Mechanism of Replication-Coupled DNA Interstrand Crosslink Repair

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SUMMARY

DNA interstrand crosslinks (ICLs) are toxic DNA lesions whose repair occurs in the S phase of metazoans via an unknown mechanism. Here, we describe a cell-free system based on Xenopus egg extracts that supports ICL repair. During DNA replication of a plasmid containing a site-specific ICL, two replication forks converge on the crosslink. Subsequent lesion bypass involves advance of a nascent leading strand to within one nucleotide of the ICL, followed by incisions, translesion DNA synthesis, and extension of the nascent strand beyond the lesion. Immunodepletion experiments suggest that extension requires DNA polymerase ζ . Ultimately, a significant portion of the input DNA is fully repaired, but not if DNA replication is blocked. Our experiments establish a mechanism for ICL repair that reveals how this process is coupled to DNA replication.

INTRODUCTION

DNA interstand crosslinks (ICLs), which covalently link the two strands of the double helix, are highly cytotoxic because they block DNA replication and transcription (Niedernhofer et al., 2005). While ICL-forming agents are commonly used in cancer chemotherapy, ICLs are also formed by endogenous cellular metabolites (Scharer, 2005). The mechanism by which ICLs are repaired in eukaryotes is unknown, but important clues come from genetic studies, which have shown that various classes of proteins are required to confer cellular resistance to ICL-inducing agents (Dronkert and Kanaar, 2001; Niedernhofer et al., 2005). One such class is the structure-specific endonucleases, which include Mus81-Eme1 and the nucleotide excision repair (NER) factor, Xpf-Ercc1, both of which are thought to perform incisions near the ICL. Another class is the translesion DNA polymerases. Among these, Rev1 and DNA polymerase ζ are particularly important (Niedzwiedz et al., 2004; Simpson and Sale, 2003). Proteins involved in homologous recombination (HR), such as Rad54, Xrcc2, and Xrcc3, are also required (Nojima et al., 2005). Finally, thirteen Fanconi anemia gene products (FA proteins) are essential for resistance to ICLs and suppression of chromosomal instability (Niedernhofer et al., 2005; Wang, 2007). Eight FA proteins form a nuclear protein complex that is thought to monoubiquitylate FancD2 and FancI, an event that is crucial for cellular resistance to ICL agents. Mutations in FA proteins give rise to Fanconi anemia, a cancer predisposition syndrome.

Indirect evidence suggests that in metazoans, the principal ICL repair pathway occurs in S phase. First, regardless of when they are treated with ICL agents, mammalian cells arrest late in Sphase with 4N DNA content (Akkari et al., 2000). Second, ICL agents lead to the formation of dsDNA breaks, but only after passage through S phase (Akkari et al., 2000; De Silva et al., 2000; Rothfuss and Grompe, 2004). Third, activation of the Fanconi anemia pathway occurs exclusively in S phase (Rothfuss and Grompe, 2004; Taniguchi et al., 2002), it is DNA replication dependent (Sobeck et al., 2006), and FA mutant cells display a prolonged S phase arrest (Akkari et al., 2001). Based on these observations and the known enzymatic properties of relevant DNA repair enzymes, a model for ICL repair in S phase has been proposed (reviewed in McHugh et al., 2001; Niedernhofer et al., 2005; Figure S1 available online). Accordingly, ICL repair is initiated by collision of a DNA replication fork with the ICL. Mus81-Eme1 and/or Xpf-Ercc1 then perform dual incisions on one parental strand surrounding the ICL, causing a dsDNA break and release of one of the replicated sister chromatids. The 3' end generated in the parental strand via incision is extended by translesion DNA polymerases past the remaining mono-adduct, which is then removed, likely via NER. Finally, the replication fork is re-established using HR. Notably, cells deficient for NER factors other than Xpf-Ercc1 are not particularly sensitive to ICL agents (De Silva et al., 2000 and references therein). Therefore, in the absence of NER, the adducted chromosome shown in Figure S1D is envisioned to be the homology donor used to re-establish the replication fork. Although the above model is attractive, it is largely hypothetical.

To elucidate the molecular mechanism of ICL repair in S phase, a biochemical system that supports this process will be vital. Extracts derived from mammalian cells have been shown



Figure 1. DNA Replication Forks Converge on an Interstrand Crosslink

(A) Structure of a nitrogen mustard-like ICL (postulated).

(B) Structure of a cisplatin ICL (based on Huang et al., 1995).

(C) Replication of pICL^{Nm} in *Xenopus* egg extracts. pCtr or pICL^{Nm} was incubated sequentially with HSS and NPE/³²P- α -dATP. At the indicated times after NPE addition, replication products were analyzed on a native agarose gel.

(D) The average replication efficiency of three independent experiments was plotted with error bars.(E) Model for replication of pICL.

(F) pICL^{Nm} was replicated as in (C). Thirty minutes after NPE addition, DNA was analyzed by electron microscopy. The predominant species, a "Figure 8" structure, is shown.

crosslinking agents (Dronkert and Kanaar, 2001). A short duplex oligonucleotide containing a nitrogen mustard-like ICL (Figure 1A) or cisplatin ICL (Figure 1B) was ligated into a 5.6 kb plasmid, generating pICL^{Nm} and pICL^{Pt}, respectively. The sequences of pICL^{Nm} and pICL^{Pt} were identical, except for a small region of about 20 base pairs surrounding the ICL (Figure S2A). For each construct, we also prepared a matched,

to promote incisions on plasmids containing ICLs (e.g., Bessho, 2003; Li et al., 1999; Mu et al., 2000), but the reactions are generally not performed in the context of S phase. To improve upon these approaches, we employed a soluble extract system derived from Xenopus eggs, which supports efficient replication of plasmid DNA templates using a physiological mechanism (Walter et al., 1998). Here, we show that Xenopus egg extracts support replication-dependent repair of plasmids containing a single, site-specific ICL. Using this system, we establish a detailed mechanism of ICL repair that includes several unexpected features. Thus, we observe that repair involves convergence of two DNA replication forks on the lesion. Subsequently, a multistep lesion bypass reaction is initiated before incisions are detected near the ICL. Finally, in contrast to prevailing models, lesion bypass involves extension of a nascent leading strand, illustrating how ICL repair is coupled to DNA replication. In addition, our data suggest that ICL repair requires DNA polymerase ζ, and that it is accompanied by activation of the ATR checkpoint and Fanconi anemia pathways.

