Dissecting the role of p53 phosphorylation in homologous recombination provides new clues for gain-of-function mutants

Anja Restle¹, Martin Färber¹, Cindy Baumann¹, Michael Böhringer¹, Karl Heinz Scheidtmann², Carsten Müller-Tidow³ and Lisa Wiesmüller¹,²,*

¹Department of Obstetrics and Gynecology, University of Ulm, 89075 Ulm, ²Institute for Genetics, University of Bonn, 53113 Bonn and ³Department of Medicine, Hematology/Oncology, University of Münster, 48129 Münster, Germany

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ABSTRACT

Regulation of homologous recombination (HR) represents the best-characterized DNA repair function of p53. The role of p53 phosphorylation in DNA repair is largely unknown. Here, we show that wild-type p53 repressed repair of DNA double-strand breaks (DSBs) by HR in a manner partially requiring the ATM/ATR phosphorylation site, serine 15. Cdk-mediated phosphorylation of serine 315 was dispensable for this anti-recombinogenic effect. However, without targeted cleavage of the HR substrate, serine 315 phosphorylation was necessary for the activation of topoisomerase I-dependent HR by p53. Moreover, overexpression of cyclin A1, which mimics the situation in tumors, inappropriately stimulated DSB-induced HR in the presence of oncogenic p53 mutants (not Wtp53). This effect required cyclin A1/cdk-mediated phosphorylation for stable complex formation with topoisomerase I. We conclude that p53 mutants have lost the balance between activation and repression of HR, which results in a net increase of potentially mutagenic DNA rearrangements. Our data provide new insight into the mechanism underlying gain-of-function of mutant p53 in genomic instability.

INTRODUCTION

Because of the central role of p53 as a gatekeeper and a caretaker, the protein must be subject to complex control mechanisms that orchestrate the multiple functions of p53 in transcription, cell-cycle control, apoptosis induction and DNA repair (1,2). Posttranslational modification of p53 by protein phosphorylation has been the most extensively studied potential functional switch mechanism, as it occurs at multiple serine and threonine residues in response to genotoxic stress (3,4). Modification of p53 on serine 15 by ATM and ATR was demonstrated to trigger the cascade of damage-induced phosphorylation and acetylation events that have been implicated in protein stabilization and enhancement of transcriptional transactivation (3,4). However, observations made with knock-in mouse models (5,6) indicated a role for serine 18 in apoptosis, but not in Mdm2-governed protein stability. Moreover, in several studies, no evidence was found for an essential role of the N-terminal casein kinase 1 (CK1) and ATM/ATR phosphorylation sites in damage-induced transcriptional transactivation (7–9). In addition, when DNA replication was blocked, p53 became phosphorylated on serine 15, but this was not accompanied by a rise in key target gene products such as p21 (10–12). This suggested that after replication fork stalling, p53 phosphorylated on serine 15 (p53pSer15) may serve additional functions unrelated to transcriptional transactivation. In support of this hypothesis, colocalization studies indicated that p53pSer15 forms a component of RAD51-specific repair assemblies (11–13).

Over the last few years, a large body of evidence has emerged indicating that p53 is directly involved in DNA repair, particularly in homologous double-strand break (DSB) repair. First, p53 recognizes three-stranded heteroduplex and four-way Holliday junctions and DNA lesions involving mismatches, gaps or DNA ends. The core domain is required for junction DNA-binding and also stimulates these activities upon mismatch recognition (2,14,15). Second, p53 physically and functionally interacts with critical enzymes and surveillance factors of homologous recombination (HR), namely with RAD51, RAD54, the MRE11 complex, BRCA1, BRCA2 and BLM, and counteracts strand exchange catalyzed by RAD51. Third, using different cell-based test systems, several groups concurrently found that Wtp53 represses...
inter- and intra-molecular HR, when triggered by DSBs or replication blocking agents. In contrast, hotspot mutants failed to downregulate these HR activities. The identification of separation-of-function mutations, which had lost p53's transcriptionsal transactivation and cell-cycle regulatory capacity, but retained HR inhibition, and vice versa, provided further evidence for p53's direct role in HR control (2,14,15). A recent report describes transcriptional repression of RAD51 by direct binding of Wtp53 to a response element within the promoter region (16). This mechanism can only partially explain the role of p53 in HR, because mutations within the p53 interaction site of the RAD51 protein abrogate HR repression by p53 (13). Moreover, p53(138V), which is defective in sequence-specific DNA binding, retains the HR-downregulatory effect (17). The biological meaning of this, at first sight, paradoxical activity directed against a fairly safe DNA repair pathway was unveiled by systematic substrate variation, which indicated a fidelity control mechanism directed against DNA exchange processes between divergent sequences (in 15). Unexpectedly, Wtp53 was more recently found to stimulate recombination in the absence of targeted substrate cleavage in a manner depending on topoisomerase I (topo I) (18,19). Spontaneous recombination events are coupled to the normal DNA metabolism in proliferating cells such as during the bypass of low level, endogenous lesions at replication forks, which are insufficient to activate stress signalling.

