The Myc-evoked DNA damage response accounts for treatment resistance in primary lymphomas in vivo

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In addition to the ARF/p53 pathway, the DNA damage response (DDR) has been recognized as another oncogene-provoked anticancer barrier in early human tumorigenesis leading to apoptosis or cellular senescence. DDR mutations may promote tumor formation, but their impact on treatment outcome remains unclear. In this study, we generated ataxia telangiectasia mutated (Atm)– proficient and -deficient B-cell lymphomas in Eu-myc transgenic mice to examine the role of DDR defects in lymphomagenesis and treatment sensitivity.

Introduction

Cancer development is a process against which normal cells appear to be well protected.1 As an early cellular barrier against imminent transformation, normal cells monitor latent oncogenic signals, which may result in activation of the ARF/p53 axis as an antiproliferative constraint, ultimately leading to apoptosis or cellular senescence.2 In turn, genetic defects in these oncogene-induced fail-safe programs are prerequisites for malignant conversion and thus are selected for in manifest malignancies.3–5

In addition, oncogene-related cellular stress enacting a DNA damage response (DDR) has been proposed as an ARF-independently constrained to limit aberrant cell division in early tumorigenesis via induction of apoptosis or senescence.6–10 Indeed, prototypic oncogenes such as Myc and Ras have been reported to cause DNA damage, possibly via the production of reactive oxygen species (ROS).11,12 Atm inactivation accelerated development of lymphomas, and their DNA damage checkpoint defects were virtually indistinguishable from those observed in Atm−/−-derived lymphomas that spontaneously inactivated the pro-apoptotic Atm/p53 cascade in response to Myc-evoked reactive oxygen species (ROS). Importantly, acquisition of DDR defects, but not selection against the ARF pathway, could be prevented by lifelong exposure to the ROS scavenger N-acetylcysteine (NAC) in vivo. Following anticancer therapy, DDR-compromised lymphomas displayed apoptotic but, surprisingly, no senescence defects and achieved a much poorer long-term outcome when compared with DDR-competent lymphomas treated in vivo. Hence, Atm eliminates preneoplastic lesions by converting oncogenic signaling into apoptosis, and selection against an Atm-dependent response promotes formation of lymphomas with pre-determined treatment insensitivity. (Blood. 2007;110:2996-3004)
regarding the relationship between Atm inactivation and clinical outcome remain controversial, 28, 29

A number of studies indicated a link between oncogenic Myc activation with Atm inactivation during tumorigenesis in different entities, including human B-cell lymphomagenesis. 30 In mice, extra copies of chromosome 15, to where the c-myc gene maps, were found in Atm−/− thymic lymphomas, 31 and recent studies demonstrated the collaboration of Atm loss and Myc activation in murine models of skin and lymphoid malignancies. 32, 33 Despite the increasing evidence for DDR impairment following oncogenic activation, the underlying mechanism by which oncogenes provoke spontaneous DDR defects and their subsequent clinical implications for treatment outcome remains elusive.

In this study, we used the Eμ-myc transgenic mouse model to comprehensively analyze the functional implications of genetically defined—as compared with sporadic—DDR defects on tumor biology and, in particular, treatment sensitivity in a large series of primary B-cell lymphomas. Our findings dissect Myc-driven activation of the DDR as a ROS-initiated and Atm-governed proapoptotic process that is selected against during lymphoma formation and demonstrate the profound impact of predefined or spontaneously acquired DDR lesions on the efficacy and long-term outcome to anticancer treatments in vitro and in vivo. Furthermore, we explore in vivo how pharmacologic intervention might preserve an intact DDR to subsequently improve the efficacy of DNA-damaging chemotherapy.

