The role of the DNA damage response protein kinase ataxia telangiectasia-mutated (ATM)- and Rad-3-related (ATR) in the cellular response to DNA damage during the replicative phase of the cell cycle has been extensively studied. However, little is known about ATR kinase function in cells that are not actively replicating DNA and that constitute most cells in the human body. Using small-molecule inhibitors of ATR kinase and over-replicating DNA and that constitute most cells in the human body. Using small-molecule inhibitors of ATR kinase and over-expression of a kinase-inactive form of the enzyme, I show here that ATR promotes cell death in non-replicating/non-cycling cultured human cells exposed to N-acetoxy-2-acetylaminofluorene (NA-AAF), which generates bulky DNA adducts that block RNA polymerase movement. Immunoblot analyses of soluble protein extracts revealed that ATR and other cellular proteins containing SQ motifs become rapidly and robustly phosphorylated in non-cycling cells exposed to NA-AAF in a manner largely dependent on ATR kinase activity but independent of the essential nucleotide excision repair factor XPA. Although the topoisomerase I inhibitor camptothecin also activated ATR in non-cycling cells, other transcription inhibitors that do not directly damage DNA failed to do so. Interestingly, genetic and pharmacological inhibition of the XPB subunit of transcription factor IIH prevented the accumulation of the single-stranded DNA binding protein replication protein A (RPA) on damaged chromatin and severely abrogated ATR signaling in response to NA-AAF and camptothecin. Together, these results reveal a previously unknown role for transcription factor IIH in ATR kinase activation in non-replicating, non-cycling cells.

As one of the major DNA damage response signaling kinases in mammalian cells, the ATM- and Rad3-related (ATR) kinase is primarily thought to respond to DNA polymerase stalling and uncoupling from DNA helicase activity as a result of template lesions or dNTP shortage (1–3). These replicative stress events are characterized by regions of single-stranded DNA (ssDNA) and a junction of ssDNA and dsDNA (5′-primer-template junction), which together serve to recruit ATR and accessory proteins to ultimately activate ATR kinase signaling (2, 4). The functional outcomes of ATR activation in response to replication stress generally involve processes that ultimately promote cell survival, such as replication fork stabilization, cell cycle delay, inhibition of replication origin firing, DNA repair, and homologous recombination (2, 5, 6).

These pro-survival functions of ATR in cells containing replication stress likely limit the therapeutic efficacy of anticancer drugs that damage DNA, and thus small molecule inhibitors of the ATR kinase are being developed as adjuvants in chemotherapy regimens (7–10). Preliminary studies using mouse models of tumor progression have indeed suggested that ATR kinase inhibition can exacerbate the antiproliferative effects of radiation and cisplatin to more effectively slow tumor growth and shrink tumor volume (11, 12).

However, the majority of cells in the body that are exposed to DNA-damaging agents through environmental, dietary, or therapeutic means are in a non-replicating and/or differentiated state. Thus, it is important to determine whether the ATR kinase has any function in the DNA damage response in non-replicating cells, and, if so, how ATR becomes activated to carry out these activities. Using purified proteins and DNA substrates in vitro or growth-arrested, confluent populations of cultured cells, several reports have indicated that ATR may become activated by the direct recognition of bulky DNA adducts by ATR or its interacting proteins (13–18), via ssDNA gaps generated by excision repair (19–23), or by transcription stress caused by RNA polymerase stalling (24–26). Unfortunately, these studies have often restricted their analyses to phosphorylation of substrate proteins, such as H2AX and p53, which are not unique to ATR (27, 28). Moreover, many of these proteins exert complex or undefined roles in cell fate following DNA damage (29–32). Thus, the actual functions of ATR in non-cycling cells have remained largely unexplored.

Nevertheless, a recent report using small molecule inhibitors of ATR kinase activity revealed a pro-apoptotic function for ATR in non-cycling cells exposed to UV light, UV mimetics, and the topoisomerase I poison camptothecin (28). Here I have further extended this finding through the use of a genetic approach in which a kinase-inactive form of ATR is overexpressed in non-cy-
ATR kinase activation in non-cycling cells

Figure 1. Pharmacological and genetic inhibition of ATR kinase protects non-replicating cells from the lethal effects of the UV mimetic NA-AAF. A, cycling and non-cycling HaCaT cells were pulsed with 10 μg/ml BrdU for 15 min. Genomic DNA was then purified and analyzed by immunodot blotting with the indicated antibodies. The graph shows the relative level of BrdU incorporation into genomic DNA (normalized to cycling cells) from three independent experiments. B, non-cycling HaCaT cells were treated with the indicated concentration of the ATR inhibitor VE-821 for 30 min prior to treatment with 15 μM NA-AAF. Cells were stained with crystal violet 24 h later to determine relative survival. C, cells were treated with the indicated concentration of AZD6738 and analyzed as described in B, D, cells were treated with caffeine and analyzed as described in B. E, non-cycling U2OS cells containing either a WT or KD FLAG-tagged ATR transgene under the control of a tetracycline-inducible promoter were left untreated or treated with 1 μg/ml tetracycline for 48 h before analysis by immunoblotting. F, non-cycling U2OS cells containing the FLAG-ATR-KD transgene were left untreated (− TET, no tetracycline) or treated with tetracycline (+ TET) for 48 h before exposure to the indicated concentration of NA-AAF. After an additional 48 h, cells were stained with crystal violet to determine relative survival. G, non-cycling U2OS cells induced to express the indicated form of ATR were treated with NA-AAF as in E to determine relative cell survival, *, p < 0.05; indicating a significant difference in survival between the two treatments or cell lines.

cycling cells. Moreover, using the autophosphorylation of ATR and the phosphorylation of SQ motif-containing proteins as biochemical markers of ATR kinase activation, I show that ATR is indeed robustly activated in non-cycling cells exposed to DNA-damaging agents, even at levels of DNA damage that do not yield appreciable cell death. Interestingly, this mode of ATR kinase signaling appears to require overt DNA damage because general inhibitors of RNA polymerase function during transcription failed to induce a significant response. Characterization of the activation mechanism of ATR in non-cycling cells unexpectedly revealed a major role for the XBP DNA translocase subunit of transcription factor IIH (TFIIH) in ATR signaling. This phenotype was correlated with failure to properly load the single-stranded DNA-binding protein RPA on damaged chromatin. Because the DNA unwinding activity of TFIIH is important for transcription and RNA polymerase function, these results implicate a novel function for TFIIH and, specifically, its XBP subunit in ATR activation. Given that the majority of cells in the body are in a quiescent or non-replicating state, these findings have important implications for understanding the physiology of ATR-dependent DNA damage signaling responses in vivo.