Upon exposure to ionizing radiation and generation of highly recombinogenic DNA lesions such as DSBs, the serines 6, 15 and 315 represent the most prominently phosphorylated p53 residues (3). On the other hand, Subramanian and Griffith (20) demonstrated that recognition of Holliday junction DNA by p53 is particularly sensitive to posttranslational phosphorylation at serine 392 as compared to serines 6 or 15. To define the role of phosphorylation in p53-dependent regulation of recombinative repair, we applied an EGFP-based recombination assay in combination with cells expressing the corresponding phosphorylation site mutants. Candidate kinases were manipulated by pharmacological or shRNA-mediated inhibition and by expression of regulatory subunits. We find that serine 15 phosphorylation by ATM/ATR is one contributing mechanism to channel functions of p53 towards DSB repair surveillance. Serine 315 and cdk2 are required for topo I-dependent recombination, which involves dynamic interactions between p53 and topo I in the nucleus. Together, our data indicate that site-specific phosphorylation of p53 plays a critical role in the fine-tuning of repair pathways with important implications for oncogenic p53 mutants.

MATERIAL AND METHODS

Cells and cell-cycle analysis

For HR-measurements KMV clones derived from the human myeloid leukemia cell line K562 and KMV (HR/3') derivatives with stably integrated HR-EGFP/3'EGFP substrate were raised in RPMI 1640 supplemented with 10% FCS and 2 mM L-glutamine (Biochrom, Berlin, Germany) (21). Stable WTK1(HR/3') clones for measurements of HR between HR-EGFP and 3'EGFP on chromosomes were established from the lymphoblastoid cell line WTK1 (21) and cultivated in RPMI 1640 with 12% FCS and 3 mM L-glutamine. When using pSVp53her expression plasmids, cells were propagated in estradiol-free medium before transfection and, then, cultivated in the presence of estradiol at a final concentration of 200 nM to fully activate the p53 moiety within the fusion protein of 90 kD (21).

For the analysis of cell-cycle phases and apoptosis induction at the time point of HR measurement, cells were collected by centrifugation, resuspended in 1 ml HBSS (25 mM HEPES, pH 7.4; 173.3 mM NaCl; 5.6 mM KCl; 0.44 mM KH2PO4; 0.34 mM Na2HPO4; 5 mM glucose; 4 mM NaHCO3) and fixed in 10 ml fixing solution (50% acetone; 40% ethanol; 10% dD2O; at −20°C) on ice for 15 min. For subsequent staining, fixed cells were washed with 2.5 ml ice-cold PBS supplemented with 0.2% EDTA, resuspended in 100 µl propidium iodide staining solution (0.25 mg/ml RNAse A; 50 µg/ml propidium iodide from Sigma in PBS) and incubated for 15 min in the dark. After diluting the suspension by 200 μl PBS with 0.2% EDTA, the stained cells were analyzed in a FACSc flow cytometer (Becton Dickinson, Heidelberg, Germany). The percentage of cells with sub-G1 DNA content, which is indicative of apoptosis, was determined, and viable cells were subdivided into populations in the cell-cycle phases G1, S and G2.

Plasmids

The following plasmids were previously described: pSUPER-TopoI (19), pSUPER-p53 (22), pcDNA3-cyclin A1 and pCMV-cyclin A1 (2,14,15). SMRAD51 chimera expression plasmid pcDNA3.1-Rad51SM (24), pKEX expression constructs for rat p53 (7), pCMV-I-SceI, pCMV-Wtp53, pCMV-p53(248W), pSVp53her and pSVp53(1-333)her (21), pCMV-p53(315A) was constructed from pCMV-p53 (BD Biosciences Clontech, Heidelberg, Germany) by use of Quickchange (Stratagene Europe, Amsterdam, The Netherlands) to introduce the point mutation. pKD-ck2-v5, pKD-E2F-1-v1 and empty vectors were purchased from Upstate. Plasmid pSUPER-WRN directs the synthesis of siRNA, which had previously been demonstrated to specifically target WRN-mRNA (25). pSUPER-WRN was generated as described for pSUPER-TopoI in Baumann et al. (19) by use of the following oligonucleotides (ThermoHybaid, Ulm): anti-WRNsiRNA-fwd, 5'-GATCCCCCTTCAGTGATCCATGTGTGCCAGAGAAAGACACAAATGTAGATCAGTGAATTTTGGAAGAA-3'; anti-WRNsiRNA-rev, 5'-AGCTTTCACAAAAATTCACTGTGATCCATGTGTGCCAGAGAAAGACACAAATGTAGATCAGTGAATTTTGGAAGAA3'. Additionally, we newly created pSV53(15A) her for the expression of p53her fusion protein mutated at serine 15 using PCR. PCR amplification relied on pSV53her template and the primer 5'-CTGTTACATGAGGAGCGCGAGCTCAGATCCTAGGCTCAGGGGCCCCTGCTGCTGGCTAGGGAAATGTTTCGACC-3', covering the 5'-end of the p53 cDNA (mutating bases underlined), and 5'-AGATGGCCATGGCAGCGG-3', covering an internal...
Neo1 restriction site. The Neo1 digested PCR product was inserted into the pUC-subcloned Xbal fragment from pSV53her, which was retransferred into pSV53her after mutagenesis. The complete sequence encoding p53(15A)her was verified by use of the ABI PRISM Big Dye Ready Reaction Cycle Sequencing kit and an ABI 377 sequencer.

