DNA damage tolerance permits bypass of DNA lesions encountered during S-phase and may be carried out by translational modification (PTM). Recent studies revealed that replication protein A (RPA), the major ssDNA-binding protein, is involved in the regulation of PCNA monoubiquitination and interacts directly with Rad18 on chromatin and in the nucleoplasm. However, it is unclear how RPA regulates this critical PTM and what functional role(s) these interactions serve. Here, we developed an in vitro assay to quantitatively monitor PCNA monoubiquitination under in vitro scenarios. Results from extensive experiments revealed that RPA regulates Rad6/Rad18 activity in an ssDNA-dependent manner. We found that “DNA-free” RPA inhibits monoubiquitination of free PCNA by directly interacting with Rad18. This interaction is promoted under native conditions when there is an overabundance of free RPA in the nucleoplasm where Rad6/Rad18 and a significant fraction of PCNA reside. During DNA replication stress, RPA binds the ssDNA exposed downstream of stalled primer/template (P/T) junctions, releasing Rad6/Rad18. Rad6 restricted the resident PCNA to the upstream duplex regions by physically blocking diffusion of ssDNA along dsDNA, and this activity was required for efficient monoubiquitination of PCNA on DNA. Furthermore, upon binding ssDNA, RPA underwent a conformational change that increased its affinity for Rad18. Rad6/Rad18 complexed with ssDNA-bound RPA was active, and this interaction may selectively promote monoubiquitination of PCNA on long RPA-coated ssDNA.

In eukaryotes, the replicative DNA polymerases (pols) anchor to PCNA sliding clamps encircling DNA to achieve the high degree of processivity required for efficient DNA replication. Under native conditions, the replicative pols are rate-limited by unwinding of double-stranded DNA (dsDNA) such that template bases are replicated as soon as they are available. This tight coupling limits exposure of ssDNA. However, the stringent replicative pols cannot accommodate distortions to the native DNA sequence. Prominent examples of these are modifications (lesions) to the native template bases from exposure to reactive metabolites and environmental mutagens such as UV radiation. Consequently, DNA synthesis on the afflicted template abruptly stops upon encountering these lesions but unwinding of dsDNA continues. These “uncoupling” events expose long stretches of the damaged template that are immediately coated by RPA, a heterotrimeric complex comprised of RPA70, RPA32, and RPA14 subunits. RPA is the major eukaryotic ssDNA-binding complex and protects exposed ssDNA from degradation and prevents formation of alternative DNA structures that are refractory to DNA synthesis. dsDNA unwinding eventually stalls after an uncoupling event and failure to re-couple DNA synthesis and dsDNA unwinding often results in dsDNA breaks that may lead to gross chromosomal rearrangements, cell-cycle arrest, and cell death. These replicative arrests may be overcome by TLS where specialized TLS pols bind to the resident PCNA and replicate the damaged DNA, allowing DNA synthesis by a replicative pol to resume. In humans, TLS requires monoubiquitination of PCNA encircling stalled P/T junctions and this critical PTM is catalyzed by Rad6/Rad18.

Rad6 is an E2 ubiquitin-conjugating enzyme that catalyzes covalent attachment of ubiquitin to a lysine residue within a target protein. On its own, Rad6 does not bind PCNA and is incapable of modifying PCNA. Furthermore, Rad6 alone can monoubiquitinate other Rad6 molecules and form mixed ubiquitin chains. The latter requires the noncovalent ubiquitin-binding site on Rad6. Selection of a target protein for Rad6 is dictated by an E3 ubiquitin ligase, such as Rad18, that simultaneously binds Rad6 and a target protein. Rad18 only functions as a homodimer and dimerization occurs through an N-terminal RING domain. A Rad18 homodimer interacts with a Rad6 through two independent binding sites. A distinct region of a Rad18 RING domain recognizes Rad6 through a canonical interface that is conserved among all E2/E3 RING complexes. The C-terminal Rad6-binding (R6B) domain of a Rad18 binds the noncovalent ubiquitin-binding site of Rad6, preventing mixed ubiquitin chain formation. In addition to the RING domain, the N-terminal half of Rad18...
contains multiple independent binding domains, one of which directly interacts with PCNA. Thus, Rad18 activates and targets Rad6 toward PCNA and only permits monoubiquitination (4).

Recent cellular studies revealed that RPA is involved in the regulation of PCNA monoubiquitination by Rad6/Rad18 and interacts with Rad18 on chromatin and in the nucleoplasm (i.e. nonchromatin associated/soluble fraction). Interestingly, the latter are quite prominent in nonperturbed human cells (8–10). However, it is unclear how RPA regulates this critical PTM and what functional role(s) these interactions serve. To address this, we developed an *in vitro* fluorescent assay to quantitatively monitor the effect of RPA on PCNA monoubiquitination under conditions that mimic *in vivo* scenarios.