Results
Pharmacological and genetic inhibition of the ATR kinase in non-cycling cells demonstrates a pro–cell death function for ATR in response to DNA damage

To better characterize the functions of the ATR kinase in non-replicating cells, the human keratinocyte-derived cell line HaCaT was grown to confluence and maintained in a low concentration of serum before treatment with several commonly used small-molecule ATR inhibitors and exposure to a DNA-damaging agent. Immunodot blotting of genomic DNA from cells pulsed with BrdU verified that the cells were not actively replicating DNA (Fig. 1A) and, hence, can be referred to as being in a non-cycling state. The non-cycling cells were then treated with a small-molecule ATR inhibitor for 30 min prior to treatment with the carcinogenic UV mimetic NA-AAF, which was employed as a model DNA-damaging agent here because it generates bulky adducts on the C8 position of guanines that block RNA polymerase movement when not removed by the nucleotide excision repair system (33–35). Cell survival was then measured 1 day after treatment by crystal violet staining of the remaining adherent cells. As shown in Fig. 1, B–D, in the absence of an ATR inhibitor, nearly 70% of the cells were killed by NA-AAF. In contrast, the highly selective ATR inhibitors VE-821 and AZD6738 provided protection to NA-AAF–treated cells and limited the extent of cell death in a dose-dependent manner, with an EC_{50} of 3.6 μM and 1.1 μM, respectively (Fig. 1, B and C).

Although relatively non-selective, caffeine has also been widely used to study ATR signaling, which is based in part on its ability to inhibit the activity of the purified enzyme (36, 37) and abrogate cell cycle checkpoints (38). However, other studies have questioned its utility for studying ATR kinase signaling in cells with DNA damage (39). When caffeine-treated, non-cycling cells were exposed to NA-AAF, I observed that, unlike the specific ATR inhibitors VE-821 and AZD6738, caffeine instead sensitized the cells to the DNA-damaging agent (Fig. 1D).
**ATR kinase activation in non-cycling cells**

Because the pharmacological ATR inhibitors may target other kinases besides ATR, I next took a genetic approach to mimic ATR kinase inactivation in non-replicating cells. I therefore took advantage of two U2OS cell lines that can be induced to express either a WT or kinase-inactive (kinase-dead, KD) form of ATR in a tetracycline-inducible manner (40–42). As shown in Fig. 1E, 48-h induction with tetracycline led to a modest increase in total ATR protein levels in the two cell lines.

Two experimental approaches were then used to determine whether the expression of the kinase-inactive form of ATR protects non-cycling cells from NA-AAF in a manner similar to that of the two highly specific ATR kinase inhibitors. In the first method, U2OS cells with the FLAG-ATR KD transgene were grown to confluence and serum-starved prior to induction with tetracycline for 2 days. Non-induced and induced cells were then exposed to increasing concentrations of NA-AAF, and cell survival was measured 2 days later. Although U2OS cells are more resistant to NA-AAF than HaCaT cells (28), increasing concentrations of the drug nonetheless yielded a decreasing fraction of surviving cells (Fig. 1F). Furthermore, cells induced to overexpress ATR-KD with tetracycline were more resistant to NA-AAF than the non-induced cells, which indicates that NA-AAF–induced cell death is dependent in part on ATR kinase activity.

In a second, related approach, I induced ATR-WT and ATR-KD expression with tetracycline in the appropriate cell lines and then monitored cell survival after NA-AAF treatment. The ATR-KD cells were found to be less susceptible to cell death than ATR-WT cells (Fig. 1G). These findings are consistent with the effects of the small-molecule inhibitors of the ATR kinase in non-cycling cells and are strikingly different from the effect of ATR kinase inhibition in asynchronous populations of cells, in which sensitization to DNA-damaging agents has been routinely observed (28, 40, 42).

Together, the pharmacological and genetic approaches for inhibiting ATR kinase function in non-cycling cells complement one another and demonstrate that one function of ATR in non-cycling cells is to promote cell death following NA-AAF treatment. Recent work showed that this pro-death function of ATR occurs in part through stimulation of apoptotic signaling (28). This function of ATR in non-cycling cells is therefore distinct from the pro-survival functions of ATR in replicating, cycling cells.

**ATR autophosphorylation on Thr-1989 in non-cycling cells**

Although the cell survival assays presented in Fig. 1 suggest that ATR is activated in non-cycling cells containing DNA damage, there are currently no established biochemical readouts for ATR activation in non-cycling cells. The checkpoint kinase CHK1 is a canonical substrate for ATR in replicating cells exposed to DNA-damaging agents. However, in non-cycling cells, CHK1 protein is not present (19, 27, 28), and, therefore, no CHK1 phosphorylation is observed following exposure to NA-AAF (Fig. 2A).

ATR has been shown to phosphorylate itself on Thr-1989 in asynchronous populations of cells exposed to inducers of replication stress (43, 44). To determine whether this residue becomes phosphorylated in non-replicating cells, I exposed both cycling and non-cycling cells to NA-AAF and then monitored Thr-1989 phosphorylation by immunoblotting. As

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**Figure 2. ATR autophosphorylation on Thr-1989 is a marker of ATR activation in non-cycling cells exposed to the UV mimetic NA-AAF.**

A, cycling and non-cycling HaCaT cells were treated with 20 μM NA-AAF for 1 h. Cell lysates were analyzed by immunoblotting with antibodies targeting the indicated proteins and phosphorlated residues (P). Quantitation of ATR autophosphorylation (average and standard error) from three independent experiments is provided below a representative immunoblot. The phospho-ATR signal was normalized to total ATR, and this ratio was set to an arbitrary value of 100 for NA-AAF–treated cycling cell sample in each experiment (set to an arbitrary value of 100). B, non-cycling HaCaT cells were pretreated with DMSO or 10 μM VE-821 (an ATR inhibitor) for 30 min prior to treatment with 10 μM NA-AAF. Cells were harvested at the indicated time point and analyzed by immunoblotting. C, quantitation of results from at least two independent experiments performed as in B. D, U2OS cells induced with tetracycline for 2 days to express either wild-type or kinase-dead forms of ATR were left untreated or treated with 60 μM NA-AAF for 4 h, and then cell lysates were analyzed by immunoblotting. Quantitation of three independent experiments is provided below the representative immunoblot data. The phospho-ATR signal was normalized to total ATR, and, p < 0.05; indicating a significant difference in NA-AAF–induced ATR phosphorylation in WT and KD cells.
A DNA damage-dependent increase in ATR phosphorylation was observed in both cycling and non-cycling cells. Although the levels of both phosphorylated ATR and total ATR were reduced in non-cycling cells in the absence and presence of NA-AAF, quantitative analyses from several independent experiments comparing the induction of ATR phosphorylation (normalized to total ATR) by NA-AAF showed a similar 2.5-fold increase in ATR phosphorylation in both cycling and non-cycling cells (Fig. 2A, bottom panel).

I next examined the kinetics of ATR phosphorylation and its dependence on ATR kinase activity. As shown in Fig. 2B, ATR became phosphorylated on Thr-1989 in a time-dependent manner following NA-AAF treatment. Importantly, this DNA damage-dependent response was prevented by treatment with the ATR kinase inhibitor VE-821 (Fig. 2, B and C). To further validate the effects of the pharmacological inhibitor on ATR phosphorylation on Thr-1989, the inducible U2OS cell lines expressing either the WT or KD forms of ATR were exposed to NA-AAF. Although an increase in ATR phosphorylation was observed in cells expressing ATR-WT, both background and NA-AAF–dependent Thr-1989 phosphorylation were significantly reduced in cells expressing ATR-KD (Fig. 2D). Moreover, expression of ATR-KD also largely abrogated NA-AAF–induced phosphorylation of the ATR/ATM substrate p53.