Antibody probes

We used the following mAbs: DO1 directed against the N-terminus of p53, C21 against topo I, B88-2 against cyclin A1, BD611168 against WRN and SX118 against p21 from Becton Dickinson, Pab421 against the C-terminus of p53 (Calbiochem, Darmstadt, Germany), 1D6 against topo I (Novocastra Laboratories Ltd, Newcastle upon Tyne, UK), E 67.1 recognizing Cyclin A2 from Cell Signaling, FPS315 against p53pSer315 (MoBiTec/MBL, Göttingen, Germany) and KH20&KH95 against E2F1 (Upstate Biotechnology Incorporated, Lake Placid, NY, USA). For immunodetection of rat p53, we used rabbit serum CM1 against p53 (Novocastra). Peroxidase-conjugated affinity purified antibodies against goat, rabbit, human and mouse IgG were obtained from Dianova, Hamburg, Germany or Biomol.

Extraction of cells, immunoprecipitation and immunoblot analysis

Total homogenates from 2 x 10⁵ KMV(HR/3') cells were prepared, subjected to SDS–PAGE (8–15%) and western blot transfer on Immobilon-P membranes (Millipore, Bedford, MA, USA) as described (21). Filters were stained with antibodies in PBS containing 0.2% Tween-20 and 5% nonfat dried milk, followed by incubation with peroxidase-conjugated secondary antibodies and detected by the ECL reagents SuperSignal® West Pico Chemiluminescent Substrate and West Dura Extended Duration Substrate (Pierce, Rockford, IL, USA). Cellular lysates for immunoblot analysis were prepared in lysis buffer (50 mM Tris, pH 7.4; 150 mM NaCl; 2 mM EGTA; 2 mM EDTA; 25 mM NaF; 25 mM β-glycerophosphate; 0.1 mM NaV; proteinase inhibitor, Roche Diagnostics, Mannheim, Germany) including phosphatase inhibitor cocktail (Roche) in the experiment shown in Figure 6B and 100 μg of total protein per sample resolved by SDS–PAGE. Filters were probed with antibodies against actin, tubulin or TBP, or Ponceau stained to control for loading differences. Alternatively, signals for a nonspecific band served to verify equal loading.

For immunoprecipitation KMV(HR/3') cells were lysed 24 h after electroporation with the indicated plasmids in 50 mM HEPES (pH 7.8), 200 mM NaCl, 0.1% Triton X-100, 20% glycerol and protease inhibitor cocktail (Roche). For isolation of topo I-Wtp53 complexes in vivo cross-linking was performed as described in Zink et al. (26). Centrifuged extracts were precleared with protein A-sepharose (PAS) for 1 h, incubated with anti-topo I or anti-GAPDH antibodies and PAS for 4 h and further processed as in Restle et al. (12).

Densitometric quantification of band intensities was performed using a ChemiImager 5500 with software (Alpha Imunotech Corporation, San Leonardo, California, USA). Values for the protein of interest were corrected with the values obtained for the corresponding loading control.

HR-analysis, drug treatment and knockdown

For the determination of HR frequencies with extrachromosomal plasmid substrate, 10⁵ KMV cells were electroporated with a total amount of 40 μg plasmid DNA (10 μg HR-EGFP/3'EGFP, 10 μg pCMV-I-SceI, 10 μg pBS, 10 μg p53 expression plasmid or empty vector controls). For chromosomal HR measurements, 4 x 10⁶ KMV(HR/3') or 3 x 10⁶ WTK1(HR/3') cells were electroporated with 30 μg plasmid DNA (10 μg pBS, 10 μg p53 expression plasmid, cyclin A1 or cyclin A2 expression plasmid or empty vector controls) with or without 10 μg pCMV-I-SceI. 48–72 h after electroporation flow cytometric analysis was performed in a FACS® (Becton Dickinson) to determine the fraction of green fluorescent cells within a population of nonfluorescent cells as described previously (21). Each measurement was paralleled by the same coelectroporation including a WtEGFP control plasmid instead of a pBS filler plasmid. The fraction of EGFP-positive cells in these controls was determined each and was used to normalize each single recombination frequency. This calculation procedure also corrects for possible rate deviations related to growth and death regulatory effects. For drug treatment, KMV(HR/3') cultures were split and treated with kinase inhibitor or solvent starting from 4 h after electroporation until measurements were performed (72 h). Caffeine (Sigma, Deisenhofen, Germany) was dissolved in H₂O and included at a final concentration of 400 μM, CGK (Sigma) in DMSO and applied at 2 μM; olomoucine (Calbiochem) was dissolved in DMSO and added at 80 μM. In topo I, p53, WRN, cdk2 and E2F1 knockdown experiments 20 μg of pSUPER-TopoI, pSUPER-p53, pSUPER-WRN, pKD-ckd2-v5, pKD-E2F-1-v1 or empty vector were added to the coelectroporation mixture (transfection efficiency: 40–80%). The statistical significance of differences was calculated by use of the t-test.