Materials and methods
Mice, cells, and drug treatments
All animal protocols used in this study were approved by the governmental review board (Landesamt für Gesundheit und Soziales, Berlin, Germany). Lymphomas harboring loss-of-function lesions were generated by intercrossing Eμ-myc transgenic mice 34 with mice harboring targeted deletions at the Atm, 35 p53, 36 ARF, 37 or bim 38 locus (all genotypes in Bl6/129 background). Monitoring of lymph nodes, processing and fixation of tissues, isolation of lymphoma cells or splenic B-lymphocytes (purified via immunobead separation [Miltenyi Biotec, Bergisch Gladbach, Germany]), and short-term cultivation and transplantation of lymphoma cells were performed as described. 39 Preparative samples are derived from mice of approximately 25 days of age with no signs of lymph node or spleen enlargement and absence of leukemia. For in vitro treatments, tumor-bearing mice received a single dose of either 4 Gy γ-irradiation or 300 mg/kg body weight of cyclophosphamide (CTX) intraperitoneally. 4 N-acetyl cysteine (NAC; Hexal, Holzkirchen, Germany) was applied as a 0.5% wt/vol based solvent.

Analysis of chromosomal aberrations, V(DJ) recombination, and genomic mutations
Chromosomal aberrations in primary lymphomas were assessed in stained metaphase spreads by fluorescence microscopy and spectral karyotyping. 5 To detect V(DJ) recombinations, IgH loci were amplified in a first genomic polymerase chain reaction (PCR) using 9 VH- or DH-specific and 1 JH-specific primers, followed by a second-round PCR containing a single VH or DH primer together with a nested JH primer. 3 A productive PCR indicates an accomplished rearrangement. Primer sequences are available upon request. 55 mutations within exons 5 to 8 were assessed by reverse transcriptase (RT)-PCR—based sequencing and homozgyous gross deletions at the INK4a/ARF locus by a genomic multiplex PCR. 4 Using primers spanning exon 1β to exon 2, RT-PCR—based expression analysis of ARF transcripts (and TATA-box binding protein [TBP] as an internal control) was carried out as described. 5

Protein analysis by immunophenotyping, immunostaining, and immunoblot
Immunophenotyping by flow cytometry as well as antigen detection by immunohistochemistry and by immunofluorescence of permeabilized cyto-}

Retroviral gene transfer, siRNA-mediated knockdown, and inducible gene expression
Retroviral gene transfer using MSCV-bcl2-ires-GFP has been described. 39 Knockdown of specific transcripts was achieved by stable introduction of retroviral siRNA cloned into the pSuper Retro system (Oligo Engine, Seattle, WA; [Atm-siRNA: 5′-GCTAGACTCTAATACTG-3′; PP5-siRNA: 5′-GACAGAGAGATTAGTG-3′]). MycERTAM cells, generated by infecting mouse embryo fibroblasts (MycERTAM-MEFs) or (ARF-deficient) NIH3T3 fibroblasts (MycERTAM-3T3) with the MSCV-MycERTAM-blasticidin retrovirus, were treated with 1 μM 4-hydroxytamoxifen (OHT; Sigma-Aldrich) or the equivalent volumes of the ethanol-based solvent.

Analysis of cellular integrity, DNA strand breaks, and metabolic condition
Cellular viability and DNA content were assessed by 7-amino-actinomycin D (Sigma-Aldrich) dye exclusion or propidium iodide–based DNA staining measured by flow cytometry and quantified using ModFit LT 2.0 Software (Verity Software House, Topsham, ME). 5 Cellular senescence was detected by staining for the senescence-associated β-galactosidase activity in cytospin preparations at pH 5.5 as described. 5 ROS levels were measured by flow cytometry in freshly isolated cells after incubation in 10 μM 2′, 7′-dichlorodihydrofluorescein diacetate (Gibco, Karlsruhe, Germany). Oxidatively modified DNA was quantified by the formamidopyrimidine-DNA glycosylase (Fpg)–COMET assay. 40 Briefly, cells were embedded in low-melting-point agarose, lysed, and exposed to Fpg (New England Biolabs, Frankfurt am Main, Germany). After gel electrophoresis and DAPI staining, mean tail moments of randomly chosen comets were quantified as the relative increment following Fpg treatment using the CometScore (TriTek, Summerduck, VA) software V1.5 for analysis. γ-H2AX foci, detected by immunofluorescence, were quantified by counting at least 25 cells per sample. Apoptosis-related DNA fragmentation was visualized using a fluorescence-based terminal deoxy-nucleotidyl transferase dUTP nick end labeling (TUNEL) assay (Roche Diagnostics, Mannheim, Germany) with DAPI as a counterstain in some experiments. Cells were mounted using VECTASHEILD (Vector Laboratories, Burlingame, CA) mounting medium for fluorescence. Images of cells were captured by the Axioplan fluorescence microscope with Plan-Neofluar objectives (100×/1.30 NA oil or 40×/0.75 NA) (Carl Zeiss, Jena, Germany) attached to a SPOT RT camera model 2.3.1 and imported into SpotAdvanced v3.5 software (Diagnostic Instruments, Sterling Heights, MI).

