DNA-dependent protein kinase promotes DNA end processing by MRN and CtIP

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The repair of DNA double-strand breaks occurs through nonhomologous end joining or homologous recombination in vertebrate cells—a choice that is thought to be decided by a competition between DNA-dependent protein kinase (DNA-PK) and the Mre11/Rad50/Nbs1 (MRN) complex but is not well understood. Using ensemble biochemistry and single-molecule approaches, here, we show that the MRN complex is dependent on DNA-PK and phosphorylated CtIP to perform efficient processing and resection of DNA ends in physiological conditions, thus eliminating the competition model. Endonucleolytic removal of DNA-PK–bound DNA ends is also observed at double-strand break sites in human cells. The involvement of DNA-PK in MRN-mediated end processing promotes an efficient and sequential transition from nonhomologous end joining to homologous recombination by facilitating DNA-PK removal.

INTRODUCTION

DNA-dependent protein kinase (DNA-PK) consists of a catalytic kinase subunit (DNA-PKcs) and the DNA end-binding heterodimer of Ku70 and Ku80 (Ku). Together, these proteins form an end recognition complex (DNA-PK) that binds to DNA double-strand breaks (DSBs) within seconds of break formation (1). DNA-PK promotes the ligation of two DNA ends by ligase IV, aided by accessory factors, and in some cases, accompanied by limited end processing (1, 2). The recognition and repair of DNA breaks by the DNA-PK–associated nonhomologous end-joining (NHEJ) machinery is generally considered to be the first and rapid phase of DNA repair, occurring within ~30 min of DNA damage, while homologous recombination is specific to the S and G2 phases of the cell cycle and occurs over a longer time frame (3, 4). DNA-PK is present at micromolar concentrations in cells (5) and binds DNA ends in all cell cycle phases, even during S phase at single-ended breaks (6).

The Mre11/Rad50/Nbs1 (MRN) complex regulates the initiation of DNA end processing before homologous recombination by catalyzing the initial 5′ strand processing at blocked DNA ends and by recruiting long-range nucleases Exo1 and Dna2 (4, 7). At single-ended breaks during replication, the nuclease activity of Mre11 was shown to be important for removal of Ku (6), suggesting that MRN catalytic processing of ends during S phase is important for recombination-mediated repair of single-ended breaks. In addition, the presence of protein blocks on DNA ends has been shown to stimulate MRN(X) endonuclease activity in vitro (8–12), consistent with the idea that chromatin-bound proteins stimulate MRN activities. Here, we considered the possibility that the presence of DNA-PK on DNA ends might regulate MRN-dependent end processing and that this interaction may be a critical component of the choice between NHEJ and homologous recombination pathways in human cells.

RESULTS

To test for the effect of DNA-PKcs on end processing by the MRN complex, we performed a nuclease assay with purified human MRN, CtIP, Ku, and DNA-PKs (fig. S1) on a 197–base pair (bp) linear double-stranded DNA (dsDNA) containing a single radiolabel at the end of one of the 5′ strands. Addition of all the protein complexes together to this substrate resulted in the appearance of a single major cleavage product approximately 45 nucleotides (nt) in length (Fig. 1A, lane 2). We previously reported MRN-mediated removal of the Ku heterodimer alone, showing that MRN makes an endonucleolytic cleavage at approximately 30 bp from the DNA end (13); here, we observe that the inclusion of DNA-PKcs in the reaction increases the efficiency of the cutting by approximately 50-fold as well as changing the position of the nuclease cleavage site to a location approximately 45 nt from the end (Fig. 1A, lanes 2 and 3). Efficient formation of the cleavage product requires CtIP, which is essential for DNA end resection in human cells (14). Mre11 nuclease activity in vitro is strictly manganese dependent (8, 15, 16); however, Mre11/Rad50 (MR) complexes from archaea and from T4 bacteriophage also exhibit nuclease activity in the absence of manganese when physiologically relevant protein cofactors are present (17, 18). Here, we observe that human MRN makes endonucleolytic cuts in magnesium-only conditions and that this requires the presence of both Ku and DNA-PKs (Fig. 1A, lane 10). Because human cells do not contain high levels of manganese (19), we conclude that the physiologically relevant endonuclease activity of MRN is thus dependent on DNA-PK.

Mre11 and CtIP both exhibit endonucleolytic activity (8, 20, 21), so we asked which protein is responsible for the catalysis. We found that substituting wild-type Mre11 with a nuclease-deficient form of Mre11, H129N (22, 23), completely abolished DNA-PK–dependent cutting by MRN, whereas we observed robust activity with nuclease-deficient N289A/H290A CtIP (Fig. 1B, lanes 4 and 5); thus, we conclude that the processing is done by the catalytic activity of Mre11.

