Resistance to DNA-damaging treatment in non-small cell lung cancer tumor-initiating cells involves reduced DNA-PK/ATM activation and diminished cell cycle arrest

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Increasing evidence suggests that tumor-initiating cells (TICs), also called cancer stem cells, are partly responsible for resistance to DNA-damaging treatment. Here we addressed if such a phenotype may contribute to radio- and cisplatin resistance in non-small cell lung cancer (NSCLC). We showed that four out of eight NSCLC cell lines (H125, A549, H1299 and H23) possess sphere-forming capacity when cultured in stem cell media and three of these display elevated levels of CD133. Indeed, sphere-forming NSCLC cells, hereafter called TICs, showed a reduced apoptotic response and increased survival after irradiation (IR), as compared with the corresponding bulk cell population. Decreased cytotoxicity and apoptotic signaling manifested by diminished poly (ADP-ribose) polymerase (PARP) cleavage and caspase 3 activity was also evident in TICs after cisplatin treatment. Neither radiation nor cisplatin resistance was due to quiescence as H125 TICs proliferated at a rate comparable to bulk cells. However, TICs displayed less pronounced G2 cell cycle arrest and S/G2-phase block after IR and cisplatin, respectively. Additionally, we confirmed a cisplatin-refractory phenotype of H125 TICs in vivo in a mouse xenograft model. We further examined TICs for altered expression or activation of DNA damage repair proteins as a way to explain their increased radio- and/or chemotherapy resistance. Indeed, we found that TICs exhibited increased basal H2AX (H2A histone family, member X) expression and diminished DNA damage-induced phosphorylation of DNA-dependent protein kinase (DNA-PK), ataxia telangiectasia-mutated (ATM), Krüppel-associated protein 1 (KAP1) and monoubiquitination of Fanconi anemia, complementation group D2 (FANCD2). As a proof of principle, ATM inhibition in bulk cells increased their cisplatin resistance, as demonstrated by reduced PARP cleavage. In conclusion, we show that reduced apoptotic response, altered DNA repair signaling and cell cycle perturbations in NSCLC TICs are possible factors contributing to their therapy resistance, which may be exploited for DNA damage-sensitizing purposes.

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Lung cancers (LCs) are among the most common human malignancies worldwide. Despite the use of aggressive multimodal chemo- and radiotherapy (CT/RT), the 5-year survival of LC patients remain as low as 15%. Resistance to CT/RT is a commonly encountered problem in both treatment-naive LC patients and during the course of CT/RT. Yet, the molecular mechanisms underlying CT/RT refractoriness are complex and still largely undefined. Identification of such mechanisms could reveal crucial tumor-specific aberrations that are amenable to targeted intervention, as well as generating biomarker profiles that will enable individualized therapy.

Emerging evidence indicates the existence of a rare fraction of tumor cells endowed with properties characteristic of somatic/embryonic stem cells, cancer stem cells or, as hereafter called, tumor-initiating cells (TICs). TICs typically show elevated expression of the stem cell-associated markers CD133, SRY-related HMG-box 2 (Sox2), octamer-binding transcription factor 4 (Oct4) and Nanog, display high tumorigenic potential and have been associated with a CT/RT-refractory phenotype. TICs can be enriched by surface marker expression or their ability to form spheres in culture, as demonstrated in different tumor types including non-small cell lung cancer (NSCLC). One of the most widely used...
markers is CD133, a cell surface transmembrane glycoprotein that has been postulated as a useful marker for stem cells in a variety of normal and tumor tissues.8–10

