Correction

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The authors note that the author name Oscar Fernadez-Capetillo should have appeared as Oscar Fernandez-Capetillo. The corrected author line appears below. The online version has been corrected.

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Nuclear phosphoinositide 3-kinase β controls double-strand break DNA repair

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Class I phosphoinositide 3-kinases are enzymes that generate 3-phosphoinositides at the cell membrane following transmembrane receptor stimulation. Expression of the phosphoinositide 3-kinase β (PI3Kβ) isoform, but not its activity, is essential for early embryonic development. Nonetheless, the specific function of PI3Kβ in the cell remains elusive. Double-strand breaks (DSB) are among the most deleterious lesions for genomic integrity; their repair is required for development. We show that PI3Kβ is necessary for DSB sensing, as PI3Kβ regulates binding of the Nbs1 sensor protein to damaged DNA. Indeed, Nbs1 did not bind to DSB in PI3Kβ-deficient cells, which showed a general defect in subsequent ATM and ATR activation, resulting in genomic instability. Inhibition of PI3Kβ also retarded the DNA repair but the defect was less marked than that induced by PI3Kα deletion, supporting a kinase-independent function for PI3Kβ in DNA repair. These results point at class I PI3Kα as a critical sensor of genomic integrity.

cancer | genomic integrity

Class I phosphoinositide 3-kinases (PI3Ks) are enzymes composed of a p85 regulatory and a p110 catalytic subunit that generate 3-phosphoinositides (PIP3) following growth-factor receptor stimulation at the cell membrane (1, 2). p110α and p110β are expressed ubiquitously, and regulate cell division and embryonic development (3–9). p110α is mainly nuclear (7); we tested whether endogenous p110β concentrated at laser tracks early and in DSB repair. To determine whether p110β activity is stimulated by DNA damage, we exposed NIH 3T3 cells to UVC or IR, then immunopurified p110β using a specific antibody (Ab) (7), and perform an in vitro PI3K assay. Both treatments increased p110β activity (Fig. 14). Considering that defects in the DDR often result in impaired cell-cycle checkpoints, we examined the consequences of interfering with p110β on DNA damage-induced G2 arrest. We determined the proportion of cells that progress into M phase by staining the cells with an SI0-phospho-histone H3 (pH3)-specific Ab (22). We used shRNA to reduce p110β expression and the selective inhibitor TGX221 to inhibit p110β (7, 23). Inhibition of p110β moderately increased the proportion of pH3+ cells (Fig. SL4). Nonetheless, a larger proportion of p110β knockdown cells progressed into M phase after IR when compared with controls, indicating that p110β deletion inhibits G2 arrest (Fig. S1B). We also examined immortalized p110β−/− murine embryonic fibroblasts (MEF), and p110β−/− MEF reconstituted with WT- or with the kinase-dead (KR)-p110β (8). p110β deletion, but not kinase inactivation impaired G2 arrest, permitting entry into mitosis (Fig. S1C).

DSB repair begins with formation of large protein complexes (foci) that contain many repair proteins (12). A large fraction of p110β localizes in the nucleus (7); we tested whether endogenous p110β formed foci after DNA damage. IR induced p110β localization in large nuclear foci (after 1 h) (Fig. S2A and B). Moreover, simultaneous staining of p110β and γ-H2AX showed partial colocalization of endogenous p110β with γ-H2AX at DSB (Fig. S2C). To confirm p110β localization at DNA damage sites, we examined GFP-p110β translocation to DSB. Cells were irradiated with an UV laser that generates DSB in defined nuclear volumes (24, 25). GFP-p110β concentrated at laser tracks early and remained associated for the entire recording period (∼300 s) (Fig. 1B, Fig. S2D, and Movie S1).

To determine whether the PI3K product PIP3 concentrates at the site of DNA damage, we performed immunofluorescence analysis using anti-PIP3 Ab. This Ab stained the cell membrane, endomembranes, and the nuclei in exponentially growing NIH 3T3 cells (Fig. S3A). Ly294002 (pan-PI3K inhibitor) reduced the cellular PIP3 signal, whereas TGX221 inhibitor was more potent in

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and ATR kinases. We examined phosphorylation of different substrates of ATM (pS3C1, pChk2) and ATR (pRad17, pChk1). γ-IR induces more markedly ATM activation while UVC triggers principally the ATR route (Fig. 2A) (19, 24). Whereas reduction of p110β levels markedly diminished ATM and ATR pathways, p110β inhibition only partially reduced ATR route (Fig. 2A).