RESULTS

Convergence of DNA Replication Forks at a Site-Specific ICL

To develop a homogeneous substrate for ICL repair, we prepared plasmids containing a single, site-specific and chemically defined ICL. This approach avoids the diverse spectrum of lesions generated when cells are treated with DNA interstrand undamaged plasmid of identical sequence termed pCtr. At least \sim 99% of pICL plasmids contained an intact interstrand crosslink (see Experimental Procedures and below). The nitrogen mustard-like ICL was designed to fit in the major groove of DNA without distorting the double helix (Figure 1A; T.A. and O.D.S., unpublished data). The cisplatin ICL (Hofr and Brabec, 2001) caused a significant helical distortion (Figure 1B; Huang et al., 1995). By comparing pICL^{Nm} and pICL^{Pt}, we sought to address whether the chemical structure of an ICL influences the mechanism of its repair.

We first analyzed how pICL^{Nm} was replicated in Xenopus egg extracts (Walter et al., 1998). In this system, plasmids are first incubated in a high-speed supernatant (HSS) of egg cytoplasm, which supports the assembly of prereplication complexes (pre-RCs) via the sequential recruitment of ORC, Cdt1, Cdc6, and MCM2-7 to DNA in a sequence nonspecific manner. Subsequently, a highly concentrated nucleoplasmic egg extract (NPE) is added. NPE triggers Cdk2-dependent replication initiation from pre-RCs, and a single, complete round of DNA replication occurs, which can be monitored via the incorporation of ³²P-α-dATP. During replication of the undamaged control plasmid (pCtr), partially replicated plasmids and fully replicated, catenated daughter molecules appear within 15 min of NPE addition (Figure 1C, lane 1) (Walter and Newport, 2000). By 30 min, resolved daughter molecules appear as nicked and supercoiled circles (Figure 1C, lane 2). In contrast, during replication of pICL^{Nm}, the appearance of nicked and supercoiled products was severely inhibited (Figure 1C, lanes 5-8). Importantly,



incorporation of ${}^{32}P-\alpha$ -dATP was identical for both plasmids (Figure 1D). These data suggest that pICL^{Nm} was fully replicated but failed to undergo decatenation (Figure 1E). In support of this interpretation, electron microscopy showed that during replication of pICL^{Nm} but not pCtr, there was a massive accumulation of "Figure 8" structures, in which each hemisphere was equivalent to the size of pICL^{Nm} (Figures 1F and S3). Together, the data demonstrate that replication initiation and elongation are highly efficient on pICL, but that forks stall after converging on the ICL.

A Stepwise Mechanism for Bypass of an ICL Lesion

To determine where replication forks arrest relative to the ICL, we mapped the ends of their leading and lagging strands. pICL^{Nm} was replicated in the presence of ³²P- α -dATP and digested with AfIIII, which cleaves 149 nucleotides (nt) to the left and 540 nt to the right of the ICL (Figure 2A). Nascent products were analyzed on a sequencing gel. Figure 2B shows that by 18 min after NPE addition, leading strand products of the *left-ward* fork formed a prominent cluster of bands ~500–520 nt in length (lane 1, black arrow; see also red strand in Figure 2A), indicating that their 3' ends were located ~20–40 nt from the ICL. Lagging strand products of the same fork were smaller and more heterogeneous (Figure 2B, lane 1, blue bracket), placing their 5' ends between ~70 and ~290 nt from the crosslink (see Figure S4

Figure 2. Multistep Lesion Bypass of an Interstrand Crosslink

(A) Structure of the replicated AfIIII fragment that includes the ICL. S, primer used to generate the sequencing ladder shown in (B).

(B) Mapping of nascent strands during replication of pICL^{Nm} or pCtr (final concentration 1.2 ng/μ). At the indicated times after NPE addition, reaction products were digested with AfIIII and analyzed on a sequencing gel alongside a sequencing ladder derived from extension of primer S on pCtr (see A). Numbers to the left indicate the sizes of the sequencing products. Leading and lagging strands for the *rightward* and *leftward* forks are indicated. Square brackets show the positions of leading strands after the initial pausing, whereas red and green arrowheads show their location after advancing toward the ICL. Open arrowhead, extension product.

(C) Enlarged and darker exposure of the bottom part of the autoradiogram shown in (B). The most prominent species observed at 45 min are indicated on the right, with the predicted last nucleotide denoted by a single letter. The exact sizes of species T147-A150 were confirmed in Figure S9. (D) Cartoon-form depiction of the results in (C).

for the identification of leading and lagging strand products). Leading strand synthesis subsequently resumed, and by 45 min, \sim 50% of the \sim 520 nt long leading strands were extended to a length of \sim 540 nt (Figure 2B, lane 4, red arrowhead), indicating that they were very close

to the ICL. By 4 hr, the \sim 540 nt species disappeared and a new product of ~690 nt accumulated (Figure 2B, lane 8, open arrowhead). This ~690 nt "extension" product can only be generated when the AfIIII restriction fragment has been completely replicated, demonstrating that the lesion was fully bypassed. No discrete replication intermediates were detected at any time using pCtr (Figure 2B, lane 9 and data not shown). By cutting out a smaller piece of replicated pICL^{Nm} surrounding the lesion and running the nascent DNA on a high-resolution sequencing gel, we found that lesion bypass occurred without significant levels of deletions or insertions (Figure S5A). Importantly, by 4 hr, the ratio of radioactivity in the 689 bp AfIIII fragment to the 4.9 kb vector backbone was essentially the same for pICL^{Nm} as for pCtr, indicating that virtually all replicated molecules that remained in the reaction had undergone lesion bypass (Figure S6B, compare lanes 8 and 9). Using pICL^{Pt}, virtually identical results for lesion bypass were observed (Figures S7, S8, and S5B). Our results indicate that the leading strand of a DNA replication fork bypasses an ICL in a series of temporally resolved steps that are essentially identical for both substrates.