These results indicate that the phosphorylation of ATR on Thr-1989 can be used as a biomarker for ATR kinase activation in non-cycling cells exposed to DNA-damaging agents and further argue that ATR activation can occur in the absence of canonical replication stress.

**ATR plays a major role in DNA damage-induced protein phosphorylation events in non-cycling cells**

Mass spectrometric studies have demonstrated that hundreds of proteins become phosphorylated by ATM and/or ATR in response to DNA damage in proliferating cells (45, 46). To determine whether ATR makes a significant contribution to protein phosphorylation events in non-cycling cells, I made use of a mixture of monoclonal antibodies targeting phosphorylated SQ motifs common to ATR and ATM kinase substrates (47) in immunoblotting experiments of lysates from NA-AAF–treated cells. As shown in Fig. 3A, NA-AAF induced the phosphorylation of many SQ motif-containing proteins in both cycling and non-cycling cells. Consistent with earlier results, CHK1 phosphorylation was only observed in cycling cells. Quantification of the total SQ motif phosphorylation from several independent experiments demonstrated a 5- to 10-fold increase in protein phosphorylation and a slightly stronger total.
response in non-cycling cells than in cycling cells (Fig. 3A, bottom panel).

Additional analyses demonstrate that the degree of SQ motif phosphorylation in non-cycling cells was dependent on NA-AAF concentration and occurred at low doses of NA-AAF that do not lead to detectable cell death (28, 48, 49) (Fig. 3B). Similarly, analysis of SQ motif phosphorylation kinetics revealed robust signaling as early as 15 min after drug administration and continued phosphorylation over the course of at least 4 h (Fig. 3C).

To clarify the dependence of SQ motif phosphorylation on ATR kinase activity, cells were pretreated with specific inhibitors of ATR and the related DNA damage response kinases ATM and DNA-PK. ATR inhibition reduced the level of SQ motif phosphorylation by ~50% (Fig. 3D). However, the ATR inhibitor alone also induced modest protein phosphorylation, which indicates that ATR inhibition may induce genomic stress that activates other DNA damage response kinases in non-replicating cells. Nevertheless, and consistent with previous evidence that ATM is activated in non-replicating cells exposed to UV mimetics (23, 27, 28, 50), ATM inhibition also partially reduced the extent of SQ motif phosphorylation. In contrast, the DNA-PK inhibitor failed to significantly affect protein phosphorylation. These results demonstrate that both ATR and ATM contribute to SQ motif phosphorylation in non-replicating cells exposed to NA-AAF.

USOS cells expressing the WT and KD forms of ATR were next employed to validate that ATR kinase activity contributes to DNA damage-dependent SQ motif phosphorylation in non-replicating cells. As shown in Fig. 3E, NA-AAF–induced protein phosphorylation was significantly attenuated in ATR-KD cells in comparison with the cells expressing ATR-WT.

In summary, these results demonstrate that SQ motif phosphorylation occurs rapidly in non-cycling cells containing bulky, transcription-stalling DNA adducts at non-lethal concentrations of NA-AAF and is mediated in large part via the ATR kinase. Thus, in addition to ATR autophosphorylation on Thr-1989, SQ motif phosphorylation can also serve as a convenient marker for DNA damage-dependent ATR signaling in non-cycling cells.

**Reduced expression of the essential nucleotide excision repair factor XPA does not significantly affect NA-AAF–induced ATR kinase signaling in non-replicating cells**

The nucleotide excision repair system can excise NA-AAF-induced DNA lesions from the genome (51), and experiments with cultured cells and defined in vitro assays with purified proteins have indicated that excision gaps enlarged by the endonucleolytic action of ExoI are stimuli for ATR kinase activation (19, 20, 52, 53). However, these analyses of ATR activation have utilized a rather limited number of protein substrates, such as p53 and RPA, which are not necessarily specific to ATR. Indeed, I recently showed that the simultaneous inhibition of both the ATR and ATM kinases was necessary to eliminate p53, H2AX, and KAP-1 phosphorylation in non-cycling human cells exposed to either UV light or the UV mimetic NA-AAF (28). Thus, the extent to which excision gaps versus other stimuli activate ATR in non-replicating cells is not known.

To determine whether ATR kinase signaling in non-cycling cells is dependent on nucleotide excision repair, expression of the core excision repair factor XPA was reduced by RNA interference. As shown in Fig. 4A, the use of a lentivirus shRNA targeting XPA mRNA efficiently lowered XPA protein levels by ~95%. Importantly, this degree of knockdown was sufficient to modestly sensitize non-cycling shXPA-expressing HaCaT cells to the short-term, toxic effects of NA-AAF (Fig. 4B).

I next examined the induction of ATR kinase signaling in NA-AAF-treated cells by monitoring the phosphorylation status of ATR and SQ motif-containing proteins in the cell lysates. As shown in Fig. 4C, strong phosphorylation was observed for both readouts of ATR activation in both control and shXPA-expressing cells. Quantitation of four independent experiments failed to detect a significant difference in protein phosphorylation between the two cell lines (Fig. 4D). These results suggest that the bulk of ATR kinase signaling in NA-AAF-treated non-replicating cells may not be the result of gaps generated by...
ATR kinase activation in non-cycling cells

nucleotide excision repair. Consistent with this interpretation, ATR kinase inhibition was shown previously to protect cells depleted of XPA from the lethal effects of NA-AAF (28). Thus, some other stimulus is likely responsible for the majority of ATR kinase signaling in non-replicating cells exposed to NA-AAF.

Direct DNA damage, but not general transcription stress, leads to robust activation of ATR kinase signaling in non-cycling cells

To further examine the mechanism of ATR kinase activation in non-replicating cells, non-cycling HaCaT cells were treated with various compounds that interfere with transcription. Camptothecin (CPT) causes direct DNA damage through the stabilization of transient topoisomerase I–DNA cleavage complexes that normally help to resolve superhelical tension that is generated in DNA during gene transcription. As shown in Fig. 5A, camptothecin induced a nearly 3-fold increase in ATR phosphorylation on Thr-1989 in a manner similar to NA-AAF. Treatment with the ATR inhibitor VE-821 largely blocked this response. In contrast, additional transcription inhibitors that ultimately cause cell death (28) but do not directly cause DNA damage failed to stimulate ATR autophosphorylation. These transcription inhibitors included triptolide (TPL), 5,6-dichloro-1-β-d-ribofuranosyl-1H-benzimidazole (DRB), and actinomycin D (ActD). Importantly, TPL, DRB, and ActD act via different mechanisms and at different stages of transcription (54). As a DNA intercalator, ActD directly inhibits the movement of RNA polymerases (54). In contrast, DRB specifically inhibits the CDK9 kinase activity of the positive transcription elongation factor P-TEFb, which normally phosphorylates RNA polymerase II and facilitates the transition of the polymerase from its initiated to its elongating state. Lastly, TPL forms a covalent complex with the XPB subunit of TFIIH, which inactivates the ATPase activity of the enzyme and prevents the initiation of transcription (55).