We previously observed that adenosine 5′-triphosphate (ATP) binding by Rad50 promotes Nbs1-dependent endonucleolytic cutting by Mre11 (8, 15). Exclusion of ATP or substitution of the non-hydrolyzable adenylyl-imidodiphosphate (AMP-PNP) analog for ATP blocks MRN nuclease activity in the presence of DNA-PK...
Fig. 1. Nucleolytic removal of DNA-PK by MRN and its stimulation by CtIP. (A) Nuclease reactions were performed with a 197-bp DNA substrate, 5′ labeled with $^{32}$P (asterisk), with MRN (50 nM), CtIP (80 nM), Ku (10 nM), and DNA-PKcs (10 nM) as indicated, in the presence of both magnesium and manganese (lanes 1 to 8) or magnesium only (lanes 9 to 16). All reactions contained the DNA-PKcs inhibitor NU7441. Products were visualized by denaturing PAGE and visualized by phosphorimager. Red and black arrows indicate the predominant product in the presence of DNA-PKcs and Ku, or Ku alone, respectively. (B) Nuclease assays were performed as in (A) in the presence of both magnesium and manganese with wild-type MRN or MRN containing nuclease-deficient Mre11 (H129N). Reactions without ATP or with AMP-PNP instead of ATP are indicated, as is the reaction without NU7441. (C) Nuclease assays were performed as in (A) in the presence of magnesium, manganese, ATP, NU7441, CtIP (C), and DNA-PK (D) with MRN (M) containing wild-type Rad50 (WT) or adenosine triphosphatase (ATPase)-deficient Rad50 K42A (KA) or D1231A (DA). (D) Nuclease assays were performed as in (A) in the presence of both magnesium and manganese with wild-type MRN in the presence of 25, 50, and 100 μM Mre11 inhibitors and NU7441. (E) Quantitation of the MRN endonuclease observed in the presence of Mre11 inhibitors expressed as percentage of the activity in the absence of inhibitors. Error bars represent SD from two replicates.
Phosphorylation of Ku has been shown to result in the removal of Ku from DNA (25) and autophosphorylation of DNA-PKcs also removes DNA-PKcs from the Ku-DNA complex (26–30); therefore, in reactions shown in Fig. 1 (A and B), we have also included an inhibitor of DNA-PKcs, NU7441, to block its kinase activity. Removal of the inhibitor reduced the efficiency of the reaction (Fig. 1B, lane 8), suggesting that reduced stability of DNA-PK on DNA reduces the nucleolytic activity of MRN on the DNA-PK–bound substrate. However, the endonucleolytic product is still observed in the absence of the inhibitor (also see fig. S2); thus, blocking DNA-PKcs phosphorylation is not absolutely essential.

Small-molecule inhibitors that specifically block either the exonuclease (PFM39) or the endonuclease (PFM01 and PFM03) activity of Mre11 have been developed (30). We found that PFM01 and PFM03 reduced MRN endonuclease activity by 60 and 98%, respectively, compared to 25% inhibition by PFM39 (Fig. 1, D and E). Efficient inhibition by PFM03 and PFM01 is consistent with Mre11 endonucleolytic activity acting on DNA ends bound by DNA-PK in this assay.

CtIP is phosphorylated by several different kinases in human cells (Fig. 2A) (31). Some of these phosphorylation events have been shown to promote DNA end resection (8, 9, 32–38), most notably the cyclin-dependent kinase (CDK)–dependent phosphorylation of T847. In addition, Ataxia telangiectasia Mutated (ATM) and Ataxia telangiectasia and Rad3 related (ATR) phosphorylation of T859 also plays an important role in CtIP binding to chromatin and CtIP-mediated resection (38, 39). To test the function of CtIP phosphorylation in the MRN endonuclease assay, we purified recombinant CtIP proteins containing phospho-blocking or phospho-mimic mutations at the T847 and T859 sites (fig. S1). Addition of these mutant proteins to the MRN endonuclease assay with DNA-PK shows that T847 and T859 phosphorylation are critical, as the phospho-deficient T847A and T859A mutants resulted in loss of CtIP stimulation (15- and 3-fold stimulation of MRN, respectively, compared to 43-fold by wild type) (Fig. 2, B (lanes 5 to 10) and C). We know that there is substantial constitutive phosphorylation of CtIP expressed and purified from insect cells (20) and infer from this result that T847/T859 phosphorylated species in the recombinant protein preparation are essential for stimulation of the MRN endonuclease activity. In contrast to these sites, phosphorylation at S327 or the ATM sites S231, S664, and S745 appears not to be important for the endonucleolytic activity (Fig. 2B, lanes 11 to 13). We previously reported that the ATM phosphorylation sites are important for CtIP intrinsic nuclease activity (21). Here, we show that these sites do not regulate DNA-PK–dependent Mre11 endonuclease activity, consistent with our observation that the N289A/H290A mutations that reduce CtIP nuclease activity also do not affect Mre11 function in this assay.

While mutation of the CtIP T847 residue to phospho-blocking alanine inhibited its stimulation of Mre11 nuclease activity, the phospho-mimic T847E mutant completely restored this ability in vitro (Fig. 2B, lane 6). This is important, as the T847E allele of CtIP was previously shown to restore localization of CtIP to sites of DNA damage as well as the resection of DSBs and promoted efficient CDK-dependent processing of these breaks (36). Thus Mre11 endonuclease activity is linked with the cell cycle–dependent processing of DNA damage via the phosphorylation of T847 in CtIP and will therefore only occur efficiently during the S and G2 phases of the cell cycle. The T859E phospho-mimic version of CtIP only partially restores Mre11 activity (Fig. 2B, lane 8), also consistent with the initial study of T859 phosphorylation by ATR in Xenopus extracts, which reported that T859E (T818E in Xenopus CtIP) is only weakly active in supporting DSB resection in CtIP-depleted extracts (39). Overall, however, the combined data suggest that CDK- and ATR/ATM-mediated phosphorylation of CtIP on T847 and T859 is critical for stimulation of MRN endonuclease activity on DNA-PK–bound ends.