Few cellular signaling networks that confer resistance to conventional CT/RT have been unequivocally demonstrated in TICs. Nevertheless, enhanced DNA repair capacity has been implicated as a likely cause of increased CT/RT resistance in this cell population in gliomas and breast cancer.11–13 With respect to NSCLC, few studies have addressed if a TIC phenotype contribute to DNA-damaging treatment resistance and if it involves altered DNA repair signaling. To address this issue, we cultured tumor spheres from a panel of LC cell lines. We show that this approach enabled enrichment of sphere-forming TICs with increased CD133 expression compared with bulk cells. The TICs were more resistant to RT- and cisplatin-induced cell death and show diminished apoptotic response in vitro and in vivo in mouse tumor xenografts. On the molecular level, our analyses of NSCLC TICs demonstrate aberrant DNA damage response (DDR) due to inadequate activation of ataxia telangiectasia-mutated (ATM), DNA-dependent protein kinase, catalytic subunit (DNA-PKcs), Krüppel-associated protein 1 (KAP1) and Fanconi anemia, complementation group D2 (FANC D2), leading to compromised cell cycle checkpoints. As a proof of principle, ATM inhibition in bulk cells increased their cisplatin resistance, as demonstrated by reduced poly (ADP-ribose) polymerase (PARP) cleavage. This is the first report showing that reduced activation of DDR can contribute to CT/RT resistance in TICs. Such differences in DDR signaling between TICs and bulk NSCLC cells may reveal important targetable pathways and allow for novel combination regimen with the potential to improve therapeutic outcome of LC disease.

Results

NSCLC cell lines contain cells with sphere-forming capacity indicative of a tumor-initiating phenotype. A panel of eight NSCLC cell lines were cultured for 7–14 days under conditions favoring stem cell growth and analyzed for their sphere-forming capacity. Four formed spheres (A549, H23, H1299 and H125), while one cell line (U-1752) could grow in stem cell media despite a lack of sphere-forming capacity (Figure 1). The time required for sphere formation varied among the cell lines as A549, H23 and H1299 cells developed distinct large spheres after 7 days, while H125 spheres still had adherent clusters along with detached spheres.

Expression of stem cell markers are increased in TICs. Expression levels of stem cell markers were characterized in the sphere-forming NSCLC TICs after 7–14 days of culture in stem cell media. The CD133 protein was expressed in 4% of H125 TICs and 10% of A549 TICs, a clear enrichment compared with the corresponding bulk cells, arbitrarily set to 1% (Figure 2a). Accordingly, mRNA levels of CD133 were increased in TICs from H125, A549 and H1299 cells (Figure 2b). The stem cell markers Sox2, Nanog and Oct4 all displayed increased mRNA expression at day 14 in A549 and H1299 TICs, whereas only Sox2 was increased in H23 TICs (Figure 2c). By contrast, increased expression of these markers was not observed in H125 TICs.

NSCLC TICs display resistance to RT. In breast cancer, glioma and medulloblastoma tumor cells with a TIC phenotype are resistant to RT.14 Here we addressed if this also is evident for NSCLC TICs. As expected, irradiation (IR) resulted in a dose-dependent reduction in the clonogenicity of H125 bulk cells, in contrast to H125 TICs, which were highly refractory to IR at 2 and 4 Gy and only sensitive to 8 Gy (Figures 3a and b). Similarly, A549 TICs were more resistant than A549 bulk cells to IR at 2, 4 and 8 Gy, even though A549 bulk cells were less sensitive than H125 bulk cells. The SF2 (surviving fraction at 2 Gy) value was increased from 0.36 to 0.66 in H125, and 0.53 to 0.71 in A549 TICs compared with bulk cells.

NSCLC TICs are more resistant to cisplatin compared with bulk cells. We next examined if NSCLC TICs were more resistant to CT using cisplatin. Notably, H125 and A549 TICs were significantly more resistant to cisplatin treatment compared with bulk cells with a relative difference in cell viability of 24 and 29% at $1\mu M$ cisplatin, respectively ($P<0.05$; Figure 3c). The area under the curve after cisplatin treatment was 1.5-fold higher in H125 TICs and 2.3-fold in A549 TICs. Transporter proteins may influence the intracellular concentration and response to cisplatin;15 however, equal mRNA expression of the major uptake transporter, copper transporter 1 (CTR1, SLC31A1 (solute carrier family 31, member 1)) and only moderately increased levels of the efflux protein ATPase, Cu$^{2+}$ transporting, β polypeptide (ATP7B) were found in H125 TICs (Supplementary Figure 1A), indicating the lack of large differences in uptake or export of cisplatin. In addition, cisplatin adduct formation was comparable in bulk cells and TICs (cisplatin 40 µM, 6 h and 10 µM, 24 h), hence ruling out inefficient drug accumulation as the main cause for the altered cisplatin sensitivity (Supplementary Figure 1B).