Because some of these agents have been shown to lead to either p53 phosphorylation and/or stabilization in fibroblasts (24, 56–59), I also monitored the phosphorylation of p53 on Ser-15 (an ATM/ATR target site) in non-cycling HaCaT cells. Strong ATR-dependent p53 phosphorylation was observed in cells treated with CPT and NA-AAF (Fig. 5A). However, the extent of p53 phosphorylation induced by DRB, TPL, and ActD was relatively weak in comparison with that induced by CPT and NA-AAF. These data, which are quantified in Fig. 5B, suggest that direct damage to the DNA template is necessary for robust ATR activation in non-cycling cells and that general inhibition of transcription is insufficient to elicit a significant response.

Inhibition of the XPB subunit of TFIIH abrogates ATR kinase activation and prevents RPA loading onto chromatin

Although the general transcription inhibitors failed to induce significant ATR kinase signaling in non-cycling cells, the results allowed me to examine whether the collision of RNA polymerases with a DNA lesion may be required to induce ATR activation (24–26, 60). Thus, according to this hypothesis, the stalling of RNA polymerase movement prior to exposure to NA-AAF or CPT should prevent the activation of ATR.

Therefore, HaCaT cells were treated with different transcription inhibitors before exposure to NA-AAF. As shown in Fig. 6A, NA-AAF treatment resulted in a clear increase in ATR, p53, and SQ motif protein phosphorylation in cells treated with DMSO, ActD, or DRB. Furthermore, quantitation of several independent experiments revealed that ActD and DRB actually caused modest potentiation of NA-AAF–induced ATR signaling (Fig. 6B). These results indicate that interfering with transcription elongation prior to DNA damage formation does not negatively impact the subsequent activation of the ATR kinase.

In striking contrast, TPL treatment instead caused strong inhibition of ATR, p53, and SQ motif phosphorylation following exposure to NA-AAF (Fig. 6, A and B), which indicates that the effect of TPL on DNA damage processing and ATR activation occurs through a distinctly different mechanism than that of the transcription inhibitors DRB and ActD. Nevertheless, to further confirm this finding, I next examined the effect of TPL on CPT-induced ATR kinase activation. Quantitation of these results is provided in Fig. 6C and shows that the activation of ATR signaling in non-cycling cells treated with CPT is largely attenuated by prior treatment with the TFIIH inhibitor TPL.

Triptolide forms a covalent complex with the XPB subunit of TFIIH (55), which inhibits the ATPase activity of XPB that is
required for TFIIH to unwind DNA during the initiation of transcription so that previously loaded RNA polymerase II can synthesize mRNA. Thus, TPL acts in a different manner than ActD or DRB, which instead inhibit transcriptional elongation via intercalation into DNA and by preventing RNA polymerase II phosphorylation (54), respectively. Given that ssDNA coated with RPA is generally considered to be a prerequisite for ATR kinase recruitment and activation in response to replication stress and other genotoxic stimuli (2, 3, 61, 62), I next examined whether TPL affected the accumulation of RPA on the chromatin fraction of cells following generation of DNA damage.

Although NA-AAF treatment led to a 2.5- to 3-fold increase in RPA protein levels on chromatin in cells treated with DMSO, DRB, or ActD (Fig. 6D), TPL instead completely blocked the DNA damage–dependent enrichment of RPA on chromatin. These findings therefore complement the ATR kinase signaling defects induced by TPL and indicate an important role for TFIIH in generating a most widely recognized signal for ATR kinase activation in non-replicating cells.

Although the use of TPL specifically implicates a role for the XBP subunit of TFIIH in ATR activation, TFIIH is a multisubunit enzyme with several distinct biochemical activities that are potentially relevant to transcription and associated genotoxic stress responses (63). Thus, to further determine whether the XBP subunit of TFIIH is specifically required for ATR kinase activation in response to DNA damage in non-replicating cells, I next examined how two additional small-molecule inhibitors of TFIIH affected DNA damage-induced ATR activation in non-replicating cells. These compounds included spironolactone and THZ1. Spironolactone (SP) induces rapid and specific proteolytic degradation of the XBP subunit of TFIIH while leaving the remaining subunits of TFIIH largely intact (64). THZ1 is
a specific inhibitor of the CDK7 kinase component of TFIIH (65) that phosphorylates RNA polymerase II to promote transcription elongation. As shown in Fig. 7A, TPL and SP abrogated SQ motif phosphorylation by 55–70% in non-replicating cells treated with NA-AAF. In contrast, THZ1 modestly stimulated SQ motif phosphorylation following NA-AAF exposure. This potentiation is analogous to the effect of DRB (Fig. 6B), which similarly inhibits RNA polymerase II phosphorylation (54). Importantly, similar results were obtained when cells were treated with the DNA-damaging agent CPT (data not shown).

To further validate that XPB is important for ATR activation in response to DNA damage in non-replicating cells, RNA interference was used to reduce XPB protein levels prior to exposure of cells to NA-AAF. Similar to the effects of TPL and SP, the XPB siRNAs partially blocked NA-AAF–induced SQ motif phosphorylation in non-replicating cells (Fig. 7B). Together, the genetic and pharmacological approaches reveal an unanticipated role for the XPB DNA translocase component of TFIIH in the activation of ATR in non-replicating cells exposed to DNA-damaging agents.

**Discussion**

The functions of the ATR kinase in promoting cell survival in response to replication stress are well documented (2, 3, 5, 6). In contrast, little is known regarding the role of ATR in response to DNA damage in cells that are not actively replicating DNA. Here I have extended the previous finding that ATR can promote an apoptotic form of cell death in response to UV light, UV mimetics, and other transcriptional stressors (28) by providing complementary pharmacological and genetic data showing that inhibition of ATR kinase activity protects non-cycling cells from DNA damage–induced lethality (Fig. 1). Thus, ATR kinase inhibition can have completely opposite functional effects on cell survival that depend not on the DNA-damaging agent but, instead, on the proliferation status of the cell and the related genomic stress that is encountered. Because the overwhelming majority of cells in the human body are in a differentiated, quiescent, or slowly cycling state, this phenomenon has important implications regarding our understanding of physiological responses to DNA damage, including in epithelial cells, which are at greatest risk of exposure to dietary, occupational, and environmental carcinogens.

These findings are also relevant to the use of ATR kinase inhibitors in cancer chemotherapy regimens (7, 8, 10). In addition to facilitating cell death of rapidly proliferating cancer cells, the results suggest that ATR inhibitors may provide protection to other cell types that are not actively undergoing DNA synthesis. Consistent with this hypothesis, a recent study indicated that the ATR inhibitor AZD6738 may be radioprotective in certain contexts within intestinal crypt cells in mice exposed to total-body ionizing irradiation (66).