RESULTS

Changes in the regulation of homologous DSB repair after mutation of p53 phosphorylation sites

Recombinative repair represents a well-established direct DNA repair function of p53 (2,14,15) and had been characterized by use of an EGFP-based assay in K562 derived KMV cell clones, which are devoid of endogenous
p53 (19,21). In this assay, HR between mutated EGFP sequences, located on the same or the sister chromatid, restores a functional EGFP gene, which can be monitored by flow cytometry via quantitation of green fluorescent cells (Figure 1A). DNA-exchange processes are triggered by DSB formation in vivo, when the rare-cutting meganuclease I-SceI is expressed and introduces a single DSB at the recognition sequence within the repair substrate.

To examine the role of p53 phosphorylation in the modulation of HR, we chose to express a series of rat p53 phosphorylation mutants, which in a previous comparative study had been shown to retain transcription activity (7). Thus, we cotransfected KMV(HR/3') cells, carrying the stably integrated HR-substrate HR-EGFP/3'EGFP (Figure 1A), with expression plasmids for I-SceI (pCMV-I-SceI) and for Wtp53, oncogenic p53(174Y) or phosphorylation mutants p53(15A), p53(4/6A), p53(390) or p53(310/313A), or empty vector. HR was reduced down to 47% by Wtp53, when compared with the p53-negative control (P = 0.034) (Figure 1B). We found similar HR repression by the mutants p53(390A) (P = 0.012) and p53(310/313A) (P = 0.005). p53(174Y), which is devoid of any functions in transcriptional and HR regulation (21), showed loss of function. After expression of p53(15A) or p53(4/6A) minor, statistically not significant, changes in HR frequencies compared to p53-negative controls were noted (P = 0.307 and P = 0.193, respectively). When we analyzed p53 expression, we noticed lower levels for p53(15A) and p53(4/6A) as compared to Wtp53 (Figure 1C). Nevertheless, regarding the products of genes, which are subject to transcriptional transactivation or repression by p53, we observed comparable p21 increases in the presence of Wtp53 as well as for p53(15A), p53(4/6A), p53(390) and p53(310/313A), and no decrease of RAD51.

Repression of homologous DSB repair by p53 is lost after ATM/ATR kinase inhibition

Phosphorylation of p53 at serine 15 by ATM and ATR is thought to initiate the cascade of DNA damage-induced posttranslational modification events on p53 (3,4). To further delineate a potential role of serine 15 phosphorylation in the regulation of DSB repair, we cotransfected KMV(HR/3') cells with pCMV-I-SceI and constructs for comparable expression of human Wtp53 and the phosphorylation mutant p53(15A) (21) (Figure 2C). Wtp53 caused a 36–51% (P = 0.017–0.046) decrease of chromosomal HR (Figure 2A and B). p53(15A) expressing cells exhibited an intermediate phenotype with a recombination frequency reduction of 10–31% (P = 0.047–0.517). For comparison, the HR regulation defective tetramerization mutant, p53(1-333), caused a statistically not significant 25% decrease (P = 0.126). To study the possible involvement of ATM and/or ATR, we next treated the cells with the ATM and ATR inhibitor caffeine (27). Western blot
analysis verified reduced phosphorylation of p53 at serine 15 after caffeine treatment (Figure 2C). Caffeine treatment completely abrogated HR-repression by Wtp53 as well as the residual influence by p53(15A) and p53(1-333) (Figure 2A).

To rule out a major contribution of the cell cycle to the effects shown in Figure 2A, we analyzed by flow cytometry the cellular distribution of G1-, S- and G2-phases. Under the conditions of HR measurements (72 h treatment with 400 μM caffeine, purposefully chosen after titration analysis to exclude cytotoxic effects), all cell types, with or without p53 expression and with or without inhibitor treatment, showed comparable cell-cycle profiles (Supplementary Figure 1A). In addition, the fraction of apoptotic cells identified by sub-G1 DNA content after propidium iodide staining, was neither significantly different in this experiment nor in the experiments described below (Supplementary Figure 1B and data not shown). This is in agreement with the fact that K562 cells are defective in apoptosis induction after p53 expression (28). Since RAD51 was demonstrated to represent the immediate target of p53-dependent HR inhibition (2,14,15), we wished to compare the influence of Wtp53 with inactivation of RAD51. For that purpose, we expressed SMRAD51, a yeast–mouse chimera, with a dominant-negative effect but which does not modify cell viability in mammalian cells (24). We found repression of DSB-induced HR in KMV(HR/30) cells down to 63% (P = 0.001) (Figure 2A). For comparison, p53(273H) showed residual chromosomal downregulation activity, however, without statistical significance (P = 0.096). Protein levels of p21 were comparably elevated in Wtp53, p53(315A), p53(392A) and p53(392D), but not p53(273H) producing cells, whereas RAD51 protein levels remained constant (Figure 3B). Thus, phosphorylation of serines 315

Figure 2. Influence of ATM/ATR-mediated phosphorylation. (A) Homologous DSB repair of chromosomal substrates was assayed in KMV(HR/3) cells as in Figure 1 using the following expression plasmids for human p53 fused to the human estrogen hormone-binding domain: pSVp53her (Wt), pSVp53(15A)her, pSVp53(1-333)her or pBS (control) in the absence or presence of 400 μM caffeine (A) or 2 μM CGK (B). Recombination frequencies in cells without p53 were set to 100% [absolute mean value in (A) and (B): 1.0 × 10⁻³]. Columns, mean (n = 3–9); bars, SEM. (C) Equal expression of p53her and p53(15A)her and decreased phosphorylation of p53her at serine 15 after caffeine and CGK treatment, respectively.