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α-tubulin (Sigma-Aldrich), γ-H2AX (Upstate, Temecula, CA), Ki67 (Dako, Hamburg, Germany), caspase-3 (Cell Signaling Technology), cleaved caspase-3/Asp175 (Cell Signaling Technology), CD45/B220 (BD Pharmingen), CD43 (BD Pharmingen), and CD90.2/Tyrl.2 (BD Pharmingen) were used, if required, with corresponding peroxidase- and fluorescence-conjugated secondary antibodies (GE Healthcare, München, Germany; Molecular Probes, Karlsruhe, Germany; Santa Cruz Biotechnology), and visualized as described.5,39

Results

Myc provokes an Atm-dependent DDR leading to apoptosis in preneoplastic B cells in vivo

NIH3T3 fibroblasts stably expressing an OHT–inducible c-Myc/estrogen receptor fusion protein (MycER<sup>TAM</sup>-3T3) confirmed that acute induction of Myc—like exposure to the DNA-damaging anticancer agent adriamycin (ADR)—elicits a PP5/Atm-dependent acute induction of Myc—like exposure to the DNA-damaging agent adriamycin (ADR), resulting in apoptosis (Figure S1, available on the Blood website; see the Supplemental Materials link at the top of the online article). To examine the role of the Atm-governed DDR in response to oncogenic Myc signaling in vivo, B-cell lymphoma–prone <i>Equi</i>-myc transgenic mice heterozygous for germ-line deletions at the Atm allele were intercrossed with Atm<sup>E</sup>-deficient nontransgenic and preneoplastic B lymphocytes consisting of Atm, Atm-P-S1987, p53, p53-P-S18, and caspase-3 with its proapoptotic cleavage product in lysates of immunobead-selected Atm-proficient and Atm-deficient cells or B cells lacking constitutive Myc signaling, indicating that Atm is required for Myc-provoked apoptosis in vivo. Accordingly, when the steady state amount of spontaneous apoptosis was determined as a fraction of DNA-fragmented TUNEL-positive cells, much more apoptosis was detected in the <i>Equi</i>-myc transgenic compared with the nontransgenic population, whereby the amount of Myc-induced cell death was significantly reduced in the absence of Atm (P = .037; Figure 1B). Thus, the activated Atm/p53 signature in the oncogene-exposed murine B-cell compartment recapitulates the occurrence of a proapoptotic DDR as previously detected in early hyperproliferative stages of human (pre)carcinomatous tissue samples.7,8

Atm inactivation collaborates with constitutive Myc expression in lymphomagenesis

Myc-driven lymphomas arising in Atm<sup>+/−</sup> mice, hereafter referred to as controls, became detectable as a palpable lymphadenopathy with associated leukemia at a median age of 101 days, whereas Atm<sup>−/−</sup> mice displayed lymphomas with a median onset of 50 days, significantly earlier (P < .001) (Figure 1C). Lymphoma development in Atm<sup>+/−</sup> mice was virtually indistinguishable from that observed in the Atm<sup>++</sup> group. Notably, all Equi-myc–initiated tumors were B220<sup>+</sup> and Thy1.2<sup>−</sup> (data not shown), indicating that Atm-deficient Equi-myc mice succumbed to B-cell lymphomas before thymic T-cell lymphomas that typically arise in nontransgenic Atm<sup>−/−</sup> mice could form and disseminate.35 Manifest lymphomas displayed reduced amounts of spontaneous apoptosis in the absence of Atm, as seen during tumor development for preneoplastic stages, while the proliferation rate appeared to be unaffected by the Atm status, indicating that the Atm-governed S-phase checkpoint is not challenged or became dysfunctional in constitutive Myc-driven lymphomagenesis (Figure S2A,B). Furthermore, Atm<sup>−/−</sup> lymphomas grew more aggressively than controls, as reflected by their dissemination into nonlymphoid organs such as lung and pancreas (Figure 1D).

