BNIP3-dependent mitophagy safeguards ESC genomic integrity via preventing oxidative stress-induced DNA damage and protecting homologous recombination

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Embryonic stem cells (ESCs) have a significantly lower mutation load compared to somatic cells, but the mechanisms that guard genomic integrity in ESCs remain largely unknown. Here we show that BNIP3-dependent mitophagy protects genomic integrity in mouse ESCs. Deletion of Bnip3 increases cellular reactive oxygen species (ROS) and decreases ATP generation. Increased ROS in Bnip3−/− ESCs compromised self-renewal and were partially rescued by either NAC treatment or p53 depletion. The decreased cellular ATP in Bnip3−/− ESCs induced AMPK activation and deteriorated homologous recombination, leading to elevated mutation load during long-term propagation. Whereas activation of AMPK in X-ray-treated Bnip3+/− ESCs dramatically ascended mutation rates, inactivation of AMPK in Bnip3−/− ESCs under X-ray stress remarkably decreased the mutation load. In addition, enhancement of BNIP3-dependent mitophagy during reprogramming markedly decreased mutation accumulation in established iPSCs. In conclusion, we demonstrated a novel pathway in which BNIP3-dependent mitophagy safeguards ESC genomic stability, and that could potentially be targeted to improve pluripotent stem cell genomic integrity for regenerative medicine.

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INTRODUCTION

Pluripotent stem cells (PSCs), including embryonic stem cells (ESCs) and induced pluripotent stem cells (iPSCs), can undergo unlimited self-renewal and maintain potency to differentiate into every cell type of an organism [1–6]. PSCs can be cultured and propagated indefinitely in vitro with a high proliferation rate, thus potentially providing an unlimited renewable source for various functional cells to serve regenerative medicine. Existing studies have shown that intrinsic characters like DNA-replications stress and environmental culture conditions both endorse PSCs easy to acquire genetic variations, which pose a serious risk for genomic instability [7–11]. Acquired DNA lesions are detrimental to PSC maintenance and function, and also raise safety concerns about the development of PSC-based cellular therapies.

In nature, ESCs, the equivalent to the inner cell mass of developing blastocysts, have a significantly lower mutation load and a higher proliferation rate with a short G1 phase compared to their differentiated progeny cells, suggesting that ESCs have an effective DNA damage response (DDR) to control their pronounced propensity to acquire DNA defects [12–20]. ESCs generate lower amounts of the genotoxic byproducts reactive oxygen species (ROS) and have increased expression of antioxidants compared to somatic cells [19, 21, 22]. In addition, DNA repair efficiency of ESCs in response to DNA damage is enhanced to protect their genomic stability relative to somatic cells. Somatic cells use error-prone non-homologous end joining (NHEJ) to repair most DNA double strand breaks (DSBs), whereas in PSCs error-free homologous recombination (HR) predominates and allows repair of DSBs with increased fidelity [23]. Compared to somatic cells, ESCs have higher levels of ATM/ATR-dependent phosphorylation of H2AX (γH2AX) and chromatin loading of single-stranded DNA (ssDNA)-binding proteins PRA32 and RAD51, which are both important for HR at DNA DSBs and stalled forks, indicating that ESCs could maintain a sensitized DDR state to mitigate DNA damage risks [24–26]. When DNA damage is not repaired, ESCs avoid the propagation of genetic lesions by undergoing either apoptosis or differentiation [18, 27–32]. Despite these multiple reports showing that PSCs have more efficient DNA repair capacity than somatic cells, the mechanisms responsible for these processes are poorly defined.

Mitochondria in human ESCs maintain a high mitochondrial priming state that sensitizes hESCs to DNA-damage-induced apoptosis, and favors the elimination of cells with DNA damage from self-renewing cell pools [27]. This finding indicates that mitochondria are involved in the maintenance of ESC genomic integrity. Mitochondria are highly dynamic, double-membrane
organelles that maintain homeostasis through autophagy [33]. In mouse ESCs, dysfunction of autophagy by depletion of ATG3 or ATG5 results in accumulation of abnormal mitochondria, leading to compromised self-renewal, pluripotency, and differentiation [34, 35]. Bcl-2 nineteen-kilodalton interacting protein (BNIP3) is an atypical member of the pro-apoptotic Bcl-2 subfamily that contains a Bcl-2 homology domain (BH3) and transmembrane domain (TM) that targets BNIP3 to mitochondria, a PEST domain that controls BNIP3 degradation, and a conserved domain [36–38]. In contrast to typical BH3-only proteins that bind to anti-apoptotic Bcl-2 family proteins via their BH3 domain, BNIP3 interacts with Bcl-2 and Bcl-XL specifically through its TM domain and N-terminus in cancer and non-cancer somatic cells [39]. The BNIP3 TM domain mediates the formation of homodimers that induce pro-survival mitophagy or cell death resembling both necrosis and apoptosis [39-43], but the interplay between BNIP3-induced apoptosis and survival functions remains obscure. Recently, we found that BNIP3-dependent mitophagy maintains mitochondrial homeostasis and pluripotency in mouse ESC [44]. However, how does BNIP3-dependent mitophagy regulate pluripotency remains obscure.