## Results

### Rad6/Rad18 monoubiquititates free PCNA

Rad6/Rad18 is active throughout the cell cycle and does not require PCNA to encircle DNA for monoubiquitination of PCNA to occur (4, 6, 7, 11–18). Interestingly, PCNA protein levels are overabundant and essentially maintained throughout the cell cycle such that at least 30% of the PCNA pool resides in the nucleoplasm (i.e. free in solution) at any time. Particularly, free PCNA accounts for more than 80% of the pool throughout G1 and G2/M, which collectively last ~16 h in rapidly proliferating human cells (19–23). Accordingly, we initially focused on monoubiquitination of free PCNA in solution over a biologically relevant time scale that approaches G1 phase (~11 h) of rapidly proliferating human cells (19). To comprehensively and quantitatively study PCNA monoubiquitination, we utilized fluorescence to monitor the attachment of ubiquitin to all proteins present. Human ubiquitin is devoid of cysteine and residue Glu-24 (Fig. S2A) is far removed from both the “hydrophobic patch” where ubiquitin-binding domains dock and the C-terminal glycine residue where target proteins are attached (24). Residue Glu-24 was mutated to cysteine (E24C) and labeled with fluorescein (Fig. S2B). All lysine residues of ubiquitin remain available for conjugation into polyubiquitin chains (7).

The attachment of ubiquitin to proteins requires Rad6/Rad18 (Fig. 1A, compare lanes 1–5 to lanes 8 and 9) and increases over time such that 2690 nM ± 61.7 nM ubiquitin is conjugated to proteins at 8 h (Fig. 1B). At this concentration, ~90% of the total ubiquitin is conjugated to proteins and the fluorescence signal remains constant (Fig. S2C). Thus, attachment of fluorescein-labeled ubiquitin to proteins does not affect the fluorescence output of the dye, permitting quantitative analysis. The ubiquitinated proteins range in size from ~20–150 kDa (Fig. 1A). In human cells, Rad18 is monoubiquitinated by Rad6 at four conserved lysine residues (Fig. S1) (25). This process has previously been reconstituted *in vitro* with recombinant human proteins and is indicated by the appearance of bands greater than 75 kDa in size (6, 7). As expected, these bands are only observed in the presence of Rad6/Rad18 (Fig. 1A, compare lanes 1–5 to lanes 8 and 9) but are independent of PCNA (Fig. 1A, compare lanes 1–5 to lanes 6 and 7; Fig. S3A, compare lanes 1 and 2 to lanes 3 and 4). Western blotting confirmed the identity of these bands as ubiquitinated Rad18 (Fig. S3A, lanes 13–16). Rad18 is clearly the predominant ubiquitinated species and increases in fluorescence intensity over time (Fig. 1A, lanes 1–5). 2610 ± 63.6 nM of ubiquitin is conjugated to Rad18 over the incubation (Fig. 1B), indicating that each Rad18 contains approximately two ubiquitin moieties on average.

A distinct band appears between 20–25 kDa in size and increases in fluorescence intensity over time (Fig. 1A, lanes 1–5). This band is only observed in the presence of Rad6/Rad18 (Fig. 1A, compare lanes 1–5 to lanes 8 and 9) and is independent of PCNA (Fig. 1A, compare lanes 1–5 to lanes 6 and 7; Fig. S3A, compare lanes 1 and 2 to lanes 3 and 4). The molecular weight of Rad6 increases from ~15 kDa to ~24 kDa by the attachment of ubiquitin and Rad6 will monoubiquitinate itself in the absence of functional interactions with Rad18 (7). Western blotting confirmed the identity of this band as monoubiquitinated Rad6 (Fig. S3A, lanes 9–12). Only ~17.7 ± 1.62 nM of ubiquitin is conjugated to Rad6 at 8 h (Fig. 1B). Thus, monoubiquitinated Rad6 is minimal, accounting for less than 3% of the total Rad6 present.

In the presence of PCNA and Rad6/Rad18 (Fig. 1A, lanes 1–5), a distinct band appears between 40 and 50 kDa and increases in intensity over time. This band disappears when PCNA is removed (Fig. 1A, compare lanes 1–5 to lanes 6 and 7; Fig. S3A, compare lanes 1 and 2 to lanes 3 and 4). The molecular weight of a PCNA monomer increases from ~36 kDa to ~45 kDa by the attachment of ubiquitin (Fig. S3C) and Rad6/Rad18 will monoubiquitinate PCNA in solution (6, 7, 17, 18). Western blotting confirmed the identity of this band as monoubiquitinated PCNA (Fig. S3A, lanes 5–8). 71.1 ± 6.79 nM ubiquitin is conjugated to PCNA monomers over the incubation (Fig. 1B). At this concentration, 23.6 ± 2.25% of PCNA monomers are monoubiquitinated (Fig. 1C), which equates to 70.7 ± 6.76% of all PCNA homotrimers containing a single ubiquitin, on average. Altogether, these results indicate that 1) ubiquitinated Rad18 (which accounts for all Rad18 present) binds to and is functional with Rad6, 2) Rad6 and Rad18 maintain active/ productive interactions throughout the incubation, and 3) Rad6/Rad18 monoubiquitinates free PCNA and this activity is prominent over a biologically relevant time scale.