Increased RT/CT resistance in NSCLC TICs is not solely attributed to differences in proliferative capacity of TICs. Stem cells are known to be in a quiescent state and it has been assumed that TICs as their malignant counterpart may be similarly less proliferative, and hence not as susceptible to conventional CT and RT.16 However, few studies have experimentally addressed the cell cycle distribution of TICs after DNA damage, and for NSCLC no such data are at hand; therefore, proliferation rate and cell cycle distribution was assessed (Figure 4). As judged by the dilution of carboxyfluorescein diacetate N-succinimidy ester (CFSE)-associated fluorescence signals after 72 h, untreated H125 TICs divided at the same rate, whereas A549 TICs proliferated at roughly one-third of the growth rate of their bulk cells (Figure 4a). While the growth of H125 and A549 bulk cells was reduced after IR and cisplatin treatment, the post-treatment proliferation of TICs showed no marked changes apart from a small reduction after the higher dose of cisplatin (10 µM) in H125 TICs, further confirming the therapy-resistant phenotype shown in Figure 3.
NSCLC TICs show impaired cell cycle arrest after DNA-damaging treatment. Reduced checkpoint response was observed in TICs after DNA-damaging treatment (Figures 4b and c). IR (8 Gy) arrested both H125 and A549 bulk cells in the G2/M phase at 24 h. By contrast, almost no (H125) or a less prominent G2 block (A549) was elicited in TICs (Figure 4b). Cisplatin treatment arrested bulk cells in the S phase after 24 h, which is typical of DNA replication-blocking agents, whereas both H125 and A549 TICs had a significantly ($P < 0.05$ versus bulk) reduced proportion of cells in the S phase after exposure to 1 mM cisplatin, and 10 mM cisplatin for A549 (Figure 4c). As expected, this initial S-phase arrest was relatively transient and followed by a more persistent G2/M arrest. After 48 h, H125 bulk cells treated with 1 mM cisplatin had progressed into G2/M, but were still blocked in the S phase after 10 mM cisplatin. In H125 TICs, we observed a trend towards less G2/M accumulation ($P = 0.09$). In A549 bulk cells, which were less cisplatin-sensitive than H125 cells, the G2/M arrest after 48 h was most evident after 10 mM cisplatin. Again, significantly less G2/M accumulation of TICs was observed ($P < 0.05$). Altogether, NSCLC TICs displayed shortened G2 checkpoint response after IR as well as reduced S and G2/M checkpoint response after cisplatin treatment, which might contribute to their refractory phenotype.

NSCLC TICs show diminished apoptotic response after DNA-damaging treatment. DNA damage-induced apoptosis following IR and cisplatin treatment was evaluated by comparing PARP cleavage and caspase 3 activation in TICs and bulk cells (Figure 5). Consistent with their increased resistance, H125 TICs showed less PARP cleavage after 24 h after either IR (Figure 5a) or cisplatin treatment (Figure 5b). Moreover, reduced caspase 3 activation was observed in H125 ($P = 0.06$) and A549 ($P < 0.05$) TICs after cisplatin treatment (Figure 5c). Thus, active caspase 3 was detected in about 30% of both H125 and A549 bulk cells after 48 h treatment with 10 mM cisplatin, whereas in comparison, only 15 and 10% of H125 and A549 TICs showed caspase-3 activation, respectively.