Exo1 is one of the critical long-range exonucleases that perform DNA end resection in human cells (4), and we have previously shown a positive effect of MRN on Exo1-mediated degradation of DNA (13, 41, 42). Despite the well-known importance of CtIP in resection in human cells, however, we have not observed any effects of recombinant CtIP on Exo1 activity in previous experiments. Here, we examined the effect of CtIP on Exo1 on a plasmid DNA substrate in the presence of MRN and DNA-PK (Fig. 2D). As expected, Exo1 is substantially blocked by the presence of DNA-PK in the reaction, an inhibition that is relieved by the inclusion of MRN and requires CtIP (Fig. 2D, lanes 4 to 6). This finding that CtIP stimulates the activity of Exo1 sevenfold in the presence of DNA-PK is consistent with the effect of CtIP on Mre11 nuclease activity we observe on the labeled substrates (Fig. 1).

We recently showed that MRN can remove DNA end blocks by sequential endo-exo-endo activities, generating a DSB adjacent to the block (8, 13), a model first suggested by processing of meiotic DNA breaks (43). In the presence of DNA-PK bound to DNA ends, we found that the MRN complex executes similar end processing of the radiolabeled substrate to generate a new DSB. We detected a ~45-bp dsDNA product using native polyacrylamide gel electrophoresis (PAGE) (Fig. 2E, lane 2), corresponding to the major product seen on denaturing gels (Fig. 1). This activity was dependent on MRN and was enhanced when both Ku and DNA-PKcs were present in the reaction (Fig. 2E, lane 2). However, the formation of the dsDNA product was less efficient (approximately 1 to 2% of the substrate cleaved at site proximal to the radiolabel) compared to the single-strand endonucleolytic product seen on denaturing gels (approximately 27 to 34% of the substrate cleaved adjacent to the labeled 5’ end) (Fig. 1, A and B).

The unexpected dependence of Mre11 nuclease activity on DNA-PK suggests that there may be specific protein–protein interactions underlying the cooperative behavior. We tested this by immobilizing purified, human CtIP on magnetic beads and monitoring binding of DNA-PK or MRN. Both MRN and DNA-PK bound to CtIP in an ATP-independent manner (Fig. 2F). Binding analysis of Ku and DNA-PKcs separately showed that each of the components of DNA-PK has affinity for CtIP (Fig. 2G). Binding of Nbs1 (added to this reaction separately from the MR complex) to CtIP is mostly independent of DNA-PK, while binding of MR to CtIP improves seven- to eightfold with DNA-PK present. Addition of EtBr or benzonase to the reaction did not affect the interaction, confirming that the binding is independent of DNA (fig. S3A). Although the stimulation of Mre11 nuclease activity requires CtIP phosphorylation on T847 and T859, we did not observe any obvious deficiency in binding between CtIP, MRN, and DNA-PK with the T847A/T859A CtIP mutant protein.
Fig. 2. MRN, DNA-PK, and CtIP promote dsDNA end resection. (A) A linear map of CtIP indicating a subset of known phosphorylated residues as well as residues important for DNA binding and catalytic activity as discussed in the text (orange, sites required for nuclease activity; green, ATM-dependent phosphorylation sites; blue, CDK-dependent phosphorylation sites; red, ATR/ATM-dependent phosphorylation site). (B) Nuclease assays were performed with MRN (12.5 nM), CtIP (40 nM), Ku (10 nM), DNA-PKcs (10 nM), and NU7441 as in Fig. 1A but with various mutants of CtIP, as indicated, in the presence of both magnesium and manganese. The red arrow indicates the predominant product formed in the presence of DNA-PKcs. (C) Nuclease assays were performed as in (B) with titrations of CtIP (10, 20, and 40 nM). (D) DNA end resection on a plasmid substrate (3.6 kb) was performed with MRN, CtIP, DNA-PK, and Exo1, as indicated, in the presence of a DNA-PKcs inhibitor. Reaction products were separated in a native agarose gel, which was stained with SYBR Green; molecular weight (MW) ladder migration is shown (kb). (E) dsDNA cleavage products from the MRN nuclease assay with DNA-PK and CtIP were detected on a 12% native polyacrylamide gel. The red arrow indicates the ~45-bp product. (F) Protein-protein interactions between CtIP, MRN, and DNA-PK were measured by IP with anti-CtIP antibody in the presence or absence of ATP followed by Western blotting of bound proteins, as indicated. (G) Interactions between CtIP, MRN, DNA-PKcs, and Ku were measured with CtIP IP as in (F) in the presence of ATP.
Deshpande et al., Sci. Adv. 2020; 6 : eaay0922     8 January 2020

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(fig. S3B); thus, the effect of the phosphorylation does not appear to be manifested through CtIP association with the other factors.

**Single-molecule observations show MRN/CtIP-mediated removal of DNA-PK and MRN from DNA ends**

To examine the interactions between DNA-PK and the MRN complex in greater detail, we turned to single-molecule microscopy. In this "DNA curtains" assay, arrays of lambda DNA molecules (~48 kb long) are attached via a biotin-streptavidin linkage to a microfluidic flow cell that is also passivated with a fluid lipid bilayer (Fig. 3A) (44). Microfabricated barriers align the DNA molecules and organize them for high-throughput analysis of bound proteins, which we have previously used to examine the interplay between the MRN complex and Ku at individual DNA ends (13). Here, we adapted this assay to observe the behavior of the DNA-PK complex, with DNA-PKcs labeled using a DNA-PKcs-specific antibody. Injection of fluorescently labeled DNA-PKcs led to only transient association with the DNA, as indicated by nonspecific sliding in the direction of buffer flow and subsequent release from the DNA ends [half-life (t1/2) = 5 ± 3 s, N = 17; Fig. 3, B and C]. Given that formation of the DNA-PK complex requires Ku and DNA (45), we considered that preloaded Ku may stabilize DNA-PKcs on the DNA ends. To test this, we injected hemagglutinin (HA)–tagged Ku heterodimer onto the DNA curtains, labeled with anti-HA antibody. As we observed previously, Ku bound only to DNA ends and did not slide on the DNA with buffer flow (13). Subsequent injection of DNA-PKcs did result in more stable association of the kinase with the Ku-bound ends; however, the inclusion of ATP in the buffer promoted rapid loss of ~80% of the bound Ku molecules (Fig. 3, D to F). These are single-turnover reactions that we observe with constant buffer flow, so even transient release of DNA-PKcs from the DNA results in loss of association. We considered that the ATP-dependent release