### RPA regulates monoubiquitination of free PCNA in a ssDNA-dependent manner

Rad6/Rad18 complexes are prominent in nonperturbed human cells and predominantly observed in the soluble fraction (i.e. nonchromatin associated) (8–10). Next, we repeated the assays described in Fig. 1 in the presence of free RPA. At the highest RPA concentration (6 μM), additional ubiquitinated species do not appear in fluorescence scans (Figs. S4 and S5) and ubiquitination of RPA subunits is not observed by Western blotting (Fig. S5). Thus, Rad6/Rad18 does not conjugate ubiquitin to free RPA and, hence, any observed effects are not attributable to RPA ubiquitination. Monoubiquitination of proteins (total) is slightly reduced over the range of RPA concentrations (Fig. 2) and this is primarily because of the minor reduction in Rad18 monoubiquitination. However, RPA dramatically stimulates monoubiquitination of Rad6 and such behavior is independent of PCNA (Fig. S4). Monoubiquitination of free PCNA is significantly reduced over the range of RPA concentrations,
and identical results are observed by Western blotting (Fig. S4). Thus, RPA directly inhibits monoubiquitination of free PCNA by Rad6/Rad18. It should be noted that RPA was limited to /H11349/6/H9262 to maintain physiological ionic strength. At this concentration, RPA is only present at a 5-fold excess over Rad18. RPA is the most abundant ssDNA-binding protein in human cells and this high concentration is maintained throughout the cell cycle such that RPA is at least 100-fold in excess of Rad18 in unper- turbed human cells (26, 27). RPA:Rad18 ratios that approach physiological conditions are likely to further inhibit monoubiquitination of free PCNA.

Rad6/Rad18 activity is imperative during DNA damage tolerance when RPA coats persistent ssDNA regions exposed by uncoupling events at replication-blocking lesions (2). Hence, the observed inhibition of Rad6/Rad18 activity by free RPA must be relieved in the presence of ssDNA. To test this, we monitored the effect of ssDNA on the RPA-dependent inhibition of free PCNA monoubiquitination. We repeated the assays described above except RPA was first pre-incubated with a 33-mer poly(dT) ssDNA template (poly(dT)$_{33}$) (Fig. S6). At physiological ionic strength, human RPA binds ssDNA with extremely high affinity ($K_D$ of fM to pM), low cooperativity, and an occluded binding site size of 30 nt (28, 29). Furthermore, nearly all ssDNA (24 ± 1 nt) occluded by a single RPA directly interacts with the protein (28–30). In contrast, human Rad18 has relatively weak affinity ($K_D$ of nM to $\mu$m) for purely ssDNA at minimal ionic strength (∼5 mM) and requires at least 49 nucleotides for binding to be observed (17). Thus, occupation of the poly(dT)$_{33}$ ssDNA by Rad6/Rad18 will be minimal, if at all. Indeed, binding of RPA to poly(dT)$_{33}$ is stoichiometric at phys-
RPA regulates monoubiquitination of PCNA on and off DNA

Four oligonucleotide-binding (OB) folds mediate binding of an RPA to ssDNA. OB-A and OB-B within the RPA70 subunit are the primary ssDNA-binding sites and together comprise an occluded ssDNA-binding site of 8 nt. Interestingly, these OB folds also comprise one of the two independent Rad18-binding sites (8, 27, 32). OB-C and OB-D reside in the RPA70 and RPA32 subunits, respectively, and extend the occluded ssDNA-binding site to 30 ± 2 nt (27–29, 32). The RPA32 subunit also contains a winged helix (WH) domain that is largely responsible for the interaction of RPA with cellular proteins, and the accessibility of this domain is enhanced upon ssDNA binding (33, 34). The second independent Rad18-binding site within RPA resides in RPA32 but has yet to be mapped (8, 10). The results from Fig. 2 indicate that RPA directly mediates Rad6/Rad18 activity toward free PCNA in a ssDNA-dependent manner. To gain insight into how this occurs we analyzed the interaction of RPA and Rad6/Rad18 as well as the composition of Rad6/Rad18 by analytical gel filtration.

Rad6/Rad18 alone peaks at 21.6 min retention time (Fig. 3A), and the appearance and disappearance of Rad18 and Rad6 are intimately correlated, indicating a stable binding interaction between these proteins. RPA alone peaks at 25.1 min retention time (Fig. 3B). When RPA and Rad6/Rad18 are pre-incubated, the Rad6/Rad18 peak undergoes a small shift to a higher molecular weight and a small increase in intensity (Fig. 3C, top panel). Furthermore, the RPA70 and RPA32 subunits elute in earlier fractions (Fig. 3C, bottom panel). Together, this indicates that RPA interacts directly, albeit weakly, with Rad18 at physiological ionic strength in the absence of any nucleic acid, in agreement with previous in vivo and in vitro studies (8). In the absence of ssDNA, both Rad18-binding sites within RPA are potentially available and, hence, it cannot be deciphered if one or both contribute to the observed interaction.