Additionally, when we tested the more potent and selective ATM/ATR inhibitor CGK (30), which also caused reduced phosphorylation of p53 at serine 15 (Figure 2C), we again observed loss of HR repression by Wtp53 and no further change for p53(15A) (Figure 2B). Altogether, our findings suggest that serine 15, which is phosphorylated by ATM and ATR, is required for fully efficient regulation of DSB-induced HR. They also show that cellular ATM and/or ATR kinase activities are needed to uncover differences in HR frequencies between p53-positive and -negative cells.

Stimulation of I-SceI-independent HR by p53 requires the cdk phosphorylation site, serine 315

Given that mutation of phosphorylated amino acids on the C-terminus of rat p53 did not alter DSB repair regulation (Figure 1B), we confirmed these observations for human p53. Therefore, we performed HR measurements on chromosomally integrated substrates after expression of human Wtp53 and cancer-related mutant p53(273H), p53(315A), p53(392A) and p53(392D). p53(315A) and p53(392A) are resistant to phosphorylation on the respective serine, whereas p53(392D) mimics phosphorylation on serine 392. p53(315A), p53(392A) and p53(392D) downregulated homologous DSB repair similar to Wtp53 (P ≤ 0.001) (Figure 3A). For comparison, p53(273H) showed residual chromosomal downregulation activity, however, without statistical significance (P = 0.096). Protein levels of p21 were comparably elevated in Wtp53, p53(315A), p53(392A) and p53(392D), but not p53(273H) producing cells, whereas RAD51 protein levels remained constant (Figure 3B). Thus, phosphorylation of serines 315
and 392 does not play a significant role in p53-mediated repression of DSB-induced HR.

Since amino acid 315 is located within the topo I-binding site of p53 (31), we next sought to determine the involvement of serine 315 in the stimulation of I-SceI-independent HR, which was shown to be mediated by Wtp53 and topo I (19). HR was monitored using KMV(HR/30) cells in combination with expression of human Wtp53, p53(315A), or the hotspot mutants, p53(273H) and p53(248W), in the absence of substrate cleavage by I-SceI. Consequently, we scored an average recombination frequency two orders of magnitude below I-SceI-induced HR (see legend to Figure 4A and B). Hence, we scored an average recombination frequency two orders of magnitude below I-SceI-induced HR (see legend to Figure 4A and B).

I-SceI-independent HR was increased ~2-fold by Wtp53 (P = 0.023), p53(273H) (P = 0.113) and p53(248W) (P = 0.001), but not by p53(315A) (Figure 4A). To determine whether HR enhancement is topo I-dependent, we repressed endogenous topo I expression by transfecting with pSUPER-TopoI (19) (Figure 5D). In these knockdown assays, HR stimulation was abolished for Wtp53, p53(273H) and p53(248W), reaching statistical significance for Wtp53 (P = 0.030) and p53(248W) (P ≤ 0.001). To exclude the possibility that topo I represents an essential component of HR, we performed the same HR measurements with simultaneous I-SceI expression for targeted substrate cleavage. Under these conditions, topo I knockdown had no effect on HR activities, regardless of the p53 status (Figure 4B). FACS analysis of the fraction of cells with G1, S or G2 DNA content or of apoptotic cells failed to detect significant differences between samples without and with expression of Wtp53, p53(273H) or p53(248W), arguing against HR stimulation being indirectly caused by the classical tumor suppressor activities (Supplementary Figure 1B). These results indicated that both Wtp53 and oncogenic mutant p53 stimulate I-SceI-independent HR, topo I-mediated recombination and revealed that serine 315 is essential for the stimulatory effect.

Oncogenic p53 mutants stimulate cyclin A1-induced, topo I-dependent homologous DSB repair

Cyclin A1, an alternative, cdk2 associated A-type cyclin, has been linked to DSB repair, and its promoter represents a transcriptional transactivation target of p53 (23).
To test a potential contribution of cyclin A1 in the regulation of homologous DSB repair by p53, we simultaneously expressed cyclin A1, p53 proteins and I-SceI in KMV (HR/3') cells. In parallel, we examined the influence of cyclin A2. We found a 69–83% enhancement of homologous DSB repair after cyclin A1 expression in cells with p53(273H) (P ≤ 0.004) and of 65–148% for p53(248W) (P ≤ 0.006), whereas in p53-negative controls frequencies were only 7–18% higher (P > 0.05) (Figures 5A, 6A and 7A). In cells expressing mutant p53(175H), we measured 78% higher HR frequencies (P = 0.013). After cyclin A2 expression, relative increases were 47–83% for p53(273H).
(P = 0.009–0.025), 42–60% for p53(248W) (P = 0.003–0.014), and 35–49% for controls (P ≥ 0.019). As expected, in cells with Wtp53 and p53(315A), DSB-induced HR was strongly suppressed, namely on average to 20% of control levels (Figure 3A), with increments of 35–41% and 17–33% upon cyclin A1 and of 13–48% and 46–47% upon cyclin A2 production (P > 0.05), respectively (Figures 5A, 6A and 7A). Flow cytometric analysis revealed that cells expressing cyclin A1 or cyclin A2 showed similar cell-cycle distributions compared with controls (Supplementary Figure 1C). Moreover, we measured recombination in the presence of the pancaspase inhibitor zVAD-fmk (50 μM) for Wtp53, p53(315A) and p53(273H) versus control (± cyclin A1 each). Recombination results did not differ from the initial results, thus, arguing against an involvement of apoptosis in the HR regulation (data not shown).