Determining the mechanism of ATR activation in non-cycling cells and its downstream functional targets are therefore important issues for improving cancer chemotherapy protocols and for understanding how DNA damage promotes mutagenesis and carcinogenesis. The use of ATR autophosphorylation and SQ motif phosphorylation shown here (Figs. 2 and 3)
should facilitate such analyses of ATR signaling by providing readily employable biochemical readouts for ATR activation in cells that are not actively replicating DNA.

The predominant model for ATR kinase activation involves its recruitment to ssDNA coated by RPA (62), which, during the replicative phase of the cell cycle, is thought to occur when DNA damage or a lack of dNTP precursors causes DNA helicase and DNA polymerase activities to become uncoupled (1). Whether a similar scenario takes place in non-replicating cells in response to RNA polymerase stalling is not clear, and it was therefore somewhat surprising that the inhibition of transcriptional elongation with DRB and ActD did not lead to robust ATR activation (Fig. 6). This finding indicates that the structural requirements for ATR recruitment and activation (4) are not satisfied in non-replicating cells in the absence of overt DNA damage caused by compounds such as NA-AAF and CPT.

Interestingly, the dramatic abrogation of DNA damage-dependent ATR signaling by the TFIIH/XPB inhibitors triptolide and spironolactone (Fig. 6 and 7) was also surprising. However, given the role of TFIIH, and specifically its XPB subunit, in unwinding DNA during transcription initiation (63, 67), its apparent function in promoting ATR kinase activation may therefore be analogous to that of the minichromosome maintenance helicase when DNA damage is encountered during DNA synthesis (1). TPL is known to form a covalent complex with Cys-342 of XPB and inhibits its ATPase activity (55, 68), which is required for DNA translocation. Thus, the demonstration that TPL interferes with RPA accumulation on damaged chromatin (Fig. 6D) is consistent with such a role for XPB/TFIIH in generating ssDNA necessary for ATR recruitment and/or activation. However, further studies are needed to test this hypothesis. Although XPB ATPase activity is also necessary for nucleotide excision repair (67), the demonstration that reduced expression of the essential excision repair factor XPA does not significantly impact ATR kinase signaling in non-cycling cells (Fig. 4) and the fact that topoisomerase I inhibition also activates ATR (Fig. 5 and 6) suggest that a transcription-associated function of TFIIH is more relevant to ATR signaling here. Whether this is strictly a transcription initiation event or is instead associated with TFIIH acting at RNA polymerases stalled at DNA lesions to generate ssDNA in other contexts will need to be more clearly resolved. Detailed biochemical studies with purified protein components and defined DNA substrates will therefore be critical to characterizing this new mode of ATR kinase activation. Finally, interfering with XPB expression and function does not completely eliminate DNA damage–dependent ATR signaling, and thus there are likely other modes of ATR activation in non-cycling cells.

Furthermore, although the use of non-cycling cells in this report was borne out of a desire to uncover replication- and cell cycle–independent functions of the ATR kinase, this TFIIH-dependent mode of ATR activation probably also occurs in cycling and replicating cells to some extent. Consistent with this hypothesis, TFIIH subunits were found by mass spectrometry to accumulate on nascent DNA at stalled replication forks (69). Preliminary studies using a sensitive assay for detecting the small excised oligonucleotide products of nucleotide excision repair (70–73) have so far indicated that TFIIH function in this context is likely independent of nucleotide excision repair.3 Thus, I speculate that TFIIH may help to resolve transcription problems or replication–transcription collisions in replicating cells in part through activation of a specific ATR signaling cascade.

In summary, the results presented here indicate that ATR activation occurs in non-cycling cells through a mechanism that may be analogous to the one that takes place in response to replication stress, in which ssDNA generated by DNA unwinding and coated by RPA plays a crucial role in recruiting and activating ATR (1, 2, 4, 61). So far, the only known function for ATR in non-replicating cells is promotion of an apoptotic form of cell death. However, it is also possible that the reduced apoptosis and cell death that is observed in cells exposed to ATR inhibitors is an indirect consequence of an abnormal DNA damage response that causes alterations to global gene expression. Along these lines, a recent report demonstrated that ATR regulates alternative splicing in UV-irradiated cells and that more than 80 genes involved in apoptosis undergo significant alternative splicing in response to UV–induced cyclobutane pyrimidine dimers (74). Nevertheless, given the broad diversity of ATR and ATM substrate proteins that have been identified by phosphoproteomics (45, 46), other functions for ATR in non-cycling cells are expected. Additional studies will therefore be necessary to fully define this new mechanism of ATR activation and its functions in non-replicating cells.

**Experimental procedures**

**Cell lines**

U2OS cell lines expressing wild-type and kinase-dead forms of FLAG-ATR (GW33 and GK41) in a tetracycline-inducible manner were obtained from Paul Nghiem (40–42). The U2OS cell lines and HaCaT keratinocytes were maintained in DMEM supplemented with 10% FBS, 6 mm L-glutamine, 100 units/ml penicillin, and 100 µg/ml streptomycin at 37 °C in a 5% CO₂ humidified incubator. Subconfluent cells grown under these conditions are referred to as cycling cells throughout this manuscript. Cells were brought to a non-cycling state following plating at 40–60% confluence, growth for 2 days in normal medium until the cells reached confluence, and then a medium change to DMEM containing 0.5% FBS for 2–3 days prior to experimentation. Measurement of BrdU incorporation into the genomic DNA of the cells grown under these two conditions was performed as described previously (28) and was used to validate that the cells were either cycling (replicating DNA) or non-cycling (not replicating DNA). FLAG-ATR expression was induced in GW33 and GK41 U2OS cell lines by addition of 1 µg/ml of tetracycline to the culture medium for 48 h.

**Chemicals and reagents**

NA-AAF was purchased from the MRIGlobal Chemical Carcinogen Repository and resuspended in 95% ethanol. BrdU, TPL, DRB, CPT, ActD, caffeine, and tetracycline were obtained from Sigma. Inhibitors of the DNA damage response kinases ATR (VE-821 and AZD6738), ATM (KU55933), and DNA-PK

