DNA-PKcs and ATM epistatically suppress DNA end resection and hyperactivation of ATR-dependent G\textsubscript{2}-checkpoint in S-phase irradiated cells

Emil Mladenov, Xiaoxiang Fan, Katja Paul-Konietzko, Aashish Soni & George Iliakis

We previously reported that cells exposed to low doses of ionizing radiation (IR) in the G\textsubscript{2}-phase of the cell cycle activate a checkpoint that is epistatically regulated by ATM and ATR operating as an integrated module. In this module, ATR interphases exclusively with the cell cycle to implement the checkpoint, mainly using CHK1. The ATM/ATR module similarly regulates DNA end-resection at low IR-doses. Strikingly, at high IR-doses, the ATM/ATR coupling relaxes and each kinase exerts independent contributions to resection and the G\textsubscript{2}-checkpoint. DNA-PKcs links to the ATM/ATR module and defects cause hyper-resection and hyperactivation of G\textsubscript{2}-checkpoint at all doses examined. Surprisingly, our present report reveals that cells irradiated in S-phase utilize a different form of wiring between DNA-PKcs/ATM/ATR: The checkpoint activated in G\textsubscript{2}-phase is regulated exclusively by ATR/CHK1; similarly at high and low IR-doses. DNA end-resection supports ATR-activation, but inhibition of ATR leaves resection unchanged. DNA-PKcs and ATM link now epistatically to resection and their inhibition causes hyper-resection and ATR-dependent G\textsubscript{2}-checkpoint hyperactivation at all IR-doses. We propose that DNA-PKcs, ATM and ATR form a modular unit to regulate DSB processing with their crosstalk distinctly organized in S- and G\textsubscript{2}-phase, with strong dependence on DSB load only in G\textsubscript{2}-phase.

Cellular responses to DNA damage (DDR) and particularly to DNA double strand breaks (DSBs)

Despite clear functional differentiations among DNA-PKcs, ATM and ATR, their activation at the DSB and their contributions to DSB processing generate opportunities for crosstalk and functional integration. Indeed, regulatory connections between DNA-PKcs and ATM have been characterized and reviewed in classical non-homologous end-joining (c-NHEJ) and ATM is the apical kinase of global cellular responses initiated by DSBs. ATM phosphorylates the checkpoint kinase, CHK2, on multiple sites including T68 and initiates chromatin-based DDR signaling by phosphorylating the histone variant H2AX to generate γ-H2AX. ATM regulates DNA-end resection (to be referred to from now on, simply as “resection”) and promotes DSB repair by homologous recombination repair (HRR). It is also thought that ATM regulates the processing of a subset of DSBs via c-NHEJ. ATR is the apical DNA replication-stress-response kinase activated through ATRIP-mediated recruitment to tracts of ssDNA coated with replication protein A (RPA). Full activation of ATR requires several factors and is complex and context dependent. ATR phosphorylates and activates CHK1, which in S-phase inactivates the CDC25A and in G\textsubscript{2}-phase the CDC25C phosphatase to enforce the checkpoint response.

Despite clear functional differentiations among DNA-PKcs, ATM and ATR, their activation at the DSB and their contributions to DSB processing generate opportunities for crosstalk and functional integration. Indeed, regulatory connections between DNA-PKcs and ATM have been characterized and reviewed. Also ATM and...
ATR jointly regulate resection at DSBs\(^{24-26,34}\). Furthermore, AT cells activate an ATR-dependent G\(_2\)/M checkpoint\(^{35,36}\) and show responses suggesting crosstalk between DNA-PKcs and ATR\(^{37}\). There is also evidence that DNA-PKcs is phosphorylated by ATR in vitro\(^{27}\) and that it facilitates ATR-CHK1 signaling\(^{38}\). This brief outline suggests a modular integration of DNA-PKcs, ATM and ATR to sustain PIKK-signaling during DSB processing.

In a recent paper\(^{39}\), we presented experiments supporting such a modular integration of DNA-PKcs, ATM and ATR in the activation of the G\(_2\)-checkpoint and the regulation of resection. The design of this study had two unique characteristics: First, it examined kinase crosstalk as a function of DSB load in the cellular genome, i.e. as a function of the IR-dose administered. Second, it specifically and exclusively analyzed in the G\(_2\)-phase of the cell cycle the response of cells irradiated also in the G\(_2\)-phase of the cell cycle\(^{33}\). These experiments showed that at low IR-doses, ATM and ATR regulate epistatically as a module the G\(_2\)-checkpoint\(^{33}\). In this module, ATR is located at the output-node and interfaces with the cell-cycle through CHK1\(^{33}\). ATM/ATR module regulates epistatically also resection at low IR-doses. On the other hand, at high IR-doses, the modular coupling between ATM and ATR relaxes and the two kinases independently contribute to G\(_2\)-checkpoint and resection. Notably, DNA-PKcs appears to also integrate to the ATM/ATR module and DNA-PKcs defects cause hyper-resection and G\(_2\)-checkpoint hyperactivation\(^{35}\).

Here, we extend these studies to cells specifically exposed to IR during the S-phase of the cell cycle and analyze resection and checkpoint activation in the following G\(_2\)-phase. Surprisingly, we discover a different regulatory organization in the outputs of the DNA-PKcs/ATM/ATR module. The results provide important insights into the cell cycle regulation of DDR that have implications for DSB processing.

**Results**

**ATR fully controls the G\(_2\)-checkpoint in cells sustaining DSBs in the S-phase of the cell cycle.**

DSBs activate the G\(_2\)-checkpoint, which manifests as an arrest in G\(_2\)-phase and which is experienced first by cells already in G\(_2\). Cells sustaining DSBs in S-phase must first complete replication and enter G\(_2\)-phase to experience the checkpoint. The recently reported\(^{35}\) intriguing crosstalk between PIKKs in the regulation of the G\(_2\)-checkpoint for cells irradiated in G\(_2\)-phase, led us to inquire how this regulatory network operates when cells are irradiated in the S-phase of the cell cycle.

The G\(_2\)-checkpoint experienced by cells sustaining DNA damage in S-phase can be studied by single-parameter flow cytometry using propidium iodide (PI) staining. Exposure of actively proliferating hTert immortalized normal human fibroblasts 82-6 (82-6 hTert) to a low IR dose (2 Gy) impairs cell division without overly inhibiting ongoing DNA replication and causes a transient, time-dependent increase in the fraction of cells in G\(_2\)-phase (Fig. 1A). This increase reflects predominantly the arrest of cells irradiated in S-phase, as only progression of cells from S-phase can increase the fraction of cells in G\(_2\), above the levels of non-irradiated cells. Cells irradiated in G\(_2\)-phase, will initially remain blocked in G\(_2\), as they also experience an arrest\(^{39}\). Thus, checkpoint quantification for cells irradiated during S-phase is possible as increase in the G\(_2\) fraction over the levels of non-irradiated controls. Accumulation in G\(_2\) continuous in these cells for approximately 6h, dropping subsequently and approaching pre-irradiation values 15 h post irradiation (Fig. 1A).

In order to explore the contribution of ATR in the investigated response, we have utilized a specific small molecule inhibitor, VE-821, (to be referred as ATRi) (Fig. 1A)\(^{35}\). Notably, all stages of the G\(_2\)-checkpoint (initiation, maintenance and recovery) are abolished, as there is no evidence for notable increase in the fraction of cells in G\(_2\). The human lung carcinoma cell line, A549, (Fig. 1B) shows a similar response, as do also additional cell lines that are discussed below. We conclude that the G\(_2\)-checkpoint activated in S-phase cells exposed to low IR doses is under complete control of ATR and that this is a rather general response.