We noticed that exogenous p53(273H) exerted residual HR-repression activities to a varying extent (Figures 5A, 6A and 7A) (21,32). Therefore, we wished to compare cyclin A1-mediated HR stimulation in cells stably expressing endogenous mutant p53 and cell derivatives without p53. Consequently, we analyzed HR activities in the human B-lymphoblastoid cell line WTK1(HR/3) carrying p53(237I) (21) and chromosomally integrated HR

Figure 6. Influence of cdk inhibition on HR regulation by cyclin A1 and mutant p53. (A) DSB-induced HR in KMV(HR/3) cells treated with DMSO (−) or 80 μM olomoucine (+). Recombination frequencies in controls −p53−/−cyclin A1−/−olomoucine were set to 100% (absolute mean value: 1.1 × 10⁻³). Columns, mean (n = 12–18); bars, SEM. (B) Immunoblot analysis for p53pSer315 and p53. (C) Cell-cycle analysis. Columns, mean (n = 2); bars, SD.

Figure 7. Regulation of HR by cyclin A1 and mutant p53 after knockdown of cdk2. (A) DSB-induced HR in KMV(HR/3) cells cotransfected with empty vector or pKD-ckd2-v5 (pKD-ckd2) for cdk2-specific RNA interference. Recombination frequencies in controls −p53−/−cyclin A1−/−pKD-ckd2 were set to 100% (absolute mean value: 1.2 × 10⁻³). Columns, mean (n = 6); bars, SEM. (B) Western blot analysis of cdk2.
substrate. HR assays were performed with and without cyclin A1 overexpression and concomitantly with or without knockdown of endogenous p53 by pSUPER-p53 (Supplementary Figure 2B). This approach revealed a statistically significant (25%, \( P = 0.001 \)) rise in HR frequencies from mutant p53-deficient (+pSUPER-p53) to -proficient cells (+pSUPER) only after cyclin A1 overexpression (Supplementary Figure 2A). Cell-cycle changes were not observed (Supplementary Figure 2C).

We next reapplied pSUPER-TopoI to analyze the involvement of topo I in the cyclin A1 and cyclin A2-dependent stimulation of homologous DSB repair. Interestingly, after topo I knockdown in KMV(HR/3) cells we did not find significant differences in HR frequencies except for the cells expressing cyclin A1 in combination with the oncogenic p53 mutants (Figure 5B and C). Thus, with p53(273H) HR was reduced by 28% \((P = 0.004)\), with p53(248W) by 43% \((P = 0.001)\). We verified topoisomerase downregulation as depicted in Figure 5D for cultures with and without cyclin A1 expression. In the same samples, p53 levels did not follow the topo I expression pattern when compared to the TBP control. Taken together, both cyclin A1 and cyclin A2 overexpression (Figure 5E) caused an enhancement of DSB-induced HR, particularly in cells expressing a p53 hotspot mutant. When compared to p53-negative controls, mutant p53-mediated HR stimulation was much more pronounced in cells with cyclin A1 rather than cyclin A2. This indicated a cyclin A1- and mutant p53-specific component. Most importantly, HR enhancement was abolished by topo I knockdown solely in cells with cyclin A1 and mutant p53.

**Mutant p53-mediated stimulation of homologous DSB repair requires cdk2**

We next used two independent approaches to analyze the role of cdk2 in regulating HR after cyclin A1 and mutant p53 expression. First, we included the cdk inhibitor, olomoucine, into KMV(HR/3') cells we did not find significant differences in HR frequencies except for the cells expressing cyclin A1 in combination with the oncogenic p53 mutants (Figure 5B and C). Thus, with p53(273H) HR was reduced by 28% \((P = 0.004)\), with p53(248W) by 43% \((P = 0.001)\). We verified topoisomerase downregulation as depicted in Figure 5D for cultures with and without cyclin A1 expression. In the same samples, p53 levels did not follow the topo I expression pattern when compared to the TBP control. Taken together, both cyclin A1 and cyclin A2 overexpression (Figure 5E) caused an enhancement of DSB-induced HR, particularly in cells expressing a p53 hotspot mutant. When compared to p53-negative controls, mutant p53-mediated HR stimulation was much more pronounced in cells with cyclin A1 rather than cyclin A2. This indicated a cyclin A1- and mutant p53-specific component. Most importantly, HR enhancement was abolished by topo I knockdown solely in cells with cyclin A1 and mutant p53.