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**Fig. 3. Single-molecule visualization of DNA-PK on DNA.** (A) Schematic of the DNA curtains assay for DNA-PK. (B) Illustration and kymograph (time series of one molecule over the course of the reaction) of DNA-PKcs injection onto DNA curtains. White arrow indicates a single DNA-PKcs binding event. This molecule then slides along the DNA in the direction of buffer flow to reach the DNA end (green arrow). The molecule then associates for a short time at the end (t) before dissociation (red arrow). (C) Lifetime of DNA-PKcs on DNA curtains in the presence of ATP. (D) Illustration and kymographs of DNA-PKcs colocalizing with an end-bound Ku and Ku(5A) in the presence or absence of ATP as indicated. (E and F) Lifetime of Ku (WT or 5A) on DNA curtains in the presence or absence of DNA-PKcs or ATP as indicated. Table shows half-life of Ku(WT) or Ku(5A) under various conditions and the number of molecules observed (N).
might be associated with the phosphorylation of Ku70 by DNA-PKcs, which inhibits DNA binding by Ku (25). To prevent Ku phosphorylation and release by DNA-PKcs, we purified a phospho-blocking mutant of Ku(T305A/S306A/T307A/S314A/T316A, "5A"), which inhibits DNA-PKcs–induced release of Ku (25). Ku(5A) bound to DNA for >2000 s, consistent with our previous observations of extremely stable binding of wild-type Ku (Fig. 3, E and F) (13). Ku(5A) stabilized DNA-PKcs at the DNA ends even in the presence of ATP (Fig. 3, D to F), confirming that phosphorylation of the Ku70 S/TQ 305–316 cluster controls the association of DNA-PK with DNA ends.

We next observed the interplay between MRN and DNA-PK [DNA-PKcs with Ku(5A)] using the DNA curtains platform. Consistent with our ensemble biochemical assays (Fig. S2), injection of dark (unlabeled) MRN and CtIP onto DNA curtains with both fluorescent DNA-PKcs and fluorescent Ku in the presence of a physiologically relevant cleavage buffer (5 mM MgCl2) led to removal of both DNA-PKcs and Ku from the ends (Fig. 4A).

To probe the dynamics of this activity, we used fluorescently labeled MRN complex together with fluorescent DNA-PKcs (Ku is unlabeled in this case) and measured the occupancy of DNA-PK on ends that are also bound by MRN. Injection of MRN alone led to stable colocalization of the two complexes (DNA-PK 1/2, >2000 s), but injection of MRN and CtIP together led to removal of both Ku and DNA-PKcs from the DNA ends (1/2 = 324 ± 15 s, N = 37; Fig. 4, B to D). In contrast, an MRN nuclease–deficient mutant (H129N) with CtIP did not remove DNA-PK; neither did CtIP added in the absence of MRN. Furthermore, injection of MRN with CtIP containing phospho-blocking mutations T847A and T859A also failed to remove DNA-PK (Fig. 4, B to D). These data suggest that colocalization of MRN with the DNA-PK complex is not sufficient to facilitate removal and that, consistent with our ensemble assays, phosphorylated CtIP is required for DNA-PK removal by MRN nuclease activity. It is notable that the rate of DNA-PK removal by MRN/CtIP under these conditions (1/2 = 324 ± 15 s) is faster than the removal of Ku by DNA-PK–mediated phosphorylation (1/2 = 780 ± 30 s); thus, the MRN-mediated pathway is likely to occur in a time frame comparable to other pathways of Ku removal in cells.

Our previous studies showed that MRN could remove Ku from DNA ends via a manganese-dependent reaction (8, 13). Here, we observe approximately the same kinetics of DNA-PK release catalyzed by MRN and CtIP as we did previously for Ku, but in magnesium–only conditions (all of the single-molecule experiments shown here were performed in the absence of manganese). These results demonstrate that DNA-PK stimulates Mre11 nuclease activity in the absence of manganese in physiologically relevant conditions and that CtIP is an essential component of this reaction.

These observations suggest a conserved mechanism for removal of blocks from DNA ends by the MRN complex (model in Fig. 4E). At two-ended DSBs, NHEJ would first engage the ends, and in cases where the ends are compatible, the ends would undergo productive end-joining. In situations where NHEJ fails and the DNA-PK complex remains at the DSB ends, Ku could be removed by DNA-PK–mediated phosphorylation (25). Alternatively, MRN can remove the complex through endonucleolytic processing, a reaction that is greatly enhanced by phosphorylated CtIP and is dependent on DNA-PK, as we show in this study. MRN activity nick either one or both the strands close to the DNA-PK complex in a sequential or simultaneous manner. The nicked or freed end is then used for extensive 5′ to 3′ resection by exonucleases, allowing damage repair through homologous recombination or homology-dependent mechanisms.