Here we uncovered that BNIP3-mediated mitophagy maintains ESC identity by preventing ESCs from oxidative stress-induced DNA damage and the resultant p53-dependent differentiation, while also maintains ESC genomic stability via protecting DNA repair by homologous recombination.

**MATERIALS AND METHODS**

**Animals**

GFP-LC3 mice (RBRC00806) [45] and Mito-RFP mice (RBRC03743) [46] were purchased from the Riken BioResource Centre. Dr. Zhuhua Zhang provided Bnip3 heterozygous mice. All protocols used for animal manipulation were approved by the Institutional Animal Care Committee.

**Reagents**

MitoTracker Red (40743ES50) was purchased from Yeasen. DCFH-DA (D6883) was purchased from Sigma Aldrich. AICAR (M9281) and Baf-A1(B1793) were obtained from Sigma Aldrich. 3-MA (D6883) was purchased from Sigma Aldrich. AICAR (S1802), Compound C (S7306), and Oligomycin A (S1478) were purchased from Selleck. 3-MA (D6883) was purchased from Sigma Aldrich. AICAR (S1802), Compound C (S7306), and Oligomycin A (S1478) were purchased from Selleck. 3-MA (D6883) was purchased from Sigma Aldrich. AICAR (S1802), Compound C (S7306), and Oligomycin A (S1478) were purchased from Selleck. 3-MA (D6883) was purchased from Sigma Aldrich. AICAR (S1802), Compound C (S7306), and Oligomycin A (S1478) were purchased from Selleck.

**Antibodies**

The following antibodies were used: anti-Bnip3 polyclonal antibody (3769, Cell Signaling Technology), anti-Actin monoclonal antibody (A5441, Sigma Aldrich), anti-LC3B antibody (PM036, Medical and Biological Laboratories, Co.), anti-Sox2 monoclonal antibody (ab29494, Abcam, for western blotting), anti-Sox2 polyclonal antibody (AB908, R&D, for IF), anti-Oct4 polyclonal antibody (ab19857, Abcam), anti-Nanog polyclonal antibody (ab106464, Abcam), anti-H2AX polyclonal antibody (CST7918, Cell Signaling Technology), anti-s3BP1 polyclonal antibody (NB100-304, Bio Techne), anti-p-ATM (Ser1891) monoclonal antibody (CST13050, Cell Signaling Technology). anti-ATM monoclonal antibody (CST2873, Cell Signaling Technology), anti-p-p53(Ser15) monoclonal antibody (CST9284, Cell Signaling Technology), anti-p-p53 monoclonal antibody (CST2537, Cell Signaling Technology), anti-Rad51 monoclonal antibody (H00009888, Abnova), anti-p-AMPAK (Thr172) (CST2531, Cell Signaling Technology), anti-p-PK (CST2532, Cell Signaling Technology), anti-SSA-E monoclonal antibody (SC-21702A647, Santa Cruz Biotechnology), Alexa Fluor 488 donkey anti-rabbit IgG (A21206, In vitrogen thermo Fisher Scientific), Alexa Fluor 594 donkey anti-mouse IgG (A11034, In vitrogen thermo Fisher Scientific). Alexa Fluor 488 donkey anti-goat IgG (A32104, In vitrogen thermo Fisher Scientific).

**Plasmids**

pMXs-Oct4, pMXs-Sox2, pMXs-Klf4, and pMXs-cMyc were purchased from Addgene (13366, 13367, 13370, 13375; Deposited by the Shinya Yamanaka lab). Mito-Keima vectors were shared by Dr. Quan Chen. Bnip3 and its mutants were cloned into pMXs and pCDH-CAG-RFP lentivectors as described previously [47]. ShRNA were designed and cloned into pSicor-GFP or pSicor vectors. The NHEJ vector, HR vector, and I-SceI-expressing plasmid were shared by Dr. Lingyi Chen.