To examine the consequences of interfering with p110β expression or activity on ATM chromatin loading, we γ-irradiated cells, fractionated them as in ref. 7, and determined ATM content in the chromatin fraction; for the ATR pathway we analyzed Rad17. ATM was present in the chromatin fraction of WT- and KR-p110β MEF, but was severely reduced in p110β−/− MEF; similarly, Rad17 loading onto chromatin was greatly impaired in p110β−/− MEF (Fig. 2B). Thus, p110β regulates ATM and ATR-pathway members binding to chromatin.

Reducing nuclear PIP3 (Fig. S3A). γ-Irradiation of NIH 3T3 cells induced formation of foci (30 min) that were PIP3-positive; some were also γH2AX-positive (Fig. S3B). DSB foci also concentrated the enzyme substrate PI(4,5)P2 (Fig. S3C). To confirm that PIP3 concentrates at damage sites, we used the GFP-Btk-PH domain, which binds selectively to PIP3 (26). We microirradiated cells with a UV laser and examined real-time Btk-PH translocation to laser tracks; Btk-PH concentrated very early and remained in this region for the entire recovery period (Fig. 1 C and D). GFP fused to the R25C mutant of Akt-PH domain (27) remained invariant (Fig. 1D and Fig. S3D). Therefore, PIP3 localizes at damaged DNA.

**Fig. 1.** p110β and PIP3 localize to damaged DNA foci. (A) NIH 3T3 cells were exposed to UV or IR and harvested at different times. p110β was immunoprecipitated (IP) (800 μg) from cell extracts and tested by in vitro PI3K assay. Control IP was with protein A. Graphs show PI3P signal intensity quantitation in arbitrary units (AU) (mean ± SD, n = 3). (Right) Western blot analysis of p110α or p110β IP (300 μg). Graphs show signal intensity quantitation in arbitrary units compared to maximum (mean ± SD, n = 3). (B) NIH 3T3 cells expressing GFP-p110β were microirradiated with an UV laser. We examined real-time GFP-p110β translocation to the DNA damage region. Dotted lines indicate laser path. (C) GFP-Btk-PH-transfected U2OS cells (24 h) were microirradiated with an UV laser and examined as in B. (Scale bars in B and C, 15 μm.) (D) The graphs show normalized fluorescence units (NFU) of the mean of n = 3 experiments performed as in B and C and plotted as a function of time. Controls are GFP and GFP fused to the R25C mutant of the Akt PH domain.

**Fig. 2.** Defective ATM and ATR pathways activation in p110β-deficient cells. (A) NIH 3T3 cells were transfected with p110β or control shRNA (48 h); other cells were treated with the p110β inhibitor TGX221 (30 μM, 4 h). Cells were irradiated (UVC or IR) and extracts collected after 1 h. Lysates were examined by Western blot using indicated Ab. ATM effectors are grouped with a square; ATR effectors are grouped with a dashed-line square. Graphs show band signal intensity (AU) (mean ± SD, n = 3). (B) p110β−/− immortalized MEF alone or reconstituted with WT- or KR-p110β were exposed to IR (10 Gy), incubated (1 h), and fractionated. Extracts were examined by Western blot with indicated Ab. The graph shows the quantitation of the signal in the chromatin fraction (mean ± SD, n = 3). (C) NIH 3T3 cells transfected with p110β shRNA or treated with TGX221 were exposed to IR (3 Gy) and processed 15 min later for immunofluorescence (IF) using anti-phospho-Rad17 or -phospho-ATM Ab. Graphs show the integrated nuclear fluorescence intensity (nuclear density, Left) and the number of foci with signal intensity > 40 AU (Right) on a representative set of cells of n > 100 examined. (Scale bars, 15 mm.) (*) Student t test P < 0.05.

**p110β Deletion Inhibits ATM and ATR Repair Pathways.** We next analyzed the impact of p110β depletion on the activity of ATM and ATR kinases. We examined phosphorylation of different substrates of ATM (pS3C1, pChk2) and ATR (pRad17, pChk1). γ-IR induces more markedly ATM activation while UVC triggers principally the ATR route (Fig. 2A) (19, 24). Whereas reduction of p110β levels markedly diminished ATM and ATR pathways, p110β inhibition only partially reduced ATR route (Fig. 2A).

