Tight regulation of ubiquitin-mediated DNA damage response by USP3 preserves the functional integrity of hematopoietic stem cells

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Abbreviations used: CLP, common lymphoid progenitor; CMP, common myeloid progenitor; DDR, DNA damage response; DSB, DNA double-strand break; DUB, deubiquitinating enzyme; ES, embryonic stem; GMP, granulocyte-monocyte progenitor; HSC, hematopoietic stem cell; IR, ionizing radiation; IRIF, IR-induced foci; IRIF, IR-induced focus; LSK, Lin−Sca1+c-Kit+ long-term HSC; LTC-IC, long-term culture–initiating cell; MPP, multipotent progenitor; MNC, mononuclear cell; MEP, megakaryocyte-erythrocyte progenitor; MNC, mononuclear cell; MPP, multipotent progenitor; SCE, sister chromand exchange; ST-HSC, short-term HSC; Ub, ubiquitin; USP3, Ub-specific protease 3; WBC, white blood cell; wpt, weeks post transplantation.

Histone ubiquitination at DNA breaks is required for activation of the DNA damage response (DDR) and DNA repair. How the dynamic removal of this modification by deubiquitinating enzymes (DUBs) impacts genome maintenance in vivo is largely unknown. To address this question, we generated mice deficient for Ub-specific protease 3 (USP3; Usp3Δ/Δ), a histone H2A DUB which negatively regulates ubiquitin–dependent DDR signaling. Notably, USP3 deletion increased the levels of histone ubiquitination in adult tissues, reduced the hematopoietic stem cell (HSC) reserves over time, and shortened animal lifespan. Mechanistically, our data show that USP3 is important in HSC homeostasis, preserving HSC self-renewal, and repopulation potential in vivo and proliferation in vitro. A defective DDR and unresolved spontaneous DNA damage contribute to cell cycle restriction of Usp3Δ/Δ HSCs. Beyond the hematopoietic system, Usp3Δ/Δ animals spontaneously developed tumors, and primary Usp3Δ/Δ cells failed to preserve chromosomal integrity. These findings broadly support the regulation of chromatin ubiquitination as a key pathway in preserving tissue function through modulation of the response to genotoxic stress.

Maintenance of genome integrity is crucial for organism physiology, including stem cell and tissue homeostasis and tumor avoidance (Hoeijmakers, 2009; Jackson and Bartek, 2009). To protect the genome from the continuous challenge of exogenous as well as endogenous genotoxic agents, cells have evolved highly effective mechanisms, collectively termed the DNA damage response (DDR; Lukas et al., 2011; Polo and Jackson, 2011). Evidence of an essential role for the DDR in tissue homeostasis has been provided in the hierarchically organized process of hematopoiesis. In fact, mouse models with defective genome maintenance mechanisms suffer from severe hematopoietic phenotypes and hematopoietic stem cell (HSC) deficiencies (Ito et al., 2004; Nijnik et al., 2007; Rossi et al., 2007).

In these animals, the accumulation of DNA damage correlates with a progressive decline of HSC proliferation and self-renewal capacity (Rossi et al., 2007; Niedernhofer, 2008). Age-dependent accrual of DNA damage occurs also in WT murine and in human HSCs, emphasizing the impact of unresolved DNA stress on the functional decline of adult stem cells, with direct consequences on tissue function (Rossi et al., 2007; Rübe et al., 2011).

DNA damage can elicit different cellular responses in HSCs, depending on the nature and extent of the lesions. For instance, unrepair DNA double-strand breaks (DSBs) can drive cell cycle arrest and senescence or apoptosis, thus contributing to impairment in stem cell function during aging. When imprecisely repaired, DNA
damage can persist as a fixed mutation in HSCs, which can be propagated to daughter cells through self-renewal and differentiation and may ultimately lead to oncogenic transformation (Rossi et al., 2008; Sperka et al., 2012). Specific properties distinguish HSCs from more restricted progenitors. Among these features is cell cycle regulation, as HSCs are mostly in a quiescent state, whereas restricted progenitors divide more frequently (Kiel et al., 2007a). Furthermore, HSCs reside in hypoxic niches in the BM. Both conditions limit the chance of HSCs to be exposed to physiological DNA damage by replication errors or reactive oxygen species, respectively (Morrison et al., 1995; Morrison and Spradling, 2008; Rossi et al., 2008).

Finally, a different DDR and the preferential use of certain DNA repair pathways may also play a role in determining how DNA damage is sensed by stem cells, as well as an altered lineage potential (Rossi et al., 2008; Pang et al., 2011; Geiger et al., 2013). Recent data point to a contribution of DDR to this phenomenon. In fact, genotoxic stress/telomere attrition can activate an HSC-specific differentiation checkpoint that drives premature lymphoid differentiation, limiting the self-renewal potential of damaged HSCs and concomitantly depleting lymphoid-competent HSCs (Wang et al., 2012). Despite the recent advances, much remains to be learnt about the molecular pathways that limit the function of aging HSCs.

Ubiquitin (Ub) modification plays a central role in the functional consequence of aging on the HSC pool is a reduced capacity in replenishing the blood system with mature cells, as well as an altered lineage potential (Rossi et al., 2008; Pang et al., 2011; Geiger et al., 2013). Recent data point to a contribution of DDR to this phenomenon. In fact, genotoxic stress/telomere attrition can activate an HSC-specific differentiation checkpoint that drives premature lymphoid differentiation, limiting the self-renewal potential of damaged HSCs and concomitantly depleting lymphoid-competent HSCs (Wang et al., 2012). Despite the recent advances, much remains to be learnt about the molecular pathways that limit the function of aging HSCs.

Ubiquitin (Ub) modification plays a central role in the response to DSBs, ensuring efficient DNA damage recognition, signaling, and repair (Lukas et al., 2011; Jackson and Durocher, 2013). After phosphorylation of the histone variant H2AX (yielding γH2AX), the coordinated activity of the RNF8 (Huen et al., 2007; Kolas et al., 2007; Mailand et al., 2007; Wang et al., 2007) and RNF168 E3 ligases leads to histone H2A and H2AX ubiquitination (Doi et al., 2009; Pinato et al., 2009; Stewart et al., 2009), which is required for efficient and stable accumulation of DDR factors, such as 53BP1 and the tumor suppressor BRCA1 (Lukas et al., 2011; Jackson and Durocher, 2013). DSB-induced ubiquitination engages several E3 ligases (Lukas et al., 2011; Jackson and Durocher, 2013). Among these, the Polycomb Ring1b/Bmi1 mono-ubiquitinate histone H2A on lysine 119 (K119; Ismail et al., 2010; Ginjala et al., 2011; Vissers et al., 2012).

Under physiological conditions, this is a prevalent modification that can constitute ~10–15% of nuclear H2A and it acts as a repressive mark for gene transcription (Goldknopf et al., 1977; Simon and Kingston, 2013). RNF8-RNF168 ligases target H2A on a distinct site, lysine 13/15 (Gatti et al., 2012; Mattiroli et al., 2012), and this K15ub mark is read by 53BP1 (Fradet-Turcotte et al., 2013). The interplay between the two Ub marks on H2A is still poorly understood (Shanbhag et al., 2010; Ginjala et al., 2011; Mattiroli et al., 2012).

Ubiquitination is dynamic and reversible (Komander et al., 2009), and to date only a few deubiquitinating enzymes (DUBs) have been implicated in negative regulation of the RNF8–RNF168 pathway (Lukas et al., 2011; Jackson and Durocher, 2013; Mosbech et al., 2013; Panier and Durocher, 2013). Among these, our group has identified the Ub-specific protease 3 (USP3). USP3 is a chromatin-associated DUB strongly active in reverting histone H2A and H2B mono-ubiquitination (Nicassio et al., 2007). Several observations link USP3 activity to DDR. First, transient USP3 silencing led to spontaneous DNA breaks, enhanced 53BP1 nuclear...
More generally, our findings indicate that deubiquitination events are as critical as ubiquitination for preventing chronic genotoxic stress in vivo.