The tight correlation of the Rad18 and Rad6 elution profiles (Fig. 3A, bottom panel) is unaffected by RPA (Fig. 3C, bottom panel), and RPA does not decrease the amount of Rad6 that co-elutes with Rad18 (Fig. 3F), indicating that RPA does not preclude Rad18 from binding Rad6. Furthermore, mixed ubiquitin chain formation by Rad6 is not observed under any experimental condition described above. This activity requires noncovalent binding of ubiquitin to Rad6 and is prohibited by the Rad18 R6B, which effectively outcompetes ubiquitin for binding to Rad6 (Fig. S9) (7). Collectively, these results indicate that the R6B domain of Rad18 remains bound to the noncovalent ubiquitin-binding site of Rad6 throughout all incubations. Next, we repeated these assays in the presence of ssDNA.

Binding of Rad6/Rad18 to ssDNA does not occur when the concentrations of poly(dT)_{13} and RPA are stoichiometric (Fig. S7 and Table S1). Rather, all RPA and ssDNA reside in a complex (Fig. 3D, top panel; Fig. S7 and Table S1) in which RPA occupies the entire length of the ssDNA (28, 29). Thus, any retention of Rad6/Rad18 by the RPA/ssDNA complex is through a direct interaction between RPA and Rad18. The Rad6/Rad18 peak (Fig. 3A, top panel) undergoes a small shift to a higher molecular weight and a significant increase in intensity (Fig. 3E, top panel). Furthermore, greater amounts of Rad18 are observed in the early fractions with RPA and ssDNA (Fig. 3E, bottom panel) compared with RPA alone (Fig. 3C, bottom panel). Finally, greater amounts of the RPA70 and RPA32 subunits are observed in the early fractions with Rad6/Rad18 and ssDNA (Fig. 3E, bottom panel) compared with Rad6/Rad18 alone (Fig. 3C, bottom panel) or ssDNA alone (Fig. 3D, bottom panel). This indicates that ssDNA-bound RPA directly interacts with Rad6/Rad18 at physiological ionic strength and ssDNA enhances the affinity of RPA for Rad6/Rad18, confirm-
Previous studies revealed that RPA is required for monoubiquitination of PCNA at ssDNA regions generated by uncoupling events during S-phase (8, 9). Similarly, Rad6/Rad18-catalyzed monoubiquitination of PCNA occurs outside of S-phase.
Western blotting confirmed complete monoubiquitination of PCNA (Fig. S10). In contrast, monoubiquitination of ~20% of PCNA monomers requires at least 8 h when PCNA is free in solution, indicating that stabilization of loaded PCNA on P/T DNA enhances monoubiquitination of PCNA by Rad6/Rad18 at least 8-fold. To gain further insight, we repeated these assays by selectively omitting individual components (Fig. 4C).

When RFC is omitted, PCNA is not assembled onto the DNA substrate (Fig. 4A, Condition 2) and only 10% of PCNA monomers are monoubiquitinated at 8 h incubation (Fig. 4C, Condition 3). This level of PCNA monoubiquitination is observed in 30–45 min when RFC is included (Fig. 4B), suggesting that loading of PCNA onto P/T DNA by RFC significantly enhances monoubiquitination of PCNA by Rad6/Rad18. In the absence of either NeutrAvidin or biotin, PCNA is loaded onto the DNA substrate but immediately diffuses off the duplex end upon release from RFC (Fig. 4A, Conditions 4 and 5) (35). Under these conditions, only 33–43% of PCNA monomers are monoubiquitinated at 8 h incubation (Fig. 4C, Conditions 5 and 6). This level of PCNA monoubiquitination is observed in ~1.25 h when the NeutrAvidin/biotin block is intact (Fig. 4B). Thus, loading and stabilization of PCNA on P/T DNA are both required for the observed enhancement of PCNA monoubiquitination on DNA (Fig. 4, B and C). When RPA is omitted, PCNA is loaded onto the DNA substrate but immediately dif-
RPA regulates monoubiquitination of PCNA on and off DNA