**Cyclin A1 stimulates association of topo I with p53**

Given that cyclin A1-mediated stimulation of homologous DSB repair depends on topo I and mutant p53, we examined whether formation of protein complexes containing oncogenic mutant p53 and topo I is involved in cyclin A1-mediated recombination stimulation. For this purpose, we performed coimmunoprecipitation analysis using ScI70-antiserum, which had previously been established for specific precipitation of topo I complexes (41). Topo I-bound proteins were recovered from extracts of cells transfected and treated for DSB-repair measurements as in Figure 6. p53(248W) bound to topo I, particularly in cells overexpressing cyclin A1 (Figure 8). Under these conditions, we were unable to detect Wtp53 signals in topo I precipitates, unless chromatin complexes were cross-linked in vivo (Figure 8), indicating weak or transient interactions with topo I. Additional treatment of these cells with olomoucine reduced the association of Wtp53 and particularly of p53(248W), with topo I, supporting the hypothesis that cyclin A1/cdk-mediated phosphorylation, plays an important role in the stable association of oncogenic p53 with topo I.
DISCUSSION

The p53 is phosphorylated on several residues in response to genotoxic stress, in particular to insults such as DSBs (2,4,15). Because of the well-established regulatory functions of p53 in DSB repair pathways, we and others previously proposed that damage-induced phosphorylation may modulate repair-related p53 activities. In this study, we systematically examined this hypothesis, demonstrating that phosphorylation of p53 at different positions modulates distinct DNA repair-related functions of p53.

Functional proof of a partial involvement of serine 15 in DSB repair

Phosphorylation of p53 at serine 15 by ATM represents the initial p53 modification induced by DSBs (4). In chromatin immunoprecipitation experiments ATM and p53 were found to localize to DSBs preceding repair (42). Moreover, we and others previously demonstrated that p53pSer15 represents the p53 subpopulation colocalizing with the MRE11 complex, RAD51 and RAD54 after replication blockage (11–13). However, so far, only indirect evidence suggested HR-related functions for this subpopulation of posttranslationally modified p53. In this work, we demonstrate that serine 15 of p53 is necessary for fully efficient repression of homologous DSB repair. On the other hand, with C-terminal phosphorylation site mutants, changes in HR downregulation were not detectable. We noticed that the phosphorylation-resistant mutant p53(15A) displayed residual HR-repression activities, whereas chemical inhibition of ATM and ATR completely abrogated p53-dependent HR downregulation. This can be explained by the involvement of additional N-terminal sites which, like serines 20 and 37, are directly or indirectly modified after ATM/ATR activation (3,4). However, we cannot exclude that ATM and/or ATR are required for p53-mediated HR repression independently of p53 phosphorylation.

Under the conditions of the DSB repair test, we observed no alterations in transcriptional transactivation nor in repression by p53(15A), when monitored via p21 and RAD51 expression. Additionally, we largely excluded that the influence of serine 15 phosphorylation on HR was caused by major p53 activities in cell cycle control or apoptosis induction. Our observations strengthen earlier conclusions of a direct involvement of p53 in RAD51-dependent gene conversion (2,14,15). Physical interactions with heteroduplex DNA recombination intermediates and RAD51 do not involve the N-terminus of p53 (15). RPA binding, which like RAD51 binding is required for HR repression by p53, was mapped around aa 48–54, and, therefore, is more likely to be modulated by N-terminal phosphorylation (43). Importantly, colocalization and immunoprecipitation experiments showed that BLM facilitates physical interactions between p53 and the sensor proteins 53BP1 and H2AX, and thereby leads to efficient localization of p53 to RAD51 foci and to a net increase in p53pSer15 (11,44). Therefore, phosphorylation of N-terminal p53 amino acids by ATM, ATR and downstream kinases may stabilize one or multiple contacts with recombination factors, enabling the HR repression function of p53.

Links between p53 and cyclin A1 in recombination

Consistent with our previous observations (18,19), we showed here that Wtp53 activates HR in the absence of targeted substrate cleavage in a topo I-dependent manner. We have now discovered the same to be true for the hotspot mutants p53(273H) and p53(248W). Serine 315 turned out to be necessary for topo I-dependent HR stimulation, whereas it was dispensable for the repression of DSB-induced HR.

Cyclin A-cdk2 was previously shown to phosphorylate serine 315 of human p53 (45,46). Interestingly, we found that even DSB-induced HR was stimulated by mutant p53 in a manner depending on topo I, when cyclin A1 was overexpressed (up to 20-fold further increase as compared to the p53-negative control). Under the same conditions, Wtp53 caused repression of homologous
DSB repair. In contrast to cyclin A1, overexpression of cyclin A2 was accompanied by a mostly p53- and topo I-independent increase in homologous DSB repair. This general increase in HR is in line with a report describing that efficient DSB-induced ATM/ATR-signaling to Chk1 requires cdk kinase activity (47). Since Chk1 is required for RAD51 activation, this implies that cdks promote HR (48).