Leading Strands Initially Stall 20-24 nt from the ICL

We next examined where exactly the leading strand stalls upon initial collision of a replisome with an ICL. Close inspection of the autoradiogram in Figure 2B showed that the earliest leading strand products to accumulate were 110 to 126 nt long (Figure 2C, lane 1), indicating that the leading strand of the *rightward* replication fork advanced to within 24 nt of the crosslink (see green strand in Figure 2A). Notably, the *leftward* fork generated a similar cluster of leading strands, with the largest prominent product also stalling 24 nt from the ICL (Figures S4C and S4D). Since the leading strands of both replication forks stalled at precisely the same distance from the lesion on either side of the ICL, the initial pause site of DNA polymerase appears to be dictated by the inherent size of the replisome's footprint on DNA, rather than the nucleotide sequence surrounding the ICL. Interestingly, there was a range of initial pause sites on either side of the ICL. Perhaps the first fork to arrive obstructs the approach of the fork coming from the other side.

Notably, we found that on pICL^{Pt}, leading strand polymerases of both forks initially paused 4 nt closer to the ICL than on pICL^{Nm} (20 versus 24 nt; Figures S7 and S8). This could be explained if the DNA surrounding the distorting cisplatin ICL was easier to unwind than the DNA surrounding the nondistorting nitrogen mustard-like ICL.

After Initial Fork Stalling, Leading Strands Advance to within 1 nt of the ICL

Shortly after the initial stalling of the replisome, DNA synthesis resumed, and by 45 min ~50% of the leading strands had advanced to the site of the crosslink (Figure 2C, lane 4). Near the nitrogen mustard-like ICL, DNA synthesis came to a second stop, generating four major species of 147–150 nt (Figure 2C, lane 4). The longest product in this cluster (150 nt) ends 1 nt before the template base that participates in the ICL (the "-1" position; Figure 2D). A similar result was observed for pICL^{Pt}, except that at 45 min, most of the products stalled at the -1 position (see Figures S7C, lane 4 and S7D). The pausing at the -1 position on pICL^{Nm} and pICL^{Pt} may reflect the time required to recruit a translesion DNA polymerase such as Rev1.

The question arises whether one or both leading strands advance to the ICL from the initial -20 pause site. If both strands advanced, complete conversion of the -20 products to the -1products should occur. However, the results for both pICL^{Nm} and pICL^{Pt} showed that a significant number of leading strand products remained paused at the initial -20 position, even after the signal at the -1 position had peaked at 45 min (e.g., Figures 2B and 2C, lane 4; Figures S4 and S7). This observation suggests that on any given DNA template, only one of the two leading strands initially advances to the ICL. In summary, lesion bypass involves an initial pausing of two leading strands on either side of the crosslink ~ 20 nt from the lesion (24 nt for pICL^{Nm}; 20 nt for pICL^{Pt}) before one of the two forks advances to the -1 position. Subsequently, the growing leading strand is extended beyond the lesion, generating the complete 689 nt AfIIII restriction fragment.

Notably, as leading strands advanced toward the ICL, there was a loss of lagging strand products, which began between 30 and 45 min (e.g., Figure 2B compare lanes 3 and 4; see also Figures S4, S7, and S8). Since this loss was observed *before* leading strands were extended past the ICL (at which time they could be ligated to downstream Okazaki fragments), it is best explained by 5' to 3' exonuclease activity, which has previously been reported in *Xenopus* egg extracts (Toczylowski and Yan, 2006).

Evidence for Uncoupling of Replicated Sister Chromatids

Current models for ICL repair hypothesize that dual incisions occur on either side of the ICL on one strand of the parental duplex ("unhooking"), allowing sister chromatid separation and lesion bypass (Dronkert and Kanaar, 2001; Niedernhofer et al., 2005; Wang, 2007; see Figures S1B and S1C). To seek evidence of unhooking in our system, replicated pICL^{Nm} was digested with HincII, which converts the "Figure 8" structure into an 11.2 kb X-shaped species with 2.3 and 3.3 kb arms (Figure 3A). In the presence of a single incision near the ICL, HincII is predicted to generate an arm fragment and a Y structure, whereas dual incisions should liberate two arms and a linear 5.6 kb fragment (Figure 3A, bottom).