To examine the consequences of interfering with p110β expression or activity on ATM chromatin loading, we γ-irradiated cells, fractionated them as in ref. 7, and determined ATM content in the chromatin fraction; for the ATR pathway we analyzed Rad17. ATM was present in the chromatin fraction of WT- and KR-p110β MEF, but was severely reduced in p110β−/− MEF; similarly, Rad17 loading onto chromatin was greatly impaired in p110β−/− MEF (Fig. 2B). Thus, p110β regulates ATM and ATR-pathway members binding to chromatin.
Immunofluorescence studies confirmed definitive activation of ATM and ATR pathways in cells with reduced p110β expression upon irradiation. p110β inhibition reduced the signal intensity of pATM and pRad17 foci, whereas p110β knockdown greatly diminished pATM and pRad17 foci number (Fig. 2C). Histone H2AX is a substrate for ATM and ATR (as well as for DNA-PK) (14, 28), p110β inhibition reduced γH2AX signal intensity in DSB foci and p110β deletion nearly eliminated γH2AX signal (Fig. S4A). We also examined 53BP1, an ATM pathway effector that regulates chromatin structure at DSB (24). p110β inhibition delayed—but p110β knockdown virtually blocked—53BP accumulation at laser tracks (Fig. S4B and Movie S2). These results show that p110β expression is critical for the association of DDR proteins (ATM, Rad17, γH2AX, and 53BP1) to DSB foci.

**Endogenous p110β Associates to Nbs1.** We used mass spectrometry to detect potential DDR proteins that interact with p110β. We transfected cells with GST-fused-p110β and performed a pull-down assay to identify nuclear proteins that might associate with p110β. We identified Rad50 (an MRN complex component), Rad17 (an ATR effector), and Rad9B (Fig. 3A), in complex with GST-p110β. We confirmed the association of Rad17 with p110β in intact cells following UVC or IR (Fig. 3A). In agreement the MRN component Rad50 association with p110β (Fig. 3A), recombinant p110α and β were found to associate to endogenous human (h)Nbs1 (29). We tested whether endogenous p110β and Nbs1 formed a complex. Endogenous p110β, but not endogenous p110α (cytosolic, ref. 6), associated constitutively with endogenous Nbs1 (Fig. 3B and C) and vice versa (Fig. 3C).

To define whether p110β regulates Nbs1 recruitment to damaged DNA, we examined translocation of GFP-murine-Nbs1 (30, 31) to laser tracks in p110β/−/− MEF. In WT-p110β-MEF, Nbs1 accumulated early (at ~15 s) and remained associated throughout the recording period (~270 s); in p110β-KR cells, Nbs1 accumulated more slowly and in smaller amounts; in contrast, p110β deletion nearly abrogated Nbs1 accumulation at laser tracks (Fig. 3D and Movie S3). Indeed, 50% of p110β/−/− MEF showed no Nbs1 accumulation and ~50% showed very low intensity and unstable Nbs1 binding at laser tracks. Results were similar in NIH 3T3 cells. p110β expression is thus necessary for Nbs1 recruitment to DSB, whereas p110β activity enhances or stabilizes Nbs1 recruitment to these sites.

**p110β Association Is Required for Nbs1 Binding to Damaged DNA.** No viable Nbs1 mutant has yet been reported to disrupt the initial recruitment of MRN to DNA (15). To test whether p110β-Nbs1 complex formation is necessary for Nbs1 binding to DSB, we assayed which residues in Nbs1 mediate association with p110β. We examined residues 653 to 669 of hNbs1, which mediate interaction of recombinant p110α and Nbs1 (29). We transfected cells with WT GFP-hNbs1, or A653-669-hNbs1, or with A670-hNbs1, and examined Nbs1/endogenous p110β association. Endogenous p110β associated efficiently with WT, but very poorly with mutant forms of Nbs1 (Fig. 4A). p110β expression is required for Nbs1 translocation to laser tracks (Figs. 3D and 4B). In addition, GFP-WT-hNbs1 but not A653-669-Nbs1 or A670-hNbs1 concentrated at laser tracks (Fig. 4C and D and Movie S4). These results show that p110β/Nbs1 association is necessary for Nbs1 recruitment to damaged DNA.

**Proliferating Cell Nuclear Antigen Concentrates at DNA-Damaged Zones Downstream of Nbs1.** Some of the proteins controlling DNA replication also regulate DNA repair. MRN complexes localize at replication forks (32), and proliferating cell nuclear antigen (PCNA), which controls DNA replication (33), also regulates NHEJ and HR in yeast (34, 35). We described that p110β associates with PCNA, controlling DNA replication (7). We explored whether PCNA also localizes to DSB in mammals, and whether this process is controlled by p110β. Although the net amount of chromatin-bound PCNA did not increase after irradiation, DNA damage induced RFP-PCNA translocation to laser tracks (Fig. 5A–C). PCNA chromatin binding diminishes by April 20, 2010 | vol. 107 | no. 16 | 7493
p110β inhibition, and more markedly upon p110β knockdown (Fig. S5A). In addition, p110β inhibition and, more strikingly, p110β deletion reduced PCNA localization at laser tracks (Fig. S5 and Movie S5). Thus, PCNA localizes to laser tracks in a p110β-dependent manner.