**MRN nuclease–mediated cutting of DNA-PK–bound ends occurs in human cells**

To determine whether nucleolytic removal of DNA-PK occurs in human cells, we performed a chromatin immunoprecipitation (ChIP) assay using the inducible AsiSI restriction enzyme system in U2OS osteosarcoma cells (46). We modified the standard ChIP protocol (fig. S4) by reducing the duration of formaldehyde crosslinking and eliminating the extensive sonication used to fragment DNA. This gentle lysis procedure, which we are naming "Gentle Lysis And Size Selection-ChIP," or GLASS-ChIP, allowed us to specifically separate small (50 to 300 bp) DNA-PK–associated DNA fragments from the bulk chromatin. The fragments were then further purified and used for quantitative polymerase chain reaction (qPCR) analysis. We monitored four genomic AsiSI sites (47) and found that this procedure generated DNA-PK–associated DNA fragments at all four sites that were strongly dependent on 4-hydroxytamoxifen (4-OHT) addition (Fig. 5A).

When cells were exposed to the DNA-PKcs inhibitor NU7441 during AsiSI induction, quantitation of DNA located very close (~30 nt) to the AsiSI genomic sites showed a 25- to 250-fold increase over background levels (i.e., levels of product formed with NU7441 but in the absence of 4-OHT induction) (Fig. 5A), consistent with the nucleolytic removal of DNA-PK–bound DNA ends we observed in vitro. These levels dropped substantially when measuring sites located farther away (~300 nt) from the AsiSI cut site (fig. S5), and no signals above background were observed at representative locations distant from AsiSI sites. With a DNA-PKcs inhibitor present as it is here, we expect that NHEJ is blocked and MRN cleavage of DNA-PK–bound ends is maximal, as we observed in purified protein reactions (Figs. 1 and 2).

Induction of AsiSI with 4-OHT in the absence of a DNA-PKcs inhibitor also generated DNA in the GLASS-ChIP assay, approximately 3- to 160-fold higher than background depending on the site, measured with primers 30 nt from the AsiSI location (Fig. 5A, inset). Under these conditions, NHEJ is not blocked; thus, the release of DNA-PKcs with associated DNA is expected to occur only as a consequence of DSB processing. The observation of these products in the absence of a DNA-PKcs inhibitor shows that processing of DNA-PKcs–bound ends occurs in human cells under normal physiological conditions.

To determine whether the DNA-PK–bound products arise through Mre11 nuclease activity, we treated the cells with the endonuclease inhibitor PFM03 based on our in vitro observations (Fig. 1, D and E). In preliminary experiments, we found that addition of 4-OHT to induce AsiSI activity in cells also exposed to PFM03 (100 μM) and the DNA-PKcs inhibitor NU7441 resulted in complete cell death within 1.5 hours. To circumvent this, we limited the treatment with 4-OHT and both DNA-PKcs and Mre11 inhibitors to 1 hour. In the absence of the Mre11 inhibitor, we observed short DNA fragments at three of four of the genomic loci tested that were dependent on 4-OHT (Fig. 5B), albeit at 10- to 20-fold reduced levels compared to the 4-hour 4-OHT induction (Fig. 5A). With the addition of PFM03, there was a significant reduction in the recovery of these fragments, indicating that Mre11 nuclease activity is responsible for the creation of DNA-PK–bound DSB products (Fig. 5B).
DISCUSSION
Here, we have shown that DNA-PK plays an important role in DNA end processing through its stimulation of Mre11 endonuclease activity. The extremely high concentration of DNA-PK in mammalian cells, particularly in human cells (5), and the high affinity of Ku for DNA ends (48) mean that DSB ends will be immediately bound by DNA-PK (see model in Fig. 4E). Removal of this block can be promoted by several distinct mechanisms: DNA-PKcs phosphorylation of Ku70, which reduces its DNA binding affinity (25); ubiquitination of Ku and degradation by proteasome-dependent or proteasome-independent mechanisms (49–51); or single-strand or double-strand endonucleolytic incision of DNA by MRN, as we show in this study. Single-strand DNA cutting by Mre11 followed by 3’ to 5’ Mre11 exonuclease activity was demonstrated as a

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Fig. 4. Single-molecule visualization of DNA-PK removal by MRN and CtIP. (A) Illustration and kymographs of DNA-PKcs (magenta) and Ku (green) upon injection of MRN and CtIP (both unlabeled [dark]). (B) Kymographs of DNA-PKcs (magenta) with Ku (unlabeled), with injection of MRN (green) alone (top), or MRN with CtIP (middle), or a nuclease-dead mutant of MRN (H129N) with CtIP (bottom). (C and D) Lifetimes and associated half-lives of the DNA-PK complex (as observed by DNA-PKcs occupancy) upon no injection (green), injection with MRN and CtIP (black), MRN alone (red), nuclease-deficient H129N MRN and CtIP (magenta), CtIP only (blue), or MRN with phospho-blocking T847A/T859A CtIP (purple). Table shows half-life of DNA-PKcs under various conditions and the number of molecules observed (N). (E) Schematic model for DNA-PK removal from DSB ends, as discussed in the text. DNA-PKcs binds to the Ku heterodimer bound to DNA at the DSB. Transition from NHEJ to homologous recombination (HR) results from (i) DNA-PKcs phosphorylation of Ku, resulting in dissociation of Ku as well as DNA-PKcs from ends, (ii) DNA-bound DNA-PK stimulation of MRN single-strand endonucleolytic cleavage followed by 5’ to 3’ resection, or (iii) DNA-PK stimulation of MRN double-stranded endonucleolytic cleavage resulting in DNA-PK loss and 5’ to 3’ resection. Long-range 5’ to 3’ resection ensues from the nick or a new DSB, creating 3’ single-stranded DNA that is used for homology search. The DNA-bound DNA-PK complex (dashed box) is the species isolated in the modified ChIP protocol (Fig. 5 and fig. S4).
mechanism for Spo11 removal by Mre11-Rad50-Xrs2 (MRX) in yeast (43).