To confirm that the effects noted above specifically reflect the response of cells irradiated in S-phase, we pulse-labeled A549 cells with BrdU (10 \* 10\(\text{cells} \text{M}, 30\text{min}) just before IR. BrdU is incorporated into DNA and labels S-phase cells, allowing thus their specific follow-up as they progress to G\(_2\) by two-parameter flow cytometry (PI vs. BrdU) (Fig. S1A). Figure 1C shows that A549 cells, labelled and irradiated in S-phase, arrest in G\(_2\) with kinetics similar to those measured using only PI-staining (Fig. 1B). Notably, treatment with ATRi abrogates the G\(_2\)-checkpoint in BrdU-labelled, S-phase A549 cells exposed to 4 Gy (Figs 1C, S1B).

To confirm the full control of ATR on the G\(_2\)-checkpoint uncovered using small molecule inhibitors, we examined available genetic systems. GM847-ATRkd cells\(^{39}\) have integrated a Tet-On promoter controlled expression cassette, which upon administration of Doxycycline (DOX) causes the expression of an ATR fragment with inactivated kinase domain (ATRkd), (Figs 1D and S4), resulting in dominant negative inhibition of ATR activity\(^{33}\). Before expression of ATRkd, GM847-ATRkd cells irradiated in S-phase develop a G\(_2\)-arrest that is completely abrogated by ATRi (Fig. 1E). Notably, treatment with DOX causes complete abrogation of the G\(_2\)-arrest in cells irradiated in S-phase (Fig. 1F), and treatment with ATRi has no additional effect.

The dominant role of ATR in the activation of the G\(_2\)-checkpoint in cells irradiated during S-phase, raises questions regarding the contribution of a key component of the G\(_2\)-checkpoint in cells exposed to low IR doses in G\(_2\)-phase, the ATM protein kinase\(^{33}\). ATM deficient AT5BIVA cells irradiated in S-phase with 4 Gy, show nearly normal activation of the G\(_2\)-checkpoint followed by a markedly prolonged arrest in G\(_2\) (Fig. 2A) as compared to wild-type cells (Fig. 1A). This response remains unchanged, as expected, after incubation with a specific ATM inhibitor (KU55933, to be referred to as ATMi). BrdU-labeling of AT5BIVA cells confirms that the observed effect reflects the response of S-irradiated cells (Figs 2B, S1C,D). This divergent contribution of ATM to the G\(_2\)-checkpoint between cells irradiated in S- and G\(_2\)-phase has generated in the past apparently contradictory results that were lively debated, until Kastan et al.\(^{40}\) explained these contradictions as the radically different cell cycle dependent ATM contribution shown here.

Notably, the sustained arrest in G\(_2\) of irradiated ATM-deficient S-phase cells depends on ATR, and is largely reversed after treatment with ATRi (Fig. 2A,B). S-phase-irradiated 82-6 hTert and A549 cells develop a very similar response after treatment with ATMi (Fig. 1A,B)\(^{35}\). An ATR contribution to this response was not considered.
in earlier studies but has been seen without performing a cell cycle specific analysis in our previous report. The contribution of ATR to the G2-checkpoint has been recently investigated by others, but interpreted as contributory to ATM function and rather specific for high-LET IR-induced DNA damage. Here, we show for the first time that ATR is entirely and exclusively responsible for the G2-checkpoint induced by low LET IR in cells exposed to low IR doses; both in a wild type, as well as in an ATM mutant genetic background.

The G2-checkpoint is implemented by suppressing the activity of Cdc25C, a phosphatase that activates CDK1 and drives cells into mitosis. ATM and ATR convey inhibitory signals to CDC25C through CHK2 and CHK1,
respectively4. The dominant role noted above for ATR in the checkpoint response suggested CHK1 as the bridge to the cell cycle machinery. In line with this expectation, UCN-01, an inhibitor of CHK1 (to be referred to as CHK1i), causes a nearly complete suppression of the G2-checkpoint in S-phase irradiated, 82-6 hTert and A549 cells (Fig. 1A,B), as well as in non-induced GM847-ATRkd cells (Fig. 1E). A nearly complete abrogation of the G2-checkpoint is also observed in S-phase AT5Biva cells after treatment with CHK1i (Fig. 2A). These results demonstrate that ATR is regulating the G2-checkpoint in this setting by activating, practically exclusively, CHK142. We note that for cells irradiated in G2 phase the G2-checkpoint only partly depends on CHK133.

Since ATM inactivation strengthened/prolonged the checkpoint, we examined whether CHK2 somehow contributes to this response. Treatment of 82-6 hTert or A549 cells with a CHK2 inhibitor (Chk2 inhibitor II, BML-277, to be referred to as CHK2i) fails to generate measurable effects on the induction and recovery of the checkpoint (Fig. 1A,B), as already reported before16,24,42–44. The same inhibitor is also ineffective, as expected, in AT5Biva cells (Fig. 2A).

Collectively, our experiments thus far suggest that activation of the G2-checkpoint in cells exposed to low doses in S-phase, exclusively requires ATR that operates through CHK1 and that ATM is not contributing to checkpoint activation. On the contrary, ATM defects have in some cell lines no consequences in the activation of the checkpoint and actually often cause its prolongation. Thus, ATM suppresses in S-phase irradiated cells a G2-checkpoint hyperactivation that remains entirely and exclusively ATR-dependent41,42, and which develops independently of CHK2. This is diametrically different from the regulation established for cells irradiated in the G2–phase, where both ATM and ATR are equally required for G2-checkpoint activation33.

Because our previous work showed that the function of the ATM/ATR module is strongly IR-dose-dependent in G2–phase irradiated cells33, we also studied the effect of higher IR doses on the G2–checkpoint in S-phase irradiated cells. 82-6 hTert cells exposed to 10 Gy initiate a strong checkpoint that blocks over 40% of the cells in G2–phase for the period of observation (Fig. 2C). Remarkably, treatment with ATRi eliminates almost fully the checkpoint response and CHK1i treatment has a similar effect (Fig. 2C)

Exposure of AT5Biva cells to 10 Gy (Fig. 2D) also generates a substantially stronger arrest in G2–phase than that measured in ATM proficient cells. Strikingly, this checkpoint remains also dependent on ATR and treatment with ATRi practically abrogates the response. The same response is also confirmed in another AT cell line, AT hTert, after exposure to 10 Gy (Fig. S1E). We conclude that in contrast to G2–phase irradiated cells, S-phase irradiated cells show no IR-dose-dependent modulation in the regulation of the G2–checkpoint; which is prolonged

Figure 2. ATM deficiency results in hyperactivation of the G2–checkpoint when cells are irradiated in S-phase. (A) As in Fig. 1A for ATM deficient, AT5Biva cells exposed to 4 Gy and treated with indicated inhibitors. (B) As in Fig. 1C for AT5Biva cells exposed to 4 Gy and treated with indicated inhibitors. (C) As in Fig. 1A for 82-6 hTert cells exposed to 10 Gy and treated with the indicated inhibitors. (D) As in (C) for AT5Biva cells. All data points represent means and standard deviations (only visible when are larger than the symbol) estimated from three independent experiments.
with increasing IR dose but remains always territory of ATR. Because in G2–irradiated cells, DNA-PKcs affected the checkpoint in ways that resemble the effect we observe here for ATM, we inquired on the checkpoint response of DNA-PKcs deficient S-phase cells.

**Figure 3.** DNA-PKcs deficiency results in hyperactivation of the G2-checkpoint in cells irradiated during the S-phase of the cell cycle. (A) As in Fig. 1A for the DNA-PKcs deficient cell line, M059J, exposed to 2 Gy and treated with indicated inhibitors. (B) As in (A) for DNA-PKcs proficient M059K cells. (C) As in Fig. 1A for 82-6 hTert cells exposed to 2 Gy and treated with the indicated inhibitors. (D) As in (A) for M059J cells, irradiated with 10 Gy and treated with the indicated inhibitors. All data points represent the means and standard deviations (only visible when are larger than the symbol) estimated from three independent experiments. (E) Western blot analysis of pCHK1-S345, a marker of ATR activation, after treatment with the indicated inhibitors of parental A549 (A549<sup>wt</sup>) and DNA-PKcs knock-out, A549 (A549<sup>DNA-PKcs<sup>−/−</sup></sup>) cells, enriched in S-phase by a single thymidine block, exposed to 10 Gy of IR and collected for analysis 9 h after irradiation. The levels of DNA-PKcs confirm the successful knockout, while the levels of CHK1 and KU70 serve as loading controls. (F) As in (E) for ATM deficient AT hTert cells analyzed 3 h after exposure to 10 Gy IR. CHK1 and RPA32 serve as loading controls. Original non-cropped images of the scanned western blot membranes are shown in Figs S4B,C.