Because both PCNA and Nbs1 binding to DSB were controlled by p110β, to define the primary event regulated by p110β in DDR, we simultaneously examined the translocation of Nbs1 and PCNA to DSB in MEF (Movies S6 and S7). p110β deletion affected both Nbs1 and PCNA loading. Nonetheless, Nbs1 translocated slightly earlier than PCNA to damaged DNA in control cells (Fig. S5). In addition, in some p110β-deficient cells in which GFP-Nbs1 did not mobilize to laser tracks, we detected PCNA translocation; finally, p110β inhibition induced a more pronounced defect in Nbs1 than PCNA translocation (Fig. S5). These findings show that interference with p110β affects the earliest sensor of DNA damage (Mre11/Rad50/Nbs1) more severely than PCNA loading and that these processes might be independent.

p110β Deletion Induces Genomic Instability. We focus our study on DSB repair, as p110β associates with HR components (Fig. 3). To demonstrate defective DSB repair in p110−/− cells, we searched for the presence of DSB in untreated cells. In addition, we irradiated these cells (γ-IR, 10 Gy) and incubated them for 1 h to permit DNA repair; we quantitated DSB using the neutral comet assay (S6). This assay showed that p110-/- MEF, but not WT or KR-p110β MEF, already had spontaneous comets in untreated cultures (~20% of cells), and showed a large proportion of comets (indicative of DSB) 1 h after IR (Fig. S4). Accordingly, p110β-depleted NIH 3T3 cells were radiation-sensitive, as they showed a higher rate of DNA damage-induced apoptosis than controls (Fig. S6). Because p110β-deficient cells fail to repair DSB and do not stop at G2/M following damage (Fig. S1), they might accumulate DNA defects. We examined chromosome breaks and chromosome numbers by DAPI staining of MEF metaphases. p110−/− MEF, but not WT or KR-p110β MEF, had abnormal chromosome numbers, chromosome breaks, and disjunction figures (Fig. 5 B and C), showing that p110β deletion causes genomic instability.

Discussion

Class I PI3K were thought to act mainly by increasing PIP3 production at the cell membrane. Here we report a unique function for PI3Kβ in the control of DSB repair. PI3Kβ was required for the binding of the first DNA damage sensor protein Nbs1 to double-strand breaks. Indeed, endogenous PI3Kβ bound to endogenous Nbs1, and this complex was necessary for efficient concentration of Nbs1 at DSB. Both PI3Kδ deletion and mutation of Nbs1 at the site of association with p110β resulted in highly defective Nbs1 localization at DSB. Because of this function in DNA damage sensing and subsequent DDR (Fig. 5D), p110β deletion resulted in defective G2 arrest, radiation sensitivity, DSB accumulation, and genomic instability.

These findings and our description on the role of p110β in DNA replication (7) suggest that there is a fundamental difference between the function of the other ubiquitous class I PI3K, p110α, which is mainly cytosolic (6, 7), and p110β. Whereas p110α regulates cell growth, as well as G0 > G1 and G1 > S phase transitions, p110β cooperates with p110α in G1 progression (6) but also concentrates in the nucleus, where it regulates DNA homeostasis.

We show that p110β is activated following DNA damage and concentrates at DSB. Both p110β kinase activity and a kinase-independent p110β function regulate DNA repair. Whereas PI3Kβ inhibition delayed activation of the DDR, PI3Kδ deletion almost abrogated it. The kinase-independent capacity of p110β to associate Nbs1 is critical for its recruitment to damaged DNA and, in turn, for amplification of the DDR. In the absence of p110β, Nbs1 was not recruited to laser tracks and, after IR, p110β knockdown cells had a very small number of pATM and pRad17-containing foci and showed defective ATM and ATR pathway activation. These defects explain the failure in G2 arrest of p110β-deficient cells, as well as their radiation sensitivity, DSB accumulation, and DNA instability.