**Figure 5.** The stability of PCNA encircling a stalled P/T junction drives PCNA monoubiquitination. A, retention and orientation of PCNA on DNA. Top, schematic representation of the FRET experiments. Cy5-PCNA was assembled on the Cy3P/BioT70 DNA substrate (Fig. 5A) in the presence of SSB, and FRET was monitored at equilibrium as in Figs. 4A, 4B, and 4C, Bottom, FRET in the presence of SSB. FRET is only observed for the Cy3P/BioT70 DNA substrate when SSB is included (Condition 2); excess Rad6/Rad18 will not compensate (Condition 3). Identical results are observed for RPA with the Cy3P/BioT33 DNA substrate (Conditions 4–6). As observed in a previous report, the FRET measured in the presence of SSB (Condition 2, 0.540 ± 0.0327) and RPA (Condition 5, 0.502 ± 0.0327) are within experimental error, indicating that the same amount of PCNA is loaded onto and stabilized at a P/T junction in the same FRET state (i.e. orientation) when the adjacent ssDNA is bound by either RPA or SSB. B, stability of PCNA encircling DNA. Top, schematic representation of the experiment. Cy5-PCNA is pre-assembled onto unlabeled P/BioT70 DNA in the presence of SSB and this solution is rapidly mixed in a stopped-flow instrument with a solution containing Cy3P/BioT70 DNA pre-bound by SSB, and FRET is monitored. Under these conditions, all Cy5-PCNA is pre-loaded onto the unlabeled P/BioT70 DNA substrate and stabilized by SSB prior to mixing, and the only pathway for dissociation of Cy5-PCNA into solution is through spontaneous opening of PCNA. Hence, RFC-catalyzed loading of Cy5-PCNA onto Cy3P/BioT70 DNA is rate-limited by spontaneous opening of PCNA. The loading trace was fit to a single-exponential and the rate constant is reported. Very similar values are obtained with RPA on the P/BioT33 P/T DNA substrates, indicating that the stability of PCNA encircling a P/T junction is the same when the adjacent ssDNA is bound by either RPA or SSB. C, monoubiquitination of loaded PCNA in the presence of a ssDNA-binding protein. Top, schematic representation of the experiment. PCNA was pre-assembled onto a P/BioT33 P/T DNA substrate in the presence of a stoichiometric amount of a ssDNA-binding protein and then monoubiquitination of target proteins was monitored as described in Figs. 1 and 2. Bottom, extent of PCNA monoubiquitination. The percentage of PCNA monomers that are monoubiquitinated is plotted as a function of time. Symbols are indicated in the figure legends and the data for each represent the average ± S.D. of three independent experiments.

Fuses off the ssDNA end upon release from RFC (Fig. 4A, Condition 3) (35) and excess Rad6/Rad18 does not compensate (Fig. 4A, Condition 6). Under these conditions, 63% of PCNA monomers are monoubiquitinated at 8 h incubation (Fig. 4C, Condition 4), similar to that observed when either NeutrAvidin or biotin are omitted. Altogether, this confirms that RPA significantly enhances Rad6/Rad18-catalyzed monoubiquitination of PCNA encircling a P/T junction by physically blocking diffusion of the sliding clamp along the adjacent ssDNA (36). SSB is a homotetramer that binds ssDNA with an occluded site size of 65 nt at 200 mM ionic strength where the ssDNA is fully wrapped around SSB, contacting each subunit within the heterotetramer. This differential mode of ssDNA-binding has no effect on the amount of PCNA loaded onto and maintained at P/T junctions (36), the orientation of PCNA encircling P/T junctions (Fig. 5A) (36), or the intrinsic stability of PCNA encircling P/T junctions (Fig. 5B). However, unlike RPA, SSB does not interact with Rad6/Rad18 (8, 10). These unique behaviors were exploited to decipher any effects of ssDNA-RPA-Rad6/Rad18 interactions (Fig. 3E) on monoubiquitination of PCNA encircling a P/T junction (Fig. 4E).

First, PCNA was pre-assembled on a P/T DNA substrate in the presence of a stoichiometric amount of a ssDNA-binding protein; SSB for P/BioT70 and RPA for P/BioT33 (Fig. 5C, top). Monoubiquitination of PCNA was then monitored over time as described above. Under these conditions, a single ssDNA-binding protein is adjacent to a P/T junction that is encircled by PCNA. On each DNA substrate, the same amount of PCNA is loaded onto the P/T junction and stabilized in the same orientation. This results in a decrease in monoubiquitination as previously suggested (8, 10). To directly test this, we utilized the *Escherichia coli* ssDNA-binding protein, SSB.

SSB is a functional homolog of human RPA and binds ssDNA noncooperatively and with very tight affinity (in pM range) at 200 mM ionic strength (28, 29, 37–39). We previously demonstrated that, like RPA, SSB binds tightly to ssDNA adjacent to a P/T junction and restricts PCNA to the upstream duplex region by physically blocking diffusion of the sliding clamp along the adjacent ssDNA (36). SSB is a homotetramer that binds ssDNA with an occluded site size of 65 nt at 200 mM ionic strength where the ssDNA is fully wrapped around SSB, contacting each subunit within the heterotetramer. This differential mode of ssDNA-binding has no effect on the amount of PCNA loaded onto and maintained at P/T junctions (36), the orientation of PCNA encircling P/T junctions (Fig. 5A) (36), or the intrinsic stability of PCNA encircling P/T junctions (Fig. 5B). However, unlike RPA, SSB does not interact with Rad6/Rad18 (8, 10). These unique behaviors were exploited to decipher any effects of ssDNA-RPA-Rad6/Rad18 interactions (Fig. 3E) on monoubiquitination of PCNA encircling a P/T junction (Fig. 4E).