**Defects in DNA-PKcs hyperactivate, epistatically to ATM, an ATR-dependent checkpoint in cells sustaining DSBs in S-phase.** Exposure in S-phase of DNA-PKcs deficient M059J (Fig. 3A) and HCT116 DNA-PKcs<sup>−/−</sup> deficient cells (Fig. S2A) to 2 Gy causes a markedly stronger arrest in G2 as compared to M059K cells, the wild-type counterpart of M059J (Fig. 3B), or wild-type HCT116 (Fig. S2B) cells. Notably, here again, ATRi abrogates the arrest (Figs 3A and S2A), as does also CHK1i. Interestingly, caffeine, a non-specific...
inhibitor of ATR, causes complete suppression of the checkpoint (Fig. 3A,B, as well as S2A and S2B). When DNA-PKcs proficient 82-6 hTert (Fig. 3C) or A549 (Fig. S2C) cells are irradiated during S-phase in the presence of the specific DNA-PKcs inhibitor NU7441 (DNA-PKcs), they also show prolonged activation of the G2-checkpoint with only slight signs of recovery in 82-6 hTert cells after 15 h. Thus, genetic ablation of DNA-PKcs and treatment with a specific DNA-PKcs inhibitor generate similar effects.

Since the hyperactivation of G2-checkpoint under conditions of suppressed DNA-PKcs activity is very similar to the response of ATM deficient cells, we investigated the effect of ATM inhibition in a DNA-PKcs deficient background. Strikingly, inhibition of ATM in M059J cells has no detectable additional effect on the checkpoint measured in the absence of the inhibitor and causes only a small delay in the activation of the checkpoint in HCT116 DNA-PKcs−/− cells (Figs 3A and S2A). Also, in DNA-PKcs proficient 82-6 hTert (Fig. 3C) cells, combined treatment with DNA-PKcsi and ATMi generates a response similar to single inhibitor treatment. Under all above described conditions, the hyperactivated G2-checkpoint relies almost fully and exclusively on ATR and CHK1 and is abrogated by ATRi or CHK1i (Figs 3A and S2C). Notably, in 82-6 hTert or A549 cells, combined treatment with DNA-PKcsi + ATMi + ATRi practically eliminates the checkpoint response (Figs 3C and S2C).

The practically complete dependence on ATRi and CHK1i of the checkpoint activated in G2 in S-irradiated DNA-PKcs deficient cells is also observed after exposure to 10 Gy, although here treatment with ATMi causes a slightly earlier initiation of the G2-checkpoint (Figs 3D and S2D). Collectively, the results of this section demonstrate that DNA-PKcs exerts pronounced inputs into the G2-checkpoint, subject to relatively small qualitative but not quantitative adjustments with IR-dose. Thus, DNA-PKcs activity is required to suppress checkpoint hyperactivation, or, in an alternative view, checkpoint recovery in the range of doses examined. In this function, DNA-PKcs operates epistatically with ATM, suggesting that in S-phase irradiated cells the two kinases function as a module to regulate processes that prevent the hyperactivation (or recovery) of an ATR-dependent checkpoint.

To directly demonstrate hyperactivation of ATR signaling under conditions of DNA-PKcs and ATM deficiency, we analyzed phosphorylation of CHK1 at Serine 345 (pCHK1-S345). This event is ATR-specific and is commonly used as surrogate marker of ATR activation. To generate results relevant to S-phase irradiated cells we employed chemical synchronization protocols that generate populations enriched in S-phase cells, which we exposed to IR. We selected A549 cells for these experiments because we had synchronization protocols, as well as a DNA-PKcs knockout mutant generated using CRISPR/Cas9 technology. Figure S2E (upper panels) shows the cell cycle distribution of irradiated (10 Gy) S-phase-enriched cells, while the lower panels their distribution 9 h later, for wild-type (left panels) and DNA-PKcs deficient (right panels) cells. Analysis of pCHK1-S345 in wild-type cells (Fig. 3E, left lanes) shows the expected increase in signal documenting ATR activation that is further potentiated by inhibition of ATM. Formation of pCHK1-S345 is enhanced in DNA-PKcs-deficient A549 cells and this effect is further amplified by inhibition of ATM (Fig. 3E, right lanes). Figure S2F shows S-phase enriched populations generated in AT hTert cells and exposed to 10 Gy. Analysis carried out 3 h later when a significant proportion of cells reach G2-phase shows strong production of pCHK1-S345 that is almost completely abrogated following treatment with ATRi, while it remains robust after treatment with DNA-PKcsi (Fig. 3F). Quantitative comparisons between the results obtained with different cell lines is confounded by differences in the initial distribution throughout the cell cycle and the subsequent progression to G2-phase of the irradiated cells. Despite this limitation, the results demonstrate ATR hyperactivation under conditions of ATM and DNA-PKcs deficiency.

The mechanistic reliance of the G2-checkpoint on ATR in S-phase irradiated cells, places at the forefront the mechanism of ATR activation which has resorption at its center. Therefore, in the following sections, we analyze in detail this endpoint under conditions similar to those employed in the checkpoint experiments.

**ATR exerts limited control on G2-phase-resection in S-phase irradiated cells.** We employed immunofluorescence (IF) to evaluate DNA end-resection by measuring RPA70 (the largest subunit of the RPA complex) retention at chromatin as a function of time after irradiation. For a specific analysis in G2-phase for cells irradiated in the S-phase, we labeled cells with EdU (30 min) before IR exposure and quantitated RPA70 signal in EdU positive cells (Figs 3A, B, C) and EdU negative cells (Figs S2A, B, C) in different times thereafter. Figure 4A outlines this form of analysis and the gates adopted to measure RPA70 signal in such cells (gate marked in red). Figure 4B shows representative IF images of the analyzed population of 82-6 hTert cells, treated or not with ATRi, exposed to 0 or 2 Gy and processed at different times thereafter. Figure 4C shows quantification at the indicated times of the integral RPA70 signal in this cohort of cells (100–150 EdU+, G2-phase cells analyzed for each data point). Results of irradiated cells are plotted together with results obtained in an identical manner from non-irradiated cells treated similarly and measured within the same gates (background signal) (Fig. 4C). We use integral RPA signal intensity per cell as a parameter in the IF analysis, arguing that it reflects the level of ATR activation better than scoring of individual foci. In addition, it allows comparison with integral signal analysis carried out to measure resection using flow cytometry (see below).

It is evident (Fig. 4C,D) that at 3 and 6 h after exposure to IR a significant increase in RPA70 signal over background is observed in EdU−, G2-phase cells, suggesting resection at DSBs that sustains the G2-checkpoint (Fig. 1A). ATRi treatment leaves in irradiated cells RPA70 signal practically unchanged (Fig. 4C). Notably, in non-irradiated cells treated with ATRi, RPA70 signal is markedly elevated (Fig. 4C). This increase likely reflects binding of RPA complex to ssDNA persisting in cells from the S-phase that have entered G2-phase; it may be generated as a result of problems encountered during DNA replication and which are enhanced after treatment with ATRi. Indeed, it is known that even under normal replication conditions, late replicating loci in heterochromatin and loci with fragile sites and repetitive elements, suffer replication fork stalling and may complete replication in G2-phase. The said effects are exaggerated after treatment with ATRi and likely cause the increase in RPA70 signal observed in non-irradiated cells. If we consider this increased signal as the “legitimate” background of the corresponding
irradiated samples and subtract it, the net RPA70 signal increase shown in Fig. 4D is obtained. Although these results appear to show a signal reduction in ATRi treated cells after IR exposure, the effect fails to reach statistical significance.