**ESC isolation and iPSC induction**

Bnip3 knockout and wild-type ESCs were isolated at E3.5 and seeded on feeder layers for selection using 2i medium. Then ESCs were cultured for 3 to 5 passages and maintained in ESC medium (knockout DMEM with 15% fetal bovine serum, 2 mM glutamine, 1 mM sodium pyruvate, 0.1 mM nonessential amino acids, 100 mg/ml streptomycin, and 100 U/ml penicillin, 0.055 mM β-mercaptoethanol, and 1000 U/ml leukemia inhibitory factor). For iPSC induction, MEFs were seeded at 50,000 cells/well in a 6-well plate and infected with a retrovirus cocktail expressing Oct4, Sox2, Klf4, and Myc; iPSC colonies were picked 14 d after infection as described previously [47].

**Generation of Bnip3 & ATM knockout and Bnip3 & p53 knockout ESCs**

ATM and Bnip3 double knockout and p53 and Bnip3 double knockout ESCs were generated using a CRISPR-Cas9 system. ATM and p53 gRNA were cloned into the px330 vector, and transfected into Bnip3 knockout ESCs with a Gene pulser Xcell II (Bio-Rad) according to the manufacturer's protocol. Then, knockout ESCs were identified by digestion with PstI for ATM and Ncol for p53 and sequencing. All knockout ESCs were further identified by western blot.

**Western blot**

Cells were lysed in RIPA buffer (50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 0.5% sodium deoxycholate, 1% Nonidet P-40, 2 mM EDTA, 10 mM NaF) with protease inhibitor cocktail (04693116001, Roche). Equivalent protein quantities (20 µg) were subjected to SDS-PAGE and transferred to nitrocellulose membranes (Millipore), which were probed with the indicated primary antibodies followed by the appropriate HRP-conjugated secondary antibodies (Santa Cruz). Immunoreactive bands were detected with a Luminata Forte Western HRP Substrate Kit (WBLUF0100, Millipore).

**Immunofluorescence microscopy**

For LC3-GFP and Mito-RFP immunostaining, we used GFP-LC3 and Mito-RFP MEFs. For Bnip3 immunostaining, cells were cultured on gelatin-coated glass slides and fixed with 4% PFA for 20 min. They were then washed with DPBS, permeabilized with 0.2% Triton-X100 for 0.5 h and blocked with 2% BSA for 1 h before staining with the appropriate primary antibody overnight at 4 °C and incubation with secondary antibody for 2 h at 37 °C. Cell nuclei were then stained with DAPI. For Mito-Keima immunostaining, ESCs were transfected with Mito-Keima and directly monitored by confocal microscopy.

**Measurement of HR**

HR measurement was carried out as described [48]. We transfected the vector into Bnip3+/− ESCs, Bnip3−/− ESCs and rescued ESCs, then selected stably expressing clones with puromycin. To induce DSB, we transfected 1 µg I-SceI-expressing plasmid into HR reporter ESC lines. ESCs were harvested for FACS analysis 48 h after infection.

**Mito-Keima assay**

Mito-Keima was used to monitor mitophagy as described [49, 50]. Mito-Keima was transfected into Bnip3+/− ESCs, Bnip3−/− ESCs, and rescued ESCs. Stably expressing clones were sorted by FACS. The bnip3-rescued vectors were transfected into the above cell lines and single clones were picked for western blot identification. ESCs were cultured under normal conditions or treated with FCCP for 3 h, and then we detected the above cell lines by confocal laser scanning microscopy for Mito-Keima imaging [51].

**Transmission electron microscopy**

Cells were fixed in 2.5% glutaraldehyde for 2 h at 4 °C and immersed in 2% osmium tetroxide. The samples were then sequentially dehydrated with 50, 70, 90, 95, and 100% ethanol. Copper grids were used for ultrathin section collection and uranyl acetate and lead citrate were used for counterstaining. Images were taken using a FEI Tecnai Spirit transmission electron microscope.
Inhibition of autophagy increases DNA damage in mouse ESCs.