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3 M. G. Kemp, unpublished data.
Cells were pelleted by gentle centrifugation. Cells were then lysed for 20 min with low serum–containing medium 6 h after the second transfection and then again 24 h later. Culture medium was replaced 1:2000 dilution in 1× PBS, and the RPA70 antibody was from Bethyl Laboratories. The phospho-ATR (Thr-1989, GTX128145) antibody was from Cell Signaling, and phospho-ATM/ATR substrate (SQ, 9607). Antibodies included antibodies against ATR (sc-1887), CHK1 (Ser-15, 9284), and phospho-ATM/ATR substrate (Thermo Scientific), or ECL Prime Western blotting substrate. Equal amounts of cell lysates were separated by SDS-PAGE, transferred to nitrocellulose, and then probed by immunoblotting using standard procedures. All blots were stained with Ponceau S, and images were obtained for later quantitative purposes. Primary antibodies included antibodies against ATR (sc-1887), CHK1 (sc-8408), actin (I-19), and XPA (sc-853) from Santa Cruz Biotechnology. The soluble cell lysates were transferred to new tubes. Chromatin-associated proteins were enriched from cells following two extractions with a modified cytoskeletal buffer (10 mM Tris-HCl (pH 7.4), 100 mM NaCl, 3 mM MgCl2, 1 mM EDTA, 1 mM Na3VO4, 10 mM NaF, and 0.1% Triton X-100). Equal amounts of cell lysates were separated by SDS-PAGE, transferred to nitrocellulose, and then probed by immunoblotting using standard procedures. All blots were stained with Ponceau S, and images were obtained for later quantitative purposes. Primary antibodies included antibodies against ATR (sc-1887), CHK1 (sc-8408), actin (I-19), and XPA (sc-853) from Santa Cruz Biotechnology and phospho-CHK1 (Ser-345, 2348), phospho-p53 (Ser-15, 9284), and phospho-ATM/ATR substrate (SQ, 9607). The phospho-ATR (Thr-1899, GTX128145) antibody was from GeneTex, and the RPA70 antibody was from Bethyl Laboratories (A300-421A). All primary antibodies were used at 1:1000 or 1:2000 dilution in 1× TBST (50 mM Tris-HCl (pH 7.4), 135 mM NaCl, and 0.1% Tween 20). Secondary antibodies included horseradish peroxidase–linked anti-rabbit IgG, anti-mouse IgG, and anti-goat IgG. Chemiluminescence was visualized with Clarity Western ECL substrate (Bio-Rad), West Femto substrate (Thermo Scientific), or ECL Prime Western blotting detection reagent (GE Healthcare/Amersham Biosciences) using the Molecular Imager Chemi-Doc XRS+ or MP imaging systems (Bio-Rad). Ponceau-stained membranes and chemiluminescent signals within the linear range of detection were quantified using Image Lab (Bio-Rad) or ImageQuant software (GE Healthcare). For each immunoblot, the phosphoprotein signal was quantified and normalized to ATR or the total Ponceau S stain. The maximum signal for each blot was set to an arbitrary value of 100, and all other phosphoprotein/total protein ratios were then normalized to this value for each immunoblot. All experiments analyzing DNA damage response signaling were repeated two to four times, as indicated, and the average (and standard error) of the phosphoprotein/total protein ratios were determined and plotted. t tests were used to determine statistically significant differences between treatment groups.

**Cell survival assays**

Cell survival assays were performed as described previously using crystal violet staining and quantitation of the solubilized dye with a spectrophotometer (28). The absorbance values of the untreated samples were set to an arbitrary value of 1 for each experiment, and all treatment samples were normalized to this value. All cell survival experiments were performed at least three times.

**Author contributions**—M. G. K. conceived the idea for this project, carried out the experiments, and wrote the paper.

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**References**

1. Byun, T. S., Pacek, M., Yee, M. C., Walter, J. C., and Cimprich, K. A. (2005) Functional uncoupling of MCM helicase and DNA polymerase activities activates the ATR-dependent checkpoint. *Genes Dev.* 19, 1040–1052

2. Cimprich, K. A., and Cortez, D. (2008) ATR: an essential regulator of genome integrity. *Nat. Rev. Mol. Cell Biol.* 9, 616–627

3. Nam, E. A., and Cortez, D. (2011) ATR signalling: more than meeting at the fork. *Biochem. J.* 436, 527–536

4. MacDougall, C. A., Byun, T. S., Van, C., Yee, M. C., and Cimprich, K. A. (2007) The structural determinants of checkpoint activation. *Genes Dev.* 21, 898–903

5. Sancar, A., Lindsey-Boltz, L. A., Unsal-Kaçmaz, K., and Linn, S. (2004) Molecular mechanisms of mammalian DNA repair and the DNA damage checkpoints. *Annu. Rev. Biochem.* 73, 39–85

6. Ciccia, A., and Elledge, S. J. (2010) The DNA damage response: making it safe to play with knives. *Mol. Cell.* 40, 179–204

7. Karnitz, L. M., and Zou, L. (2015) Molecular pathways: targeting ATR in cancer therapy. *Clin. Cancer Res.* 21, 4780–4785

8. Toledo, L. I., Murga, M., and Fernandez-Capetillo, O. (2011) Targeting ATR and Chk1 kinases for cancer treatment: a new model for (new and old) drugs. *Mol. Oncol.* 5, 368–373

9. Llona-Minguez, S., Höglund, A., Jacques, S. A., Koolmeister, T., and Helleday, T. (2014) Chemical strategies for development of ATR inhibitors. *Expert Rev. Mol. Med.* 16, e10

10. Fokas, E., Prevo, R., Hammond, E. M., Brunner, T. B., McKenna, W. G., and Muschel, R. J. (2014) Targeting ATR in DNA damage response and cancer therapeutics. *Cancer Treat. Rev.* 40, 109–117

11. Fokas, E., Prevo, R., Pollard, J. R., Reaper, P. M., Charlton, P. A., Cornellis, B., Valls, K. A., Hammond, E. M., Oclina, M. M., Gillies McKenna, W., Muschel, R. J., and Brunner, T. B. (2012) Targeting ATR in vivo using the novel inhibitor VE-822 results in selective sensitization of pancreatic tumors to radiation. *Cell. Death Dis.* 3, e441

12. Vendetti, F. P., Lau, A., Schamus, S., Conrads, T. P., O’Connor, M. J., and Bakkenist, C. J. (2015) The orally active and bioavailable ATR kinase inhibitor AZD6738 potentiates the anti-tumor effects of cisplatin to resolve ATM-deficient non-small cell lung cancer in vivo. *Oncotarget* 6, 44289–44305

**ATR kinase activation in non-cycling cells**

(nu7441) were purchased from Selleckchem. SP and THZ1 were obtained from APEXBio.
ATR kinase activation in non-cycling cells

13. Unsal-Kacmaz, K., Makarov, A. M., Griffith, J. D., and Sancar, A. (2002) Preferential binding of ATR protein to UV-damaged DNA. *Proc. Natl. Acad. Sci. U.S.A.* **99**, 6673–6678

14. Jiang, G., and Sancar, A. (2006) Recruitment of DNA damage checkpoint proteins to damage in transcribed and nontranscribed sequences. *Mol. Cell. Biol.* **26**, 39–49

15. Choi, J. H., Lindsey-Boltz, L. A., and Sancar, A. (2007) Reconstitution of a human ATR-mediated checkpoint response to damaged DNA. *Proc. Natl. Acad. Sci. U.S.A.* **104**, 13301–13306

16. Choi, J. H., Lindsey-Boltz, L. A., and Sancar, A. (2009) Cooperative activation of the ATR checkpoint kinase by TopBP1 and damaged DNA. *Nucleic Acids Res.* **37**, 1501–1509