Both control and Atm<sup>−/−</sup> lymphomas accomplished clonal V(D)J recombination and progressed to a more mature, CD43<sup>−</sup>B-cell state, indicating that differentiation was not blocked in the absence of Atm (Figure S2C and data not shown). Spectral karyotyping of Atm<sup>−/−</sup> samples confirmed no signs of gross aneuploidy but revealed a higher frequency of nonrecurrent translocations (6 of 7 Atm<sup>−/−</sup> compared with 2 of 7 control samples tested; Figure S2D-E), highlighting the role of Atm in proper DNA double-strand break repair.

Hence, Atm resides in a proapoptotic signaling pathway that is activated during Myc-driven lymphomagenesis to counter malignant transformation, and inactivation of Atm alleles promotes development of apoptotically compromised lymphomas in a nonhaploinsufficient fashion.

The Atm/p53 and Arf/p53 axes cooperate as proapoptotic growth restraints in Myc-driven lymphomagenesis

Inactivation of Arf, the Myc-inducible tumor suppressor controlling p53 protein levels via interaction with Mdm2,42,43 has been
that became evident after exposure to ADR (Figure 2D). Hence, the Atm-mediated antioncogenic response complements ARF signaling to the common downstream effector p53 to counter onco-
genic transformation via execution of apoptosis.

A subset of manifest Myc lymphomas displays functional defects in the Atm-governed DDR

Because the Myc-initiated DDR signature is present in preneoplastic B cells (Figure 1A), we hypothesized that full-blown malignancies might have selected for Atm-related DDR defects during lymphomagenesis. We therefore scanned primary control lymphomas for the presence or absence of activated DDR components by immunoblot analysis. Beside detectable basal levels of phosphorylation-activated Atm (ie, Atm-P-S1987) in some cases, 6 of 12 control lymphomas tested exhibited considerable amounts of p53-P-S18, whereas these phospho-proteins were not detectable in normal B cells (Figure 3A) and Atm−/− lymphomas (Figure 3B). No activated DDR mediators could be visualized in any of the remaining control lymphomas tested despite comparable Myc expression levels, indicating either lack of basal activation or structural inactivation of this pathway in about half of the samples.

Therefore, we tested whether stimulation by exogenous DNA damage may either result in a detectable DDR or might unmask checkpoint defects such as impaired apoptotic activation or “drug damage–resistant cell cycle progression into S phase” reminiscent of the entirety of lymphomas presented in panel A.

Figure 3. DDR-compromised Myc lymphomas share checkpoint defects with Atm-deficient lymphomas. (A) Immunoblot analysis of Myc, Atm, Atm-P-S1987, p53, p53-P-S18, and tubulin as a loading control in immunobead-isolated normal B cells and 12 randomly selected control lymphomas. (B) Immunoblot analysis of the presence or absence of activated DDR components by immunoblot analysis. Beside detectable basal levels of phosphorylation-activated Atm (ie, Atm-P-S1987) in some cases, 6 of 12 control lymphomas tested exhibited considerable amounts of p53-P-S18, whereas these phospho-proteins were not detectable in normal B cells (Figure 3A) and Atm−/− lymphomas (Figure 3B). No activated DDR mediators could be visualized in any of the remaining control lymphomas tested despite comparable Myc expression levels, indicating either lack of basal activation or structural inactivation of this pathway in about half of the samples.

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of the “radiosensitive DNA synthesis” checkpoint defect known from Atm−/− cells. Control lymphomas that formed with detectable basal p53-P-S18 levels consistently produced strongly induced p53-P-S18 levels accompanied by high amounts of apoptosis-related cleaved caspase-3 after exposure to ADR, a signature that was entirely absent in control lymphomas with undetectable basal p53-P-S18 levels and in Atm−/− lymphomas prior to and after ADR (Figure 3B top). Thus, control lymphomas with detectable and ADR-inducible p53-P-S18 levels were considered DDR competent (“DDR+,” 6 of 12 cases tested), whereas those not producing any p53-P-S18 band (despite wild-type p53 genes by sequencing analysis [of 12 control and 6 Atm−/− lymphomas each]; data not shown) formed the DDR-impaired (“DDR−,” 6 of 12 cases tested) group. Importantly, when the same individual lymphoma cell populations were exposed to ADR after stable infection with a bcl2 retrovirus to block apoptosis, Atm−/− and DDR− control lymphomas but not DDR+ lymphomas exhibited considerable DNA synthesis activity as indicated by cells in S phase (Figure 3B bottom). Hence, reduced basal levels of DDR components selected for in control lymphomas reflect genetic lesions that produce a drug damage-resistant and apoptotically compromised phenotype virtually indistinguishable from the checkpoint defects detectable in Atm−/− lymphomas.