**Measurement of ATP and ROS**

A CellTiter-Glo Luminescent Cell Viability Assay kit (Promega Corporation, 0000092970) was used to assess cellular ATP content. Cellular ROS was measured by flow cytometry using HDCF-DA from Sigma Aldrich (D6883).

**Measurement of OCR and ECAR**

Seahorse XF24 analyzer (Seahorse Bioscience Asia, Shanghai) was used for detecting OCR and ECAR in ESCs or iPSCs. Briefly, ESCs or iPSCs were seeded at 60,000 cells/well for 6 h, and measurements were made strictly according to the standard protocol in the manual. ATP contribution was calculated under the manual.

**AP staining and clone formation assay**

For AP staining, a BCIP/NBT Alkaline Phosphatase Colour Development Kit (Beyotime) was used according to the manufacturer’s instructions. The clone formation assay was used as described. Briefly, ESCs were trypsinized into single cells that were seeded into 6-well plates (coated with feeder) at 1000 cells/well and cultured for one week. The number of AP-positive cells was counted after AP staining.

**RESULTS**

Inhibition of autophagy increases DNA damage response in mouse ESCs

Previous study reported that loss of autophagy caused a synthetic lethal deficiency in DNA repair in mouse embryo fibroblasts (MEFs) via a checkpoint kinase 1 (Chk1) dependent manner [52]. ESCs failed to activate the ATR-Chk1 signaling axis under alkylation-induced DNA damage [53]. To test whether autophagy contributes to genomic stability regulation in ESCs, we inhibited autophagy by 3MA or Baf1 treatment in ESCs. Treatment of ESCs with either 3MA or Baf1 resulted in increased ROS production and decreased ATP generation (Fig. 1A, B). Correspondingly, DNA damage response in ESCs was dramatically enhanced as indicated by the elevation levels of 53BP1 and γH2AX upon 3MA or Baf1 treatment (Fig. 1C, D). Importantly, the enhanced DNA damage response in ΔLIR Bnip3 ESCs could be rescued by gain expression of wide-type but not LC3-binding dead mutant V8D Atg3. This result supports the ATG3-mediated autophagy contributes to DNA damage response in ESCs.

BNIP3-dependent mitophagy safeguards ESC identity by preventing dysfunctional mitochondrion accumulation-induced oxidative stress and corresponding p53-mediated differentiation

Recently we have uncovered BNIP3-mediated mitophagy maintains mouse ESC pluripotency by regulating mitochondrial homeostasis [44]. Knockout of Bnip3 leads to enhanced ROS generation, decreased ATP production and impaired mitochondrial cristae without affecting cellular viability and apoptosis (Supplementary Fig. 1). Since Bnip3 knockout is associated with significant increases in ROS generation, and excessive intracellular ROS levels can cause DNA damage, we next asked whether Bnip3 depletion resulted in increased amounts of ROS-mediated DNA damage. In bnip3−/− ESCs, γH2AX and 53BP1 expression, as well as levels of the oxidative stress-induced DNA damage indicator 8-OHdG, were significantly increased compared to wild-type ESCs (Fig. 2A, B and Supplementary Fig. 2A). These defects could be partially restored by gain expression of wild-type but not ΔLIR Bnip3 or addition of the antioxidant NAC (Fig. 2A–E and Supplementary Fig. 2B). Together, these data indicated that increased ROS generation induced by dysfunction of BNIP3-dependent mitophagy in ESCs directly caused genomic DNA damage.

We then assessed whether deterioration of self-renewal and pluripotency in bnip3−/− ESCs was caused by ROS accumulation that occurred with Bnip3 depletion. As expected, the decreased expression of pluripotency genes and compromised colony...
BNIP3-dependent mitophagy protects ESC from oxidative stress-induced DNA damage. A Western blot of γH2AX and 53BP1 in bnip3−/− ESCs and rescued ESCs with actin as the loading control. B Increased cellular 8-OHdG in bnip3−/− ESCs was rescued by gain of expression of wild-type but not ΔLIR Bnip3. Data shown are the mean ± SD, n = 3; *, P < 0.05; **, P < 0.01; Student’s t-test. C NAC treatment (100 μM for 12 h) suppressed elevated oxidative stress in bnip3−/− ESCs. Data shown are the mean ± SD, n = 3; *, P < 0.05; **, P < 0.01; NS no significant difference, Student’s t-test. D NAC treatment antagonized increased DNA damage in bnip3−/− ESCs. Data shown are the mean ± SD, n = 3; *, P < 0.05; NS no significant difference, Student’s t-test. E Compromised pluripotency gene expression and enhanced DNA damage in bnip3−/− ESCs can be rescued by NAC treatment. Actin served as the loading control. F Deteriorated self-renewal of bnip3−/− ESCs can be restored by NAC treatment. Data shown are the mean ± SD, n = 3; *, P < 0.05; **, P < 0.01; NS no significant difference, Student’s t-test. G Activation of ATM and p53 in bnip3−/− ESCs can be rescued by wild-type but not ΔLIR Bnip3. Actin served as the loading control. H Decreased expression of pluripotency genes in bnip3−/− ESCs can be rescued by knockout of either ATM or p53. Data shown are the mean ± SD, n = 3; P < 0.01; Student’s t-test.