17. Liu, Y., Fang, Y., Shao, H., Lindsey-Boltz, L., Sancar, A., and Modrich, P. (2010) Interactions of human mismatch repair proteins MutSα and MutLα with proteins of the ATR-Chk1 pathway. *J. Biol. Chem.* **285**, 5974–5982

18. Yilmaz, S., Sancar, A., and Kemp, M. G. (2011) Multiple ATR-Chk1 pathway proteins preferentially associate with checkpoint-inducing DNA substrates. *PloS ONE* **6**, e22986

19. Lindsey-Boltz, L. A., Kemp, M. G., Reardon, J. T., DeRocco, V., Iyer, R. R., Modrich, P., and Sancar, A. (2014) Coupling of human DNA excision repair and the DNA damage checkpoint in a defined *in vitro* system. *J. Biol. Chem.* **289**, 5074–5082

20. Sertic, S., Pizzi, S., Cloney, R., Lehmann, A. R., Marini, F., Plevani, P., and Muzzi-Falconi, M. (2011) Human exonuclease 1 connects nucleotide excision repair (NER) processing with checkpoint activation in response to UV irradiation. *Proc. Natl. Acad. Sci. U.S.A.* **108**, 13647–13652

21. Willis, J., Patel, Y., Lentz, B. L., and Yan, S. (2013) APE2 is required for ATR-Chk1 checkpoint activation in response to oxidative stress. *Proc. Natl. Acad. Sci. U.S.A.* **110**, 10592–10597

22. Vrouwe, M. G., Pines, A., Overmeert, R. M., Hanada, K., and Mullenders, L. H. (2011) UV-induced photodegradations elicit ATR kinase-dependent signaling in non-cycling cells through nucleotide excision repair-dependent and -independent pathways. *J. Cell Sci.* **124**, 435–446

23. Ray, A., Blevins, C., Wani, G., and Wani, A. A. (2016) ATR- and ATM-mediated DNA damage response is dependent on excision repair assembly during G2 but not in S phase of cell cycle. *PloS ONE* **11**, e0159344

24. Derheimer, F. A., O’Hagan, H. M., Krueger, H. M., Hanasoge, S., Paulsen, M. T., and Ljungman, M. (2007) RPA and ATR link transcriptional stress to p53. *Proc. Natl. Acad. Sci. U.S.A.* **104**, 12778–12783

25. Ljungman, M. (2007) The transcription stress response. *Cell Cycle* **6**, 2252–2257

26. Lindsey-Boltz, L. A., and Sancar, A. (2007) RNA polymerase: the most specific damage recognition protein in cellular responses to DNA damage? *Proc. Natl. Acad. Sci. U.S.A.* **104**, 13213–13214

27. Wakisugi, M., Sasaki, T., Matsumoto, M., Nagaoka, M., Inoue, K., Inobe, M., Horibata, K., Tanaka, K., and Matsunaga, T. (2014) Nucleotide excision repair-dependent DNA double-strand break formation and ATM signaling activation in mammalian quiescent cells. *J. Biol. Chem.* **289**, 28730–28737

28. Kemp, M. G., and Sancar, A. (2016) ATR Kinase Inhibition protects non-cycling cells from the lethal effects of DNA damage and transcription bly during G1 but not in S phase of cell cycle. *Cell Cycle* **15**, 2879–2887

29. Cortez, D. (2003) Caffeine inhibits checkpoint responses without inhibiting the ataxia-telangiectasia-mutated (ATM) and ATM- and Rad3-related (ATR) protein kinases. *J. Biol. Chem.* **278**, 37139–37145

30. Ninghui, P., Park, P. K., Kim, Y., Vaziri, C., and Schreiber, S. L. (2001) ATR inhibition selectively sensitizes G1 checkpoint-deficient cells to lethal premature chromatin condensation. *Proc. Natl. Acad. Sci. U.S.A.* **98**, 9092–9097

31. Casper, A. M., Ninghui, P., Arlt, M. F., and Glover, T. W. (2002) ATR regulates fragile site stability. *Cell* **111**, 779–789

32. Ninghui, P., Park, P. K., Kim, Y. S., Desai, B. N., and Schreiber, S. L. (2002) ATR is not required for p53 activation but synergizes with p53 in the replication checkpoint. *J. Biol. Chem.* **277**, 4428–4434

33. Liu, S., Shiotani, B., Lahiri, M., Maréchal, A., Tse, A., Leung, C. C., Glover, J. N., Yang, X. H., and Zou, L. (2011) ATR autophosphorylation as a molecular switch for checkpoint activation. *Mol. Cell.* **43**, 192–202

34. Nan, E. A., Zhao, R., Glick, G. G., Bansbach, C. E., Friedman, D. B., and Cortez, D. (2011) Thr-1989 phosphorylation is a marker of active ataxia telangiectasia-mutated and Rad3-related (ATR) kinase. *J. Biol. Chem.* **286**, 28707–28714

35. Matsuoka, S., Ballif, B. A., Smogorzewska, A., McDonald, E. R., 3rd, Hurov, K. E., Luo, J., Bakalarski, C. E., Zhao, Z., Solimini, N., Lerenthal, Y., Shiloh, Y., Gly, S. P., and Elledge, S. J. (2007) ATM and ATR substrate analysis reveals extensive protein networks responsive to DNA damage. *Science* **316**, 1160–1166

36. Stokes, M. P., Rush, J., Macneill, J., Ren, J. M., Sprott, K., Nardone, J., Yang, V., Beausoleil, S. A., Gly, S. P., Livingstone, M., Zhang, H., Polakiewicz, R. D., and Comb, M. J. (2007) Profiling of UV-induced ATM/ATR signaling pathways. *Proc. Natl. Acad. Sci. U.S.A.* **104**, 19855–19860

37. Traven, A., and Heierhorst, J. (2005) SQ/TQ cluster domains: concentrated AT/ATR kinase phosphorylation site regions in DNA-damage response proteins. *BioEssays* **27**, 397–407

38. van Oosterwijk, M. F., Filon, R., Kalle, W. H., Mullenders, L. H., and van Zeeland, A. A. (1996) The sensitivity of human fibroblasts to N-acetoxy-2-acetylaminofluorene is determined by the extent of transcription-coupled repair, and/or their capability to counteract RNA synthesis inhibition. *Mol. Cell. Biol.* **16**, 3779–3788

39. van Oosterwijk, M. F., Filon, R., De Groot, A. J., van Zeeland, A. A., and Mullenders, L. H. (1998) Lack of transcription-coupled repair of acetylaminofluorene DNA adducts in human fibroblasts contrasts their efficient inhibition of transcription. *J. Biol. Chem.* **273**, 13599–13604

40. Tresini, M., Warmerdam, D. O., Kolovos, P., Snijder, L., Vrouwe, M. G., Demmers, J. A., van IJcken, W. F., Grosveld, F., Medema, R. H., Hoeijmakers, J. H., Mullenders, L. H., Vermeulen, W., and Marteiij, J. A. (2015) The core spliceosome as target and effector of non-canonical ATM signaling. *Nature* **523**, 53–58