NSCLC TICs exhibit diminished activation of DDR after DNA-damaging treatment. We hypothesized that differences in the cellular capacity to repair DNA damage could contribute to the increased resistance of NSCLC TICs. For
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these mechanistic studies, we included the IR-mimetic drug bleomycin in addition to IR and cisplatin. As shown in Figures 6a and b (for quantification data, see Supplementary Figure S2), there were no major differences in total levels of the key DDR regulators ATM and the non-homologous end-joining protein DNA-PKcs comparing H125 bulk cells and TICs, regardless of the presence of DNA damage. Next, we analyzed their activation pattern and in H125 bulk cells, pDNA-PK (pS2056), pATM (pS1981), as well as the DNA double-strand break marker γH2AX (H2A histone family, member X; pS139) were all increased after IR (Figure 6a) and after bleomycin treatment (Figure 6b). A higher basal level of γH2AX was noted in TICs compared with bulk cells, therefore the fold induction 1 h after DNA damage (IR, 1.8- versus 2.7-fold; bleomycin, 3.9- versus 5.2-fold) was lower in TICS than in bulk cells. In line with this, untreated H125 TICS had longer comet tails than bulk cells, suggesting increased levels of spontaneous DNA damage (Supplementary Figure 3). Notably, DNA damage-induced phosphorylation of DNA-PK, ATM and the ATM substrate KAP1 (pS824) after IR, bleomycin and cisplatin was reduced in H125 TICS compared with bulk (Figures 6a and b). Similar patterns of reduced pATM and pKAP1 response after IR were also seen in A549, H1299 and H23 TICS.

Because of the observed differences in the ATM pathway, we addressed if the activation pattern of the ATM substrate checkpoint kinase 2 (Chk2) was altered in H125 TICS. Irradiated TICS mounted a more robust initial Chk2 (pT68), whereas Chk2 activation in bulk cells was less prominent. Coherent with the ATM and KAP1 data, bleomycin activated Chk2 and structural maintenance of chromosomes 1 (SMC1), another ATM substrate, to a lower extent in H125 TICS. By contrast, the Chk2 target cell division cycle 25 homolog A (Cdc25A) did not show marked differential regulation in TICS versus bulk cells (Figure 6c). Both bleomycin and cisplatin treatment induced the monoubiquitination of FANCD2, a DNA damage-responsive protein, in bulk cells, whereas the induction was less pronounced in H125 TICS and absent in A549 TICS (Figures 6b and c). Rad51, a central player in homologous recombination, was unchanged by bleomycin treatment. Taken together, our data suggest that NSCLC TICS have increased levels of spontaneous γH2AX-decorated DNA lesions and yet were unable to mount an efficient DDR after treatment with exogenous DNA-damaging agents.

To determine if reduced DDR activation is related to therapy resistance, we evaluated cisplatin-induced apoptotic response in H125 bulk cells in the presence or absence of DNA-PKcs (NU7026) or ATM (KU55933) inhibitors (Figure 6d). Incubation with the ATM inhibitor before and during cisplatin treatment reduced PARP cleavage and the DNA-PK inhibitor in combination with cisplatin caused a minor reduction in PARP cleavage compared with cisplatin alone, suggesting that ATM and its substrates are of major importance for the resistant phenotype of TICS.

Xenografts derived from NSCLC TICS show impaired cisplatin response. To verify that NSCLC TICS are refractory to cisplatin also in vivo, we injected severe-combined immunodeficiency (SCID) mice with H125 bulk cells and TICS, respectively, and assessed tumor growth (median tumor size) over time. As shown in Figure 7, xenografts derived from H125 bulk cells (bulk tumors) grew slower during cisplatin treatment (P = 0.067 versus mock treatment), but regained full proliferative capacity when cisplatin treatment was ended (indicated by an arrow), and eventually reached similar sizes as the mock-treated bulk tumors. On the contrary, xenografts derived from TICS were highly resistant to cisplatin treatment and grew equally well regardless of whether the mice were treated with cisplatin (Figure 7). Thus, the in vivo data further support a DNA damage treatment-refractory phenotype of NSCLC TICS.