To study resection at higher IR doses, we employed a quantitative flow cytometry-based method\(^3\)\(^3\)\(^1\)\(^1\). Cells are incubated, with EdU to label cells in S-phase and resection is measured by detecting RPA70 in EdU\(^{+}\), G2-phase cells, identified by co-staining of DNA with propidium iodide (PI). The upper panels in Fig. 5A show as an example raw data as dot plots and the gates used to quantitate RPA, EdU and PI signals using results obtained from independent experiments. A student t-test was used for statistical analysis and the individual p-values are indicated.
Figure 5. ATR plays no role in the regulation of DNA end-resection in cells irradiated with high IR doses during S-phase when analyzed in the subsequent G2-phase of the cell cycle. (A) Summary of the three-parametric flow cytometry analysis utilized to quantitate DNA end-resection in cells exposed to high IR doses in S-phase. Plots illustrating RPA70 vs. PI signals (upper panels), or EdU vs. PI signals (middle panels). On the middle panels the gates applied to quantify DNA end resection in G2-cells irradiated in S-phase (EdU+) are also indicated. The lower part of the figure shows histograms of RPA70 signal evaluated in non-irradiated (0 Gy, green) or irradiated (10 Gy, red) EdU+, G2-phase cells. (B) Histograms of RPA70 signal intensity measured in EdU+, G2-phase, 82-6 hTert cells exposed to 5, 10 and 15 Gy. (C) Upper panels: Histograms of the intensity of RPA70 signal as a function of time in EdU+, G2-phase, 82-6 hTert cells irradiated with 10 Gy in the presence or absence of ATR inhibitor. Lower panels: Quantitative analysis of three independent experiments, showing the arithmetic means for RPA70 signal intensity. The error bars represent standard deviations. A student t-test was utilized for statistical analysis. The individual p-values are indicated on top. (D) Left panel: Histograms showing the RPA70 intensity, 3 h after exposure of GM847-ATRkd cells to 10 Gy after pretreatment (+DOX) or not (−DOX) with doxycycline. Right panel: Quantitative analysis of two independent experiments, showing the arithmetic means of RPA70 intensity. The error bars represents standard deviations.
3 h after irradiation of 82-6 hTert cells with 0 or 10 Gy. The histograms in the lower panel of Fig. 5A show intensity distribution of RPA70 signal in the defined gates in irradiated and non-irradiated cells. The robust RPA70 signal increase observed in cells exposed to 10 Gy indicates extensive resection at DSBs. Figure 5B shows that IR-induced resection can be conveniently quantitated in a range of doses between 5 and 15 Gy using this method.

We employed the above protocols to analyze resection as a function of time after exposure of 82-6 hTert cells to 10 Gy. The results in Fig. 5C show robust resection at 3 h, 6 h and 9 h after IR, in line with the activation of ATR required to sustain the checkpoint. Notably, and in line with the IF analyzes, inhibition of ATR has little effect on the overall IR-induced resection (Fig. 5C, upper panels). However, here again slight increase in RPA70 signal is measured in non-irradiated cells. The lower panels in the Fig. 5C summarize the results of three experiments using as parameter the arithmetic mean of RPA70 signal distribution. Here again, ATRi has no effect on net signal intensity, while non-irradiated controls show higher background signal. Plotting of the same results using the median of RPA70 signal distribution as a parameter instead, leads to similar conclusions (Fig. S3A).

A similar analysis in non-induced GM847-ATRkd cells exposed to 10 Gy also shows robust resection at 3 h in G2 for S-phase irradiated cells (EdU+) (Fig. 5D, left panel). Inhibition of ATR by administration of DOX causes no detectable suppression of IR-induced resection but causes also a marked increase in the RPA70 signal in non-irradiated cells (Fig. 5D, left panel). The bar graphs in Fig. 5D (right panels) show the compiled results of two experiments. Similar trends are also observed in AS49 cells, although the double peaks in the RPA70 signal distributions, particularly in the non-irradiated samples, complicate the quantitative analysis of the results (Fig. S3B).

Collectively, the above results show robust resection in G2-phase in cells irradiated during the S-phase of the cell cycle, in line with the ATR dependent checkpoint documented in the previous section. ATRi treatment failed to change the IR-induced RPA70 signal, but increased the background signal in non-irradiated cells. As a result the background-corrected net increase in RPA70 signal in irradiated ATRi-treated cells could be interpreted as showing partial suppression of resection by ATRi. While final answer for this ambiguity will require alternative methods of resection analysis in cells irradiated in S-phase (see below), we suggest that for cells irradiated in S-phase, ATR exploits resection for activation and implementation of the G2-checkpoint, but that it exerts itself only limited regulation on resection. This is in contrast to the strong regulatory inputs ATR has in cells irradiated in G2-phase, where ATR inhibition suppresses resection: completely at low and partially at high IR-doses33.

Inhibition of CHK1 leaves G2-phase resection unchanged in S-phase irradiated 82-6 hTert and AS49 cells (Fig. 6A,B). As for ATRi, albeit less pronounced, CHK1i increases RPA70 signal intensity in non-irradiated cells generating the same issues in the analysis of CHK1i effect on resection as discussed above for ATRi. Since CHK1i failed to suppress resection in G2-irradiated cells33, we infer that the present set of data also suggests no effect of CHK1 on IR-induced resection. If this is indeed true it means that the effect of ATR/CHK1 inhibitors on non-irradiated cells may not be fully transmitted to irradiated cells. We present next results that further support this notion.

To validate the above effects on resection with an independent method, we grew 82-6 hTert cells for 24 h in the presence of BrdU, labeled them with EdU for 30 min and exposed them to IR. At different times after IR, we analyzed cells by flow cytometry to measure, in a cell cycle dependent manner, ssDNA by staining non-denatured DNA with BrdU-specific antibody. The overall analysis is very similar to that described for RPA, except that BrdU signal on ssDNA is instead detected to assess resection. Figure 6C (left panels) shows that exposure to 10 Gy robustly increases BrdU signal in EdU+ cells analyzed in G2-phase at 3, 6, and 9 h post-irradiation, confirming robust resection that sustains the checkpoint. Treatment with ATRi leaves here again the overall BrdU signal intensity practically unchanged (Fig. 6C, right panels). Non-irradiated samples show a smaller increase in BrdU signal after treatment with ATRi as compared to the RPA70 measurements presented above. Furthermore, background subtracted analysis of such distributions from three experiments (Fig. 6D) supports the notion that ATRi does not affect resection in S-phase irradiated cells. Similar conclusions are drawn when the results are analyzed using the median of the BrdU signal distribution instead (Fig. S3C).

DNA-PKcs and ATM defects lead to persistent resection in an epistatic manner in subsequent G2-phase in S-phase irradiated cells. When AT5BIVA cells are irradiated in S-phase, resection measured in G2-phase is overall reduced as compared to ATM proficient cells, but persistent over time (Fig. 7A) explaining the persistent checkpoint developing under these conditions (Figs 2 and S1E). Treatment of these cells with ATRi has only a small effect on resection (Fig. 7B). Notably, ATM deficiency partly rescues the effect of ATRi on DNA replication in non-irradiated cells, mitigating thus some of the above outlined complications in the interpretation of the results obtained. Similar trends are also seen in AT hTert cells, before and after treatment with ATRi (Fig. S3D). When ATM proficient, 82-6 hTert cells are treated with ATMi, marked and persistent resection develops (Fig. 7C). Thus, in S-phase irradiated cells, ATM suppresses late hyperresection.