Lymphomas developing under ROS scavengers preserve an intact DDR machinery

Given our observation that Myc signaling—like treatment with a DNA-damaging compound—provokes PPS/Atm-dependent p53 activation (Figure S1), we aimed to assess marks of DNA damage in Myc-driven lymphomas with different Atm background. ROS—a potential cause of DNA lesions that have been associated with Myc activation in vitro—were detectable at significantly increased levels in Eμ-myc lymphoma cells compared with nontransgenic Atm+/− and Atm−/− B cells (Figure 4A). Consequently, quantification of DNA damage as a result of oxidative stress using the Fpg-COMET assay unveiled higher numbers of DNA strand breaks in Myc-driven lymphomas compared with nontransgenic B cells irrespective of their Atm status (Figure 4B). Phosphorylation of the histone H2A variant, H2AX, at serine 139 is known as an early, Atm-mediated response to DNA double-strand breaks, resulting in γ-H2AX foci at the sites of DNA damage. The number of γ-H2AX foci was found markedly increased under constitutive Myc expression compared with nontransgenic B cells and appeared to be independent of Atm (Figure 4C). Most γ-H2AX foci measured in manifest Eμ-myc lymphomas reflect sites of endogenous DNA damage rather than marks at apoptotic strand breaks, because similar levels of γ-H2AX foci were also detected in Atm−/− and control lymphomas stably overexpressing the antiapoptotic Bcl2 protein. Because γ-H2AX foci may form in an Atm-independent fashion via substituting action of the related Atr (ataxia-telangiectasia Rad3-related) kinase, it is conceivable that comparable steady state levels of γ-H2AX foci in control versus Atm−/− lymphoma cells may result from the extended persistence of unresolved DNA strand breaks in the absence of proper Atm function. Indeed, immunoblot analysis confirmed the presence of serine 431-phospho–activated Atr (Atr-P-S431) in Atm−/− and, less profoundly, in control lymphomas but not in nontransgenic B cells (Figure 4D). Importantly, LPS-enforced proliferation of nontransgenic primary B cells to an extent comparable with Myc-driven lymphoma cell proliferation produced a much smaller increase of ROS levels (1.5-fold versus about 5-fold) and was not associated with enhanced numbers of γ-H2AX foci (Figure 4E). Thus, the profound increase in marks of DNA damage in Myc-driven cells with or without Atm is predominantly linked to genotoxic but not to hyperproliferative signaling emanating from the oncogene.

These findings indicated that Myc-evoked oxidative stress acts as the upstream signal that instates the selective pressure against a functional DDR machinery as observed in a subset of control lymphomas (Figure 3A). Therefore, we examined lymphoma development when oxidative stress is pharmacologically blunted by the ROS scavenger NAC, which prevents DNA damage and protects cells from a subsequent PPS/Atm-dependent DDR elicited