Discussion

TICS display resistance to CT and RT in several tumor types. Although a few studies have suggested the existence of a therapy-resistant phenotype of NSCLC TICS, most of the underlying molecular events remain elusive. Here we demonstrate that NSCLC TICS are refractory to both RT and the DNA-damaging CT agent cisplatin in vitro, and cisplatin in vivo. Importantly, we show that NSCLC TICS exhibited inadequate activation of DDR signaling, impaired checkpoint...
To confirm a TIC phenotype of the NSCLC spheres formed, we analyzed several stem cell markers and revealed that three out of four sphere-forming cell lines (A549, H125 and H1299) had increased the proportion of cells expressing CD133. However, H23 spheres did not, indicating that CD133 may not be a uniform marker for NSCLC TICs, as reported earlier. Aside from CD133, NSCLC TICs have been reported to show increased expression of several stem cell markers, including Sox2, Oct4, Nanog, aldehyde dehydrogenase and CD44. The increased expression of Sox2, Nanog and Oct4 in A549 TICs was consistent with but not another report, indicating some variance in the TIC phenotype. We did not detect increased expression of Sox2/Nanog/Oct4 in H125 TICs or Nanog/Oct4 in H23 TICs despite their clear sphere-forming capacity. Importantly, this heterogeneity in the expression of putative stem cell markers in NSCLC TICs did not affect their response pattern following DNA-damaging treatment.

A novel finding in this study is that NSCLC TICs enriched as spheres display apparent radiation resistance compared with bulk cells. This supports studies of TICs in other tumor types such as glioma where isolated primary CD133+ cells were more radioresistant than CD133− cells. We could also confirm the previously reported cisplatin-refractory phenotype of NSCLC TICs. Although an increased expression of ATP7B in TICs could partly contribute to platinum resistance, the increase was relatively minor and is unlikely to account for the large differences in cytotoxicity that we observed. Using a combination of quantitative real-time PCR, immunofluorescence, flow cytometry and comet assay, we conclusively show that cisplatin can be transported into TICs where it forms adducts and induce DNA strand breaks at comparable levels as in bulk cells. Likewise, both bleomycin and IR caused similar amounts of DNA damage in both bulk cells and TICs, as shown by the comet assay (Supplementary Figure 3).

TICs have been shown to possess several features associated with resistance to DNA damage, including slower cell cycle kinetics, increased DNA repair capacity and resistance to apoptosis. Here, we present novel data demonstrating reduced cell cycle arrest after IR or cisplatin treatment of NSCLC TICs. This is mechanistically important as alterations in the checkpoint response might allow TICs to evade the proapoptotic effects of RT/CT and thereby contribute to resistance. CFSE confirmed that the growth of TICs was essentially unchanged following DNA-damaging treatment. As untreated H125 TICs and bulk proliferated at comparable rates, the activation of cell cycle checkpoints in TICs was unlikely to simply be delayed and their increased therapy resistance was not purely due to quiescence. Although slower growth of untreated A549 TICs may partly account for their resistance, both H125 and A549 TICs showed impaired apoptotic response following DNA-damaging treatment, which indicates that the rate of proliferation was not crucial for the differences in chemo- or radio-sensitivity. To our knowledge, only one other study has reported decreased cell cycle arrest in CD133+ cells, albeit observed in liver cancer cells at a much higher dose of radiation (15–20 Gy). In concordance with our findings, CD133+ liver cancer cells also showed diminished PARP cleavage and Annexin V-staining after IR. Taken together,
these data suggest that transient checkpoint responses may represent a hitherto unappreciated mechanism that allows TICs to evade DNA damage-induced apoptosis.