We also demonstrate in this work that double-strand DNA cutting by MRN is promoted by stable occupancy of DNA-PK on the ends, as shown by the increased efficiency of MRN cleavage in the presence of a DNA-PKcs inhibitor in vitro and by our observation of the release of DNA-PK-bound DNA fragments in human cells. This requirement for stable occupancy of DNA-PK ensures that MRN does not cleave ends that are undergoing successful end-joining, as this outcome also leads to DNA-PKcs dissociation (1). DNA-PK and MRN have often been viewed as competitors for DNA ends, but the data presented here show that

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**Fig. 5.** DNA-PK–associated DNA end fragments are observed at AsiSI breaks in human cells using GLASS-ChIP. (A) Small dsDNA products resulting from nucleolytic cleavage of DNA-PK–bound AsiSI–generated DNA ends (dashed line box, Fig. 4E) were isolated from U2OS cells treated with 4-OHT or vehicle for 4 hours as indicated. NU7441 (10 μM) was added as indicated for 5 hours starting at 1 hour before 4-OHT addition to induce AsiSI. DNA-PK–bound DNA was isolated using a modified ChIP protocol (GLASS-ChIP, fig. S4) and quantified by qPCR using primers located ~30 nt from the AsiSI cut site (results from primers ~300 nt from cut sites in fig. S5). Primer set U1 (solid) is upstream, whereas D1 (checkered) is downstream of the AsiSI cut sites. The DNA quantitated from U2OS cells in the presence or absence of 4-OHT and a DNA-PKcs inhibitor (NU7441) is shown for each AsiSI site (46). The inset magnifies the results for experiments performed in the absence of NU7441. Results are from three independent biological replicates, with Student’s two-tailed t test performed; *P < 0.05, **P < 0.01, ***P < 0.001, in comparison to equivalent samples without 4-OHT. (B) The GLASS-ChIP protocol was performed as in (A) using cells treated with a DNA-PKcs inhibitor (NU7441, 10 μM), a Mre11 inhibitor (PFM03, 100 μM), and 4-OHT for 1 hour as indicated. Results are from three independent biological replicates, with Student’s two-tailed t test performed; **P < 0.005 and ****P < 0.0001, in comparison to equivalent samples without PFM03.
these complexes are linked in an ordered pathway (Fig. 4E), consistent with cytological observations (52). The dual roles of DNA-PKcs ensure rapid repair by NHEJ as well as an efficient transition to homologous recombination by promoting MRN-dependent initiation of end processing.

Loss of DNA-PKcs in mice results in radiosensitivity as well as immunodeficiency due to a failure to complete V(D)J recombination, and naturally occurring mutations in the gene have generated viable but immunocompromised animals (53). In contrast, no human patients have been identified with complete loss of DNA-PKcs, and human cell lines lacking the protein exhibit a severe growth deficit in addition to DNA damage sensitivity, extreme genomic instability, and telomere shortening (54). Similarly, Ku70- and Ku80-deficient mice are radio-sensitive but viable, yet deletion of Ku subunits in human cells generates spontaneous DNA breaks, telomere defects, and cell lethality within several divisions (55). This combined set of observations shows that DNA-PK plays many roles in human cells and is more essential for the maintenance of genomic stability in human cells compared to rodents or other mammals. The function of DNA-PKcs in regulating MRN/CtIP-mediated end processing that we describe in this work is likely an important component of the responsibilities of DNA-PKcs in genome maintenance. Further structural analysis is necessary to fully understand how phosphorylated CtIP enables the MRN complex to breach the otherwise impenetrable block that DNA-PK creates on DNA ends and how the many posttranslational modifications on CtIP, MRN, and other factors present at ends can modulate these outcomes. The extreme sensitivity of U2OS cancer cells to simultaneous use of DNA-PKcs and Mre11 inhibitors points toward a potential use of these small molecule inhibitors in treatment of cancers with active homologous recombination pathways.

MATERIALS AND METHODS

Plasmid, strains, and protein purification

Recombinant human proteins [MRN wild-type, M(H129N)RN, MR, MR(K42A), MR(D123A), Nbs1, CtIP wild type, and CtIP mutants] were expressed and purified from Sf21 insect cells as described previously (8, 21, 24). 3XFLAG-MRN, 3XHA-Ku70/80 heterodimer, and Exo1-biotin were made as described previously (13). DNA-PKcs from human cells was purified as described previously (56). The Ku70(5A) mutant (T305A/S306A/T307A/S314A/T316A) was made from the baculovirus transfer vector containing wild-type Ku70 by QuikChange mutagenesis (Stratagene) to create the 5A mutant expression plasmid pTP4143 and the corresponding bacmid pTP4177, from the baculovirus transfer vector containing wild-type Ku70 by QuikChange mutagenesis (Stratagene) to create the 5A mutant expression plasmid pTP4143 and the corresponding bacmid pTP4177, which was used with wild-type Ku80 baculovirus to infect Sf21 cells and purify Ku(5A).