DNA-PKcs deficient M059J cells exposed to 10 Gy in the S-phase show in G2-phase enhanced resection that persists for longer times (Fig. 7D, left panels) as compared to M059K cells (Fig. 7D, right panels). Treatment of these cells with ATMi fails to significantly increase resection, suggesting that for this endpoint and under the conditions employed ATM and DNA-PKcs work epistatically. ATRi exerts no effect on resection in this genetic background as well, and in this particular cell line it only has a relatively small effect on RPA70 signal intensity in non-irradiated cells (Fig. 7E). Similar conclusions can be drawn from results obtained in DNA-PKcs deficient, HCT116 cells (Fig. S3E). Also, treatment with CHK1i in a DNA-PKcs deficient background does not affect the DNA end resection (Fig. S3F). Thus, for cells irradiated in S-phase, both ATM and DNA-PKcs function in an epistatic manner to suppress resection at DSBs during their processing in G2-phase, and this response mirrors closely their effects on the G2-checkpoint.
Discussion

**ATR exclusively activates the G\(_2\)-checkpoint in cells irradiated in S-phase.** While the individual functions of ATM and ATR are relatively well-characterized, aspects of their functional cooperation and cross-talk continue to emerge. We recently reported intriguing functional interactions between ATM and ATR in the regulation of the G\(_2\)-checkpoint in cells irradiated in G\(_2\)-phase, showing striking mechanistic adaptations with increasing load of DSBs in the genome. Specifically, we discovered a complete functional coupling between ATM and ATR in the regulation of the G\(_2\)-checkpoint in cells with low numbers of DSBs induced by exposure to low doses of IR (see model description in Fig. 8). Under these conditions and phase of the cell cycle, ATM and ATR are equally required for the activation of the checkpoint, and inhibition of either kinase completely abrogates this activation. We use here the term epistasis to describe this relationship, but we caution the reader that there are alternative definitions of this term. This previously unreported functional coupling between ATM and ATR appears directional, connecting to the cell cycle machinery primarily through ATR to CHK1 signaling (Fig. 8).

Strikingly, this tight modular interdependence relaxes when cells are exposed to high IR doses. Now, independent outputs from ATM and ATR activate the G\(_2\)-checkpoint and inhibition of both kinases is required to fully suppress this activation (Fig. 8). Either way, G\(_2\)-checkpoint activation remains the exclusive territory of the ATM/ATR duo. This dose-dependent modular integration between ATM and ATR raises ATR relevance to the same level as ATM, and suggests that views of an ATM-centered regulatory organization of the G\(_2\)-checkpoint need revision.

The results presented here, while confirming a modular cooperation between DNA-PKcs, ATM and ATR in the overall regulation of the G\(_2\)-checkpoint, they also uncover striking and unexpected differences in the specific wiring of the module when cells are irradiated in S-phase. Thus, in stark contrast to the epistatic and...
dose-dependent regulation of the checkpoint by both ATM and ATR in cells irradiated in G2-phase, in cells irradiated in S-phase the G2-checkpoint depends entirely on ATR that functions practically exclusively through CHK1; moreover, the mechanism of this regulation is not fundamentally altered with increasing IR-dose (see model description in Fig. 8).

This observation is highly significant mechanistically, as it suggests that with the transition of a cell from S- to G2-phase, regulatory and possibly also structural changes in genome organization occur that profoundly modify the mechanistic underpinnings of the G2-checkpoint. It is evident therefore, that only strictly cell cycle
phase-specific analysis of DDR will generate “pure” results for sound mechanistic interpretations advancing our understanding of its molecular underpinnings. When the experimental data “mix” the diametrically different responses of cells irradiated in S- and G2-phase, misleading conclusions are likely to be drawn, or apparently contradictory results may be obtained in different cell systems.

In cells irradiated in S-phase, ATM and DNA-PKcs epistatically suppress G2-checkpoint hyper-activation. Not only is in cells irradiated in the S-phase ATR the sole controller of the G2-checkpoint, but in addition ATM, the intimate partner of ATR for cells irradiated in G2-phase, assumes now a diametrically different contribution to the checkpoint. In this setting, ATM is not required in any way for the activation of the checkpoint, but supports instead its recovery; as a consequence ATM deficiency causes G2-checkpoint prolongation and hyperactivation. This is similar to the function of DNA-PKcs in cells irradiated in G2-phase33. Strikingly, in cells irradiated in S-phase, the functions of ATM and DNA-PKcs are epistatic in the sense that inhibition of either kinase generates equivalent effects and inhibition of both kinases produces no additivity. We conclude that both kinases epistatically regulate the same process related to the recovery from the checkpoint, and that defects cause ATR and checkpoint hyperactivation (Fig. 8).

Reduced control of resection by ATR in G2-phase in cells irradiated in S-phase. Resection at a DSBs is an important determinant of DSB-repair pathway choice favoring HRR, SSA, and alt-EJ10,54,55. The fundamental requirement for resection in ATR activation links thus a key aspect of checkpoint control to the specific forms of DSB processing that require resection56. Indeed, c-NHEJ is not linked to ATR signaling and the G2-checkpoint57–60; an observation fully in line with c-NHEJ’s fast kinetics. The fact that resection accompanies the activation of the G2-checkpoint in cells irradiated in S-phase provides mechanistic explanation for the involvement of ATR. Strikingly though, in contrast to G2-irradiated cells, where ATR is not only passively responding to resection, but is also actively regulating it33, in cells irradiated in S-phase ATR only responds to resection but has only limited, if any, control in the process. As for cells irradiated in G2, CHK1 is not playing a role in the regulation of resection. This immediately raises the question as to whether the ATR mediated regulatory phosphorylation of CtIP61 occurs differently in cells irradiated in S-phase versus cells irradiated in the G2-phase of the cell cycle56.

We are aware that the resection measurements in S-phase-irradiated cells have some ambiguity, as a consequence of ATRi effects on DNA replication that increase RPA signal in non-irradiated S-phase cells when they reach G2-phase. However, the lack of any notable reduction in the RPA70 signal in the irradiated population by two independent methods is in line with this conclusion.
DNA-PKcs and ATM function in an epistatic manner to suppress hyperresection. The last significant contribution of the results reported here is the epistatic suppression of hyperresection by DNA-PKcs and ATM in cells irradiated in S-phase when they reach G2-phase. This persistent and epistatic hyperresection observed in the absence of DNA-PKcs and ATM activities explains the above discussed hyperactivation of the checkpoint as hyperactivation of ATR. These results tightly link resection to ATR activation and the mounting of ATR-dependent checkpoints (Fig. 8). There is evidence for links between DNA-PKcs and ATM with the former exerting strong negative regulation on ATM through phosphorylation at multiple sites. DNA-PK also stimulates the MRN complex and CtIP for efficient endonucleolytic processing of DNA ends under physiological conditions. How these interesting observations feed into the cell cycle dependent regulation in the crosstalk between members of the PIKK kinase family (DNA-PKcs, ATM and ATR) will be subject of future investigations.

However, whereas DNA-PKcs deficiency causes increased levels of resection that persist, ATM deficiency leads to reduced initial levels of resection that also persist. This points to qualitatively similar inputs between DNA-PKcs and ATM in the regulation of resection and G2-checkpoint activation that have different quantitative manifestations.

A model integrating the activities DNA-PKcs, ATM and ATR in S- and G2-phase. In aggregate, we describe cell cycle specific contributions of PIKK family members (DNA-PKcs, ATM and ATR) to the regulation of G2-checkpoint and DNA end-resection that are consistent with crosstalk and possibly cross-regulation among these kinases. Therefore, we postulate their modular integration. Our working hypothesis is summarized in the model outlined in Fig. 8. The first opportunity of functional integration among these kinases derives from their ability to be recruited to DSBs at different stages in their processing, both for cells irradiated in S- and in G2-phase. Indeed, for DSBs that will eventually undergo resection, all three kinases will be successively recruited, and will contribute, in varying ways as we saw above, to the regulation of this resection process. The modular integration model proposed here, specifically applies to this subset of DSBs and it is also this subset of DSBs, which according to our results, has links to the checkpoint response.

While the initial recognition and binding to DSBs will likely start from DNA-PK, recruitment and activation of ATM and subsequently of ATR will ensue. Such sequential functional integration for the processing of an individual DSB explains why the substrates of DNA-PKcs, ATM and ATR are largely overlapping: to ensure sustained DDR signaling as DSBs transit through the different processing steps.