To obtain further mechanistic insight, we profiled several DNA repair proteins, and interestingly, NSCLC TICs tend to show increased basal levels of the DNA damage marker γH2AX. Taken together with data from the comet assay, we hypothesize that TICs may harbor more spontaneous DNA damages and that the increased baseline γH2AX could reflect a subpopulation primed to respond to DNA damage, as previously reported for CD133+ glioma cells. Notably, NSCLC TICs showed diminished phosphorylation of H2AX, ATM, DNA-PKcs and FANCD2 monoubiquitination after exposure to DNA-damaging agents. Given that a similar reduced activation of γH2AX after DNA damage has been reported in mammary TICs, it raises the question as to whether or not the TICs are a pre-existing population with intrinsically altered DDR or if these properties could be affected by culture and selection. This has been difficult to address in our system, as we have not sorted out the CD133+ cells before the sphere culturing process. Nonetheless, differential expression of DDR proteins was reported in sorted CD133+ versus CD133− cells shortly after isolation from the primary tumor, implying that TIC properties differ inherently from their bulk counterpart, and is not solely a result of in vitro culture. In addition, cisplatin normally binds to serum proteins, so having no serum in the stem cell media should increase its potency and TICs should incur more DNA damage; however, equal damage was revealed by the comet assay. Interestingly, while the bulk cells have repaired most of their damage 3 h after bleomycin treatment, as judged by the disappearance of comet tails, DNA strand breaks appeared to persist in TICs, suggesting that the DNA repair capacity of TICs may be diminished.

A hyper-active DDR is usually considered to enhance TICs survival, but there is also data from studies on glioma where no differences in DNA base excision, single-strand break repair or resolution of γH2AX nuclear foci were found between bulk cells and TICs. Our data clearly demonstrate reduced activation of an ATM and KAP1-mediated pathway in NSCLC TICs. Importantly, bulk H125 cells cotreated with the ATM inhibitor KU55933 decreased cisplatin-induced PARP cleavage, uncovering a causal relationship between reduced ATM signaling and resistance to DNA-damaging treatment. ATM is a key driver of the DNA damage signaling network and is required for cell cycle arrest and DSB repair through homologous recombination. Therefore, the reduced ATM signaling is in accordance with the impaired cell cycle arrest that we see. Notably, mutations of the ATM gene in B-cell chronic lymphocytic leukemia patients were associated with impaired CT response and reduced overall and treatment-free survival. ATM, ATR and DNA-PK rapidly phosphorylates the transcriptional repressor KAP1 on Ser824, leading to increased expression of KAP1 target genes involved in checkpoint control (p21\textsuperscript{WAF1/Cip1}) and apoptosis (Bcl-2-associated X protein (Bax), p53 upregulated modulator of apoptosis (Puma), Noxa). Therefore, the impaired phosphorylation of KAP1 might provide a mechanistic explanation for the reduced DNA damage-induced cell cycle arrest and apoptosis in NSCLC TICs.

Persistent checkpoint arrest is detrimental to many tumor cells. Consequently, the transient arrest followed by checkpoint recovery, especially coupled to decreased apoptosis, will favor survival of TICs. The general conclusion of our study is that TICs are able to survive both a higher basal level of DNA damage and to escape the checkpoint arrest despite the presence of DNA damage. This correlates well with data showing higher mutation rates in a subtype of glioblastoma defined on the basis of a CD133-related gene expression signature. Further studies are needed to investigate the DNA repair activity in TICs and to validate the findings using siRNA towards primarily ATM and KAP1. An increased knowledge of the specific alterations of these components in TICs may open up for therapeutic interventions as a way to improve the RT and/or CT response of NSCLC.

Materials and Methods

Cell culture. The human NSCLC cell lines A549 and H23 (adenocarcinomas), H125 (adenosquamous), H157, H661 and H1299 (large cell carcinoma) are commercially available from ATCC (American Type Culture Collection; Manassas, VA, USA), whereas U-1752 (squamous cell carcinoma) and U-1810 (mixed large cell/adeno carcinoma) were established at Uppsala University as described.