Figure 3B shows that the X-shaped structure expected from Hincll digestion was generated soon after addition of NPE (lane 1). Moreover, a very low level of all the intermediates expected from single incision events was detected. After 1 hr, there was a dramatic reduction in the abundance of the X-shaped molecules, and a concomitant increase in the 5.6 kb linear species (Figure 3B, lanes 6-8). The conversion of X-shaped molecules to linear products was not accompanied by accumulation of arm fragments (Figure 3B, lanes 6-8), as might have been expected (Figure 3A, bottom). We propose that these arms were generated by incision but are not visible due to a highly active resection activity (see Figures S10B, S10C, and S10F). This interpretation is consistent with the 5' to 3' exonuclease activity detected in Figure 2B, which degrades lagging strand products and which should also attack newly exposed 5' ends generated by incision (as depicted in Figure S10B). It is also consistent with the fact that between \sim 45 and \sim 120 min after addition of NPE, we commonly observe a 40%-70% loss of total replication products (e.g., Figure 3B) as well as a decline in total nascent strand products (Figure 2B, compare lanes 1 and 8). This loss of replication products is linked to repair since it is never observed with pCtr (Figure 2B, compare lanes 8 and 9 and data not shown), and we speculate it may reflect the activity of a 5' to 3' exonuclease that prepares the DNA template for HR. In summary, our results clearly show that after replication of pICL^{Nm}, the two sister chromatids that are initially connected by the ICL are separated, most likely via incisions near the ICL. This leaves a continuous daughter molecule that can serve as the DNA template for lesion bypass.

A Leading Strand Advances to the ICL *before* There Is Evidence of Incision

We wanted to know when the separation of sister chromatids occurs relative to the events underlying lesion bypass. We thus compared the abundance of the X-shaped species with that of the leading strand intermediates over time and plotted the results (Figure 3C). At 45 min, only ~15% of X-shaped molecules had disappeared, presumably due to incisions (Figure 3C, blue line), while the abundance of leading strands at the -1 to -4 position on pICL^{Nm} had already peaked (Figure 3C, gray dashed line). Assuming that only one leading strand on each replicated plasmid advances to the ICL, virtually every template in the reaction has undergone this process by 45 min. Similar results were observed with pICL^{Pt} (data not shown). Therefore, advance of



Figure 3. Detection of Incisions near the ICL

(A) Expected intermediates resulting from single or dual incisions near the ICL (see text). Note that the first incision could also occur to the left of the ICL, giving rise to a short arm and large Y structure.

(B) pICL^{Nm} replication products were digested with HincII and separated on a native agarose gel (lanes 1–8). Replicated pCtr was digested with HincII to generate a 5.6 kb size marker (lane 9, only 20% of the reaction loaded).

(C) Advance of the leading strand to the ICL precedes incisions. At each time point in (B), the relative abundance of X-shaped molecules was quantified using a phosphorimager (blue line). At 18 and 22 min, before replication was complete, the level was assigned a value of 100. The relative abundance of leading strand products from both forks at the -1 and -4 positions (Figure 2B) (gray dashed line) is plotted. The graph shows the average of four independent experiments with error bars.

the leading strand to within a few nucleotides of the ICL does not require prior incision.

ICLs Are Fully Repaired in a Replication-Dependent Manner

To determine whether any plasmids are fully repaired during incubation in Xenopus egg extract, we exploited the fact that one of the two Accl restriction sites in pICL^{Nm} coincides with the lesion (Figure S2A) and therefore cannot be cleaved by Accl (Figure 4A, lane 3). However, if the ICL is repaired in an errorfree manner, both Accl sites become accessible, and Accl digestion should yield two linear DNA restriction fragments (2.3 and 3.3 kb), as seen with pCtr (Figure 4A, lane 2). As shown in Figure 4B, lanes 10–17, incubation of pICL^{Nm} in NPE resulted in a gradual increase in the yield of 2.3 and 3.3 kb Accl restriction fragments, indicative of repair. However, a complication is that even on unrepaired molecules, any incisions near the ICL (shown as red arrows in Figure 4B) would result in 2.3 and 3.3 kb fragments after digestion at the distal Accl site. To identify the background level of these products, we digested an equal volume of the replicated plasmid with Hincll, whose single restriction site coincides with the distal Accl site. As shown in Figure 4B (lanes 1-8), 2.3 and 3.3 kb Hincll products arose early, then declined after 90 min, likely due to degradation. Therefore, to quantify bona fide regeneration of the proximal Accl site, we subtracted the Hincll digestion products (lanes 1-8) from the Accl digestion products (lanes 10-17) and plotted the difference (Figure 4C, red line). This analysis shows that the ICL-proximal Accl site began to be regenerated about 90 min after NPE addition and reached a plateau at 120 min, soon after the extension step of lesion bypass (Figure 4C). By 4 hr, \sim 8% of the DNA molecules that had initially undergone DNA replication were cleaved at both Accl sites. Given that there is significant degradation of the

DNA template during the reaction, the repair efficiency was higher (\sim 15%) if calculated as a percentage of molecules *remaining* in the reaction at each time point. Similar results were obtained for the pICL^{Pt} substrate (see below).

To address whether Accl cleavage at the ICL is dependent on DNA replication, we inhibited replication initiation by blocking Cdk2/Cyclin E activity with p27Kip (Walter et al., 1998). pICLNm was digested with AccI or HincII as in Figure 4B and visualized by Southern blotting. Figure 4D shows that when DNA replication was inhibited, no repair products were detected (e.g., compare lanes 3 and 9), suggesting that ICL repair is replication dependent. An alternative explanation is that the 2.3 and 3.3 kb Accl restriction fragments reflect the replication of a contaminating, uncrosslinked plasmid. However, this is unlikely for three reasons. First, Accl cleavage was significantly delayed relative to the bulk of DNA replication, which is normally complete within 20-30 min of NPE addition. Second, based on quantification of the Southern blot shown in Figure 4D (lanes 7-9 versus 13), less than 0.2% of pICL^{Nm} was cleavable by Accl at the ICL-proximal site in the absence of DNA replication, indicating that at least 99.8% of the molecules were modified at the restriction site. Third, 30 min after NPE addition, when DNA replication of pICL^{Nm} was essentially complete, only 0.6% of plasmids had undergone decatenation (data not shown), indicating that at least 99.4% of plasmids contained an ICL. Given that we observe between 3% and 8% regeneration of the Accl site, we conclude that our extracts support bona fide DNA replication-dependent repair of ICLs.

