCCTACGTGCAGAGGTTCCACTGCCCTGAG-3′. All shRNA, deletion, and mutant constructs were verified by sequencing (Biomolecular Research Core, University of Virginia).

Cell culture and treatment
MCF-10A, MCF-7, and T47D were cultured as indicated by the American Type Culture Collection. Primary HME cells were purified from tissue (McCaffrey and Macara, 2009) and cultured (Eisinger-Mathason et al., 2008). For irradiation, the cells were washed with PBS and exposed to 20 J/m² UVC. Comet assay was performed according to the manufacturer (Trevigen, Inc.) and electrophoresed under alkaline conditions (275 mA for 30 min at 4°C). Cell fractionation was performed as described previously.

Figure 8. Breast cancer cells do not have active ERK8. (A) Analysis of endogenous ERK8 levels in breast cancer cell lines. Whole cell extracts (WCE) from different breast cell lines were normalized for Ran. For comparison, MCF-10A cells were transduced with luciferase (Luc) or ERK8 shRNA, and the insoluble (I) fraction was analyzed 5 d after transduction. (B) Lysates from the various breast lines were transduced with Venus-ERK8, normalized to expression of Venus-ERK8, and immunoblotted for active ERK8 [pERK8]. (A and B) The black lines indicate that intervening lanes were removed. (C) Loss of ERK8 does not decrease PCNA levels in MCF-7 cells. MCF-7 cells were transduced with luciferase or ERK8 shRNA for 5 d. Lysates were normalized to Ran and immunoblotted. (D) Model of ERK8 regulation of PCNA levels. Active ERK8 preferentially binds to the chromatin through its binding partners. ERK8\CHROMATIN and PCNA\CHROMATIN interact via the ERK8 PIP box. Loss of ERK8 binding to PCNA via silencing or mutation of the PIP domain results in the increased recruitment of HDM2, which enhances PCNA turnover. Autoinhibition is alleviated by mutating Pro390 and Pro398. However, these mutations interfere with chromatin binding. LV, lentivirus.

Materials and methods

Plasmids and lentiviral production
Constructs used to generate lentivirus, including pSPAX2, pLVTHM, and pMD2G, were provided by D. Trono (Swiss Federal Institute of Technology, Lausanne, Switzerland; Zufferey et al., 1997). pLV-Venus and pLV-mRFP lentivirus constructs were provided by I.G. Macara (University of Virginia, Charlottesville, VA; McCaffrey and Macara, 2009). pCMV5HIS6-Ub was provided by D. Wotton (University of Virginia). pWPI was purchased from Addgene. Lentiviral production was performed as described previously (McCaffrey and Macara, 2009) but titered using MCF-10A cells.

The short hairpin sequence including the ERK8-targeting shRNAs (in bold) are (ERK8-1) 5′-GATCCCCACATTTACCTGGTGTTTGAATCAAGAGA-TCAAACACCAGGTAAATGT TTTTGGAAA-3′ and (ERK8-2) 5′-GATCCC CGACAGATGCCCAGAGAACA TTCAAGAGATGTTCTCTGGGCATCTGTC TTTTGGAAA-3′. The short hairpin sequence including the luciferase-targeting shRNAs (in bold) is 5′-GATCCCCCGTACGCGGAATACTTCGA TTCAAGAGATCGAAGTATTCCGCGTACG TTTTGGAA-3′ (Malliri et al., 2004). GST-PIP and GST-PIPm were generated by subcloning into pGEX2T (GE Healthcare) the annealed oligonucleotides with the upper strand sequences 5′-GATC CCGAGGCACTGCAGCACCCCTACGTGCAGAGGTTCCACTGCCCTGAG-3′ and 5′-GATCCCAGGCACTGGCGCACCCCTACGTGCAGAGGTTCCACT GCCCTGAG-3′. All shRNA, deletion, and mutant constructs were verified by sequencing (Biomolecular Research Core, University of Virginia).

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(Avkin et al., 2006). Transient transfection was performed with Lipofectamine 2000 (Invitrogen) reagent using nontargeting siRNA #1, human MDM2 SMARTpool siRNA, or human PCNA SMARTpool siRNA [Thermo Fisher Scientific]. The following chemicals were used: nulitin-3 (EMD); hydroxyurea, etoposide, cisplatin, and MMS (methyl methane sulfonate; Sigma-Aldrich); and H2O2 (Thermo Fisher Scientific) and M宁NG6 (N-methyl-N-Nitrosoguanidine; TCI America).