RESULTS

USP3-deficient mice show normal postnatal development and genome-wide increase in mono-ubiquitinated H2A and H2B

To investigate the biological role of USP3, we have generated conditional USP3-deficient mice (Fig. 1, A and B; Materials and methods). Germline disruption of the floxed Usp3 allele was achieved through the actin-cre deleter strain (Fig. 1, C and D). Usp3-deleted (Usp3Δ/Δ) pups were born at Mendelian ratio and with no overt phenotypes in the presence of effective USP3 deletion as measured by protein and mRNA levels (Fig. 1, E–G).

foci, and DDR activation in human cancer cells. Second, upon ionizing radiation (IR), USP3-depleted cells were unable to efficiently resolve γH2AX and Ub conjugates at DNA damage foci, with a concomitant delay in cell cycle recovery (Nicassio et al., 2007). Finally, overexpression of USP3 opposes the accumulation of RNF168 and the downstream mediator 53BP1 at IR–induced DSBs in a manner dependent on its catalytic activity (Doil et al., 2009; Mosbech et al., 2013; EC, unpublished data). Altogether, these findings support a role for USP3 in counteracting RNF8-RNF168–mediated signaling by removal of Ub marks from chromatin. However, it remains unresolved whether USP3 is key in chromatin ubiquitination in vivo. Here, we investigate the consequences of USP3 loss of function in vivo, which enabled us to discover a critical role for de-ubiquitination in maintaining tissue homeostasis under physiological conditions and stress response.

More generally, our findings indicate that deubiquitination events are as critical as ubiquitination for preventing chronic genotoxic stress in vivo.
To address the effect of loss of USP3 on nuclear ubiquitination, we performed immunofluorescence analysis with the pan anti–mono- and –poly-ubiquitinated substrate FK2 antibody. USP3Δ/Δ primary mouse embryonic fibroblasts (MEFs) displayed a significantly higher FK2 signal and a pronounced enrichment in nuclear foci compared with WT cells (Fig. 2A). Attributing increased ubiquitination to USP3 loss of function, ubiquitination levels were effectively rescued when an active USP3 protein was stably reexpressed in these cells (Fig. 2B).

Figure 3. USP3Δ/Δ mice exhibit shorter lifespan, increased tumorigenesis, and spontaneous genotoxic stress in MEFs. (A–E) Cohorts of WT (n = 26) and USP3Δ/Δ (n = 34) mice were monitored for survival for 90 wk. (A) Kaplan Meier general survival analysis. (B) Histopathological analysis of spleens from WT and USP3Δ/Δ mice and representative H&E-stained spleen sections from 5-mo-old animals. Bars: (left) 500 µm; (right) 20 µm. Low myelopoiesis in one 10-mo-old USP3Δ/Δ animal; low lymphoid compartment in a 15-mo-old USP3Δ/Δ mouse. (C) Kaplan Meier tumor-free survival analysis and distribution of tumor types in USP3Δ/Δ mice. (D and E). H&E staining of histological sections of representative malignancies in USP3Δ/Δ mice. (D) Moderately differentiated papillary carcinoma of the lung (17 mo). (E) Adenomatosis in the stomach (14 mo). Bars: (top) 500 µm; (bottom) 50 µm. (F) Constant field gel electrophoresis (CFGE) analysis of WT and USP3Δ/Δ MEFs. Results are the mean ± SD of three independent experiments. (G) Quantification of chromosomal aberrations in metaphase preparations of WT and USP3Δ/Δ MEF. A minimum of 42 cells/genotype was assessed. Mean ± SEM of one of two representative experiments is shown. Bar, 10 µm. (H) SCEs analysis in WT and USP3Δ/Δ sh-p53 MEFs. SCEs in representative metaphases are indicated by arrows. Inset, chromosome with double SCE. SCEs in a minimum of 48 cells/genotype were quantified. Mean ± SEM of one of two representative experiments is shown. Bar, 10 µm. For all panels: *, P ≤ 0.05; **, P ≤ 0.01; ***, P ≤ 0.001. P-value was assessed by Log-rank test (A and C) or by Student’s t test (F–I).
signal, we isolated core histones from MEFs. Immunoblotting revealed that the mono-ubiquitinated forms of H2A (uH2A) and H2B (uH2B) were reproducibly more abundant in Usp3Δ/Δ compared with WT MEFs (Fig. 2 C). Enhanced chromatin-bound FK2 Ub signal and uH2A/uH2B levels were also found in vivo, as exemplified by freshly isolated BM cells and by histone fractions purified from spleen and liver from Usp3Δ/Δ mice (Fig. 2, D and E). Together, our data indicate that USP3 is dispensable for normal embryogenesis and postnatal development but it is required to prevent accumulation of uH2A and uH2B in adult tissues, providing our previous in vitro data (Nicassio et al., 2007) with in vivo significance.

Loss of USP3 leads to shorter life span and increased cancer incidence

To examine the effect of USP3 deletion on animal life span, we monitored cohorts of Usp3Δ/Δ (n = 34) and WT (n = 26) mice up to 90 wk of age. Kaplan Meier survival analysis indicated that Usp3Δ/Δ mice have a significantly shorter lifespan compared with their WT littermates (Fig. 3 A).

We detected a hypocellular spleen in 7 out of 34 Usp3Δ/Δ mice (20.5%) between 6 and 18 mo, whereas only 1 out of 26 (3.8%) WT mice showed such phenotype at the age of 18 mo, suggesting a decline in adult tissue function (Fig. 3 B). Reinforcing this, clinicopathological analysis revealed that Usp3Δ/Δ mice had a significant decrease in tumor-free survival compared with their WT counterparts (Fig. 3 C). Spontaneous tumors occurred in 11 of 34 (32.3%) Usp3Δ/Δ mice compared with only 1 of 26 (3.8%) WT mice. (Fig. 3 C and Table S1). Pre-neoplastic lesions and a broad spectrum of tumor types in the lung, gastrointestinal tract, and brain were specifically associated with the Usp3Δ/Δ genotype (Table S1; Fig. 3, C–E). Collectively, our data show that Usp3Δ/Δ mice are susceptible to tissue function decline and neoplasia, a phenotype consistent with earlier studies on physiological ageing and experimentally enforced accumulation of DNA damage (Hoeijmakers, 2009; Jackson and Bartek, 2009).

USP3-deficient mice develop lymphopenia with age

Mouse models with a defective DDR display severe hematopoietic phenotypes and HSC deficiencies (Ito et al., 2004; Nijnik et al., 2007; Rossi et al., 2007). Consistently, Usp3Δ/Δ mice showed defects in hematopoietic tissues (Fig. 3 B). Therefore, we set out to determine whether USP3 deficiency functionally affected hematopoiesis.

First, we analyzed peripheral blood in 10–17- and 44-wk-old (aged) animals. We found a considerable decrease in the numbers of white blood cells (WBCs) in Usp3Δ/Δ aged mice only compared with their WT littermates, suggesting an age-dependent cellular attrition in the hematopoietic system in Usp3Δ/Δ mice. Lymphocytes were more affected by USP3 deletion (Table 1).