Persistence of an Adducted Parental Strand after Lesion Bypass

The question arises as to how the Accl site is regenerated since lesion bypass alone should yield an adducted intermediate,



Figure 4. Replication-Dependent Repair of an ICL

(A) An ICL blocks cleavage by AccI. Fifteen nanograms of pICL^{Nm} or pCtr was digested with AccI, separated on a native agarose gel, and stained with SYBR Gold. (B) ICL repair assay. At different times after addition of NPE/³²P- α -dATP, pICL^{Nm} or pCtr was recovered and equal aliquots were digested with HinclI (lanes 1–9) or AccI (lanes 10–18). (Note: lanes 1–9 represent a darker exposure of Figure 3B.) Twenty percent of the reaction was loaded in lanes 9 and 18. (C) At each time point shown in (B), the repair efficiency was calculated as explained in the text and graphed (red line). Extension products from the same ex-

(C) At each time point shown in (B), the repair efficiency was calculated as explained in the text and graphed (red line). Extension products from the same experiment shown in Figure 4B were plotted for comparison (gray dashed line).

(D) pICL^{Nm} or pCtr was replicated using NPE lacking radioactivity and optionally supplemented with p27^{Kip}. Plasmid was recovered, digested as indicated, and examined by Southern blotting using pCtr DNA as probe. Twenty percent of the reaction was loaded in lane 13. Samples were supplemented with a 1.2 kb HindIII fragment of pCtr before extraction (loading control).

which is not expected to be cleavable by Accl (Figure S10C, bottom plasmid). Although NER is not essential for higher eukaryotic cells to survive exposure to ICL agents, it has been proposed to remove the adduct after lesion bypass (see Introduction). To examine whether this occurs in our system (as depicted in Figures S10F and S10G), we used strand-specific Southern blotting to monitor the presence of an adduct in the parental strands, which would retard their mobility. To generate size markers for the unadducted strands, pCtr or pICLPt was replicated in the presence of ³²P-α-dATP. After 4 hr, the replicated DNA was digested with AfIII and Asel to cut out a small fragment encompassing the ICL (Figure 5A). As expected, DNA replication generated nascent products of 178 nt (top strand) and 176 nt (bottom strand) (Figure 5B, lanes 1 and 2). To examine parental strands, the same reaction was performed in the absence of ${}^{32}P-\alpha$ -dATP, and AfIIII/Asel digestion products were hybridized with a strand-specific probe complementary to the top strand. This analysis revealed a new band migrating roughly 1 nt above the unadducted top strand using pICLPt (Figure 5C, lane 3, Top-AD), but not with pCtr (Figure 5C, lane 4). Because this band was also not seen in the radioactively labeled nascent products (Figure 5B, lanes 1 and 2), it must correspond to a modified parental strand. Similarly, re-probing the stripped blot for the bottom strand showed a new band just above the unadducted bottom strand (Figure 5D, lane 3, Bottom-AD). We conclude that an adduct (AD) persisted in both parental strands. Since lesion bypass is expected to occur randomly using either the rightward or leftward fork (Figure 5A), 50% of each parental strand should be shifted if no adducts were removed, and this was in general agreement with our observations (Figures 5C and 5D, lane 3). The data suggest that most of the remaining adducts were resistant to repair, although we cannot rule out that



Figure 5. An Adduct Persists in the Parental Strand after Lesion Bypass

(A) Cartoon illustrating replication of an AfIIII/Asel restriction fragment harboring a cisplatin ICL. Due to the different overhangs generated by these enzymes, digestion of pICL^{Pt} yields top and bottom strands of 178 and 176 nt, respectively. Lesion bypass by the rightward fork yields a radioactively labeled nascent top strand and an adducted, parental bottom strand (Bottom-AD), while lesion bypass by the leftward fork results in a labeled nascent bottom strand and an adducted parental top strand (Top-AD). Strand-specific Southern blotting was used to detect either the top strands (blue lines) or the bottom strands (green lines). (B) Detection of nascent strands. pICL^{Pt} or pCtr was replicated in the presence (lanes 1 and 2) or absence (lanes 3 and 4) of $^{32}P-\alpha$ -dATP. After 4 hr, replication products were digested with AfIIII and Asel, separated on a 5% denaturing polyacrylamide gel and transferred to a Nylon membrane. Radioactive products were visualized using a phophorimager.

(C) Detection of the nascent top strand and the adducted parental top strand (Top-AD) on the membrane in (B) by Southern blotting using a bottom-strand probe. (D) Detection of the nascent bottom strand and the adducted parental bottom strand by stripping and reprobing the membrane in (C) using a top-strand probe. Primer S was used to generate a sequencing ladder from pCtr that serves as a size marker (see Figure 2). Green and blue arrowheads indicate the 176 nt and 178 nt sequencing products, respectively (see Figure S5C for sequence information and location of primer S). The migration of the digested DNA replication products is retarded by 1 nt with respect to the sequencing products (See Figure S9 for discussion of this effect). Adducted parental strands also persisted on pICL^{Nm} (data not shown).

a small fraction was removed by NER, giving rise to the fully repaired products detected in Figure 4. Alternatively, fully repaired products may be generated by HR (see Discussion). The small degree of gel retardation observed in Figure 5 suggests that only a single base or nucleotide, rather than an oligonucleotide, remained attached to the parental strand via the ICL. This is either because the original incisions occurred immediately adjacent to the ICL or because oligonucleotides generated during unhooking were trimmed by a nuclease (Hazrati et al., 2007).