First, PCNA was pre-assembled on a P/T DNA substrate in the presence of a stoichiometric amount of a ssDNA-binding protein; SSB for P/BioT70 and RPA for P/BioT33 (Fig. 5C, top). Monoubiquitination of PCNA was then monitored over time as described above. Under these conditions, a single ssDNA-binding protein is adjacent to a P/T junction that is encircled by PCNA. On each DNA substrate, the same amount of PCNA is loaded onto the P/T junction and stabilized in the same orientation.
RPA regulates monoubiquitination of PCNA on and off DNA

Discussion

RPA is the most abundant ssDNA-binding protein in human cells and this high concentration is maintained throughout the cell cycle (26, 27). Furthermore, human RPA binds to ssDNA with extremely high affinity \( (K_a \sim 10^9 \text{M}^{-1}) \) at physiological ionic strength (28, 29). Together, this ensures that RPA immediately coats exposed ssDNA, protecting it from degradation and preventing formation of alternative DNA structures (4). RPA also directly engages in DNA metabolism through protein-protein interactions. For example, under native conditions when ssDNA exposure is minimal, free RPA binds the p53 tumor suppressor protein in the nucleoplasm and inhibits binding of p53 to transcriptional promoters. Upon generation of persistent ssDNA regions, RPA binds tightly to the exposed templates, releasing and activating p53 (40–42). Furthermore, RPA bound to exposed ssDNA directly recruits various proteins involved in DNA repair pathways and the DNA damage checkpoint/response (43). Our studies on human Rad6/Rad18 now reveal that RPA can regulate the activity of a protein complex both in the nucleoplasm and on exposed ssDNA.

ssDNA mediates RPA-Rad18 interactions

The results from Fig. 3 indicate that 1) RPA and Rad18 interact in solution, albeit weakly, in agreement with previous in vivo and in vitro studies (8); 2) ssDNA-bound RPA directly interacts with Rad6/Rad18 at physiological ionic strength; and 3) ssDNA enhances the affinity of RPA for Rad6/Rad18. Altogether, these results confirm previous hypotheses (8, 10). When bound to ssDNA, OB folds A and B of RPA70, the primary ssDNA-binding sites of RPA, are occupied by ssDNA (28–30) and, hence, unlikely to bind Rad18. Previous independent reports revealed that transactivator proteins directly compete with ssDNA for binding to the ssDNA-binding OB folds of RPA70 (44, 45). Particularly, binding of ssDNA and p53 to RPA are mutually exclusive and such behavior is critical for p53 regulation in human cells (40–42, 46). These transactivator proteins directly bind RPA through “acidic patches” that may adopt negatively charged, amphipathic helices that structurally mimic ssDNA binding to an OB fold of RPA (45, 47–49). The portion of Rad18 that interacts with RPA contains many acidic residues (Fig. S1). Specifically, amino acid sequence 171–189 is \( \sim 30\% \) D/E and may serve as an “acidic patch” to interact directly with RPA70.

RPA inhibits monoubiquitination of PCNA in solution

For PCNA monoubiquitination to occur, Rad6/Rad18 must be in an active conformation and interact productively with PCNA. Regarding the former, Rad18 must remain a homodimer and Rad6 must bind the Rad18 homodimer at two independent binding sites (4, 5, 7, 51, 52). Rad18 monoubiquitination, which requires homodimerization of Rad18 (25), is robust in the absence of RPA and ssDNA (Fig. 1). Furthermore, Rad6 monoubiquitination is negligible and polyubiquitin chain formation is not observed. Finally, monoubiquitination of free PCNA is quite significant over a biologically relevant time course, with ubiquitin attached 70.74 ± 6.756% of all PCNA homotrimers on average. Altogether, this indicates that functional interactions between the Rad18 homodimer and Rad6 are achieved and maintained in the absence of RPA and ssDNA, and functional Rad6/Rad18 interacts productively with free PCNA in solution.

Monoubiquitinated PCNA is only observed on chromatin after exposure of human cells to agents that generate persistent stretches of RPA-coated ssDNA (4, 6, 7, 17, 18). This stringent selectivity is critical as aberrant PCNA monoubiquitination leads to increases in both spontaneous and damaged-induced mutagenesis (53). Interestingly, ubiquitin-specific protease 1 (USP1), the primary deubiquitinase for PCNA, is dramatically down-regulated throughout G1 when more than 80% of the PCNA pool resides in the nucleoplasm (13–16, 19–23). A similar scenario arises in S-phase when USP1 is degraded or inactivated in response to genotoxic agents that cause fork uncoupling (2, 54). The results presented in Figs. 3 and 4 indicate that ssDNA-free RPA directly interacts with Rad6/Rad18 and inhibits monoubiquitination of free PCNA. Such behavior may compensate for loss of USP1 by preventing monoubiquitination of free PCNA and, hence, maintain the selectivity of PCNA monoubiquitination in the absence of USP1.