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Bulk cells were cultured in RPMI-1640 supplemented with 10% heat-inactivated fetal bovine serum (HyClone, Sweden). To assess sphere-forming ability, cells were seeded at a density of 10,000 cells per ml in Dulbecco’s modified Eagle’s media/F12 (Lonza, Basel, Switzerland) supplemented with 20 ng/ml epidermal growth factor, 10 ng/ml basic fibroblast growth factor (PeproTech, London, UK), 2 μg/ml heparin (Sigma-Aldrich, Stockholm, Sweden) and B27 supplement (Invitrogen, Stockholm, Sweden) in non-adherent plates. Fresh supplements were added at day 3 or 4, and both adherent and non-adherent cells were kept when replating the cells at day 7. Experiments were started after 10–14 days of culture in stem cell media. Before experiments, spheres were trypsinized and mechanically dissociated to generate single cells. For kinase inhibitor experiments, cells were pretreated with 10 μM NU7026 (DNA-PKcs inhibitor) or KU55933 (ATM inhibitor) 1 h before DNA-damaging treatment, and the inhibitors were kept during the 24 h before cells were pelleted for western blot analysis.

**IR procedure.** TICs and bulk cells were irradiated with 2, 4 or 8 Gy. The IR source was Co60 with a dose rate of 0.4 Gy/min. At 1, 2, 4, 24 or 48 h after IR, cells were pelleted and frozen for western blot analysis or fixed for caspase 3 and cell cycle analysis.

**Clonogenic survival assay.** Following IR, 500 cells were seeded in RPMI media in 10 cm² dishes. After 9 days (A549) or 14 days (H125) depending on the cellular growth rate, the resulting colonies were stained using 0.5% crystal violet in 25% methanol and colonies containing more than 50 cells were counted. For each radiation dose duplicate samples were prepared. Evaluation of RT response was...
Reagents and random hexamer primers (Applied Biosystems, Carlsbad, CA, USA). Analyses were performed in duplicate on a 7900 Fast Real-Time PCR System with Fast SYBR Green reagents (Applied Biosystems) and Fast program, initiated at 95 °C for 20 s, followed by 40 cycles of 95 °C for 1 s and 60 °C for 20 s. The specificity of the reactions was analyzed by melting curve analysis to confirm the presence of a single amplified product. mRNA levels were normalized to TATA binding protein (TBP) to confirm equal cDNA input, using the 2^(-ΔΔCt) method for calculation of the relative expression values. The mRNA levels are expressed as fold to bulk cells in Figure 2. Primer sequences were: CD133 - forward 5'-CATCAAGCAACACCAAGCTG T-3', reverse, 5'-TGTCCAATGGAAATTTAGCC CTTT-3'; Sox2 – forward, 5'-CACCCTCAGGATGCTTCTT-3', reverse, 5'-GAGCT G GCTCCTGAGCTTGAA-3'; Nanog – forward, 5'-AAACTGTGGCCGCAATATGC AAGTG-3', reverse, 5'-TCCCTGTTGTTAGAAGATGAAGGG-3'; Oct4 – forward, 5'-AGGCCCCATTTCCAAGG CC-3', reverse, 5'-TGGACCCTCTCCGGGT TTGG-3'; CTR1 – forward, 5'-GGGTTAAGAGCTGAGAGAGGATGG-3', reverse, 5'-GCCGATGATGCCCATCCCTTGTTG-3'; ATP7B – forward, 5'-GGGTTAAGAGCTGAGAGAGGATGG-3', reverse, 5'-GCCGATGATGCCCATCCCTTGTTG-3'; and TBP – forward, 5'-CAGGAGACCGCAGCTGATTT-3', reverse, 5'-TTTTCTTG CTCCGAGTCTGGAC-3'.