Rev7-Depleted Extracts Are Defective for ICL Repair

Vertebrate cells deficient for Rev7 or Rev3, the two subunits of DNA pol ζ , are extremely sensitive to cisplatin, suggesting a defect in ICL repair. To determine whether DNA pol ζ participates in ICL repair in our cell-free system, we depleted Rev7 from egg extracts. Immunodepletion with Rev7-specific antibodies removed the vast majority of Rev7 and Rev3 from HSS and NPE (Figure 6A and data not shown). In the Mock-depleted and Rev7-depleted extracts, the leading strand of the rightward fork advanced to within 1 nt of the ICL in pICL^{Pt}, and this product then disappeared in both extracts with similar kinetics (Figure 6B, lower panel, -1 product). Strikingly, in the Rev7-depleted extract but not in the mock-depleted extract, a new species accumulated (the "0 product") that was exactly 1 nt longer than the -1 product (Figure 6B, lower

panel). The 0 product likely reflects translesion synthesis in which a nucleotide is inserted across from the adducted template G (Figure 6D). Build-up of the 0 product in the absence of Rev7 was observed in four independent experiments (Figure 6B and data not shown). Importantly, we also found that in Rev7-depleted extracts, regeneration of the ICL-proximal restriction site was reduced on average 3-fold relative to the mock-depleted control, demonstrating a defect in the final outcome of ICL repair (Figure 6C). The residual repair may be due to incomplete Rev7 depletion or redundant activities. The accumulation of the 0 product in Rev7-depleted extract agrees well with the proposed role for Pol ζ in extending leading strands beyond a DNA lesion (Washington et al., 2004). It is therefore unlikely that the repair defect observed in these extracts is caused by codepletion of unknown lesion bypass activities. Our results suggest a specific role for DNA pol ζ in ICL repair and argue that ICL repair in our cell-free system occurs via a physiological mechanism.

Collision of a DNA Replication Fork with an ICL Leads to Checkpoint Activation and Triggers the Fanconi Anemia Pathway

When mammalian cells are treated with agents such as mitomycin C that induce ICLs, FancD2 is ubiquitylated and the ATR checkpoint kinase is activated (Wang, 2007). We therefore



Figure 6. Repair of pICL Is Defective in Rev7-Depleted Extracts

(A) Rev7 immunodepletion. Undepleted, mock-depleted, and Rev7-depleted HSS and NPE were analyzed by western blotting using Rev7 antibody. A relative volume of 100 corresponds to 0.3 µl extract.

(B) Accumulation of a new lesion bypass intermediate in Rev7-depleted extracts. pICL^{Pt} was replicated in mock-depleted or Rev7-depleted HSS and NPE (4 ng/µl final DNA concentration). At the indicated times, products were digested with AfIIII and analyzed on a sequencing gel (as in Figure S7B). Numbers to the left indicate the size of the sequencing products. The new replication intermediate is indicated (0 product).

(C) The average repair efficiency in mock- and Rev7-depleted extracts in four independent experiments is plotted with error bars.

(D) Cartoon depicting the intermediate that accumulates in Rev7-depleted extracts. We infer that a C residue is inserted at position 145 since the translesion step is likely performed by the cytidyl transferase Rev1, and some of the products are digestible with Sapl.

examined whether these damage-response pathways are also activated upon replication of purified ICL-containing plasmids. As shown in Figure S11A, replication of pICL^{Pt} caused a massive induction of Chk1 phosphorylation at serine 344 (Chk1-P; lower panel, lanes 2-6) that was comparable to the level seen in the presence of aphidicolin, which induces helicase uncoupling (lanes 12-13) (Byun et al., 2005). Replication of undamaged plasmid did not induce Chk1 phosphorylation. pICL^{Pt}-induced Chk1 phosphorylation was completely dependent on replication initiation (Figure S11B, lower panel, lanes 1 and 2). The data indicate that collision of a DNA replication fork with an interstrand crosslink activates the ATR signaling pathway. Replication of pICLPt also converted FancD2 to a slower migrating form (Figure S11A, top panel, lanes 2-6), which represents an ubiquitylated species (data not shown). Like ICL repair, FancD2 ubiquitylation was critically dependent on DNA replication (Figure S11B). The replication-dependent activation of the Fanconi anemia and ATR pathways by pICL further suggests that repair of these DNA templates in Xenopus egg extracts involves a physiological mechanism.

DISCUSSION

Using Xenopus egg extracts and model DNA templates containing a single, site-specific ICL, we have established an in vitro sysuse cellular sensitivity to ICL agents to make inferences about the process of ICL repair, our approach directly examines the process of repair using a DNA template that contains no DNA lesions other than ICLs. Using this system, we show that ICL repair is directly coupled to DNA replication. Examination of repair intermediates suggests the following mechanism of ICL repair (Figure 7). Initially, two DNA replication forks converge on the ICL, generating a structure in which the leading strands of each fork stall \sim 20–40 nt from the ICL, and the 5' ends of the lagging strands are located at a greater and more variable distance from the lesion (Figure 7B). After an \sim 20 min delay, lesion bypass is initiated when the leading strand of one fork is extended to within a few nucleotides of the ICL (Figure 7C). After a further \sim 30 min, the two sister chromatids that are joined via the ICL are uncoupled, likely via dual incisions surrounding the ICL (Figure 7D). Concurrent with sister chromatid uncoupling, a nucleotide is incorporated across from the adducted base that formed part of the ICL (Figure 7E), an event termed translesion synthesis. At present, we cannot determine whether plasmid uncoupling or translesion synthesis occurs first. Soon after incision/translesion synthesis, the leading strand is extended beyond the ICL and ligated to the first downstream Okazaki fragment (Figure 7F). Immunodepletion of Rev7 suggests that this extension step is dependent on DNA polymerase ζ. Finally, fully

tem that supports ICL repair in S phase. Unlike many studies that





When DNA containing an ICL (A) undergoes DNA replication, the leading strands of two converging replication forks initially stall 20–40 nt from the lesion (B). One leading strand (indicated in red) is then extended to within 1 nt of the ICL, a step which may require prior replisome remodeling (C). Subsequently, the two sister chromatids are uncoupled via dual incisions (yellow scissors) on either side of the ICL, possibly by XPF and/or Mus81 (D). Next, a translesion DNA polymerase (possibly Rev1) inserts a nucleotide across from the adducted base (E), after which DNA polymerase ζ extends the nascent strand beyond the ICL (F). Finally, two fully repaired DNA duplexes are generated through the action of nucleotide excision repair (NER) on the top duplex and homologous recombination (HR) on the bottom duplex (G).