formation of bnip3−/− ESCs were partially rescued by NAC treatment (Fig. 2E, F), supporting that the deterioration of bnip3−/− ESCs results from dysfunctional BNIP3-dependent mitophagy induced oxidative stress.

To dissect the interplay between enhanced genomic DNA damage by Bnip3 depletion and ESC identity, we examined downstream events of ROS accumulation, particularly the DNA damage response (DDR) in bnip3−/− ESCs. We found that increased phosphorylation of ATM (Ser1981) and p53 (Ser15) in bnip3−/− ESCs, which can be rescued by gain expression of wild-type but not ΔLIR Bnip3, could be restored by NAC treatment, indicating that intracellular ROS accumulation following Bnip3 knockout contributed to activation of ATM and p53 (Fig. 2G and Supplementary Fig. 2C, D). Correspondingly, both colony formation and expression of pluripotency marker genes in bnip3−/− ESCs showed significant defects, and colonies of bnip3−/− ESCs exhibited differentiation phenotype (Fig. 2H, I and Supplementary Fig. 4A–C). Together, these data suggested that BNIP3-mediated mitophagy maintains ESC identity by preventing dysfunctional mitochondrion accumulation caused oxidative stress and subsequent p53-dependent differentiation.

BNIP3-dependent mitophagy protects genomic integrity by safeguarding homologous recombination

In addition to compromised self-renewal and pluripotency associated with DDR, Bnip3-knockout ESCs form colonies with lower efficiency, providing an ideal model to evaluate long-term effects of excessive ROS on ESC genomic stability. We next investigated whether increased levels of intracellular ROS in bnip3−/− ESCs would significantly impair genomic integrity during long-term propagation. We propagated Bnip3+/+, vector-bnip3−/−, ΔLIR-bnip3−/−, and WT-bnip3−/− ESCs routinely for 30 passages either under normal conditions or with NAC or H2O2 treatment. Then we performed whole genomic sequencing on each cell line at either initial passage or after propagation for 30 passages. Compared to Bnip3+/+ ESCs, the bnip3−/− ESC genome accumulated more mutations during long-term propagation (Fig. 3A). Mitochondrial genome of Bnip3+/+ ESCs showed similar mutation rates with that of bnip3−/− ESCs. This enhanced accumulation of mutations in
Excessive AMPK activation decreases homologous recombination

Adenosine monophosphate (AMP)-activated protein kinase (AMPK) is a cellular energy sensor that is regulated by both cellular adenine nucleotides and glucose [54-56]. As shown above, Bnip3 depletion resulted in dysfunctional mitochondrial accumulation and significantly decreased cellular ATP generation by oxidative phosphorylation (Supplementary Fig. 5A) [44]. Although mouse ESCs had higher expression levels of AMPK and enhanced AMPK phosphorylation than somatic fibroblasts constitutively [57], the lower rate of ATP generation in Bnip3−/− ESCs further enhanced AMPK phosphorylation without affecting AMPK expression, which was rescued by gain of expression of wild-type but not ΔLIR-Bnip3 [Fig. 3D and Supplementary Fig. 5B]. We next asked whether excessive AMPK activation in Bnip3−/− ESCs is responsible for the defective HR and associated increased in DNA mutation accumulation. Inhibition of excessive AMPK activation with Compound C in Bnip3−/− ESCs partially restored the compromised HR activity without affecting DNA damage (Fig. 3E, F and Supplementary Fig. 5C). In contrast, treatment of Bnip3−/− ESCs with the AMPK agonist AICAR or the mitochondrial

bnip3−/− ESCs was partially rescued by wild-type but not ΔLIR-Bnip3, indicating that Bnip3-dependent mitochondrial protects genomic stability during long-term propagation of ESCs (Fig. 3A).