Western blot analysis. Protein was extracted from cell pellets using RIPA buffer (50 mM Tris-HCl (pH 7.4), 150 mM NaCl, 0.5% Igepal, 5 mM EDTA (pH 8.0), 0.1% SDS) supplemented with PhosStop and Protease inhibitor cocktail tablets (Roche Diagnostics, Basel, Switzerland). Proteins were separated on 3–8% Tris-acetate gradient gels in NuPage Tris-acetate running buffer (Invitrogen) or 4–12% Bis-Tris gradient gels in NuPage MOPS running buffer (Invitrogen), depending on the size of the protein to be analyzed and transferred to nitrocellulose membrane (Hybond ECL; Pharmacia Biotech, GE Healthcare, Uppsala, Sweden), which was blocked in 5% BSA and probed with different primary antibodies. The following primary antibodies were used: p-tubulin and α-tubulin (ab92547 and ab5621, respectively; Abcam), total DNA-PKcs (MS-423P0; Thermo Fisher Scientific, Fremont, CA, USA), α-tubulin (ab5621), total ATM and FANCD2 (1549-1 and 2986-1, respectively; Epitomics, Burlingame, CA, USA), pKAP1 and pSMC1 (pS378; A300-776A and pSer966; A300-050A, respectively; Bethyl Laboratories, Montgomery, TX, USA), PARP, cdc25A and β-tubulin (sc-7150, sc-97 and sc-5286, respectively; Santa Cruz Biotechnology, Santa Cruz, CA, USA), pChk-2 (pThr68, no. 2661; Cell Signaling, Danvers, MA, USA), p-Chk2 (T78/S68; Sigma-Aldrich) and GAPDH (no. 22753; Trevigen, Gaithersburg, MD, USA). The membranes were incubated with primary antibodies overnight at 4 °C, followed by probing with either infrared dye-conjugated goat anti-rabbit or -mouse secondary antibodies (LI-COR, Cambridge, UK) for 45 min at room temperature. The membranes were scanned and quantified using the Odyssey S infrared Imaging System (LI-COR). A representative blot of three independent experiments is shown, unless otherwise stated in the figure legend.

Comet assay. H125 bulk cells and TICs were treated with 5 µM/µl bleomycin or 8 Gy IR for 1 and 3 h, or 40 µM cisplatin for 16 and 24 h. A total of 20,000 cells per ml in PBS were mixed with low melting agarose at a 1:10 ratio and 900 cells were added per slide to allow for solidification. The slides were washed with PBS and mounted using 1 ml 0.05% Agarose in PBS. The slides were then placed under a coverslip followed by a 30-min incubation in neutral comet running buffer before the slides were put in an electrophoresis tray to allow DNA to migrate for 45 min. Slides were immersed in DNA precipitation buffer for 30 min followed by incubation in 70% ethanol for another 30 min. The slides were dried at 37 °C for 15 min before the addition of 100 µl SYBR Gold to each sample for 30 min. Following another drying step, the comets were analyzed at a × 40 magnification using a fluorescence microscope.

Xenograft experiments. All parts of the experiment were performed according to the Swedish Ethical guidelines for animal experiments and were approved by the Animal Ethical committee in Stockholm, Sweden. Female SCID mice in a BALB/c background, at an age of 8–10 weeks at the time of inoculation, were used to establish xenografts. Xenografts were established by subcutaneously injecting one million H125 bulk or TICs in 100 µl PBS in the right-side flank of the mice. The tumor sizes were monitored twice a week and calculated according to the formula: tumor weight = length × width^2/2, as described in Gallo et al.15 When a tumor reached 100 mg, the mouse was included in the study (day 0) and allocated either to no treatment (injected with...
saline alone) or to intraperitoneal injections of 1 mg/kg cisplatin in a volume of 100 μl. The mice were treated two times a week, a maximum of nine times in total. The mice were killed if the tumor size approached 1200 mm^3 or if the health conditions of the mice were severely impaired.

**Statistics.** All values are expressed as mean ± S.D. from three independent experiments, unless otherwise stated in the figure legend. Comparison between two groups was made using an unpaired two-tailed *t* test; *P* < 0.05 was considered significant.

**Conflict of Interest**
The authors declare no conflict of interest.

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