While this modular integration in the functions of DNA-PKcs/ATM/ATR ensure optimized processing of DSBs (acting as a form of optimization module), it is functionally important that recruitment of any one of these kinases to the DSB can also occur independently of the presence of the other kinases, explaining why frequently depletion of one kinase does not eliminate the activation of the others. Thus, the function of DNA-PKcs in c-NHEJ does not require ATM or ATR, and ATM is fully activated in the absence of ATR activity.

Furthermore, our results suggest coupled functions for essential aspects of DDR such as the regulation of checkpoint activation and recovery, as well as the regulation of resection. Our results indeed point to a tight linkage between the former and the latter as already reported for the yeast. Notably, we could demonstrate that these coupled functions are wired in the different manner in cells sustaining DSBs in S-phase and in G2-phase of the cell cycle (Fig. 8). Thus, in cells sustaining DSBs in G2-phase, ATM and ATR function as a tightly and epistatically operating pair that has under its full control resection and G2-checkpoint activation. Functional uncoupling requires high loads of DSBs in the genome and may reflect a means cells employ to better cope with high loads of DSBs. In this sub-module, ATR occupies the exit node with reference to cell cycle regulation, mediated by Chk1 and possibly an additional unidentified kinase. Here, the third member of the family, DNA-PKcs, regulates resection and checkpoint recovery (Fig. 8).

On the other hand, in cells irradiated in S-phase the checkpoint is regulated entirely by ATR that is activated by resection, controlled by DNA-PKcs/ATM that now operate as a pair epistatically. The exact role of ATR in the regulation of this resection will require additional investigations. Connections between resection and checkpoint affecting recovery from the checkpoint and a defining role of KU in this response have been reported for Saccharomyces cells after inducing a single DSB. The characterization of the molecular underpinnings of DNA-PKcs, ATM/ATR interactions is a promising area for future mechanistic investigations.

Methods
Cell culture and irradiation. All cell lines employed were grown in 10–20% fetal bovine serum (FBS)-supplemented cell culture media, at 37 °C in an atmosphere of 5% CO2 in air. DNA-PKcs knock-out and parental A549 cells, parental HCT116 and DNA-PKcs knock-out HCT116 cells were maintained in McCoy’s 5 A medium; ATM deficient cell lines, AT5BIV A and AT hTert (GM2052), as well as the conditionally ATR-deficient, GM847-ATRkd, and 82-6 hTert cells, were grown in Dulbecco’s modified Eagle’s medium (D-MEM). If not stated otherwise, cells were exposed to IR at 37 °C using a 320 kV X-ray machine with a 1.65 Al filter (GE-Healthcare). The dose rate at 500 mm distance from the source was 2.7 Gy/min.

Synchronization of cells in S-phase. In order to enrich cells in S-phase of the cell cycle a single thymidine block was applied. Cells were treated with 2 mM thymidine for 18 h and were released by single washing with PBS and immediate incubation in fresh growth medium. Cells were allowed to progress through the S-phase for 3 h and were treated with PIKK inhibitors 1 h prior to irradiation. Cells were collected at 3 or 9 h after irradiation.

Generation of DNA-PKcs knock-out A549 cell line. CRISPR/Cas9 technology was utilized for the generation of DNA-PKcs knock-out cells. For this purpose, A549 cells (A549-wt) were transfected with Cas9
expression plasmids, which were constructed to express a DNA-PKcs exon specific gRNA. The plasmid also co-expresses a GFP protein, which was utilized to select transfected cells by cell sorting for further analysis. Individual A549 clones were isolated and DNA-PKcs expression was monitored by western blotting (Fig. 3E). The isolated DNA-PKcs negative clones were further confirmed by analyzing their sensitivity to IR using clonogenic assays (data not shown). A pool of the following gRNAs was applied to generate DNA-PKcs knockout, A549 cells: gRNA1-AAAAAGCCTACACTCAGGGAC, gRNA2- CAGCAAGTGCACCTGTGTA, gRNA3- ATCCGACTTT GGGCATGCGT, gRNA4- GATCAGCAGCCAGTCCTCA,gRNA5- CAGACATCTGAAACATTTA.

Treatment of cells with kinase inhibitors. Caffeine (Sigma-Aldrich) was dissolved in distilled water at 100 mM and was used at a final concentration of 4 mM. 2-morpholin-4-yl-6-thianthren-1-yl-pyran-4-one (KU55933, ATMi, Calbiochem) was dissolved in DMSO (Sigma-Aldrich) at 10 mM and was used at 10 μM final concentration. 7-hydroxyxaurasporine (UCN-01, CHK1i, Calbiochem) was dissolved in DMSO at 100 μM and was used at 100 nM final concentration. The CHK2 inhibitor (CHK2 Inhibitor-II (BML-277), CHK2i, Calbiochem) was dissolved in DMSO at 1 mM and used at a final concentration of 400 nM. 8-(4-Dibenzoethinyl)-2-(4-morpholinyl)-4H-1-benzopyran-4-one (NU7441, DNA-PKcsi, Tocris) was dissolved in DMSO at 10 mM and was used at 10 μM final concentration. 3-Amino-6-[4-(methylsulfonyl)phenyl]-N-phenyl-2-pyrazinecarboxamide (ATRi, VE-821, Haoyuan Chemexpress) was dissolved in DMSO at 10 mM concentration and was used at a 5 μM final concentration. Unless indicated otherwise, all inhibitors were added to the cells 1 h before irradiation and were maintained until collection for analysis. To induce the expression of ATR kinase-dead (ATRkd) protein that exerts a dominant-negative function on ATR, exponentially growing GM847-ATRkd cells were exposed for 4 h to doxycycline hyclate (DOX), (Sigma-Aldrich) at a concentration of 3 μg/ml. DOX was maintained in the cultures during the duration of the experiment.

Indirect immunofluorescence (IF) and image analysis. For IF analysis33, cells were grown on cover-slips coated with poly-L-lysine (Biochrom). S-phase cells were labelled with 10μM of 5-ethyl-2′ deoxyuridin (EdU) for 30 min. Further manipulations were as previously described33. Primary antibody for RPA7044 was diluted (1:300) in PBG solution. Alexa Fluor-conjugated secondary antibody, anti-mouse IgG Alexa Fluor 488 (ThermoFisher Scientific, A11001), was applied at 1:400 dilution for 1 h at RT. The EdU signal was developed using an EdU developing kit (ThermoFisher Scientific) according to the manufacturer’s instructions. Cells were counterstained with 100 ng/ml DAPI (ThermoFisher Scientific) at RT for 5 min and coverslips were mounted in ProFluor antifade reagent (PromoCell). Scanning and analysis were carried out on an automated imaging system (Metasystems). For the quantification of parameters of interest, approximately, 1600 cells were analyzed to obtain ~200, EdU+ G2-phase cells.

Flow cytometry analysis of DNA end-resection by RPA70 and BrdU signal quantification. DNA end-resection analysis using RPA70 detection were described previously33. Briefly, exponentially growing cells were pulse-labeled for 30 min with 10 μM EdU. When, BrdU detection was used, cells were grown for 24 h in presence of 10 μM BrdU and then labelled with EdU. EdU labeling was interrupted by rinsing the cells with pre-warmed PBS and fresh medium containing or not PIKK inhibitors was supplied. Cells were irradiated with indicated IR doses and were collected by trypsinization at the indicated times after irradiation. The unbound RPA was extracted by incubating in ice-cold PBS containing 0.2% TritonX-100 and cells were fixed in 3% PFA. Cells were blocked with PBG blocking buffer overnight at 4°C and incubated with a monoclonal antibody raised against RPA70 (αSSB70B, mouse hybridoma cell line kindly provided by Dr. J. Hurwitz), anti-KU70 (529) (GeneTex, GTX77607), anti-ATR (Santa Cruz Biotechnology, sc-28901), anti-CHK1 (G-4) (Santa Cruz Biotechnology, sc-8408), anti-pCHK1-S345 (Cell Signaling Technology), anti-DNA-PKcs (Merck Millipore, PC127) and were used at 1:500 to 1:4000 dilutions. The secondary antibodies were: anti-mouse IgG conjugated with IRDye680 or anti-rabbit-IgG conjugated with IRDye800 (Li-COR Biosciences, 92668020 and 92632211) at 1:15,000 dilution. Immunoblots were visualized by scanning the membranes in an Odyssey infrared scanner (Li-COR Biosciences). Digital images were processed using the brightness contrast functions of the dedicated Odyssey software. Raw non-cropped scanned membranes are presented.