41. Hess, M. T., Gunz, D., and Naegeli, H. (1996) A repair competition assay to assess recognition by human nucleotide excision repair. *Nucleic Acids Res.* **24**, 824–828
52. Giannattasio, M., Follonier, C., Tourrière, H., Puddu, F., Lazzaro, F., Pascero, P., Lopes, M., Plevani, P., and Muzi-Falconi, M. (2010) Exo1 competes with repair synthesis, converts NER intermediates to long ssDNA gaps, and promotes checkpoint activation. Mol. Cell 40, 50–62
53. Kemp, M. G., and Hu, J. (2017) Post excision events in human nucleotide excision repair. Photochem. Photobiol. 93, 178–191
54. Bensaude, O. (2011) Inhibiting eukaryotic transcription: which compound to choose? How to evaluate its activity? Transcription 2, 103–108
55. Titov, D. V., Gilman, B., He, Q. L., Bhat, S., Low, W. K., Dang, Y., Smeaton, M., Demain, A. L., Miller, P. S., Kugel, J. F., Goodrich, J. A., and Liu, J. O. (2011) XPB, a subunit of TFIIH, is a target of the natural product triptolide. Nat. Chem. Biol. 7, 182–188
56. van Gijssel, H. E., Mullenders, L. H., van Oosterwijk, M. F., and Meerman, F. (2011) XPB, a subunit of TFIIH, is a target of the natural product triptolide. Photochem. Photobiol. 95, 596–604
57. Klibanov, S. A., O’Hagan, H. M., and Ljungman, M. (2001) Accumulation of soluble and nucleolar-associated p53 proteins following cellular stress. J. Cell Sci. 114, 1867–1873
58. Ljungman, M., O’Hagan, H. M., and Paulsen, M. T. (2001) Induction of ssr15 and lvs382 modifications of p53 by blockage of transcription elongation. Oncogene 20, 5964–5971
59. O’Hagan, H. M., and Ljungman, M. (2004) Nuclear accumulation of p53 following inhibition of transcription is not due to diminished levels of MDM2. Oncogene 23, 5505–5512
60. Ljungman, M., and Lane, D. P. (2004) Transcription: guarding the genome by sensing DNA damage. Nat. Rev. Cancer. 4, 727–737
61. You, Z., Kong, L., and Newport, J. (2002) The role of single-stranded DNA and polymerase α in establishing the ATR, Hus1 DNA replication checkpoint. J. Biol. Chem. 277, 27088–27093
62. You, Z., and Elledge, S. J. (2003) Sensing DNA damage through ATRIP recognition of RPA-ssDNA complexes. Science 300, 1542–1548
63. Egly, J. M., and Coin, F. (2011) A history of TFIIH: two decades of molecular biology on a pivotal transcription/repair factor. DNA Repair 10, 714–721
64. Alekseev, S., Ayadi, M., Brino, L., Egly, J. M., Larsen, A. K., and Coin, F. (2014) A small molecule screen identifies an inhibitor of DNA repair inducing the degradation of TFIIH and the chromosensitization of tumor cells to platinum. Chem. Biol. 21, 398–407
65. Kwiatkowski, N., Zhang, T., Rahl, P. B., Abraham, B. J., Reddy, J., Ficarro, S. B., Dastur, A., Amzallag, A., Ramaswamy, S., Tesar, B., Jenkins, C. E., Hannon, N. M., McMillin, D., Sanda, T., Sim, T., et al. (2014) Targeting transcription regulation in cancer with a covalent CDK7 inhibitor. Nature 511, 616–620
66. Vendetti, F. P., Leibowitz, B. J., Barnes, J., Schamus, S., Kiesel, B. F., Abberbock, S., Conrads, T., Clump, D. A., Cadogan, E., O’Connor, M. J., Yu, J., Reumer, J. H., and Bakkenist, C. J. (2017) Pharmacologic ATM but not ATR kinase inhibition abrogates p21-dependent G1 arrest and promotes gastrointestinal syndrome after total body irradiation. Sci. Rep. 7, 41892
67. Coin, F., Oksenych, V., and Egly, J. M. (2007) Distinct roles for the XPB/p52 and XPD/p44 subcomplexes of TFIIH in damaged DNA opening during nucleotide excision repair. Mol. Cell 26, 245–256
68. He, Q. L., Titov, D. V., Li, J., Tan, M., Ye, Z., Zhao, Y., Romo, D., and Liu, J. O. (2015) Covalent modification of a cysteine residue in the XPB subunit of the general transcription factor TFIIH through single epoxide cleavage of the transcription inhibitor triptolide. Angew. Chem. Int. Ed. Engl. 54, 1859–1863
69. Dungrawala, H., Rose, K. L., Bhat, K. P., Mohini, K. N., Glick, G. C., Couch, F. B., and Cortez, D. (2015) The replication checkpoint prevents two types of fork collapse without regulating replosome stability. Mol. Cell 59, 998–1010
70. Song, J., Kemp, M. G., and Choi, J. H. (2017) Detection of the excised, damage-containing oligonucleotide products of nucleotide excision repair in human cells. Photochem. Photobiol. 93, 192–198
71. Choi, J. H., Kim, S. Y., Kim, S. K., Kemp, M. G., and Sancar, A. (2015) An integrated approach for analysis of the DNA damage response in mammalian cells: nucleotide excision repair, DNA damage checkpoint, and apoptosis. J. Biol. Chem. 290, 28812–28821
72. Choi, J. H., Gaddameedhi, S., Kim, S. Y., Hu, J., Kemp, M. G., and Sancar, A. (2014) Highly specific and sensitive method for measuring nucleotide excision repair kinetics of ultraviolet photoproducts in human cells. Nucleic Acids Res. 42, e29
73. Hu, J., Choi, J. H., Gaddameedhi, S., Kemp, M. G., Reardon, J. T., and Sancar, A. (2013) Nucleotide excision repair in human cells: fate of the excised oligonucleotide carrying DNA damage in vivo. J. Biol. Chem. 288, 20918–20926
74. Muñoz, M. J., Nieto Moreno, N., Giono, L. E., Cambindo Botto, A. E., Dujardin, G., Bastianello, G., Lavoro, S., Torres-Méndez, A., Menck, C. F., Blencowe, B. J., Irinia, M., Foiani, M., and Kornblüth, A. R. (2017) Major roles for pyrimidine dimers, nucleotide excision repair, and ATR in the alternative splicing response to UV irradiation. Cell. Rep. 18, 2868–2879
75. Yang, X., Boehm, I. S., Yang, X., Salehi-Ashtiani, K., Hao, T., Shen, Y., Lubonja, R., Thomas, S. R., Alkan, O., Bhindari, T., Green, T. M., Johannesen, C. M., Silver, S. J., Nguyen, C., Murray, R. R., et al. (2011) A public genome-scale lentiviral expression library of human ORFs. Nat. Methods 8, 659–661