Next, we quantified our results using flow cytometry. Significant absolute and relative loss of both B (B220+) and

| Cohort (n) | Age | WBC ×10^9/liter | Lymphocyte ×10^9/liter | RBC ×10^12/liter | Hgb mM/liter | Plt ×10^9/liter |
|-----------|-----|-----------------|------------------------|------------------|-------------|----------------|
| WT (9)    | 10–17 | 4.8 ± 1.3       | 4.4 ± 1.3              | 4.6 ± 0.3        | 4.5 ± 0.4   | 621 ± 184      |
| Usp3Δ/Δ (13) | 10–17 | 4.2 ± 1.4       | 3.8 ± 1.3              | 4.4 ± 0.2        | 4.3 ± 0.2   | 584 ± 192      |
| WT (10)   | 44   | 5.7 ± 1.4       | 5.1 ± 1.0              | 4.6 ± 0.3        | 4.5 ± 0.3   | 628 ± 292      |
| Usp3Δ/Δ (13) | 44   | 3.6 ± 1.4***    | 3.2 ± 0.9***           | 4.3 ± 0.1*       | 4.3 ± 0.2   | 506 ± 167      |

Hgb, hemoglobin; Plt, platelets. Significant difference between USP3Δ/Δ and WT mice as determined by Student’s t test: *, P ≤ 0.05; ***, P ≤ 0.001. Numbers indicate means ± SEM. Results are from three independent experiments.
T (CD3+) lymphoid lineages was observed in the blood of aged mice, whereas alterations in the myeloid population (CD11b+) fell below the threshold for significance (Fig. 4 A and S1). Importantly, skewed hematopoiesis was mirrored in the BM of Usp3ΔΔ mice. FACS analysis showed that the lymphoid compartment (CD19+) was substantially reduced in aged but not in 17-wk-old Usp3ΔΔ mice, whereas the myeloid (CD11b+) lineage was not considerably affected (Fig. 4 B and not depicted). During B cell development, all B cell subpopulations were uniformly depleted in aged Usp3ΔΔ mice only (Fig. 4, C and D). Thus, loss of USP3 significantly affected hematopoiesis, notably leading to age-dependent cell loss of B and T cell lineages.

Qualitative and quantitative defects in the adult hematopoietic stem and progenitor cell compartment in Usp3ΔΔ mice

Because B cell defects occurred at early stages of commitment, the USP3 phenotype may arise in the hematopoietic stem and progenitor cell compartment rather than at later stages of development such as at B cell specification or during programmed rearrangements of the IgH locus (Alt et al., 2013). To test this, we performed multiparameter flow cytometry in 17- and 44-wk-old mice on the Lin- Sca1+c-kit+ (LSK) compartment, which is enriched for HSCs. This analysis revealed a profound decrease in the LSK numbers in Usp3ΔΔ mice at both ages (Fig. 5, A and B; and Fig. S2 A). In addition, although the LSK population expanded in WT mice with age, consistent with previous results (Morrison et al., 1996), no increase in absolute numbers of LSKs was found in Usp3ΔΔ 44-wk-old animals (Fig. 5 A). These results show that deficiency in the HSC-enriched LSK compartment precedes the reduction in mature cells observed in the blood of Usp3ΔΔ mice, indicating that this compartment is dependent on USP3.

The LSK compartment includes subpopulations of both long-term HSC (LT-HSC; cells capable of long-term reconstitution of the hematopoietic system) and short-term HSC (ST-HSC), as well as multipotent progenitors (MPPs; Osawa et al., 1996; Christensen and Weissman, 2001; Yeung and So, 2009). To discriminate between these, we used the CD150 SLAM HSC surface receptor in conjunction with the LSK, flk2/CD135, and CD34 markers (Kiel et al., 2007b; Fig. S2 A).

In aged mice, USP3 loss resulted in significantly lower absolute numbers and frequencies of all three primitive populations: LT–HSCs (LSK, flk2/CD135 “CD34+CD150−”, 1.4-fold reduction), ST–HSCs (LSK, flk2/CD135 “CD34+CD150−”, 1.8-fold reduction), and MPPs (LSK, flk2/CD135 “CD150+”; two-fold reduction). In contrast, at 17 wk of age, Usp3ΔΔ animals only showed a reduction in ST–HSCs and MPPs, whereas the least proliferative LT–HSCs were less affected by USP3 deletion (Fig. 5, A and B).

In the more mature progenitor pools, USP3 loss resulted in a preferential decrease in common lymphoid progenitors (CLPs) already at 17 wk of age, with minimal changes in the numbers of common myeloid progenitors (CMPs), granulocyte-monocyte progenitors (GMPs), or megakaryocyte-erythrocyte progenitors (MEPs), indicating that the latter can functionally cope with a reduced pool of LSKs (Fig. 5 C and Fig. S2 B). Deficiency in CLP homeostasis was validated in vitro as colony formation capacity of Usp3ΔΔ BM cells in methylcellulose specifically affected lymphoid progenitors (Fig. 5, D and E).

Thus, based on phenotypic profiling, loss of USP3 had a significant effect on the size of the HSC, LSK, and CLP compartments that was aggravated upon aging, and caused quantitative and qualitative defects in colony-forming ability in lymphoid progenitor populations. Together with the notion that early embryogenesis is not affected in Usp3ΔΔ mice, the above results indicate that USP3 is functionally required for adult HSC maintenance.

USP3-deficient HSCs display cell-autonomous functional decline

To investigate the basis of the Usp3ΔΔ HSC phenotype, we performed in vivo and in vitro functional assays. To functionally test the HSC repopulation ability, we transplanted WT or Usp3ΔΔ BM (CD45.2) cells in a 1:1 ratio with WT Bl6 competitor population (CD45.1) into lethally irradiated WT Bl6 recipients (CD45.1). In this competitive transplantation experiment, we chose to use young adult mice (8 wk old), before numerical defects would affect the Usp3ΔΔ progenitor...
compartment. Consistent with our previous observation in Usp3Δ/Δ animals (Figs. 4 and 5), 2–16 wk post transplantation (wpt) peripheral blood analysis showed a moderate but significant decrease in the repopulating ability of Usp3Δ/Δ BM cells compared with WT donors (Fig. 6 A, left). In particular, Usp3Δ/Δ donor cells poorly contributed to the lymphoid B and T lineages (B220+ and CD3+, respectively). As expected, the myeloid (Gr1+) and the LSK compartments were unaffected (Fig. 6 A, middle and right, respectively).

Next, we performed a noncompetitive repopulation assay, in which 100% of WT or Usp3Δ/Δ BM from 39–42-wk-old mice was transplanted. To cope with the reduced number of HSCs in Usp3Δ/Δ mice at this age, we normalized the BM dose to the actual HSCs present at the time of transplantation (stem cell equivalent BM). Nonetheless, Usp3Δ/Δ LSKs and primitive HSCs in recipient BM at 16 wpt showed significantly less engraftment (Fig. 6 B). Finally, we performed serial transplantations. Reasoning that sequential transplantation forces LT-HSC to self-renew and to differentiate to repeatedly reconstitute the recipient hematopoietic system (Purton et al., 2006), we chose to use BM cells from young (8 wk old) donor mice. Notably, at 12 wpt, secondary recipients of Usp3Δ/Δ BM displayed a significantly reduced donor-derived LSK population compared with WT (Fig. 6 C).

To examine the impact of USP3 loss on HSC homeostasis in response to an external and temporally controlled source of hematopoietic stress, we exposed the mice to sequential injections of the pyrimidine analogue 5-Fluorouracil (5-FU). Although we did not detect BM hypocellularity in Usp3Δ/Δ mice at any age, a continuous regimen of 5-FU that induces severe myelosuppression exposed the defect in the BM reconstituting ability of aged Usp3Δ/Δ mice as estimated by the significant decrease in Usp3Δ/Δ BM cellularity compared with WT animals (Fig. 6 D). These data indicate that chronic stress exposure exacerbates the Usp3Δ/Δ phenotype in the early progenitor compartment, the LT–HSCs.