Figure 4. Myc induces DNA damage via ROS in vivo. Comparison of freshly isolated Eμ-myc lymphoma cells and immunobead-selected nontransgenic B cells derived from an Atm+/− or an Atm−/− background (n = 4 individual samples per genotype). (A) 2′,7′-Dichlorodihydrofluorescein diacetate–based flow cytometric analysis of cellular ROS levels (left); representative example visualized by fluorescence microscopy (right). Original magnification, ×200. (B) Oxidative DNA damage measured as the relative induction of mean tail moments in the COMET assay prior to and after treatment with Fpg; quantification (left); representative examples of comets compared with nontransgenic B cells (right). (C) γ-H2AX foci per cell in cytosin preparations; same cell populations stably expressing Bcl2 to block apoptosis in the right panel; quantification (left) and representative examples (right). (D) Immunoblot analysis of total Atr, Atr-P-S431, and tubulin as a loading control in nontransgenic B lymphocytes as compared with representative control and Atm−/− Eμ-myc lymphomas. (E) Proliferation (left, cells with S-phase DNA content), ROS induction (middle, as in panel A), and γ-H2AX foci (right, as in panel C) in primary nontransgenic B cells displayed as relative values at 48 hours after LPS stimulation (50 μg/ml) compared with non-LPS.
by Myc in established cell lines in vitro (Figure S3A).12,50 Eµ-myc transgenic mice were continuously exposed to NAC-supplemented drinking water starting at a midembryonic age. Importantly, when γ-H2AX foci were quantified in NAC-pretreated versus untreated lymphomas at manifestation, significantly less foci were found in NAC-protected control lymphomas (P = 0.016; Figure 5A), indicating that Myc-originated signals otherwise leading to DNA damage were blunted by the ROS scavenger. Consequently, lymphomas that formed in NAC-treated animals displayed significantly less TUNEL-positive apoptotic cells compared with untreated controls (P = 0.024; data not shown). This difference was even larger in cells isolated from Eµ-myc transgenic mice at a preneoplastic stage (P = 0.006; Figure 5B). Notably, Myc-inducible high ARF levels remained unchanged in response to NAC (Figure S3B-D), suggesting that NAC action attenuates an oncogene-evoked DDR but not oncogenic signaling to ARF. Notably, NAC-treated mice (n = 19) developed lymphomas with latencies comparable with untreated controls (data not shown), presumably due to a neutralizing net effect of the lowered induction of the DDR-initiated proapoptotic machinery and, simultaneously, the lowered selective pressure against this machinery and a slight reduction in proliferation when NAC is present. Importantly, NAC-protected control lymphomas, although lacking basal activation of DDR components (to compare with non-NAC–treated control lymphomas, see Figure 3A), consistently retained a functional DDR that became apparent only after exposure to ADR in vitro (Figure 5C, compare with Figure 3B). Therefore, NAC treatment effectively reduces marks of endogenous DNA damage and attenuates DDR-initiated apoptosis in Myc-driven lymphomagenesis, thereby uncovering ROS as critical mediators of the Myc-provoked DDR that is subsequently selected against in manifest lymphomas.

Preserving a functional Atm-governed DDR determines superior treatment outcome

Given the impaired ADR-inducible checkpoint responses in Atm⁻/⁻ and DDR-compromised control lymphomas in vitro, we tested whether a functional DDR machinery at diagnosis may predict superior short-term and long-term outcome to various DNA-damaging anticancer therapies in vitro and in vivo. Firstly, freshly isolated control lymphoma cells belonging to the DDR⁺, the DDR⁻, or the NAC-protected group as well as Atm⁻/⁻ and p53⁻/-null lymphomas (ie, Eµ-myc lymphomas that formed in p53⁻/- mice and lost the remaining p53 wild-type allele9) were exposed to increasing doses of ADR in vitro, and viability was assessed after 19 hours. DDR⁺ control lymphomas—like lymphomas that formed under NAC—displayed excellent sensitivity to ADR (Figure 6A). In contrast, Atm⁻/⁻ lymphomas and DDR⁻ control lymphomas exhibited profound drug resistance only surpassed by the drug-insensitive p53⁻/-null lymphomas. Notably, the poor sensitivity of Atm⁻/⁻ lymphomas was not due to additional genomic alterations, because preneoplastic Atm⁻/⁻ B cells were equally insensitive to ADR (Figure S4A). To further assess short-term drug-induced apoptosis in lymphomas of the respective groups growing in their natural environment, lymph node sections from mice harboring transplanted lymphomas obtained 4 hours after systemic treatment with the anticancer agent CTX were subjected to TUNEL staining. Again, DDR⁺ control and NAC-protected control lymphomas produced massive apoptosis in sit following CTX, whereas...
apoptosis was much more sparse in CTX-treated Atm−/− and DDR− control lymphomas (Figure 6B).