Polyacrylamide gel electrophoresis (SDS-PAGE) and western blotting. SDS-PAGE and western blot analysis were carried out according to the previously published protocol33. Briefly, cells were collected by trypsinization and were washed in ice-cold PBS. Approximately 3 × 106 cells were lysed in ice-cold RIPA buffer (ThermoFisher Scientific) supplemented with Hal generphosphatase and protease inhibitor cocktails (ThermoFisher Scientific), as recommended by the manufacturer and processed for SDS-PAGE and subsequently for western blotting as described33. The primary antibodies were: anti-RPA32 (mouse hybridoma cell line kindly provided by Dr. J. Hurwitz), anti-KU70 (529) (GeneTex, GTX77607), anti-ATR (Santa Cruz Biotechnology, sc-28901), anti-CHK1 (G-4) (Santa Cruz Biotechnology, sc-8408), anti-pCHK1-S345 (Cell Signaling Technology), anti-DNA-PKcs (Merck Millipore, PC127) and were used at 1:500 to 1:4000 dilutions. The secondary antibodies were: anti-mouse IgG conjugated with IRDye680 or anti-rabbit-IgG conjugated with IRDye800 (Li-COR Biosciences, 92668020 and 92632211) at 1:15,000 dilution. The secondary antibodies were: anti-mouse IgG conjugated with IRDye680 or anti-rabbit-IgG conjugated with IRDye800 (Li-COR Biosciences, 92668020 and 92632211) at 1:15,000 dilution. The secondary antibodies were: anti-mouse IgG conjugated with IRDye680 or anti-rabbit-IgG conjugated with IRDye800 (Li-COR Biosciences, 92668020 and 92632211) at 1:15,000 dilution. The secondary antibodies were: anti-mouse IgG conjugated with IRDye680 or anti-rabbit-IgG conjugated with IRDye800 (Li-COR Biosciences, 92668020 and 92632211) at 1:15,000 dilution.
3. Alt Frederick, W., Zhang, Y., Meng, F.-L., Guo, C. & Schwer, B. Mechanisms of Programmed DNA Lesions and Genomic Instability in the Immune System. Cell 152, 417–429 (2013).
4. Blackford, A. N. & Jackson, S. P. ATM, ATR, and DNA-PK: The Trinity at the Heart of the DNA Damage Response. Mol Cell 66, 801–817, https://doi.org/10.1016/j.molcel.2017.05.015 (2017).
5. Faltick, J., Coates, I. & Jackson, S. P. Conserved modes of recruitment of ATM, ATR and DNA-PKcs to sites of DNA damage. Nature 434, 605–611 (2005).
6. Zou, L. & Elledge, S. J. Sensing DNA Damage Through ATRIP Recognition of RPA-sDNA Complexes. Science 300, 1542–1548, https://doi.org/10.1126/science.11083430 (2003).
7. Gell, D. & Jackson, S. P. Mapping of protein-protein interactions within the DNA-dependent protein kinase complex. Nucleic Acids Research 27, 3494–3502 (1999).
8. Singleton, B. K., Torres-Arzayus, M. I., Rottingham, S. T., Taccioli, G. E. & Jeggo, P. A. The C terminus of Ku80 activates the DNA-dependent protein kinase catalytic subunit. Mol Biol Cell 19, 3267–3277 (1999).
9. Jette, N. & Lees-Miller, S. P. The DNA-dependent protein kinase: A multifunctional protein kinase with roles in DNA double strand break repair and mitosis. Progress in Biophysics and Molecular Biology 117, 194–205, https://doi.org/10.1016/j.pbiomolbio.2012.02.003 (2015).
10. Cencicchi, R., Rondinelli, B. & D’Andrea, A. D. Repair Pathway Choices and Consequences at the Double-Strand Break. Trends in Cell Biology 26, 52–64, https://doi.org/10.1016/j.tcb.2015.07.009 (2015).
11. Rothkamm, K., Krüger, I., Thompson, L. H. & Löbrich, M. Pathways of DNA Double-Strand Break Repair during the Mammalian Cell Cycle. Molecular and Cellular Biology 23, 5706–5715 (2003).
12. Shiloh, Y. & Ziv, Y. The ATM protein kinase: regulating the cellular response to genotoxic stress, and more. Nature Reviews. Molecular Cell Biology 14, 197–210 (2013).
13. Pauli, T. T. Mechanisms of ATM Activation. Annual Review of Biochemistry 84(12), 11–12.28, https://doi.org/10.1146/annurev-biochem-060614-034335 (2015).
14. Ahn, J.-Y., Schwarz, J. K., Pwenna-Worms, H. & Canman, C. E. Threonine 68 phosphorylation by ataxia telangiectasia mutated is required for efficient activation of Chk2 in response to ionizing radiation. Cancer Research 60, 3934–3936 (2000).
15. Brown, A. L. et al. A human Cds1-related kinase that functions downstream of ATM protein in the cellular response to DNA damage. Proceedings of the National Academy of Sciences of the United States of America 96, 3745–3750 (1999).
16. Matsuoka, S., Huang, M. & Elledge, S. J. Linkage of ATM to cell cycle regulation by the Chk2 protein kinase. Science 282, 1893–1897 (1998).
17. Melchiornia, R., Chen, X.-B., Blasina, A. & McGowan, C. H. Threonine 68 is required for radiation-induced phosphorylation and activation of Cds1. Nature Cell Biology 2, 762–765 (2000).
18. Seuly, R. & Xie, A. Double strand break repair functions of histone H2AX. Mutation Research/Fundamental and Molecular Mechanisms of Mutagenesis 750, 5–14, https://doi.org/10.1016/j.mrfmmm.2013.07.007 (2013).
19. Schwertman, P., Bekker-Jensen, S. & Mailard, N. Regulation of DNA double-strand break repair by ubiquitin and ubiquitin-like modifiers. Nature Reviews. Molecular Cell Biology 17, 379–394, https://doi.org/10.1038/nrm.2016.58 (2016).
20. Almeyr, M. & Lukas, J. To spread or not to spread—chromatin modifications in response to DNA damage. Current Opinion in Genetics & Development 23, 156–160, https://doi.org/10.1016/j.gde.2012.11.013 (2013).
21. Hustedt, N. & Durocher, D. The control of DNA repair by the cell cycle. Nat Cell Biol 19, 1–9, https://doi.org/10.1038/ncllcell.2016.352 (2016).
22. Chapman, J. R., Taylor, M. R. G. & Boulton Simon, J. Playing the End Game: DNA Double-Strand Break Repair Pathway Choice. Molecular Cell 47, 497–510, https://doi.org/10.1016/j.molcel.2012.07.029 (2012).
23. Adams, K. E., Medhurst, A. L., Dart, D. A. & Lakin, N. D. Recruitment of ATR to sites of ionising radiation-induced DNA damage requires ATM and components of the MRN protein complex. Oncogene 25, 3894–3904, https://doi.org/10.1038/sj.onc.3702450 (2006).
24. Cuadrado, M. et al. ATM regulates ATR chromatin loading in response to DNA double-strand breaks. Journal of Experimental Medicine 203, 297–303, https://doi.org/10.1084/jem.20051923 (2006).
25. Jazayeri, A. et al. ATM- and cell cycle-dependent regulation of ATR in response to DNA double-strand breaks. Nature Cell Biology 8, 37–45, https://doi.org/10.1038/ncllcell.2006.66 (2006).
26. Myers, J. S. & Cortez, D. Rapid Activation of ATR by Ionizing Radiation Requires ATM and Mre11. Journal of Biological Chemistry 281, 9346–9350, https://doi.org/10.1074/jbc.M513265200 (2006).
27. Riballo, E. et al. A pathway of double-strand break rejoining dependent upon ATM, Artemis, and proteins localizing to g-H2AX foci. Molecular Cell 16, 715–724 (2004).