Finally, to corroborate our in vivo findings with an in vitro functional assay, we performed long-term culture–initiating cell (LTC-IC) assays, a retrospective assessment of the presence of functional HSCs via ex vivo propagation (van Os et al., 2008). To this end, we purified Lineage negative (Lin−) cells from 15–17-wk-old Usp3Δ/Δ and WT mice. After long-term culturing, Usp3Δ/Δ Lin− cells showed a significant decrease in colony-forming ability in methylcellulose, indicating a reduced activity of Usp3Δ/Δ primitive hematopoietic cells (Fig. 6 E). Collectively, these findings experimentally support a predominantly cell-autonomous function of USP3 in HSCs under both homeostatic conditions and stress-induced hematopoiesis.
USP3–deficient HSCs suffer from greater DNA damage in vivo and in vitro

To test whether insufficient preservation of genome integrity underlies the USp3Δ/Δ hematopoietic phenotype, we first performed immunostaining for γH2AX, a marker of DDR activation (Rogakou et al., 1998). We quantified the number of γH2AX foci in freshly isolated FACS-sorted (sorted) HSCs from aged (44 wk old) mice. Although only rare γH2AX foci were observed in WT HSCs (>80% of LT-HSCs and >60% of ST-HSCs showed no more than two foci per cell), a high percentage of the USp3Δ/Δ LT-HSCs and ST-HSCs stained positive for γH2AX, with 45 and 85%, respectively, showing multiple foci (Fig. 7, A and B). Accumulation of γH2AX foci in USp3Δ/Δ progenitors was a predominant, although not exclusive, feature of aged USp3Δ/Δ HSCs (Fig. 7 C).

To investigate the actual presence of DNA breaks, we performed an alkaline comet assay. A significantly higher proportion of increasingly damaged comet tails was detected in sorted LSKs and in total BM cells from aged USp3Δ/Δ mice, reflecting accumulation of DNA breaks in these cells (Fig. 7, D–F). Positive detection of comet tails was not due to an increase in apoptosis (see below). Collectively, these results demonstrate that USP3 protects hematopoietic stem and progenitor cells from accumulation of spontaneous DNA damage in vivo.

To test whether DNA damage occurs in USp3Δ/Δ HSCs owing to enforced proliferation upon stress-induced expansion, we sorted LT–HSCs from aged mice and investigated DNA damage accrual in actively proliferating HSCs in vitro. Analysis of γH2AX and 53BP1 foci indicated in USp3Δ/Δ HSCs, as measured by FK2 immunofluorescence (Fig. 7 H). Consistent with our previous findings in MEFs, conjugated Ub at the chromatin was also enhanced in USp3Δ/Δ LT-HSCs, as assessed by FK2 immunofluorescence (Fig. 7 H).

Because 53BP1 accumulation at DSB sites is dependent on RNF168-mediated ubiquitination (Doil et al., 2009; Stewart et al., 2009; Fradet-Turcotte et al., 2013), we asked whether enhanced FK2 would positively correlate with 53BP1 recruitment. Notably, double immunofluorescence staining revealed that colocalizing 53BP1-conjugated Ub foci were indeed enriched in USp3Δ/Δ LT-HSCs, suggesting that Ub-dependent DNA damage signaling was enhanced in these cells (Fig. 7 I).

To determine possible functional consequences of DNA damage accumulation in the absence of USP3, we analyzed micronuclei formation as biomarkers of genotoxic stress (Fenech et al., 2011). DAPI analysis showed a higher frequency of micronuclei in USp3Δ/Δ LT–HSCs cultures, which were exacerbated upon culturing (Fig. 7 J). Together,
Increased cellular aging underlies USP3-deficient BM defects

Persistent DNA damage may lead to either DDR-induced differentiation or cell cycle restriction, both of which may account for the phenotypes we discovered in USP3-deficient mice (Rossi et al., 2008; Sperka et al., 2012; Wang et al., these findings are consistent with 17-wk-old USP3-deficient proliferating ST-HSCs being more vulnerable, and so more easily lost, than quiescent LT-HSCs (Fig. 5, A and B), and implicate an integral role for USP3 in genomic maintenance of HSCs through the control of chromatin ubiquitination.

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Figure 8. Cellular senescence in Usp3Δ/Δ HSC compartment and BM. (A) Cytospins of BM cells from Usp3Δ/Δ and WT mice were assayed for SA-β-galactosidase activity. The percentage of SA-β-Gal–positive cells was quantified by counting 100 cells on three separate fields (n = 4 mice per genotype). Bar, 20 µm. (B and C) Cytospin preparations of sorted LSKs from BM of Usp3Δ/Δ or WT mice were immunostained for HP1γ (B) and H3K9Me3 (C). Focal nuclear staining is visible in insets. Signal intensity per nucleus was quantified by ImageJ. n = 3 per genotype. A minimum of 1,000 nuclei/sample was evaluated. Data are mean ± SEM of one of two representative experiments. Bars: 75 µm; (inset) 10 µm. (D) Immunostaining for HP1γ and H3K9Me3 on BM sections from Usp3Δ/Δ and WT mice. The percentage of positive cells was quantified in 3 fields on a minimum of 1,500 cells/field per sample. n = 7.
Hematopoietic analysis revealed that... Fig. 9 A). For all panels: *, P ≤ 0.05; **, P ≤ 0.01; ****, P ≤ 0.0001; ns, not significant.

was evaluated in two replicate experiments. Bar, 20 µm. Mice were 32 wk old (A–D) or 40–44 wk old (E and F). G–J: LT-HSCs for in vitro expansion were isolated from two independent experiments is shown (mean ± SEM).

time 0 and 96 h after plating are shown. Bar, 300 µm. (H) Immunostaining of in vitro expanded LT-HSCs for H3K9Me3. Signal quantification by ImageJ as phase contrast object confluence.

USP3 protects HSCs and mice from IR

To test the impact of replicative stress on LT-HSC fitness, we exposed these cells to c-Myc using a cell-permeable protein transduction approach. Notably, a significantly higher percentage of Usp3Δ/Δ LT-HSCs displayed SA-β-gal activity upon Myc expression, suggesting that they are more sensitive to oncogene-induced senescence (Fig. 8 J). Our results indicate that Usp3Δ/Δ HSCs engage cell cycle restriction under the conditions of cytokine-induced and oncogene-induced replicative stress.
by performing colony formation assays with BM cells from mice before and after IR treatment. *Usp3*ΔΔ/ΔΔ BM cells plated 7 d after exposure to 5 Gy showed significantly reduced colony formation. Consistent with our previous results (Fig. 5 F), no differences were observed with cells from nonirradiated mice (Fig. 9 D).
To investigate the consequences of IR on HSCs in vivo, mice were exposed to 5 Gy and sacrificed 24 h later. Notably, the Usp3Δ/Δ primitive progenitor compartment (i.e., LSKs and HSCs) was acutely decreased compared with WT controls (Fig. 9 E and Fig. S2 C). Reduction in LSK cellularity upon IR was accompanied by an increase in the fraction of Annexin V+/PI+ cells in Usp3Δ/Δ mice (Fig. 9 F).

To gain mechanistic insight into USP3-mediated protection of HSCs from genotoxic stress, we sorted LT-HSCs and analyzed their response to IR in vitro. Both WT and Usp3Δ/Δ HSCs showed a similar immediate induction of γH2AX and 53BP1 IR-induced foci (IRIF) upon 2 Gy. By 24 h after IR, however, although the great majority of IRIFs were resolved in WT cells, γH2AX and 53BP1 foci persisted in a significant fraction of Usp3Δ/Δ cells, suggesting a delay in DSB repair (Fig. 9, G and H). This was accompanied by an increase in the number of micronuclei, indicative of the actual presence of broken/lagging chromosome in Usp3Δ/Δ cells (Fig. 9 I).

Together, these data demonstrate that USP3-deficient HSCs are more sensitive to genotoxic stress in vivo and in vitro and support the notion that unresolved DNA damage affects the functional integrity of Usp3Δ/Δ HSCs in homeostasis, as well as upon aging and stress. Overall, the data point to USP3 as a critical mediator of DDRs in vivo.