Recent reports on cellular senescence imposed as another tumor-suppressive DDR to oncogenic signals in early (pre)malignant lesions suggested that DDR defects may be selected for to overcome this barrier.5,9,10,14,51,54,55 Moreover, disruption of the senescence program might impact on treatment outcome as well, because drug-inducible senescence was shown to extend survival of mice harboring Myc lymphomas.39 Therefore, we tested whether drug-inducible senescence remains available in the absence of Atm. Contrasting the premature senescent phenotype of Atm−/− MEFs,52 Myc-driven Atm−/− lymphomas, like controls, apparently form by either never imposing or bypassing oncogene-provoked senescence. However, both DDR− control and Atm−/− lymphoma cells, infected with Bcl2 to protect from apoptosis, entered cellular senescence 7 days after exposure to ADR in vitro as demonstrated by senescence-associated β-galactosidase staining (Figure 6C). Hence, disruption of the DDR by spontaneous defects or loss of Atm genes compromises chemotheraphy-inducible apoptosis while leaving cellular senescence as a drug-responsive program available.

Finally, we asked whether these differences in apoptotic short-term responsiveness would translate into different long-term outcome in vivo. To this end, mice harboring transplanted primary lymphomas received a single 4 Gy dose of total body γ-irradiation when lymph node enlargements became palpable. Using this well-tolerated dose, most animals entered a remission but all eventually relapsed. Mice harboring DDR− control lymphomas achieved the most durable remissions, whereas those bearing DDR− lymphomas displayed the shortest benefit from treatment (Figure 6D). The mean time to relapse in the Atm−/− lymphoma group was significantly shorter compared with the DDR− (P = .004) but not to the DDR− control lymphoma group. The group of mice harboring NAC-protected lymphomas behaved similarly to the DDR− control lymphoma group and responded significantly better when compared with the DDR− control lymphoma group (P < .001). Hence, preservation of an intact DDR at diagnosis is critical for the execution of an apoptotic response and long-term outcome after DNA-damaging therapy in vivo.

Discussion

Our study is the first comprehensive genetic approach that mechanistically links oncogene-provoked DDR defects to treatment outcome in aggressive lymphomas in vivo. Using a well-established Myc-driven mouse model for the generation of primary lymphomas with defined genetic lesions that recapitulate genetic and histopathological features of human non-Hodgkin lymphomas,53 we identified the 2-step DDR induction by ROS initiation followed by Atm transduction as the critical anticancer restraint that complements the ARF axis in response to Myc signaling. While in nononcogene-driven scenarios the precise impact of Atm inactivation on cell growth and death remained controversial,27 the collaboration of Atm defects with oncogenic Myc action unmasks apoptosis as the key Atm-mediated effector mechanism in response to Myc. Importantly, our data not only demonstrated a significant acceleration of Myc-driven lymphogenesis in an Atm−/− background but unveiled, as a proof of relevance, spontaneous selection against a functional DDR during lymphoma formation, and this group indeed displayed a tendency toward shortened tumor latencies (mean onset, 90 days for DDR− versus 120 for DDR+ lymphomas).

We considered—in addition to ROS—other mechanisms by which oncogenes might activate the DDR machinery. Stalled DNA replication forks due to Myc-induced hyperproliferation may produce DNA damage that triggers an Atr/Nbs1/Chk1-governed checkpoint.56-58 Indeed, the increased amounts of Atr-P-S431 detected in Myc-driven lymphomas were suggestive for hyperreplication stress. Conversely, the slight decrease of proliferation in NAC-treated lymphomas might have been sufficient to no longer produce aberrant DNA replication intermediates, thereby canceling a key trigger of the Atr-mediated DDR checkpoint. However, evidence presented here argues for ROS as the critical oncogene-evoked initiator of proapoptotic DNA damage signaling. Firstly, enforced proliferation by LPS in primary nontransgenic B cells to a level reminiscent of mitogenic, constitutive Myc signaling failed to generate a profound increase in ROS and did not enhance γ-H2AX marks of DNA damage, thereby suggesting that Myc-evoked ROS and Myc-driven proliferation are largely independent effector functions. Secondly, acute induction of Myc in fibroblasts (ie, before a relevant impact on proliferation could have been observed) resulted in a DDR that was dependent on ROS, as demonstrated pharmacologically, and on Atm, as confirmed genetically. Thirdly, Eμ-myc transgenic preneoplastic B cells (ie, prior to the potential acquisition of more complex genomic alterations during transformation) already displayed activated DDR components associated with cleavage of caspase-3, and this proapoptotic signature was entirely absent in the Atm-deficient counterpart despite Atr activation at this time (data not shown). In accordance with Atr’s ability to substitute for Atm with delayed kinetics, it is not surprising to detect higher activation of Atr in Atm−/− lymphomas, but apparently not as a functionally relevant component of proapoptotic signaling.49 Fourthly, biallelic deletion of Atm dramatically shortened tumor latencies in Myc-driven tumor models irrespective of the activity of the Atr kinase. Taken together, while constitutive Myc signaling, presumably via hyperreplication stress, is undoubtedly associated with Atr activation, our data support the view that ROS are the critical mediators of proapoptotic Myc-induced DNA damage sensed and transduced primarily by Atm.