Immunodetection and immunofluorescence

Primary antibodies used were monoclonal anti-p27, rabbit anti–RF-C1, monoclonal anti-GST, rabbit anti–HA, monoclonal anti–HD72, monoclonal anti–p53, and monoclonal anti–PCNA (Santa Cruz Biotechnology, Inc.); monoclonal anti–cyclin D1, monoclonal anti–Rb, rabbit anti–phospho–Ser/Thr428 ATR, rabbit anti–phospho–Thr68 Chk2, and rabbit anti–Chk2 (Cell Signaling Technology); monoclonal anti–phospho–Thr, rabbit anti–phospho–Ser, rabbit anti–cyclin E, and monoclonal anti–γH2AX (Millipore); rabbit anti–phospho–Thr (Invitrogen); monoclonal anti–Rad9 and rabbit anti–phospho–MAPK (pThr202/pTyr204, Thermo Fisher Scientific); rabbit anti–ERK8 (Abgent); monoclonal anti–PCNA (Dako); monoclonal anti–Ran (BD); monoclonal anti–cyclin A (Novoceastra); goat anti–GFP and rabbit anti–GFP (Venus; Abcam); and monoclonal anti–HA (12CA5; lymphocyte Culture Center, University of Virginia). Secondary antibodies used were goat anti–mouse, goat anti–rabbit, and donkey anti–goat (Jackson Immunoresearch Laboratories, Inc.).

Detection of PCNA and Venus-tagged chromatin-binding proteins was adapted from Sporbert et al. (2002). In brief, cells were plated on coverslips, washed twice with ice-cold PBS, and permeabilized for 30 s with ice-cold cytoskeleton buffer (10 mM Pipes, pH 6.8, 250 mM sucrose, 50 mM NaCl, 2 mM MgCl2, 2 mM EGTA, and 0.1% Triton X-100). The permeabilized cells were washed three times with ice-cold wash buffer (PBS containing 3 mM MgCl2 and 400 U RNase-free DNase I/10 µg DNA (New England Biolabs), and then incubated with 3 mM MgCl2 and 400 U Rnase-free DNase I/10 µg DNA (Invitrogen). Cell lysates were incubated with the soluble fraction. Approximately 140 µl of Dynabeads sheep anti–mouse secondary antibody (Invitrogen) was washed with immunoprecipitation buffer and incubated with lysate for 1 h at 4°C in a thermomixer [900 rpm]. The immunoprecipitates were washed five times in the appropriate immunoprecipitation buffer, three times high salt buffer (10 mM Hepes, pH 7.4, 300 mM NaCl, 10 mM KCl, and 1 mM EDTA), and three times with the immunoprecipitation buffer. The beads were boiled in 2x SDS lysis buffer [0.12 M Tris–Cl, pH 6.8, 4% SDS, 20% glycerol, 0.2 M DTT, and 0.008% bromophenol blue], heated to 100°C for 5 min

Pulse-chase labeling

Cells were washed and preincubated for 4 h with cold labeling media (Met-free DMEM/F12 [Sigma–Aldrich], 5% dialyzed horse serum, 0.5 µg/mL hydrocortisone, 100 ng/mL cholera toxin, 10 µg/mL insulin, and 5.75 µM cold Met). [3H]Met (200 µCi/mL, 800 Ci/mmol; PerkinElmer) was added and incubated for 4 h. After washing with labeling media plus 2 mM cold Met, the cells were replaced in their normal growth media and harvested for immunoprecipitation.

Statistical analysis

Data are presented as mean ± SEM; Student’s two-tailed t test was used for comparisons, with P < 0.05 considered significant.

Online supplemental material

Fig. S1 shows that ERK8 regulates the cell cycle. Fig. S2 shows that loss of ERK8 increases DNA damage. Fig. S3 shows that ERK8 regulates PCNA turnover. Fig. S4 shows that inhibition of p53 degradation does not alter PCNA protein levels. Fig. S5 examines the activation, levels, and subcellular distribution of ERK8 in response to DNA-damaging agents. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.201002124/DC1.

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