In contrast to the consequences of deleting p110β, its inhibition delayed, but did not abrogate Nbs1 association to laser tracks; the number of IR-induced foci was roughly 50% that of normal cells and showed reduced pATM and pRad17 intermediate frequency (IF) signal intensity. In vivo imaging of cells with inhibited p110β showed unfocused, delayed, and unstable concentration of Nbs1, PCNA, and 53BP1 in laser tracks, suggesting that p110β activity stabilizes or facilitates protein recruitment to DSB. Given the high negative charge of PIP2, the reported local increase in PIP2 at DSB might help to maintain DNA (also negatively charged) in an open conformation by repelling electrostatic forces. Alternatively, PIP2 might sequester histones (positively charged), contributing to stabilization of chromatin in an open conformation at DSB. PIP2 localization at DSB sites might also recruit PH domain-containing proteins, such as PKBζ/Akt1, which concentrate at
pik3cb<sup>−/−</sup> littermate embryos were found; 70% appeared normal and 30% appeared small and moribund (10). In MEFs obtained from the latter, KR-p110β content was smaller than in the apparently normal ones, supporting the existence of p110β kinase-independent functions (10). Tissue-specific p110β-deletion mouse models have been reported; whereas PTEN<sup>−/−</sup> prostates develop tumors, p110β-deletion in PTEN<sup>−/−</sup> prostate impeded tumorigenesis (8). p110β<sup>−/−</sup> prostates did not show apparent morphological defects (8). The observation that prostate develops in conditional p110β<sup>−/−</sup> mice (8) contrasts with the requirement of p110β for DNA replication and repair, suggesting that other proteins might replace or compensate p110β function in some tissues. Conventional p110β deletion, however, impairs embryonic development very early, at E 2 to 3 (4). Because DSB repair is critical for embryonic development (11), we hypothesize that the kinase-independent function of p110β in DNA repair might contribute to cause the embryonic lethality of p110β<sup>−/−</sup> mice.

Current knowledge classifies three distal homologs of PI3K, DNA-PKcs, ATM, and ATR, as key DDR regulators. DNA-PKcs controls NHEJ; ATM and ATR control HR (12–15). This model should be expanded to integrate the nuclear class I PI3K isoform p110β in DNA repair pathway activation at the DSB-sensing step (Fig. 5D). We have examined the function of p110β on HR DSB repair, as p110β associates with HR repair proteins (Fig. 3); however, we cannot rule out the involvement of p110β in additional repair processes. The involvement of p110β in nuclear PIP3 production and DDR increases the complexity of the mechanisms by which class I PI3K pathway might regulate survival and tumorigenesis. p110β activity is essential for prostate cancer formation in the mouse (8), suggesting the use of interfering p110β-based therapy. Our results point to the importance of testing the status of the p53 gene (frequently mutated in cancer), because defects in apoptosis mechanisms in cancer cells might result in increased genomic instability following interference with p110β.

**Materials and Methods**

**Reagents and cDNA.** β-actin antibody (Ab) was from Sigma, histone Ab from Chemicon Intern, p110β (if) and Rad17 (if) from Santa Cruz, p110β from Cell Signaling, Chk1 and Chk2 from Novoceastr and Upstate, Rad17 (Western blot), pRad17, and p-SMC1 were from Abcam. Other antibodies used were PIP3 (Echelon Bioscience), PCNA (BD Transduction), p-histone H3 (Ser10; Abcam); pEGFP-C1-53BP1 (24), murine γH2AX (Millipore), p-ATM (1981; Rockland); p-Chk1 (Ser345), ATM, and Nbs1 were from Cell Signaling. p110α Ab was donated by A. Klippel (Merck, Boston, MA). [32P]ATP was from Amersham. TGX221 (used at 30 μM) was from ACCC. p110β shRNA was from Origene and others shown in Fig. 1 (6). All remaining reagents were from Sigma. The construct encoding the PKB-mutant-PhD domain in the pEGFP-C1 vector was a gift of J. Downward (Cancer Research, London, United Kingdom). pSG5-Myc-p110β (7), pEGFP-C1-35B3 (24), murine Nbs1-2GFP (30), and pEn-mRFP-PCNA2 (33) have been described. pEGFP1-p110β and pEBG-GST-p110β were constructed by subcloning p110β from pSG5-myc-p110β into pEGFP-C1 and pEBG. GFP-Btk-Ph-Domain was described (26). pEGFP-C2-WT- Nbs1b, pEGFP-C2-ΔNbs1b, and -Ifh1 were prepared by subcloning into pEGFP-C2 (mutants were kindly donated by Y. C. Chen, Yang-Ming University, Taipei, Taiwan) (29).

**Cell Culture and Transfection.** Cell lines were maintained as reported (7). Immortalized p110β-deficient MEFs were donated by J. Zhao and T. Roberts (Dana-Farber Cancer Institute, Boston, MA) (8). Cell synchronization at G1/S and metaphase arrest were described (6, 7). To examine the proportion of cells in mitosis, we stained the cells with an antibody recognizing S10–polymerizing kinase and metaphase checkpoint kinase. The signal from the antibody recognizing S10–polymerizing kinase was quantitated by flow cytometry. The proportion of cells in mitosis was calculated from the percentage of cells with high levels of the signal.

**Irradiation-Induced DNA Damage.** Cells were irradiated using UV or ionizing radiation (IR). UVC radiation was generated using a UV generator (λ = 254 nm; Vilber Lourmat) and IR was delivered by a γ-irradiation (MARK 1, Shepherd and Associates) that uses a 137Cs probe. Real-time recruitment of DNA repair proteins to microcassette-generated DNA damage sites was reported (24, 25).