repaired daughter duplexes are formed, as evidenced by Accl digestion of the DNA sequence underlying the ICL. The Accl site is likely regenerated by HR and/or NER (Figure 7G; see also Figure S10), although an involvement of base excision repair cannot be ruled out.

The question arises why more of the input DNA was not repaired since lesion bypass is highly efficient. One likely explanation is that most molecules retain an adduct in the parental strand that precludes digestion by Accl (Figure 5). This observation suggests that NER is not highly active in our system. The remaining adduct is unlikely to be lethal since it will not disrupt chromosome segregation. Another potential explanation is that the repair is error prone, although we disfavor this idea, as explained below. Finally, recombinational repair may be inefficient or aberrant. Indeed, although we observe X-shaped molecules during 2D gel electrophoresis that are consistent with Holliday junction intermediates (data not shown) and 5' to 3' resection (Figure 2B), the latter process may be so rapid as to destroy many daughter molecules before they can engage in HR (Figure S10F, top plasmid).

The mechanism of ICL repair characterized here differs in important ways from current models for this process, which have been inferred largely from genetic data and the biochemical properties of relevant DNA repair enzymes (Dronkert and Kanaar, 2001; Niedernhofer et al., 2004, 2005; Niedzwiedz et al., 2004; Wang, 2007). Thus, prevailing models of ICL repair envision that an incised parental strand is extended past the DNA lesion (Figure S1D), whereas our results clearly show that this reaction involves one of the two nascent strands (Figure 7). This observation reveals how ICL repair is coupled to DNA replication. A potential benefit of using a nascent strand is that replicative polymerases, and possibly translesion DNA polymerases, need not be recruited de novo. Moreover, in current models, lesion bypass can begin only after incision has occurred since the latter process generates a 3' end within the parental strand that will be extended past the lesion. In contrast, by extending a nascent strand, lesion bypass can begin before incisions. Indeed, our data indicate that after the initial pause at -20, the leading strand advances to within a single nucleotide of the ICL before any incisions occur. We inferred the absence of incisions at early time points from the persistence of the X-shaped molecule generated by Hincll digestion (Figure 3B) since any incisions near the ICL should cause this structure to collapse (Figure 3A). A potential exception is an incision that occurs close to the ICL, internal to a significant region of duplex DNA, since in this scenario the arm fragment may not be released (Figure S12A). However, such an incision would be revealed as soon as the replication fork advances to the incision (Figure S12B). We therefore believe that incisions are not made until after the leading strand moves to the -1 position. A beneficial consequence of initiating lesion bypass before incisions have occurred is that repair of one sister chromatid is already underway at the moment of incision. This reduces the lag period between the time that dsDNA breaks are generated and when a fully duplex sister chromatid is available for homology-directed repair of the break.

A second difference is that while current models of ICL repair propose collision of a single replication fork with the lesion, we observe convergence of two forks on the ICL. Based on the following considerations, the latter situation may occur frequently, even in somatic cells. First, since initiations at neighboring origins generally take place concurrently (Berezney et al., 2000), two converging forks will arrive at an ICL contemporaneously if the lesion is located near the midpoint between two origins. Second, if the ICL is located immediately adjacent to one origin, one of the two converging forks will reach the ICL at most 40 min before the other (assuming an average interorigin

distance of ~50 kb and a fork speed of ~1.25 kb/min; Berezney et al., 2000; Ge et al., 2007). Importantly, repair is quite slow, taking several hours, both in vitro (this paper) and in mammalian cells (Akkari et al., 2000). Moreover, it has been proposed that when one replication fork stalls, new replication forks are rapidly established from nearby, surplus pre-RCs (Ge et al., 2007). Therefore, even when an ICL is located adjacent to an origin, two DNA replication forks may still arrive at the lesion almost contemporaneously. A potential advantage of the dual fork collision mechanism is that there is no need to re-establish a processive replication fork after lesion bypass is complete. This consideration is particularly relevant given the absence of a known replication restart pathway in higher eukaryotes. In some instances, only one fork can collide with the ICL, for example when two such lesions occur between neighboring pre-RCs, but this situation should be rare. In summary, it seems safe to conclude that cells must be able to cope with single and dual collisions of replication forks with ICLs, and future work will be required to determine whether the mechanisms of repair differ in the two situations.

Our results raise important questions about the mechanism of lesion bypass. First, what happens during the 20 min between the arrival of the leading strand at position –20 and its further extension toward the ICL? It is possible that parts of the replication fork must be disassembled before the leading strand can be extended. For example, we speculate that the replicative DNA helicase, the MCM2-7 complex (Takahashi et al., 2005), must be removed to allow the leading strand to advance to the ICL. Notably, many nascent products were observed between the initial pause site and the site of the ICL (Figures 2C and S7C), suggesting that the leading strand was extended in a nonprocessive fashion. These results indicate that the mode of DNA synthesis changes substantially after the initial pause, possibly due to a reliance on strand-displacement synthesis.