Inhibition of free PCNA monoubiquitination by RPA is directly and completely reversed by binding of ssDNA to RPA (Fig. 3). Under all conditions tested, monoubiquitination of Rad18 is not significantly affected (Fig. 2), mixed ubiquitin chain formation (Fig. S9) is not observed, and the occupancy of Rad6 within Rad6/Rad18 is maintained (Fig. 3). Collectively,
this indicates that RPA does not alter the composition of Rad6/Rad18; the Rad18 homodimer remains intact and maintains contact (via R6B) with the noncovalent ubiquitin-binding site on Rad6. RPA may regulate Rad6/Rad18 activity by mediating the Rad18 RING-Rad6 interaction in a ssDNA-dependent manner. In a previous report, full-length human Rad18 containing the RING domain inhibited polyubiquitin chain formation by Rad6 as well as Rad6 monoubiquitination. However, the isolated R6B was only sufficient to inhibit polyubiquitin chain formation and had no effect on Rad6 monoubiquitination, suggesting the Rad18 RING-Rad6 interaction may inhibit Rad6 monoubiquitination and promote PCNA monoubiquitination (7). If RPA mediates the Rad18 RING-Rad6 interaction, the effects of RPA on monoubiquitination of Rad6 and PCNA should be inversely correlated. Such behavior is clearly evident in the presence and absence of ssDNA (Fig. 2), in agreement with the proposed model. However, alternative models are also possible. For example, Rad6/Rad18 must productively bind PCNA for monoubiquitination to occur (7). Hence, RPA may regulate the activity of Rad6/Rad18 by mediating the Rad18-PCNA interaction in a ssDNA-dependent manner. Future studies will decipher these and other models.

**RPA enhances monoubiquitination of PCNA encircling stalled P/T junctions**

RPA promotes monoubiquitination of PCNA at persistent ssDNA regions generated throughout the cell cycle and such regulation is required for this PTM (8, 9, 13, 16). The results presented in Fig. 4 reveal that this is achieved, at least partially, by RPA stabilizing PCNA at P/T junctions by physically blocking diffusion of PCNA along the adjacent ssDNA. The results presented in Fig. 3 confirm that RPA directly interacts with Rad6/Rad18 and this interaction is significantly enhanced by binding of RPA to ssDNA. Thus, ssDNA/RPA complexes can directly recruit Rad6/Rad18 to the vicinity of stalled P/T junctions, as previously proposed (8, 10).

The extent and lifetime of such recruitment and whether it affects monoubiquitination of PCNA on DNA depends on the relative binding affinities of Rad6/Rad18 for an RPA/ssDNA complex and PCNA encircling stalled P/T junctions. To test this, we directly compared monoubiquitination of PCNA encircling a P/T junction abutted by either a single RPA or a single SSB (Fig. 5); only the former interacts with Rad6/Rad18 (8, 10). Under these conditions, any observed differences in PCNA monoubiquitination are attributed to interactions between Rad6/Rad18 and the RPA adjacent to the P/T junction. As the reaction progresses, the fraction of PCNA that is monoubiquitinated (i.e. reaction product) increases. This mimics S-phase in human cells after UV treatment where monoubiquitinated PCNA builds up and persists on ssDNA regions for many hours post UV (9, 55, 56). A relatively high affinity of Rad6/Rad18 for the RPA/ssDNA complex would inhibit PCNA monoubiquitination over time compared with SSB as the accumulating reaction products immobilize Rad6/Rad18, impeding turnover. In contrast, a relatively weak affinity favors direct binding of PCNA by Rad6/Rad18 from solution and localizes Rad6/Rad18 to an RPA/ssDNA complex only transiently, if at all, permitting efficient turnover throughout the incubation even as products accumulate. The results presented in Fig. 5 indicate that PCNA monoubiquitination is independent of the identity of the ssDNA-binding protein. Thus, the affinity of Rad6/Rad18 for the RPA/ssDNA complex is relatively weak compared with PCNA, and direct interactions, if any, between Rad6/Rad18 and the RPA/ssDNA complex do not affect monoubiquitination of the resident PCNA. However, these results do not rule out that weak, transient interactions selectively enhance monoubiquitination of PCNA on longer RPA-coated ssDNA regions. In human cells, persistent ssDNA regions generated at UV-induced lesions are 150–1250 nt in length, with the latter representing ~65% (2). Here, the effective concentration of RPA near stalled P/T junctions is much higher (~5- to 42-fold) than the resident PCNA and, hence, would promote interactions of Rad6/Rad18 with RPA along the ssDNA (28–30). These interactions may foster “successful collisions” between Rad6/Rad18 and the resident PCNA without significantly increasing the lifetime of Rad6/Rad18 on RPA-coated ssDNA. This unique model is the focus of future studies. Altogether, the results presented in Figs. 3–5 suggest that after DNA replication stress, RPA binds the ssDNA exposed downstream of stalled P/T junctions and restricts the resident PCNAs to the upstream duplex regions by physically blocking diffusion of PCNA along ssDNA. This activity is required for efficient monoubiquitination of PCNA on DNA. Furthermore, Rad6/Rad18 directly interacts with RPA/ssDNA complexes, albeit weakly, and these interactions may promote monoubiquitination of PCNA on lengthy RPA-coated ssDNA regions.