**Fig. 5.** p110β deletion induces radiation sensitivity and genomic instability. (A) Representative p110β<sup>−/−</sup> immortalized MEF, alone or reconstituted with WT- or KR-p110β, were exposed to IR (10 Gy) and incubated (1 h), or were untreated; all were tested in neutral comet assays. The graph shows the tail moment for these MEF (mean ± SD, n = 50 cells). Tail Moment = (%DNA in tail × tail length)/100. (Scale bar = 125 μm.) (B) Representative p110β<sup>−/−</sup> MEF in metaphase or prometaphase (indicated). p110β<sup>−/−</sup> MEF showed supernumerary chromosomes and fused chromosomes (two chromosomes with one centromere, inset 2, arrowhead). Control adjacent chromosomes (with two centromeres) are shown in inset 3 and 4 (arrows). Representative images of n = 50 examined. (Scale bar = 15 μm.) (C) Percent-MEF with the indicated phenotypes. (D) Repair of DNA DSB is stimulated by a phosphorylation-based signaling cascade termed the DNA damage response. The earliest DSBR sensor for HR is the MRN complex, which binds to DNA and activates the class IV PI3K ATM. ATM permits binding of replication protein A, which assists from pSG5-myc-p110β domain in the pEGFP-C1 vector was a gift of J. Downward (Cancer Research, London, United Kingdom). pSG5-Myc-p110β (7), pEGFP-C1-35B3 (24), murine Nbs1-2GFP (30), and pEn-mRFP-PCNA2 (33) have been described. pEGFP1-p110β and pEBG-GST-p110β were constructed by subcloning p110β from pSG5-myc-p110β into pEGFP-C1 and pEBG. GFP-Btk-Ph-Domain was described (26). pEGFP-C2-WT- Nbs1b, pEGFP-C2-ΔNbs1b, and -Ifh1 were prepared by subcloning into pEGFP-C2 (mutants were kindly donated by Y. C. Chen, Yang-Ming University, Taipei, Taiwan) (29).

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DSB, associates to DNA-PK, and promotes cell survival (37). PTEN, a phosphatase that dephosphorylates PIP3, also controls DSB repair by regulating Rad51 expression (38). The potential crosstalk between PI3Kβ, PKBα/Akt1 and PTEN action in DDR requires further analysis.

We conclude that whereas p110β activity (PIP3) facilitates DNA repair, p110β expression is required for this process, supporting the concept that p110β contribution in DDR is mainly kinase-independent. The kinase-independent function of p110β was also seen in KR-p110β knock-in mice; these mice were born at lower numbers (50%) than expected. Moreover, at E 13.5, two groups of
Before laser treatment, the cell medium was changed to a phenol red-free DMEM (Invitrogen). Immediately after microirradiation, repeated images of the same field were acquired in an integrated confocal unit operated by LCS software v2.6.1. Images were recorded at ~3.2 s after DSB generation, with a gap of 3.2 s per image.

**Cell Lysis, Immunoprecipitation, Western Blotting, PI3K Assay, and GST-p110β Pull-Down.** Total cell lysates were prepared in RIPA lysis buffer (7). For protein-protein interactions, cells were extracted as cytosolic and nuclear fractions (7). Triplet fractionation (cytosol, nuclear soluble, and chromatim). IP, Western blot, and PI3K assays were as described (7).

For pull-down, NIH 3T3 cells transfected with pEBG-GST or p-p110β-NLS (48 h) were fractionated as cytoplasmic and nuclear extracts. Nuclear lysates (1 mg) were incubated with glutathione beads (2 h, 4 °C). Beads were washed twice with lysis buffer and once with 50 mM Tris (pH 7.5; bound p110β) was resuspended in 2 × Laemmli buffer and resolved by SDS/PAGE. The gel was stained using a silver staining kit (Amersham). Stained protein bands were excised and the gel cut into small pieces and analyzed by mass spectrometry. p110β-associated proteins were identified by comparison with the peptide sequence database.

**Immunofluorescence, Comet Assay, and Statistical Analyses.** Cells were fixed with 4% formaldehyde in PBS (10 min, room temperature), blocked using PBS buffer (0.3% TX-100) for 1 h (room temperature). Cells were incubated with antibodies (1 h, room temperature), followed by three washes with staining buffer. Secondary antibodies were added to samples and incubated (1 h, room temperature), followed by three washes. Mounting medium containing DAPI (VectaShield) was added and cells were visualized in an AV100 Flow-view Olympus microscope or Leitz DMRB (Leica). Neutral comet assays were performed using the Comet Assay Kit (Trevigen) according to manufacturer’s instructions.