DNA substrates

TP542 (GGTTTTCGGATCCGACGGGTTG) was 5′ [32P] radiolabeled using T4 polynucleotide kinase (New England Biolabs) and purified (New England Biolabs Nucleotide Removal Kit). The 197-bp DNA substrate labeled on the 5′ end was created by PCR using 5′ [32P] radiolabeled TP542 (GGTTTTCGGATCCGACGGGTTG) and TP4373 (GAGGATCGGAGATGGAGAATCGTCAATAATATATGACGTTT) with pTP3718 followed by purification on a native 0.7% agarose gel as described previously (8). The DNA substrate used in Fig. 2A was made by digesting pCDF-1b plasmid with SspI to create a 3.6-kb linear DNA with blunt ends and was used without further purification. The lambda DNA substrate used in single-molecule studies was prepared as described previously (13).

Nuclease assays

Ten microliters of nuclease assays contained Ku70/80 (10 nM), DNA-PKcs (10 nM), hMR (50 nM), Nbs1 (50 nM), and CtIP (80 nM) unless indicated otherwise in figure legends. Reactions were performed with 0.1 nM DNA in 25 mM 3-(N-morpholino)propanesulfonic acid (MOPS) (pH 7.0), 20 mM Tris (pH 8.0), 80 mM NaCl, 8% glycerol, 1 mM dithiothreitol (DTT), 1 mM ATP, 0.2 mM NU7441, 5 mM MgCl2, 1 mM MnCl2, and bovine serum albumin (BSA) (0.2 mg/ml), with additions or exceptions as noted in the figures and legends. hMR and Nbs1 were preincubated (10 min at 4°C) before addition to reactions, and Ku and DNA-PKcs were allowed to bind DNA in the presence of NU7441 for 5 min at 25°C followed by the addition of other components. Assays were carried out in Protein LoBind Eppendorf tubes (Millipore) at 37°C for 30 min. Reactions were stopped with 0.2% SDS and 10 mM EDTA, lyophilized, dissolved in formamide, boiled at 100°C for 4 min, loaded on denaturing polyacrylamide gels containing 16% acrylamide, 20% formamide, and 6 M urea, and separated at 40 W for 1.5 hours, followed by phosphorimager analysis. Mre11 inhibitors were added at final concentrations of 25, 50, and 100 µM in assay reactions. Mre11 inhibitor dilutions were made in 20% dimethyl sulfoxide (DMSO) in buffer A, which results in 6% DMSO in these assay reactions, and an equivalent concentration of DMSO was used in control reactions. For the detection of dsDNA products, reactions were treated with 10 µg of proteinase K at 37°C for 1 h after stopping the nuclease reaction. The samples were then separated using 12% native PAGE in 1× tris-borate-EDTA (TBE) buffer at 150 V followed by phosphorimager analysis.

DNA resection assays on plasmid substrates were carried out similarly using 0.1 nM 3.6-kb pCDF-1b plasmid linearized with SspI and 1 nM Exo1 in the presence of the DNA-PKcs inhibitor NU7026 (200 µM). Reactions were stopped with 0.2% SDS and 10 mM EDTA, and 1.5 µg of proteinase K was added and incubated at 37°C for 90 min followed by 10-min incubation at 50°C. These reactions were then separated in a 0.7% native agarose gel in 1× TBE buffer, stained with SYBR Green, and visualized with a ChemiDoc gel imager (Bio-Rad).

Protein-protein interactions

IP assays were performed with anti-CtIP antibody (Active Motif 14E1) in Protein LoBind Eppendorf tubes (Millipore). In a 40-µl reaction volume, 20 nM Ku70/80, 20 nM DNA-PKcs, 25 nM hMR, 25 nM Nbs1, and 16 nM CtIP were incubated as indicated in the absence of DNA in 25 mM Mops (pH 7.0), 20 mM tris (pH 8.0), 80 mM NaCl, 8% glycerol, 1 mM DTT, 1 mM ATP, 0.2 mM NU7441, 5 mM MgCl2, and BSA (0.2 mg/ml) at 37°C for 10 min. The reaction was then incubated with 2 µg of anti-CtIP antibody, 0.4% CHAPS, and 2 µl of protein A/G magnetic beads (Pierce) at 37°C for 30 min with shaking. Beads were isolated using a magnet and washed three times with wash buffer [buffer A [25 mM tris (pH 8.0), 100 mM NaCl, 10% glycerol] containing 0.4% CHAPS and BSA (0.1 mg/ml)]. Tubes were changed at the third wash step, and proteins bound to the beads were eluted with 1× SDS-PAGE loading dye prepared in wash buffer. Samples were heated at 100°C for 5 min and separated by 8% SDS-PAGE followed by Western blotting. Because Mre11 and Ku80 migrate very close to each other, blots were developed...
sequentially—first for FLAG-tagged CtIP, Nbs1, and Mre11, then for DNA-PKcs and His-tagged Rad50 and Ku70, and lastly for biotin-tagged Ku80. Proteins were detected using rabbit anti-FLAG (Cell Signaling), rabbit anti–DNA-PKcs (Bethyl Laboratories), and mouse anti-His (GenScript) antibodies as well as streptavidin–Alexa Fluor 680 conjugate (Invitrogen) using an Odyssey imager (Li-Cor).