RPA is critical for PCNA monoubiquitination as knockdown to undetectable levels in human and budding yeast cells effectively eliminates this PTM on DNA after treatment with replication-blocking agents (8, 9). Additional cellular factors have also been implicated to various extents (4) and new discoveries continue to emerge (57, 58). Some have been shown to bind Rad18 and/or PCNA and may promote PCNA monoubiquitination through direct interactions with these proteins. Future biochemical and cellular studies are needed to elucidate the mechanism(s) by which each factor promotes PCNA monoubiquitination and to decipher how these distinct pathways for regulation are interconnected in vivo.

**Experimental procedures**

**Recombinant human proteins**

Detailed information about plasmid construction and the expression, purification, and labeling of proteins can be found in the supporting information.

**Oligonucleotides**

Oligonucleotides were synthesized by Integrated DNA Technologies (Coralville, IA), purified on denaturing polyacrylamide gels, and the concentrations were determined from the absorbance at 260 nm using the calculated extinction coefficients. For annealing the DNA substrates (Fig. S6), the primer and corresponding complementary template strands were mixed in equimolar amounts in 1× Annealing Buffer (10 mM Tris-HCl, pH 8.0, 100 mM NaCl, 1 mM EDTA), heated to 95 °C for 5 min, and slowly cooled to room temperature.
**RPA regulates monoubiquitination of PCNA on and off DNA**

**Antibodies**

Primary antibodies for PCNA (PC5, mouse monoclonal IgG1), RPA70 (MA70-2, mouse monoclonal IgG1), RPA32 (MA34, mouse monoclonal IgG1), RPA14 (mouse monoclonal IgG1), and Rad18 (H-77, rabbit polyclonal IgG), and secondary antibodies (goat anti-rabbit IgG-HRP) were purchased from Santa Cruz Biotechnology. Rad6 primary antibody (rabbit polyclonal IgG) was purchased from Abcam.

**Ubiquitination assays**

All ubiquitination assays were performed at room temperature (23 ± 2 °C) in 1 x ubiquitination assay buffer supplemented with 1 mM TCEP, and the final ionic strength was adjusted to physiological (200 mM) by the addition of appropriate amounts of KOAc. Rad6/Rad18, RPA, and poly(dT)₃₃, or pre-incubated complexes thereof, were loaded by RFC (180 nM with 1 mM ATP) onto an unlabeled DNA substrate (200 nM either Cy3P/BioT33 or Cy3P/BioT70 with 800 nM NeutrAvidin) that was pre-incubated with an ssDNA-binding protein (400 nM of either RPA or SSB) and FRET was monitored over time as described previously (35). Loading traces were fit to a single-exponential increase and normalized to their respective ranges.

**Analytical gel filtration**

Experiments were performed at 4 °C on a ÄKTA purifier (GE Healthcare) using a Superdex 200 10/300 GL column (GE Healthcare) in 1 x ubiquitination assay buffer supplemented with 1 mM TCEP, and the final ionic strength was adjusted to physiological (200 mM) by the addition of appropriate amounts of KOAc. Rad6/Rad18, RPA, and poly(dT)₃₃, or pre-incubated complexes thereof, were loaded in a volume of 150 µl. Concentrations of each are indicated in the respective figure/figure legend. The elution profile was analyzed by UV absorbance. Where indicated, fractions of a given elution profile were collected, resolved on 4–20% Mini-PROTEAN TGX™ Gels, and visualized by Coomassie Blue staining or Western blotting with Rad6 antibody. Western blots were quantified and normalized to a Rad6 loading control.

**Fluorescence microscopy**

All experiments were performed at room temperature (23 ± 2 °C) in 1 x ubiquitination assay buffer supplemented with 0.1 mg/ml BSA and 1 mM TCEP, and the final ionic strength was adjusted to physiological (200 mM) by the addition of appropriate amounts of KOAc. For steady-state fluorescence, measurements were done in Jobin Yvon FluoroMax-4 Fluorimeter. Assay solutions contained 110 nM Cy3-labeled P/T DNA (either Cy3P/BioT33 or Cy3P/BioT70), NeutrAvidin (0 or 440 nM), 1 mM ATnP, and either an ssDNA-binding protein (0 or 110 nM of RPA or SSB) or Rad6/Rad18, RPA, and poly(dT)₃₃, or pre-incubated complexes thereof, were loaded in a volume of 150 µl. Concentrations of each are indicated in the respective figure/figure legend. The elution profile was analyzed by UV absorbance. Where indicated, fractions of a given elution profile were collected, resolved on 4–20% Mini-PROTEAN TGX™ Gels, and visualized by Coomassie Blue staining or Western blotting with Rad6 antibody. Western blots were quantified and normalized to a Rad6 loading control.

**Acknowledgment**—We thank Dr. Titia K. Sixma who generously provided the plasmids for expression of Rad6/Rad18.

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