As expected, H2O2 treatment during long-term propagation further increased genomic mutation load in Bnip3+/−, vector-bnip3−/−, ΔLIR-bnip3+/−, and WT-bnip3−/− ESCs. Unexpectedly, however, NAC treatment of vector-bnip3−/− or ΔLIR-bnip3−/− ESCs during long-term propagation did not mitigate the elevated genomic mutation load (Fig. 3A). These data suggest that increased ROS levels are likely not the direct cause of enhanced genomic mutation load observed in bnip3−/− ESCs during long-term propagation. To dissect this observation further, we assessed DNA repair by homologous recombination (HR) in Bnip3+/−, vector-bnip3−/−, ΔLIR-bnip3+/−, and WT-bnip3−/− ESCs. Surprisingly, we found that depletion of Bnip3 significantly decreased the HR rate (Fig. 3B). These defects were also observed in developing bnip3−/− blastocysts (Fig. 3C). The compromised HR in bnip3−/− ESCs was partially compensated by gain of expression of wild-type, but not the LIR-binding deficient ΔLIR-Bnip3 (Fig. 3B). These data indicate that Bnip3-dependent mitophagy protects mouse ESC genomic stability by safeguarding HR.
Bnip3 overexpression improves iPSC genomic integrity. A Overexpression of Bnip3 does not affect reprogramming efficiency. Data shown are the mean ± SD, n = 3; NS no significant difference. B Bnip3 overexpression enhanced mitochondrial autophagy in reprogramming cells. GFP: LC3; RFP: Mitochondria. C Confocal fluorescence micrographs of reprogramming cells. SSEA-1-positive cells at reprogramming day 3 were sorted for staining with γH2AX and RAD51. Green, γH2AX; Red, RAD51; Blue, DAPI. D Statistical data for γH2AX foci in (C). Cells were randomly divided into 3 groups and the foci were counted. Data shown are the mean ± SD, for RAD51 and Bnip3. E Statistical data for RAD51 and γH2AX colocalization foci/γH2AX foci in (C). F Confocal fluorescence micrographs of iPSCs. Green, γH2AX; Red, SOX2; Blue, DAPI. G Western blot detection of RAD51, γH2AX, p-AMPK, AMPK, and Bnip3 in wild-type or Bnip3-overexpressing iPSCs. Actin served as the loading control. H Quantifications of each protein expression in (G). Data are shown as mean ± SD, n = 3; ***, P < 0.001; NS no significant difference, Student’s t-test. I Acquired indels in indicated iPSC lines.

respiration inhibitor oligomycin compromised DNA repair by HR (Supplementary Fig. 5D). These data support that excessive activation of AMPK did result in impaired HR DNA repair in mouse ESCs.

We next tested whether excessive activation of AMPK could diminish ESC genomic integrity. Both Bnip3+/- and WT-bnip3−/− ESCs were treated with X-ray irradiation in the presence or absence of AICAR or oligomycin. Both the AMPK agonist and mitochondrial inhibitor significantly increased the genomic mutation load in Bnip3+/- ESCs, indicating that excessive AMPK activation caused by mitochondrial dysfunction in ESCs deteriorated genomic stability (Fig. 3G). In addition, we treated vector-bnip3−/− and ΔLIR-bnip3−/− ESCs with X-rays and the AMPK inhibitor Compound C, and found that inhibition of excessive AMPK activation in bnip3−/− ESCs dramatically decreased the genomic mutation load (Fig. 3G). At the same time, we evaluated the genome stability of Bnip3+/- and bnip3−/− ESCs after 30 passages in the presence or absence of Compound C by detecting the DNA damage maker and HR maker via western blot. We found the surviving bnip3−/− ESCs showed enhanced DNA damage with no DNA repair changes after 30 passages. Long-term culturing led to increased DNA damage in both Bnip3+/- and bnip3−/− ESCs. Compound C treatments significantly enhanced DNA repairs in both Bnip3+/- and bnip3−/− ESCs (Supplementary Fig. 5E, F). Together, these data suggested that enhanced accumulation of genomic mutations is due to excessive AMPK activation induced by deterioration of mitochondrial bioenergetics in bnip3−/− ESCs.