Our data add to the intense scientific debate on relative tumor-suppressive contributions of ARF-mediated versus DNA damage–initiated signals as upstream components of p53. Myc-driven hyperproliferation imposes an ARF response via E2F transcription factors,59 although other Atm-independent, possibly metabolic pathways might also be involved in ARF induction. Recently, ARF inactivation was shown to disrupt p53’s tumor-suppressive function in response to chemically induced oncogenic stresses in vivo, but no DDR-defective genetic model such as the Atm−/− mouse was included in this setting.50 In the Eμ-myc transgenic mouse, tumor onset was accelerated by defined genetic ablation of Atm through impairment of p53-mediated apoptosis. Although p53 protein levels remained very low in response to even drug-induced DNA damage due to lack of protein-stabilizing posttranslational modifications in the absence of Atm, p53 apparently was still susceptible to ARF-mediated oncogenic signals. This duality of oncogenic signaling to p53 turned out to be amenable to pharmacologic dissection, because antioxidative intervention ablated the oncogene-evoked DDR without affecting Myc-mediated ARF induction in fibroblasts and B cells. As a consequence, neither Atm loss nor NAC treatment alleviated the pressure to inactivate components of the ARF/p53 axis in primary lymphomas. Thus, our data support the view that ROS on one side and oncogenic
hyperreplication on the other side challenge cellular countermeasures whose tumor-suppressive potential is rather complementary than overlapping. Accordingly, human tumors—and lymphoid malignancies in particular—not only frequently exhibit INK4a/ARF mutations but may present with simultaneous defects in the DDR machinery, namely at the level of the Atm kinase.61,62

Whether selective defects in certain tumor suppressors or increased genetic instability may ultimately drive tumor development and progression remains an issue of debate. In our Eµ-myc model, compromised apoptosis as a consequence of Atm loss became already apparent in preneoplastic Atm−/− B cells, but this defect did not permit the survival of cytogenetically grossly aberrant lymphomas; we rather observed lymphomas with close to normal chromosome counts harboring translocations indicative of improper DNA double-strand break repair in the absence of Atm. Likewise, nontransgenic Atm−/− mice develop T-cell lymphomas that select for specific chromosome alterations (for example, gains that include the c-myc locus) instead of randomly accumulating genomic aberrations.31 Hence, genetic instability may occur as a byproduct of Atm dysfunction, but Myc-driven lymphoma formation relies on sufficient suppression of latent proapoptotic capabilities by targeting both the Atm/p53 and ARF/p53 cascades.

Oncogene-derived ROS have important implications for the outcome of anticancer therapy, because their excessive presence may primarily activate but ultimately select against an intact proapoptotic DDR machinery at diagnosis or later during tumor progression (Figure 3 and data not shown). Recently, a pharmacologic approach was proposed to preferentially kill cancer cells by taking advantage of their abnormal ROS levels,63 which might be of particular relevance to target tumors that failed to maintain an increased genetic instability may ultimately drive tumor development.31 Hence, genetic instability may occur as a byproduct of Atm dysfunction, but Myc-driven lymphoma formation relies on sufficient suppression of latent proapoptotic capabilities by targeting both the Atm/p53 and ARF/p53 cascades.

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The Myc-evoked DNA damage response accounts for treatment resistance in primary lymphomas in vivo

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