**Statistical analyses** were performed using StatView 5[12]. Gel bands, curve integration, and fluorescence intensity were quantitated with ImageJ software. Cell cycle profiles were analyzed with multicyle AV for Windows (Phoenix Flow Systems). Statistical significance was evaluated with the Student’s t test and the χ² test calculated using Prism5 v5.0 software. Fluorescence values in the irradiated region were annotated during recording. NFU were obtained by subtracting the fluorescence value of the first frame and comparing this value to maximal fluorescence intensity in control cells (considered 1). Tail Moment was calculated using Comet Assay IV software from Perceptive Instruments.
Fig. S1. p110β deletion impairs G2/M checkpoint. (A) NIH 3T3 cells were arrested at the G1/S border by double-Thi block and released. At 5 h after release, cells were treated with TGX221 (30 μM, 1 h), and at 6 h samples were irradiated (IR, 5 Gy). The proportion of mitotic cells was analyzed by p-histone H3* staining at 12 h after release. The graph shows the fraction (mean ± SD, n = 3) of p-histone H3* cells in each condition compared to maximum (~45% p-H3* cells in nonirradiated controls, considered 100). To control inhibitor activity, we examined extracts from exponentially growing NIH 3T3 cells alone or preincubated with TGX2221 (30 μM, 1 h) by Western blot using phospho-PKB Ab. (B) NIH 3T3 cells transfected with p110β shRNA (shRNA4, 48 h) or (C) p110β−/− immortalized murine embryonic fibroblasts (MEF), some reconstituted with WT or kinase-dead (KR) p110β, were treated as in A. Maximum p-histone H3* cells was ~15% in control NIH 3T3 G2/M cells and ~10% in control G2/M MEF. In B, to control shRNA efficiency, extracts from NIH 3T3 transfected with four different sets of p110β shRNA (48 h) were examined by Western blot. All four p110β shRNAs induced a similar defect in G2 arrest. In C, to control PI3K pathway status in MEF, we examined extracts from S phase cells by Western blot using phospho-PKB. *, P < 0.05.
Fig. S2. p110β localizes in DSB foci. (A) NIH 3T3 cells were irradiated (IR, 5 Gy) and processed after 1 h for immunofluorescence (IF) using anti-p110β Ab. Insets show DAPI staining. Graphs show IF intensity along the depicted line in arbitrary units (AU) in a representative cell (n = 50). (B) Graph illustrates the integrated area of each individual peak obtained in the IF-signal profiles performed as in A. (*), P < 0.05. The graph shows that the p110β-IF integrated signal in focus of IR cells is greater. (C) Irradiated NIH 3T3 cells (IR, 5 Gy) were processed for indirect IF upon IR (15 min), using anti-γH2AX (red) and -p110β (green) Ab. Insets show 20x magnifications. Arrows indicate some of the foci in which γH2AX and p110β colocalize. (D) NIH 3T3 cells expressing GFP were microirradiated with an UV laser. We examined real-time GFP translocation to the DNA damage region. Dotted lines indicate laser paths. (Scale bars A and D, 15 μm.)
**Fig. S3.** Endogenous PIP₃ localizes in DSB foci. (A) Exponentially grown NIH 3T3 alone or pretreated (1 h) with TGX221 or Ly294002 were stained by IF using anti-PIP₃ Ab. Insets show DAPI staining. The graphs illustrate the integrated signal from the entire cell or from the nucleus (indicated). (*), P < 0.05. (B) NIH 3T3 cells were irradiated (IR, 5 Gy); after 30 min, PIP₃ (green), γH2AX (red), and DNA (DAPI) were examined by IF. (C) NIH 3T3 cells were irradiated (IR, 5 Gy); after 30 min, PIP₂ (green) and DNA (DAPI) were examined by IF. (D) NIH 3T3 cells expressing the R25C mutant of the Akt PH domain were microirradiated with an UV laser and we examined real-time translocation of the probe to the laser path (dotted lines). (Scale bars in A–D, 15 μm.)
Fig. S4. Defective phospho-γH2AX and 53BP1 focus assembly in irradiated p110β-deficient cells. (A) NIH 3T3 cells were transfected with p110β or control shRNA (48 h); the latter cells were treated with TGX221 (30 μM) or DMSO (4 h). Cells were exposed to IR or UVC, and processed (5 min later) for indirect IF using antiphospho-γH2AX Ab. The graphs represent the integrated nuclear fluorescence intensity for a representative set of cells (n = 20) of > 100 examined. (Scale bar = 15 μm.) Unpaired χ² P values indicated. (B) NIH 3T3 cells were transfected with p110β or control shRNA (48 h) in combination with GFP-53BP1 (24 h); other cells were transfected with GFP-53BP1 (24 h) and treated with TGX221 (4 h). Cells were microirradiated with an UV laser. Normalized fluorescence units (NFU) along laser tracks was calculated and plotted as a function of time. The assembly curves represent mean ± SD (n = 6). (*), P < 0.05.
Fig. S5. p110β regulates proliferating cell nuclear antigen (PCNA) recruitment to damaged DNA. (A) p110β−/− immortalized MEF alone or reconstituted with WT or KR-p110β were irradiated (IR, 5 Gy) and collected after 1 h. Cells were fractionated into cytosol/cytoskeleton, nuclear (N) and chromatin (Chr) fractions, and the last two fractions were examined by Western blot using anti-PCNA Ab. The graph shows the percentage of chromatin-bound PCNA compared to that in the nuclear soluble fraction (considered 100%; mean ± SD, n = 3). (*), P < 0.05. (B and C) NIH 3T3 cells transfected (24 h) with RFP-PCNA were microirradiated and recorded. The RFP-PCNA concentration at double-strand break sites was determined by confocal microscopy live imaging. Dotted lines in the first image frame indicate the laser paths across the nucleus. Normalized fluorescence units along laser tracks was calculated and plotted as a function of time. Assembly curves represent the mean ± SD (n = 6), (*) , P < 0.05. (B)). (D) p110β−/− immortalized MEF alone or reconstituted with WT or KR-p110β were cotransfected with RFP-PCNA and GFP-Nbs1 (24 h). Cells were microirradiated with an UV laser. We examined real-time RFP-PCNA and GFP-Nbs1 translocation to the DNA damage region. The figures shows the RFP-PCNA and GFP-Nbs1 signals at three recording times of representative cells (indicated). (Scale bars C and D, 15 μm.)
**Fig. 56.** p110β deletion induces radiation sensitivity. NIH 3T3 cells were transfected with p110β or p110α-specific shRNA (48 h). p110α and p110β expression levels were examined by Western blot. Cell-cycle profiles in control or p110α- and p110β-shRNA-transfected cells were examined by flow cytometry at different times post-UVC exposure (80 J/m²). The percentage of cells with sub-G1, G0/G1, S, or G2/M DNA are indicated. The graph shows the percentage of cells with a sub-G1 DNA content 24h upon IR (mean ± SD, n = 5). (*), P < 0.05.