From our data and given that cyclin A1 is a target gene product of Wtp53 (23), we would like to put forward the following hypothesis (Figure 9): Wtp53 may support rare HR processes via cyclin A1 and topo I in normal, proliferating cells, particularly during S-phase, when active DNA synthesis is associated with replication fork stalling and recombinative bypass events. However, in the presence of severe damage such as DSBs, ATM-signaling may dominantly promote HR-repression functions of Wtp53, i.e. overcome HR stimulation. Oncogenic mutant p53 has lost the HR repression but retained the HR-stimulation function resulting in a net increase of recombination.

**Dynamic p53-topo I interactions as the mechanistic basis of cyclin A1-induced recombination**

Topo I is known to bind p53 between aa 302 and 321 (31), i.e. in a region containing serine 315, which is phosphorylated in response to cyclin A1 expression. Immunoprecipitation experiments revealed that overexpression of cyclin A1 promoted physical interactions between topo I and p53(248W). For Wtp53, we noticed comparatively weak or transient topo I binding, which is in agreement with observations made by Gobert et al. (31). Cyclin A1-enhanced p53–topo I interactions were reduced after pharmacological inhibition of cdks, particularly for p53(248W). Thus, we propose that cyclin A1/cdk2-mediated phosphorylation of p53 enables stable complex formation with topo I, thereby causing hyper-recombination in p53 mutant cells (Figure 9).

Topo I has been recognized as a DNA metabolizing enzyme that—once localized to DNA lesions by other proteins—is necessary for the recruitment of DNA repair enzymes. On the other hand, topo I is potentially dangerous due to its nicking–closing activity (49). Topo I facilitates initiation of genetic recombination through strand breakage after strand pairing and/or resolution of DNA junctions (50–52). p53, and oncogenic mutant p53 in particular, was previously demonstrated to stimulate topo I relaxation activity (31,53,54) and the formation of topo I double cleavage complexes causing oligonucleotide excision coupled to recombinative gap filling (55). From this, it is conceivable that stable complex formation between topo I and mutant p53 after cdk-mediated p53 phosphorylation is sufficient to inappropriately stimulate recombinase activities through recruitment to genomic DNA breaks and enzymatic activation of topo I.

**Gain-of-function in p53 mutants and genome destabilization**

It has been well-established that cancer-related p53 mutants, including p53(273H) and p53(248W), not only lost tumor suppressor functions, but also acquired additional oncogenic activities (56). In this study, we utilized cellular systems derived from p53-negative or mutant p53 cells to exclude dominant negative effects on endogenous Wtp53. p53 gain-of-function effects on genomic instability, in particular, are manifested by aberrant ploidy, increased mutation and gene amplification rates (56–58). We and others proposed that topo I mediates gain-of-function of mutant p53 (18,31,41,54). To our knowledge this is the first report that demonstrates that cyclin A1 stimulates DSB-triggered HR in cells that express mutant p53 but not in cells lacking p53, which indicates a gain-of-function. This effect is mediated by cdk2-dependent phosphorylation and by topo I. Cyclin A1
overexpression revealed a gain-of-function of mutant p53 even when directly comparing HR frequencies in cells with versus without mutant p53. This was evident after knockdown of endogenous mutant p53 (Supplementary Figure 2).

Increased topo I activities were detected in several cancer types and found to correlate with abnormal p53 function in melanomas (49). Since mutant p53 is transcriptionally defective, it is important to note that p53 is not the only regulator of cyclin A1 expression. Rather, upon mutation of TP53, alternative transcription factors, such as the oncoproteins c-myc, Six1 and PML-RARα, can induce cyclin A1 expression (59–61). In fact, elevated cyclin A1 levels have been detected in several cancers including male germ cell tumors (62), breast cancer (61) and in acute myeloid leukemia (63), whereas it is normally suppressed in most somatic cells (Figure 9). Evidence for the important role of cyclin A1 in the initiation of acute myeloid leukemia was provided by a transgenic mouse model with targeted cyclin A1 in the bone marrow (64). Most intriguingly, with respect to the observed synergistic effect of mutant p53 and cyclin A1 on recombination, Tokumaru and colleagues (65) found a negative correlation between cyclin A1 hypermethylation, with robust cyclin A1 protein expression and p53 mutation in head and neck cancer. From our results, mutant p53 is predicted to promote genome instability in tumors with cyclin A1 overexpression.

In summary, we have demonstrated that phosphorylation of p53 at distinct amino acids does differentially regulate direct DNA repair-related activities. We provide functional data, which unveil a role of the ATM/ATR-target serine 15 in fully enabling p53 to downregulate HR and of edk2-dependent phosphorylation in activating HR. Phosphorylation of p53 at serine 15 forms part of the ATM/ATR-controlled DNA damage response network, which is activated early in tumorigenesis before genomic instability occurs (66,67). p53Ser15 may, therefore, contribute at least in part through repair regulatory functions to the barrier against cancer, and indeed, recent in vivo data demonstrate that serine 15 is required for p53-mediated tumor suppression (68). On the other hand, we provide evidence that oncogenic p53 mutant proteins have lost the balance between activation and repression of recombination repair, which under conditions such as cyclin A1 overexpression may result in an increase of mutagenic DNA rearrangements such as gene amplification or loss of heterozygosity. Our data provide new answers to the longstanding question of the mechanism underlying the gain-of-function in oncogenic p53 mutants in genome instability.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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