DISCUSSION

We determine here the impact of negative regulation of DNA damage-induced Ub signaling on organism physiology, which was largely unexplored. USP3 is one of the few DUBs that have been linked to this aspect of DDR. (Nicassio et al., 2007; Doil et al., 2009; Jackson and Durocher, 2013; Panier and Durocher, 2013; Sharma et al., 2014). Using a mouse genetics approach, we provide evidence that defective deubiquitination does have profound consequences in vivo, as USP3-deleted mice have a shortened lifespan, associated with a functional decline of the hematopoietic stem and progenitor compartment and with sporadic spontaneous neoplasia. Linked to these phenotypes is chronic genotoxic stress, most likely resulting from the altered histone ubiquitination upon USP3 loss, as measured in vivo and in vitro. In support of this, we find that USP3 protects HSCs from DNA damage through modulation of Ub-dependent DDR signaling. This role in HSC maintenance is, to our knowledge, a previously uncharacterized function of the RNF8-mediated ubiquitination pathway.

USP3 and cancer

Similarly to other components of the Ub-DDR network, such as 53BP1, RNF8, and RNF168 (Ward et al., 2003; Li et al., 2010; Bohgaki et al., 2011; Halaby et al., 2013), Usp3Δ/Δ mice are mildly tumor prone, with up to 9% of the animals showing neoplasia within 1 yr of age. This is consistent with the moderate, though measurable, magnitude of chromosomal aberrations in Usp3Δ/Δ cells. Because a broad spectrum of tumor types was observed, we propose this pathology to be related to a general tissue function decline. Progressive accumulation of mutations as a consequence of unresolved DNA damage in Usp3Δ/Δ cells may underlie this phenotype. Although a genome maintenance defect is a plausible explanation, our results do not rule out the possibility that USP3’s role in tumor suppression may also involve deubiquitination of still uncharacterized substrates, besides H2A and H2B, potentially regulating distinct cellular processes. The ability to deubiquitinate multiple substrates is in fact not unprecedented within the USPs (Komander et al., 2009), and a novel target for USP3, RIG–1, has been recently reported (Cui et al., 2014).

USP3 and Ub-mediated DDR in HSC maintenance

Pathology arising from Usp3 deficiency clearly manifests in hematopoiesis. Although hematopoietic homeostasis is achieved in young adult Usp3Δ/Δ mice, there is indeed a marked decline in HSC activity over time. In physiological conditions, loss of USP3 mainly exacerbate the decline of the lymphoid progenitor compartment, a phenotype reminiscent of an aged hematopoietic system, in which functional decline of aged HSCs is associated with immunodeficiency and skewed differentiation in the elderly (Rossi et al., 2008; Pang et al., 2011). USP3 is accountable for the age-dependent cellular attrition in HSCs, because the BM from young (8 wk old) Usp3Δ/Δ mice, although functionally compromised compared with WT of the same age, performed better than BM from aged (44 wk old) mice. In turn, a gradual loss of functional HSCs is further exacerbated under conditions of stress, as demonstrated by the reduced capacity of Usp3Δ/Δ HSCs for self-renewal and by the lineage skewing upon transplantation in vivo, as well as by the reduced proliferative potential in vitro. Again, these features are characteristic of an aged HSC phenotype (Geiger et al., 2013) and directly implicate USP3 in protecting HSCs from the consequences of physiological stress in vivo.

In agreement with the enhanced DNA breaks in MEFs, USP3 loss was further associated with spontaneous DNA damage in HSCs in vivo and in vitro. Our results support the DDR to be implicated in the functional defects of Usp3Δ/Δ HSCs, as they feature increased radiation sensitivity in vivo and prolonged DDR upon IR in vitro. These findings are in agreement with several studies demonstrating the dependence of the hematopoietic system on an intact DDR (Ito et al., 2004; Nijnik et al., 2007; Rossi et al., 2007; Niedernhofer, 2008). Nornocellularity of the BM and a more pronounced skewing toward lymphoid lineages under homeostasis is shared with some DDR-deficient mouse models, such as the Ku80-deficient mice (Rossi et al., 2007). However, the Usp3Δ/Δ HSC phenotype as revealed upon BM transplantation appeared less severe compared with classical components of the HR, NHEJ, NER, or telomere maintenance pathways (Ito et al., 2004; Nijnik et al., 2007; Rossi et al., 2007; Niedernhofer, 2008). Because we observe enhanced ubiquitination in Usp3Δ/Δ mice, these data further support a role for USP3, and in turn for deubiquitination, as a modulator of DDR signaling, the alteration of which would have less severe consequences for
genome integrity compared with mutations in core DSB repair factors. In fact, neither apoptosis nor DNA damage-induced premature lymphoid differentiation significantly occurred in Usp3Δ/Δ primitive hematopoietic progenitors. Rather, we observed features of senescence. Not surprisingly, these features, along with spontaneous DNA damage, were further enhanced when we challenged Usp3Δ/Δ primitive progenitors with replicative stress in vitro, thereby implicating USP3 in fine-tuning DDR during proliferation. In this, our data on USP3 provide a framework to further investigate how lack of control of the Ub-dependent DDR pathway contributes to the functional decline observed in aged HSCs.

Histone ubiquitination by RNF8-RNF168 is central in DDR activation (Lukas et al., 2011; Jackson and Durocher, 2013). This raised the hypothesis that deubiquitination by USP3 may in turn be important for restraining/silencing the DDR. Consistently, enhanced Ub conjugates were accompanied by spontaneous 53BP1 foci as well as a prolonged retention of 53BP1 upon IR in Usp3Δ/Δ HSCs. Notably, these data extend previous observations obtained with RNAi by us and others (Nicassio et al., 2007; Gudjonsson et al., 2012). The pathologies found in Usp3Δ/Δ mice further indicate that in vivo, this aspect of DDR may be equally relevant to DDR activation in genome maintenance. We can envision at least two deleterious scenarios that could result from increased association/stabilization of genome caretakers such as 53BP1 with chromatin as a consequence of USP3 depletion. First, enhanced binding of DDR mediators might initiate unnecessary repair events, thus contributing to genomic instability. Consistent with this possibility, the spreading of chromatin ubiquitination at DSBs due to amplification of the RNF168 pool has been linked to enhanced DSB repair (Gudjonsson et al., 2012). Second, hyperubiquitination may lead to prolonged association of DDR factors at damaged DNA sites, which could in turn delay silencing of DDR signaling and cell cycle recovery (Nicassio et al., 2007). If deubiquitination is delayed, we hypothesize that persistent uH2A and possibly uH2B conjugates, which have also been implicated in regulating DSB repair (Moyal et al., 2011; Nakamura et al., 2011), may become detrimental during subsequent genetic insults. In such a situation, repair factors may get targeted simultaneously to repaired and de novo broken DNA. Over time, this scenario is likely to destabilize the genome, leading to chromosomal aberrations, stem cell depletion, and cancer.

In summary, the USP3 knockout mouse model establishes the importance of USP3 in limiting chronic genotoxic stress and HSC aging. Also, it highlights the relevance of understanding the mechanistic basis of chromatin deubiquitination, a still poorly understood aspect of DDR, and how it contributes to diseases related to chromosomal metabolism and to the pathogenesis of cancer.

MATERIALS AND METHODS

Mouse strains, monitoring of the mice, and TBL. Generation of conditional (Usp3flo; MGI:5490049; FVB;129P2-Usp3tm1.2Nki) and null (Usp3Δ; MGI:5490048) Usp3 alleles was performed as described below. LoxP sites were positioned in introns flanking the Znf-UBP domain coding region (exon 2 and 3) of Usp3. Actin-Cre was used for germine deletion. Mice in this study were in a mixed 129/Ola × FVB genetic background (MGI:5490048; FVB;129P2(B6)-Usp3tm1.1Nki). USP3-null lines generated from two independently targeted embryonic stem (ES) cell clones were analyzed in parallel for Mendelian offspring and for development and gave similar results. Mice used in BM reconstitution experiments were backcrossed into a C57BL/6j (The Jackson Laboratory) strain background (Ly5.2+) for 8 generations (MGI:5490048; B6.129P2(FVB)-Usp3tm1.1Nki). Mice were maintained in a specific pathogen-free environment. All animal experiments comply with local and international regulations and ethical guidelines and have been authorized by our local experimental animal committee at The Netherlands Cancer Institute (DEC-NKI). Mice were monitored for survival and tumor development for 22 mo. Time to death was defined as the latency between birth and unexpected death or a terminal disease stage indicated by >20% weight loss or other symptoms of severe sickness. Mice were sacrificed by CO2 asphyxiation and underwent necropsy. Organs were fixed as described below for histopathological analysis. Hypocellularity in the spleen of Usp3Δ/Δ mice was evaluated by blind examination of H&E-stained spleen sections. Sensitivity to IR was assessed with the use of 2–4-mo-old littermates. Mice were irradiated with an experimentally defined sublethal dose of IR (7 Gy; HF320 Radiobiology Constant Potential X-ray Unit, Pantak, East Haven, CT) and monitored for 28 d.