Enhancement of iPSC genomic integrity by increasing BNIP3-mediated mitophagy

Given that BNIP3 has a role in protecting genomic integrity of ESCs, we tested whether enhancement of BNIP3-dependent mitophagy during reprogramming improves genomic stability of iPSCs. Bnip3 overexpression significantly enhanced mitochondrial autophagy during reprogramming without affecting reprogramming efficiency (Fig. 4A, B). By sorting of SSEA-1 positive cells on reprogramming day 3, we found that Bnip3 overexpression significantly alleviated ROS production (Supplementary Fig. 6A, B). Correspondingly, DDR activation, as evidenced by the presence of γH2AX foci, was significantly decreased in cells overexpressing Bnip3 compared to WT reprogramming cells (Fig. 4C). Importantly, activation of DNA repair was dramatically increased in cells overexpressing Bnip3 compared with WT reprogramming cells (Fig. 4D, E). These data suggested that enhancement of BNIP3-dependent mitophagy by overexpression of Bnip3 during reprogramming lessened oxidative stress-induced DDR and increased DNA repair by HR. Thus, we would expect to see improved genetic integrity in the resulting iPSCs having Bnip3 overexpression. Correspondingly, the established iPSCs with Bnip3 overexpression showed decreased ROS levels and enhanced...
mitochondrial respiration (Supplementary Fig. 6C–G). Supportively, the iPSCs with Bnip3 overexpression exhibited decreased DNA damage and AMPK activation, while enhanced DNA repair (Fig.4F–H). Whole genomic sequencing on established iPSC lines carrying either empty vector or Bnip3 cDNA, and their parental fibroblasts, demonstrated that Bnip3 overexpression indeed dramatically reduced the mutation load in established iPSCs (Fig. 4I).

DISCUSSION

In this study, we revealed that BNIP3-mediated mitophagy maintains ESC identity by ensuring mitochondrial integrity that avoids excessive ROS generation and the corresponding p53-dependent ESC differentiation. Meanwhile, BNIP3 protects ESC genomic stability by safeguarding homologous recombination through maintenance of mitochondrial bioenergetics. In addition, reinforcement of BNIP3-dependent mitophagy during reprogramming is demonstrated to be an alternate strategy to improve iPSC genomic stability (Fig. 5).

Excessive levels of cellular ROS, the byproducts of mitochondrial oxidative phosphorylation, are detrimental to genomic DNA, proteins, and lipids [58]. Previous studies showed that ESC mitochondria are in an immature state characterized by a globular shape, blurred cristae, and limited respiration capability compared to differentiated somatic cells [59–64]. Somatic cells utilize oxidative phosphorylation, whereas ESCs are almost entirely glycolytic with low mitochondrial activity under aerobic conditions [22, 64]. In agreement with these mitochondrial characteristics, ESCs produce less ROS and have a higher antioxidant capacity compared to somatic cells [19, 65, 66]. Thus, it was proposed that lower cellular ROS serves as a defense mechanism for ESCs to maintain genomic integrity. In contrast, here we showed that increased cellular ROS levels that accompany Bnip3 depletion resulted in p53 activation that promoted ESC differentiation and corresponding compromises in self-renewal. Treatment with NAC or knockout of p53 in Bnip3−/− ESCs compensate for the compromised self-renewal, highlighting the relationships between Bnip3-knockout-induced ROS accumulation, p53 activation, and maintenance of ESC identity. Interestingly, a recent study showed that loss of Cops5, conserved COP9 signalosome subunit 5, resulted in increased ROS generation, p53 activation, G2/M arrest, and apoptosis rather than p53-dependent differentiation in mouse ESC [48]. The mechanisms for these differential effects of ROS on determination of ESC fate are unclear. Existing studies have proposed that signal transduction of ROS-induced p53 activation may depend on the extent of DNA damage, since higher levels of DNA double-stranded break (DSB) damage induce both p53-dependent and p53-independent apoptosis, whereas low levels of DNA damage suppress p53-dependent expression of critical pluripotency genes like Nanog and Oct4 [29].

Treatment of Bnip3+/+ ESCs with H2O2 further increased genomic mutation load during long-term propagation; however, NAC treatment did not decrease genomic mutation accumulation in the same timeframe (Fig. 3A). Intriguingly, treatment of Bnip3−/− ESCs with NAC had little impact on genomic mutation load during long-term propagation of these cells (Fig. 3A), although it could restore the defective self-renewal ability (Fig. 2F). These data suggest that BNIP3-dependent mitophagy does not directly rely on ROS homeostasis maintenance function to protect ESC genomic integrity. Instead, ESCs developed a fail-safe elimination mechanism to maintain genomic stability by inducing p53-dependent differentiation of cells having elevated ROS levels to remove them from self-renewing pools.