On pICL^{Nm}, the leading strand pauses again at the -4 to -1positions (Figure 2C). We propose that the pauses seen at the -4 to -2 positions result from the difficulty of unwinding DNA around the ICL. The pause at -1, which is also observed on pICL^{Pt}, probably reflects the time required to recruit a translesion DNA polymerase, a step that may involve PCNA ubiquitylation (Lehmann et al., 2007). Based on genetic experiments, the dCMP transferase Rev1 is a prime candidate for this polymerase (Niedzwiedz et al., 2004; Simpson and Sale, 2003). Since the nitrogen mustard-like and cisplatin ICLs are formed between G residues, the involvement of Rev1 suggests that the translesion step itself should be largely error free. Indeed, we infer that some products must be error free given that they can be digested with restriction enzymes. Genetic experiments suggest that the Rev1 product is subsequently extended by DNA polymerase ζ, a heterodimer of Rev3 and Rev7 (Lehmann et al., 2007). Our results provide strong support for this idea since we see accumulation of the translesion product when pICLPt is replicated in Rev7-depleted extracts. Importantly, there is generally a gap of several hundred nucleotides between the ICL and the 5' end of the first lagging strand product located downstream. Since DNA polymerase ζ is error prone, we speculate that there might be a switch back to a replicative DNA polymerase to avoid high mutation rates. Intriguingly, we observed no accumulation of the translesion product when pICL^{Nm} was replicated in Rev7-depleted extracts (data not shown). This observation suggests that a nitrogen mustard-like ICL can be handled by other translesion DNA polymerases. It also shows that removing DNA polymerase ζ from our extracts results in highly specific defects.

We have shown that replication of pICL in Xenopus egg extracts leads to activation of the ATR checkpoint pathway. This observation is interesting because replication-dependent ATR activation is normally thought to occur as a result of helicase uncoupling (Byun et al., 2005), which should not be possible in the presence of an ICL. We speculate that the checkpoint is launched on single-stranded DNA that is exposed on the lagging strand template after a fork collides with the ICL. We also found that pICL replication promotes FancD2 ubiguitylation, suggesting that the FA pathway participates in ICL repair. The recently reported replication-independent ubiquitylation of FancD2 by undamaged plasmids in egg extracts likely resulted from the use of very high DNA concentrations (Sobeck et al., 2007). The cell-free system we have developed is ideally suited to elucidate the roles of these DNA damage-response pathways, as well as known repair enzymes, in the process of interstrand crosslink repair.

EXPERIMENTAL PROCEDURES

Preparation of pICL

A cisplatin ICL duplex was prepared (Hofr and Brabec, 2001), purified on a Mono-Q column, and ligated into pSVRLuc (See Figure S2). The analysis of the crosslink-containing substrate (restriction digest, real-time PCR) confirmed the presence of ICL in \geq 99% of the plasmids. For pICL^{Nm}, oligos containing a single modified guanosine base, 7-deaza-7-(2,3-dihydroxy-pro-pyl)-guanine (G*), were annealed and processed to generate the ICL depicted in Figure 1A (T.A. and O.D.S., unpublished data). The crosslinked DNA duplex was ligated into pSVRLuc. About 99% of pICL^{Nm} contained an ICL, as judged by AccI restriction digest and quantitative PCR. Both the nitrogen mustard-like and cisplatin ICLs are stable under the neutral reaction conditions employed in this study.

Electron Microscopy

DNA was purified from replication reactions and EM was performed using the cytochrome c drop spreading method (Thresher and Griffith, 1992). The contour length of all DNA molecules was determined using image J software and converted to basepairs using the 200 nm scale bar in each micrograph, assuming 3.4 A° /basepair.

Xenopus Egg Extracts and Replication

DNA replication and preparation of *Xenopus* egg extracts (NPE and HSS) were as described (Walter et al., 1998). Briefly, plasmid was incubated with HSS for 20 min, followed by addition of two volumes of NPE containing ${}^{32}P-\alpha$ -dATP (0.25 μ Ci/µl, 3000 Ci/mmol final). Unless otherwise indicated, the final plasmid concentration in the reaction was 1.8 ng/µl. To measure DNA replication efficiency (Figures 1C and 1D), 2 µl aliquots of the reaction were analyzed on native agarose gels (Takahashi et al., 2004). For other applications, 10 µl aliquots of replication reactions were stopped with 90 µl Stop Solution A (1% SDS, 1 mM EDTA, 100 mM Tris, pH 8), treated with RNase A followed by 0.5 μ g/µl Proteinase K at 37°C, followed by phenol/chloroform extraction. DNA was ethanol precipitated in the presence of glycogen (30 μ g/µl final) and resuspended in 5–10 µl of 10 mM Tris buffer (pH 7.5).

Nascent Strand Analysis

Extracted replication products were digested with the indicated enzymes followed by addition of 0.5 volumes of Stop Solution B (95% formamide, 20 mM EDTA, 0.05% bromophenol blue, 0.05% xylene cyanol FF). Restriction fragments (3 μ l) were separated on 42 cm long, 5% polyacrylamide sequencing gels prepared with Rapidgel-XL in 0.8X GTG Buffer (USB Corporation, Cleveland, OH, USA). Gels were transferred to filter paper, dried, and visualized with a phosphorimager. Sequencing ladders were generated using the Cycle Sequencing kit from USB.

Antibodies

FancD2 antiserum was prepared against residues 1–172 of *Xenopus* FancD2 and its specificity confirmed using western blotting. Antibody against full-length *Xenopus* Rev7 protein was prepared using the same procedure. Rev7 was removed from HSS and NPE using three rounds of depletion with the Rev7 antiserum.

SUPPLEMENTAL DATA

Supplemental Data include Supplemental Experimental Procedures and twelve figures and can be found with this article online at http://www.cell. com/cgi/content/full/134/6/969/DC1/.

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