Construction of Usp3 targeting vector. Targeting construct for USP3 was generated using the pFlexible targeting vector system (van der Weyden et al., 2005). The pFlexible vector (pAG14) contains the PGK-driven positive/negative selectable marker puroDtk and loxP (L) and FRT (F) recombination target sites. This allow for efficient conversion of a single targeting event into a null or conditional mouse line.

A BAC clone (bMQ-415010; Geneservice) from 129S7/AB2.2 mouse DNA (indexed 129S7/SvEvBrl-Hprt-m2 (AB2.2) library, displayed on the Ensembl Genome Browser [http://www.ensembl.org/Mus_musculus/] under the DAS source 129S7/AB2.2) containing the USP3 locus was used to generate the USP3 targeting construct, and DNA fragments were amplified from the BAC DNA by PCR using the Expand Long range dNTPack kit (Roche). Fragments were as follows: the 3′ homology arm (3.5 kb PCR fragment) of exon 2 in the USP3 gene, flanked by Ascl restriction site: forward primer, 5′-GGCGCGCCCAAGTTGGAGATAGCCATGCC-3′; reverse primer, 5′-GGCGCGCCGAATTCTGTGTTAAAAACTGTGGTTTTCCCACA-3′; the conditional arm (4.3 kb PCR product containing exon 2 and 3, flanked by PacI sites; forward primer containing PacI-BamHI-KpnI sites, 5′-AGGATCCGGTACCTGTTGACTTCAAAAGAGAGTG-3′; reverse primer, 5′-TTAATTGAGATTTGTTGAAGTTCATCTCC-3′); reverse primer, 5′-TTAATATCTTCTCTATATATACCTTCACCTTCAAGAAAAGAGATGCTGTTGAAGATAGCCATGCC-3′; reverse primer, 5′-GGCGCGCCGCCTCGTCCCTCTGGTTCTTATTAGT-3′; reverse primer, 5′-GGCGCGCCCGCTTGGTTTTAAAAAGCAGATAACGAC-3′. PCR products were first cloned in pCR-XL-TOPO Vector (TOPO XL PRC Cloning kit; Invitrogen) using one short SfiI chemically competent Escherichia coli (Invitrogen). The PCR-XL–TOPO–cloned homology arms were fully sequenced prior subcloning into the pFlexible vector using the Ascl, PacI, and NotI restriction sites incorporated by PCR. Primers used for sequencing are available upon request. The final pFlexible-USP3CKO-FRT (conditional knockout–Flpe recombine target) construct was fully sequenced prior to electroporation.

Generation of Usp3<sup>CKO-FRT</sup> ES cells. The USP3 targeting vector was linearized with Clal and electroporated into 129/Ola-derived ES cells of the E14 Ubiline 1B10 (Robertus-Maandag et al., 1998). Cells were grown under 1.8 mg/ml puromycin selection on a lethally irradiated MEF feeder layer. After 5 d of drug selection, ~300 puromycin resistant ES clones were picked and amplified in nonselective medium. High molecular weight DNA was isolated according to standard procedures (Jonkers et al., 2001), and genomic DNA was amplified with a primer pair encompassing the LoxP1 site, primers 37 (forward, 5′-ATAATTGGCCTGAT-3′) and 38 (reverse, 5′-TATCTTGAGCTTGGATTGTC-3′), yielding a DNA fragment of 447 bp for the WT allele and of 519 bp for

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the targeted allele. For Southern blotting, the probes were generated and labeled by PCR. The 5′ probe (527 bp; forward primer, 5′-AAGG-GATTTGAGGATAGGCACGGGC-3′; reverse primer, 5′-TTTGAATCTACAGGATT-3′) hybridizes to a 10 kb BamHI restriction fragment in WT cells and a 7.2 kb fragment in homologous recombinants. The 3′ probe (600 bp; forward primer, 5′-ATAATGCTGTAAGGAAGGACGG-3′; reverse primer, 5′-AATTTCTCTTTACTCAGT-3′) recognizes a 7.6 kb EcoRI fragment in WT cells and a 10 kb fragment in recombinants. The probe recognizing the puroycin cassette (470 bp; forward primer, 5′-GAGCT- GCAAGAACCTTTCC-3′; reverse primer, 5′-ATATAAACCCCGAGTAGCC-3′) hybridizes to a 10 kb EcoRI fragment in recombinants only.

**Generation of Usp3CreERT2 conditional and Usp3-deleted (Δ) mice.** Three independent ES cell clones with homologous integration at the targeting site were injected into C57Bl/6j blastocysts to generate chimeric mice. These chimeras were subsequently crossed with FVB/N females to produce heterozygous offspring. Heterozygous mice with successful germ line transmission of the targeted allele were crossed with C57Bl/6j mice expressing Flp recombinase (B6.Cg-Tg(ACERTFLP)9205Dym/A; Rodriguez et al., 2009) to remove the neomycin resistance cassette resulting in Usp3Cre allele. To obtain Cre-mediated deletion of the floxed allele in the germline, heterozygous Usp3 conditional mice were crossed with Actin-Cre deleter (FVB/N), in which cre expression is under the human β-actin gene promoter and is expressed in all of the cells of the embryo by the blastocyst stage of development, and intercrossed to generate Usp3Δ/Δ mice.

**Genotyping of Usp3 mice.** Genotyping of F1 tail DNA was performed by Southern blot using the same DNA probes as described above for ES cells. The 3′ probe recognizes a 16.6 kb KpnI DNA fragment in the WT allele, a 9.5 kb fragment in the Usp3CreERT2 allele, and a 12.2 fragment in the deleted allele. The 3′ probe hybridizes to a 7.6 kb EcoRI fragment in WT cells, a 10 kb fragment in the Usp3CreERT2 allele, and a 6 kb fragment in the deleted allele. After founders were established, genotyping of F2 DNA was performed by PCR. Tail DNA was extracted using the Direct PCR DNA Extraction Reagents (QIAGEN) and PCR performed using Platinum PCR Supermix (Invitrogen). The Usp3 alleles were detected with the following primers: the LoxP1 site in intron 1, yielding a 447 and 519 bp product for the WT or Usp3CreERT2, respectively (forward primer 37, 5′-ATAATGCTGTAAGGAAGGACGG-3′; reverse primer 38, 5′-TCATCTGTAGCTTGTGATTGC-3′); the LoxP2 site in intron 3, yielding a 419 and a 519 bp product for the WT or the Usp3Δ, respectively (forward primer 35, 5′-GATGACAGCAACGATACTGG-3′; reverse primer 36, 5′-ATAGACAGGACTTTACTACC-3′); and the Usp3Δ allele, yielding a 544 bp product in the Usp3Δ allele (forward primer 37 and reverse primer 36). The Cre transgene (a 200 bp PCR fragment) was detected with the following primers: forward primer, 5′-GTGTTACCTGGTTACTATCAG-3′; reverse primer, 5′-ATAGACAGGACTTTACTACC-3′. The Cre transgene yields a 700 bp DNA fragment (forward primer, 5′-GGTTCAACTGAGCCCAAGCTTCC-3′; reverse primer, 5′-GTGATTACGTCCTTACACCTGC-3′; PCR for the FLIP transgene yielded a 780 bp DNA fragment (forward primer, 5′-GGTTCAACTGAGCCCAAGCTTCC-3′; reverse primer, 5′-GTGATTACGTCCTTACACCTGC-3′; Rodriguez et al., 2009).