**Single-molecule experiments**

Single-molecule DNA curtains were performed as previously described (13). For labeling of DNA-PKcs, 400 nM anti-mouse secondary Qdots (Thermo Fisher) were incubated with 360 nM anti-DNA-PKcs antibody (Abcam ab44815) on ice for 10 min in 5 μl of DNA-PKcs loading buffer [40 mM tris-HCl (pH 8.0), BSA (0.2 mg/ml), 5 mM MgCl₂, 1 mM ATP, and 2 mM DTT]. After buffer switch, buffer [40 mM tris-HCl (pH 8.0), BSA (0.2 mg/ml), 5 mM MgCl₂, 1 mM ATP, and 2 mM DTT]. DNA-PKcs was added to this mixture to a final concentration of 267 nM Qdots, 240 mM antibody, and 167 mM DNA-PKcs in 7.5 μl for another 10 min on ice. This mixture was diluted to a final volume of 200 μl (10 nM Qdots, 9 mM antibody, and 6.25 nM DNA-PKcs). Ku(5A) was loaded on DNA curtains essentially as described previously for wild-type Ku (13). Next, DNA-PKcs was injected onto DNA curtains from a 100-μl/min loop at 200 μl/min in DNA-PKcs loading buffer. For MRN cleavage experiments, after DNA-PKcs loading, the buffer was switched to MRN cleavage buffer [40 mM tris-HCl (pH 8.0), 60 mM NaCl, BSA (0.2 mg/ml), 5 mM MgCl₂, 1 mM ATP, and 2 mM DTT]. After buffer switch, mixtures of 3 nM MRN, 14 nM CtIP, or both were loaded as previously described (13).

**GLASS-ChiP assay**

Human U2OS cells with inducible AsiSI (46) were grown to 50 to 60% confluency in 150-mm dishes and treated with 600 nM 4-OHT for 4 hours at 37°C. When used, 10 μM NU7441 was added 1 hour before 4-OHT addition. After 4-OHT treatment, cells were fixed with 1% formaldehyde for 7 min at room temperature (RT) with gentle rotation. Crosslinking was stopped by addition of 125 mM glycine for 5 min, washed twice with cold phosphate-buffered saline, harvested, flash-frozen in liquid nitrogen, and stored at −80°C. For the ChiP assay, we modified a standard protocol (Abcam) with a gentle lysis procedure and minimal, low-level sonication to rupture cells without extensive DNA damage. The formaldehyde-fixed cells were thawed at RT for 5 min, resuspended in radioimmunoprecipitation assay buffer [50 mM tris-HCl (pH8.0), 150 mM NaCl, 2 mM EDTA (pH8.0), 1% NP-40, 0.5% sodium deoxycholate, and 0.1% SDS] with 1× protease inhibitors (Pierce, A32955), and sonicated using a QSonica sonicator at low power (amplitude 20) for 10 s, followed by 10 pulses after a 20-s interval. Cell lysates were then centrifuged at 3000 rpm for 3 min at RT to remove the bulk of chromatin. The supernatant was then incubated with 1.6 μg of anti-DNA-PKcs pS2056 antibodies (Abcam 124918) overnight at 4°C, followed by incubation with 25 μl of protein A/G magnetic beads (Pierce) at RT for 2 hours. Beads were then washed sequentially once in low-salt wash buffer [0.1% SDS, 1% Triton X-100, 2 mM EDTA, 20 mM tris-HCl (pH 8.0), and 150 mM NaCl], once in high-salt wash buffer [0.1% SDS, 1% Triton X-100, 2 mM EDTA, 20 mM tris-HCl (pH 8.0), and 500 mM NaCl], and once in LiCl wash buffer [0.25 M LiCl, 1% NP-40, 1% sodium deoxycholate, 1 mM EDTA, and 10 mM tris-HCl (pH 8.0)]. Beads were then resuspended in TE buffer [10 mM tris (pH 8.0) and 1 mM EDTA] and transferred to a fresh tube and finally eluted with 100 μl of elution buffer (1% SDS and 100 mM NaHCO₃). Crosslinks were reverted for the elutions (65°C for 24 hours). DNA fragments <300 bp from ChiP elution were separated by pulling down larger fragments using paramagnetic AMPure XP beads: 65 μl of AMPure XP beads was added to the uncrosslinked ChiP elution and mixed thoroughly. After 10-min incubation at RT, beads were isolated using a magnet. The supernatant was collected, and 25 μl of fresh AMPure XP beads was added and mixed thoroughly. After 10 min at RT, beads were separated and the supernatant was cleaned using a Qiagen Nucleotide Cleanup kit. At the final step, DNA was eluted in 60 μl of elution buffer and used for qPCR. Four different AsiSI sites were monitored using up- and downstream primers within 40 to 52 bp from the AsiSI cut site (table S1). %DNA was calculated using the following equation: \( \frac{2(\text{Ct( INPUT)} - \text{Ct( TEST)})}{\text{Ct( TEST)}} \times 100 \)%. DNA values obtained for IP in the absence of antibodies were subtracted from the values obtained in the presence of antibody and used for plotting the graphs in Fig. 5. To monitor the Mre11 nuclease dependence, we treated the human U2OS cells expressing inducible AsiSI with 10 μM NU7441, 100 μM PFM03, and 600 nM 4-OHT simultaneously for 1 hour at 37°C before harvesting cells.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/6/2/eay0922/DC1

Fig. S1. Purified recombinant proteins used in this study.

Fig. S2. Ku(5A) promotes MRN-dependent endonuclease activity.

Fig. S3. Interactions between CtIP, DNA-PKcs, Ku, and MRN do not depend on DNA or CtIP phosphorylation.

Fig. S4. Schematic diagram of GLASS-ChiP protocol.

Fig. S5. Quantification of DNA immunoprecipitated with anti-DNA-PKcs pS2056 using primers 300 bp away from AsiSI DBS.

Table S1. Primers used for qPCR in Fig. 5 and fig. SS.

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