**Movie S1.** A representative NIH 3T3 cell expressing GFP-p110β; the video includes 89 frames taken every 3.2 s after laser irradiation.

**Movie S1**
**Movie S2.** Three concatenated movies of NIH 3T3 cells expressing GFP-S3BP1: a representative laser-irradiated NIH 3T3 cell (first cell), a NIH 3T3 cell pretreated with TGX221 (second cell), and a NIH 3T3 cell transfected (48 h before irradiation) with p110β shRNA (third cell). For recording, frames were taken every 3.2 s after laser irradiation.

**Movie S3.** Concatenated movies of representative laser-irradiated MEF expressing GFP-Nbs1 recorded every 3.2 s. The movie includes a representative MEF expressing WT-p110β (first), a KR-p110β reconstituted p110β−/− MEF (second), and a p110β−/− MEF (third).
Movie S4. Concatenated movies of representative U2OS cells expressing WT or mutated GFP-hNbs1 upon laser irradiation (recorded every 3.2 s). Cells were transfected with WT-GFP-hNbs1 (first), GFP-A465-hNbs1 (second), or GFP-A4670-hNbs1 (third).

Movie S5. Representative RFC-PCNA-expressing NIH 3T3 cells microirradiated with an UV laser and recorded (first cell), or pretreated with TGX221 (1 h) before irradiation (second cell), or transfected (48 h before irradiation) with p110β shRNA (third cell, concatenated movies). The videos include ~50 frames taken every 3.2 s after irradiation.
Movie S6. With Movie S7, a simultaneous analysis of GFP-Nbs1 (green) and RFP-PCNA (red) concentration at laser irradiated tracks in p110β−/− MEF reconstituted with WT-p110β (first cell) or KR-p110β (second cell, concatenated movies). Video starts immediately after laser irradiation; frames were recorded every 3.2 seconds. The video was mounted with half of the Nbs1-GFP frames (every 6.4 s) and half of the RFP-PCNA frames (every 6.4 s).

Movie S6

Movie S7. A typical p110β−/− MEF examined similarly to that shown in Movie S6. Video starts immediately after laser irradiation; frames were recorded every 3.2 seconds. The video was mounted with half of the Nbs1-GFP frames (every 6.4 s) and half of the RFP-PCNA frames (every 6.4 s).

Movie S7