**Protein analysis.** Total protein lysate from MEFS and histone fractions from cells and tissues were prepared as previously described (Nicassio et al., 2007). A complete list of antibodies used in Western blotting is found in Table S5.

**Hematology.** Peripheral blood was collected into EDTA-coated microtubes, and blood cell counts were analyzed on a COUNTER Ac T dif (Beckman Coulter). For FACS analysis, blood was depleted from red blood cells by hypotonic lysis and staining was performed with fluorescent-labeled antibodies specific for B220, CD3, CD11b. Samples were analyzed on a CyAn ADP Analyzer or on a LSR, Fortessa (BD). A complete list of antibodies used in flow cytometry is found in Table S6.

**Flow cytometric analyses and cell sorting of hematopoietic subpopulations.** BM mononuclear cells (MNCs) were stained for lymphoid and myeloid lineage markers. To analyze and isolate BM LSK, LT-HSC, ST-HSC, MPP, CLP, and CMP subpopulations (Osozawa et al., 1996; Christensen and Wessman, 2001; Kiel et al., 2007b; Karumky et al., 2008; Rodrigues et al., 2008; Yeung and So, 2009), MNCs were first stained with Lineage Cell Detection Cocktail-Biotinylated mouse antibody (Macs; Miltenyi Biotec). For FACS analysis, cells were then directly stained with fluorochrome-conjugated antibodies against Sca1 (Pacific blue), c-kit (APC), CD34 (FITC), CD135 (Flk2 (PE), CD150 (PECy7), CD16/32 (PerCPcy5.5), and CD127 (PerCPcy5.5). For cell sorting of LSK, LT-HSC, and ST-HSC, depletion of lineage+ cells was performed using BioLume MicroBeads (Macs; Miltenyi Biotec) and magnetic columns (Macs; Miltenyi Biotec) before staining. Samples were analyzed with a CyAn ADP Analyzer. Cell sorting was performed with a FACSaria (BD).

All FACS data were analyzed using FlowJo Software (Tree Star). Apoptosis in Lin-, LSKs, and HSCs (defined as above) was assessed in freshly isolated BM cells using the FitC Annexin V Apoptosis detection kit II (BD). Samples were analyzed with the LRS Fortessa (BD).

**BM transplantation.** In competitive BM transplantation, 1.5 × 10^6 BM MNCs from 8-wk-old donor test animals (CD45.2, WT, or Usp3Δ/Δ) were mixed in a 1:1 ratio with WT competitor cells (CD45.1) and injected into lethally irradiated (2 × 5.5 Gy TBI) C57BL/6-CD45.1 recipient mice.Recipient mice peripheral blood was monitored by FACS analysis every 2 wk for 16 wk. Donor contribution and multilineage reconstitution were assessed based on the expression, respectively, of CD45.1/CD45.2 antibodies or B220, CD3, and Gr1 markers in the CD45.2+ fraction. At 16 wpt, primary recipients were sacrificed and the frequency of donor-derived CD45.2 LSK assayed by phenotypic profiling. In noncompetitive transplantation, BM cells from 39–42-wk-old WT or Usp3Δ/Δ mice were transplanted for each genotype, cells from 3 donor mice were pooled and 5 × 10^6 cells injected into lethally irradiated CD45.2 primary recipients. Flow cytometry staining for LSK and HSC of donors was performed to ensure that the HSC frequency in test and control BM would be comparable. Differences in HSC were corrected before transplantation to transplant stem cell equivalents. LSKs and HSCs analysis of primary recipient was performed at 16 wpt, as described above.

For serial transplantation, irradiated primary recipients were injected noncompetitively with 3 × 10^5 BM cells (stem cell equivalents) from 8-wk-old WT or Usp3Δ/Δ mice (for each genotype, BM cells from 3 mice were pooled). For secondary transplants, primary recipients were sacrificed at 12 wpt, BM cells from 3 recipients per genotype were pooled, and 3 × 10^6 of these cells were transplanted into lethally irradiated hosts. Secondary recipients were sacrificed at 12 wpt and analysis for donor-derived Lin- LSKs, and HSCs was performed as described above. In each experiment, 5 recipient mice with test or control cells were assayed. Samples were analyzed with an LRS Fortessa.

**5-FU treatment.** 5 Fluorouracil (5-FU; Sigma-Aldrich) was administered to 44-wk-old mice weekly intraperitoneally at a dose of 150 mg/kg for 4 consecutive wk. BM cellularity (two femurs) was evaluated at 44 wk after 5-FU treatment.

**LTC-IC assay.** OP9 stromal cells were grown in Iscove’s modified Dulbecco’s medium supplemented with 20% FBS and 50 mM 2-mercaptoethanol. Confluent OP9 stromal layers in 6 wells were irradiated with 15 Gy and subsequently seeded with purified Lin− cells (15,000 cells per well) in triplicates. After 5 wk of culturing (MyeloCult M5350 [STEMCELL Technologies], supplemented with 276 ng/ml hydrocortisone; replacement of 1/2 medium volume every week), cells were harvested and seeded in methylcellulose for CFU-C colony assays (as below).

**RNA-seq gene expression analysis.** For gene expression analysis, LSKs (FACS sorted from freshly isolated BM), naive splenic B cells (freshly FACS sorted), or FACS-sorted LPS-activated B cells were used. n = 4 Usp3Δ/Δ and n = 4 WT littermates (2 mo old). Cells from individual animals were pooled and total RNA was extracted. Samples were prepared using TruSeq protocols, and standard sample preparation protocols and RNA-seq was performed on a HiSeq2000 machine (Illumina) at the NKI Genomics Core Facility.
Naïve splenic B cells were obtained by CD43 depletion using biotinylated anti CD43 (Clone S7; BD), and the iMag system (BD), as described by the manufacturer. Naïve B cells were directly FACS sorted with fluorochrome-conjugated antibody specific for CD19 (APC), or cultured in vitro for 4 d in IMDM + 8% FBS and 50 µg/mL E. coli LPS (055:B5; Sigma-Aldrich) to obtain activated B cells, followed by sorting. Two independent experiments were performed. Cell sorting was performed by FACSaria (BD). The data discussed in this publication have been deposited in NCBI’s Gene Expression Omnibus (Edgar et al., 2002) and are accessible through GEO Series accession no. GSE58495.

HSC liquid culture, time-lapse imaging, and Tat-cMyc fusion protein delivery. LT-HSCs were sorted from BM of 40–44-wk-old mice (WT n = 3, Usp3Δ/Δ n = 4). Cells from individual animals were pooled and grown in liquid cultures. Cells were grown in RHB-A Stem Cell Culture Medium (STEMCELL Technologies) containing SCF, Flt-3 ligand, and TPO (50 ng/mL), and IL-3, IL-6, and IL-11 (25 ng/mL; PeproTech). Medium was replenished and cells were expanded in 3% oxygen to maintain optimal growth. To evaluate proliferation, 1,000 cells were plated in 96-well plates at day 8 or 11 of culture. 4 wells per conditions were imaged (phase-contrast) with a 4 h interval for 6 d using the IncuCyte FRL (Essen BioScience). Confluenence was determined by the IncuCyte software, based on area (confluence) metrics. Plating of 500 cells/well gave similar results. The cDNA of human transcription factor cMYC (NM_020467.3) was PCR amplified and subcloned in pGEX6P1 plasmid containing the membrane penetrating peptide YGRKKRRQRRR from HIV TAT protein. Expression of recombinant Tat-cMyc was induced in log phase growing bacteria E. coli BL21 DE3 strain (Invitrogen) upon treatment of 1 mM isopropyl-b-D-galactoside for 3 h at 30°C. Purification comprised a further step to remove GST tag from Tat-cMyc protein using PreScission Protease (GE Healthcare). Liquid cultures of HSCs were treated with 1 µg/mL of purified Tat-c-Myc for 5 d, with daily replenishing of the fusion protein.