Genomic DNA can be damaged by exogenous stresses like UV light, ionizing radiation, and chemical compounds, as well as by...
endogenous stress like ROS, and DNA replication and transcription activities [67, 68]. Mammalian cells have developed sophisticated mechanisms to protect their genomic integrity. After DNA damage, cells quickly initiate DDR to induce cell-cycle arrest, apoptosis, or senescence [15]. In contrast to somatic cells, PSCs have a short G1 phase and no G1/S checkpoint, which possibly causes constitutive replication stresses like single-stranded DNA gap accumulation and corresponding constitutive activation of DDR as characterized by elevated phospho-H2AX [24, 69]. Earlier studies showed that PSCs preferentially employ HR over NHEJ, thus repairing DNA DSBs with enhanced fidelity [23]. At the same time, expression of some DNA repair machinery components has been found to be significantly increased in PSCs compared to somatic cells [17, 65]. In addition, ESC-specific factors like ZSCAN4, FILIA, COPSS, and SALL4 have been shown to regulate pluripotency and genomic integrity coordinately [48, 70–72]. In contrast, here we demonstrated a novel pathway in mouse ESCs that connects mitochondrial integrity and genomic stability through BNIP3-dependent mitophagy that maintains genomic stability by safeguarding DNA repair. Dysfunction of BNIP3-dependent mitophagy caused the accumulation of abnormal mitochondria and in turn, decreased ATP generation. The resulting decreased intracellular ATP levels of \( \text{BNip3}^{-/-} \) ESCs further activate AMPK, leading to compromised DNA repair by HR. Interestingly, we previously showed that AMPK is highly expressed in ESCs relative to somatic fibroblasts, and that phosphorylation of ULK1 by AMPK is essential for mESC self-renewal and maintenance of pluripotency [57]. These data together suggest that appropriate activation of AMPK is required for ESCs to maintain cellular homeostasis and identity, whereas over-activation of AMPK in ESCs is detrimental to genomic stability. Indeed, chemical activation of AMPK in X-ray-stressed wild-type ESCs significantly enhanced the genomic mutation load, while AMPK inactivation in X-ray-stressed \( \text{BNip3}^{-/-} \) ESCs ameliorated genomic DNA damage (Fig. 3G).

Reprogramming of somatic cells to iPSCs is an inefficient process. Suppressed expression of the tumor suppressor gene \( p53 \) significantly enhances reprogramming efficiency, due to decreased \( p53 \)-dependent DNA damage responses and induction of apoptosis [15, 73–77]. The potential need for \( p53 \) inactivation for successful reprogramming raises concerns about genomic stability of iPSCs and genomic structure variations have, in fact, been identified in multiple iPSC lines [9, 78, 79]. More than half of the somatic coding mutations present in 22 human iPS cell lines analyzed were found to be acquired during the reprogramming process, independently of the reprogramming method used [80]. These findings concerning genomic instability of iPSCs pose safety concerns for the clinical development of iPS technology [81]. In this regard, our results suggest an alternate strategy wherein reinforcement of BNIP3-dependent mitophagy during reprogramming could be used to improve iPSC genomic stability and facilitate clinical application of these cells.

**Data deposition**

Whole genome sequencing data of Vector-Bnip3\(^{+/+}\), Vector-bnip3\(^{-/-}\), \( \Delta \text{LIR-bnip3}^{-/-} \), and WT-bnip3\(^{-/-}\) ESC were uploaded to Science Data Bank. The CSTR of our genome sequencing data is 31253.11.scienedb.02743, and the DOI is 10.5776/scienedb.02743. The details of data are available from the corresponding author on reasonable request.

**DATA AVAILABILITY**

The data and materials during the current study are available from the corresponding author on reasonable request.
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AUTHOR CONTRIBUTIONS
QZ, KL, and TZ designed and performed the experiments, analyzed the data, and wrote the paper; QZ, KL, LZ, ZL, LW, and JC performed the experiments and analyzed the data; YX, AZ, QC, and TZ coordinated and supervised the experiments.

COMPETING INTERESTS
The authors declare no competing interests.

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All protocols used for animal manipulation were approved by the Institutional Animal Care Committee.

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