28. Cimprich, K. A. & Cortez, D. ATR: an essential regulator of genome integrity. Nature Reviews. Molecular Cell Biology 9, 616–627, https://doi.org/10.1038/nrm2450 (2008).
29. Guo, Z., Kumagai, A., Wang, S. X. & Dynw, W. G. Requirement for Atr in phosphorylation of Chkl and cell cycle regulation in response to DNA replication blocks and UV-damaged DNA in Xenopus egg extracts. Genes & Development 14, 2745–2756 (2000).
30. Liu, Q. et al. Chkl is an essential kinase that is regulated by Atr and required for the G2/M DNA damage checkpoint. Genes & Development 14, 1448–1450 (2000).
31. Zhao, H. & Puwenna-Worms, H. ATR-Mediated Checkpoint Pathways Regulate Phosphorylation and Activation of Human Chkl. Nature Cell Biology 21, 4129–4139, https://doi.org/10.1038/ncllcell.2011.429 (2011).
32. Hekmat-Nejad, M., You, Z., Yee, M. C., Newport, J. W. & Cimprich, K. A. Xenopus ATR is a replication-dependent chromatin-modifying enzyme. EMBO Reports 10, 629–635 (2009).
33. Tajima, H., Lee, K.-I. & Chen, B. P. C. ATR-Dependent Phosphorylation of DNA-Dependent Protein Kinase Catalytic Subunit in Response to UV-Induced Replication Stress. Molecular and Cellular Biology 26, 7520–7528 (2006).
34. Lin, Y.-F., Shi, H.-Y., Shang, Z., Matsunaga, S. & Chen, B. P. DNA-PKcs is required to maintain stability of Chkl and Claspin for optimal replication response stress. Nucleic Acids Research 42, 4463–4473, https://doi.org/10.1093/nar/gku116 (2014).
35. Chiby, W. A. et al. Overexpression of a kinase-inactive ATR protein causes sensitivity to DNA-damaging agents and defects in cell cycle checkpoints. EMBO Journal 17, 153–159, https://doi.org/10.1093/emboj/17.1.153 (1998).
36. Xu, B., Kim, S.-T., Lim, D.-S. & Kastan, M. B. Two Moleurally Distinct G2/M Checkpoints Are Induced by Ionizing Irradiation. Molecular and Cellular Biology 22, 1049–1059, https://doi.org/10.1128/MCB.22.4.1049-1059.2002 (2002).
62. Zhou, Y.
61. Peterson, S. E.
60. Wang, H., Wang, X., Iliakis, G. & Wang, Y. Caffeine could not efficiently sensitize homologous recombination repair deficient cells
59. Wang, X., Wang, H., Iliakis, G. & Wang, Y. Caffeine-induced radiosensitization is independent of non-homologous end joining of
42. Shibata, A.
44. Rainey, M. D., Black, E. J., Zachos, G. & Gillespie, D. A. Chk2 is required for optimal mitotic delay in response to irradiation-induced DNA damage incurred in G2 phase. Oncogene 27, 896–906, https://doi.org/10.1038/sj.onc.1210702 (2008).
43. Hardcastle, I. R. et al. Discovery of Potent Chromone-4-one Inhibitors of the DNA-Dependent Protein Kinase (DNA-PK) Using a Small-Molecule Library Approach. Journal of Medicinal Chemistry 48, 7829–7846 (2005).
45. Hardcastle, I. R.
46. Labib, K. & Hodgson, B. Replication fork barriers: pausing for a break or stalling for time? EMBO Reports 8, 346–353 (2007).
47. El Achkar, E., Gerbault-Seureau, M., Muleris, M., Dutrillaux, B. & Debatisse, M. Premature condensation induces breaks at the interface of early and late replicating chromosome bands bearing common fragile sites. Proc Natl Acad Sci USA 102, 18069–18074, https://doi.org/10.1073/pnas.0506497102 (2005).
48. Visser, A. E. et al. Spatial Distributions of Early and Late Replicating Chromatin in Interphase Chromosome Territories. Experimental Cell Research 243, 398–407 (1998).
49. Saldivar, J. C. et al. An intrinsic S/G2 checkpoint enforced by ATR. Science 361, 806–810, https://doi.org/10.1126/science.aap9346 (2018).
50. Ruiz, S.
53. Lavin, M. F. Ataxia-telangiectasia: from a rare disorder to a paradigm for cell signalling and cancer. Nature Reviews. Molecular Cell Biology 15, 7–18, https://doi.org/10.1038/nrb3719 (2014).
51. Forment, J. V., Walker, R. V. & Jackson, S. P. A high-throughput, flow cytometry-based method to quantify DNA-end resection in interphase of early and late replicating chromosome bands bearing common fragile sites. Molecular and Cellular Biology 22, 6521–6532 (2002).
55. Symington, L. S. & Gautier, J. Double-Strand Break End Resection and Repair Pathway Choice.
62. Wang, H., Wang, H., Powell, S. N., Iliakis, G. & Wang, Y. ATR Affecting Cell Radiosensitivity Is Dependent on Homologous Recombination Repair but Independent of Nonhomologous End Joining. Cancer Research 64, 7139–7143 (2004).
56. Wang, H. et al. Caffeine inhibits homology-directed repair of I-sceI-induced DNA double-strand breaks. Oncogene 23, 824–834, https://doi.org/10.1038/sj.onc.1207168 (2004).
57. Wang, H., Wang, X., Wang, H., Iliaakis, G. & Wang, Y. ATR Affecting Cell RadioSensitivity Is Dependent on Homologos Recombination Repair but Independent of Nonhomologous End Joining. Cancer Research 64, 7139–7143 (2004).
58. Wang, H., Wang, H., X., Wang, H., Iliaakis, G. & Wang, Y. Caffeine-induced radiosensitization is independent of non-homologous end joining of DNA double strand breaks. Radiation Research 159, 426–432, 10.1667/0033-7587(2003)159[0426:CRIO]2.0.CO;2 (2003).
59. Wang, H., Wang, X., Iliaakis, G. & Wang, Y. Caffeine could not efficiently sensitize homologous recombination repair deficient cells to ionizing radiation-induced killing. Radiation Research 159, 420–425 (2003).
60. Peterson, S. E. et al. Activation of DSB Processing Requires Phosphorylation of CtIP by ATR. Molecular Cell 49, 657–667, https://doi.org/10.1016/j.molcel.2012.11.020 (2013).
61. Deshpande, R. A. et al. DNA-PKcs promotes DNA end processing. bioRxiv, https://doi.org/10.1101/395731 (2018).
62. Kenny, M. K., Schlegel, U., Fournex, H. & Hurwitz, J. The role of human single-stranded DNA binding protein and its individual subunits in simian virus 40 DNA replication. Journal of Biological Chemistry 265, 7693–7700 (1990).

Acknowledgements

The authors are grateful to Drs Karlene Cimprich, Penny Jeggo, Markus Lobrich, Eric Hendickson, Tej Pandita and Joan Allalunis-Turner for providing cell lines. Work supported by grants from the Federal Ministry of Education and Research, BMBF (02S8254, 02S8467, 03NUK005C, 02NUK043B) and the Deutsche Forschungsgemeinschaft, D.F.G., (GRK1739, ILS1-11-1).

Author Contributions

E.M., G.I. and X.F. designed the project and formulated the experiments. E.M. performed the IF and DNA end-resection analyzes, E.M., X.F., K.P.-K. and A.S. performed remaining experiments and analyzed results. G.I. and E.M. performed the manuscript, E.M. prepared the figures. All authors read and reviewed the manuscript.

Additional Information

Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-019-51071-6.

Competing Interests: The authors declare no competing interests.

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2019