Quantitative real-time (qRT) PCR. Total RNA was extracted using TRIzol reagent (Life Technologies) and cDNA was prepared using SuperScript II RT and oligo(dT)₅ primers (Life technologies). qRT-PCR was performed on a StepOnePlus RT-PCR system (Applied Biosystems) using SYBR Green PCR Master Mix (Applied Biosystems). The amount of target, normalized to an endogenous reference (HPRT), was calculated by 2⁻ⁿΔCt. Primer sequences were as follows: Hprt forward, 5’-CTGTTGAAAGGACCTCTTCG-3’; Hprt reverse, 5’-TGAAGTACCTATCATATGCAGGCA-3’; Usp3Δ forward 1, 5’-TCGTGTGACGTGTCAAGTGTC-3’; and Usp3Δ reverse 1, 5’-ATGGTGTAGAATGCTTCTGC-3’. The DAPI channel (Blue) was used to select the nuclei of the cells in the field, and pixel intensity in each nucleus was measured on the second channel, red channel for FK2, HP1γ, and H3K9Me3. Data are presented as fluorescence intensity per cell or percentage of positive cells. For HP1γ and H3K9Me3, cells with a pixel intensity >50 (on a 1–255 scale) were considered positive. For counting of γH2AX foci, foci in the green channel were defined as particles >0.2 µm² with intensity >80 (on a 1–255 scale), and images were counted blind. Cells were isolated from 3 mice per genotype at 17 or 44 wk of age. Sorted cells from individual animals were pooled and deposited on cytospin for IF. Two independent experiments were performed and 200–500 cells were counted per genotype per experiment. For counting γH2AX/53BP1 foci, or FK2/53BP1 foci, in vi tro expanded LT-HSCs, foci were defined in each channel as above and foci positive for both signals were scored as colocalizing. For quantification of immunohistochemistry slides, cells with pixel intensity >50 (for HP1γ) or 45 (for H3K9Me3, on a 1–255 scale) were considered positive. Data were analyzed using Excel (Microsoft) and Prism 6 (GraphPad Software).

Histopathological analysis and immunohistochemistry. For histopathological analysis, tissues and organs were collected and fixed in formalin (or ethanol-acetic acid-formalin [EAF]) for 48 h (or 24 h in EAF). Fixed tissues and organs were subsequently dehydrated, embedded in paraffin, and 2-µm sections were prepared and stained with hematoxylin and eosin (H&E) for routine examination. For immunohistochemistry, 4-µm-thick sections were prepared and stained for activated Caspase 3, H3K9Me3, or HP1γ was performed according to the manufacturers’ recommendations. Secondary antibodies were coupled to Strep–AB complex, and immunodetection was performed using Liquid DAB Substrate Chromogen System (Dako). For H3K9Me3, slides were blocked with Powervision block (DPCB blocking) and binding was detected using the Powervision peroxidase kit (DVPVR110 HRP). Counterstaining was performed with hematoxylin.

MEF isolation and retroviral transduction. MEFs were cultured in 3% oxygen in DMEM (Life Technologies) supplemented with 10% FBS (Sigma-Aldrich) and 1% penicillin/streptomycin (Life Technologies). MEFs were isolated from embryonic day 14.5 embryos. In brief, fetal tissue was rinsed in PBS, minced, rinsed twice in PBS, and incubated with trypsin/EDTA (Life Technologies) for 10–15 min at 37°C and subsequently thoroughly dissociated in medium. After removal of large tissue clumps, the remaining cells were plated out in a 75-cm² flask. After 48 h, confluent cultures were frozen down. Retrovirus carrying empty vector or short-hairpin (sh) RNA directed against p53 (pRetroSUPER sh-p53; Drac and Bernards, 2003), pBABE-Puro empty vector, or expressing WT-USP3 was used to infect MEFs overnight. The next day, infected MEFs were split in medium containing puromycin to select the infected population. Efficient p53 knockdown was verified by Western blotting.

Chromosome analysis and SCE. Isolated MEFs were grown in 3% O₂. Analysis of chromosomal aberrations and SCE was performed on Giemsa-stained metaphase spreads as previously described (Nejenhuis et al., 2010). Images of metaphase spreads were captured using a microscope (Axiovert S100; Carl Zeiss) equipped with a camera (AxioCam Color; Carl Zeiss) and AxioVision software. For SCE analysis, 3 × 10⁵ cells were plated in 10-cm dishes. After 24 h, 10 µM BrdU was added, incubated 40 h, the last 4 h of which in the presence of 0.2 µg/mL Colcemid. Metaphase spreads were prepared as above with the following exceptions. Metaphase spreads were stained with Hoechst 5 µg/ml for 12 min, rinsed with water, and washed with water
for 5 min. Next, the slides were covered with water and coverslips and exposed to UV light (365 nm) for 1 h. Coverslips were removed and slides were rinsed using 2× SSC (Sodium Citrate buffer; 300 mM NaCl and 30 mM Tri-sodium Citrate) at 65°C 15 min. Finally, coverslips were washed with water 5 min and stained with Giemsa. Imaging and analysis was as described above.

**Constant field gel electrophoresis, comet assay, and β-galactosidase activity assay.** DSBs in primary MEFs were measured by Constant field gel electrophoresis (CFGE) as previously described (Neijenhuis et al., 2010). Alkaline comet assay was performed on total BM or LSK cells freshly sorted from 10-mo-old mice using the Comet assay kit ( Trevigen). Images were taken using a microscope (AxioObserver Z1; Carl Zeiss) equipped with a CCD camera and AxioVision software; comets were scored using Comet-Score software acquiring comet metrics as described (CometScore Tutorial) and analyzed using Excel. Senescence-associated β-galactosidase activity was assessed in cytosin preparation of freshly isolated BM cells from 32-wk-old littermates or sorted and in vitro cultured LT-HSCs using the Senescence-associated β-galactosidase staining kit (Cell Signaling Technology). Phae contrast pictures were captured using a microscope (Axiovert S100; Carl Zeiss) equipped with a camera (AxioCam Color) and AxioVision software.

**Pre-B and CFU-C colony-forming assay.** BM MNCs were seeded on 35-mm culture dishes in triplicate in methylcellulose medium supplemented with cytokines: MethoCult GF M3434 (for CFU-C, 5 × 10⁴ cells or 2 × 10⁵ cells, as indicated) and MethoCult M3630 (for CFU–Pre-B, 5 × 10⁴ cells; STEMCELL Technologies). CFU-Cs (colony forming units in culture) include CFU-GEMM (granulocyte, erythroid, macrophage, megakaryocyte), multipotential progenitors and lineage-restricted progenitors of the erythroid (BFU-E, burst-forming unit–erythroid), and granulocyte, monocyte-macrophage (CFU-GM). Cultures were incubated at 37°C under 5% CO₂. Colonies were quantified at day 8.

**Statistics.** Statistical analysis was performed using an unpaired two-tailed Student’s t test (Prism 6.0). Animal survival experiments were analyzed with a Log-rank nonparametric test and expressed as Kaplan-Meier survival curves.

**Online supplemental material.** Fig. S1 shows representative FACS profiles for hematopoietic stem and progenitor subpopulations (relative to Fig. 6 and Fig. 9 E). Table S1 shows neoplastic leukaemias (relative to Fig. 6 and Fig. 9 E). Table S2 lists genes validated by RNF168 ubiquitin ligase. Table S3 shows the DNA-damage-induced H2A Lys 15 ubiquitin mark. Table S4 lists genes validated by RNF8 transduces the DNA-damage signal via histone ubiquitylation of programmed DNA lesions and genomic instability in the immune system. Cell 152:417–429. http://dx.doi.org/10.1016/j.cell.2013.01.007

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