IP6K1 Reduces Mesenchymal Stem/Stromal Cell Fitness and Potentiates High Fat Diet-Induced Skeletal Involution

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Abstract

Mesenchymal stem/stromal cells (MSCs) are the predominant source of bone and adipose tissue in adult bone marrow and play a critical role in skeletal homeostasis. Age-induced changes in bone marrow favor adipogenesis over osteogenesis leading to skeletal involution and increased marrow adiposity so pathways that prevent MSC aging are potential therapeutic targets for treating age-related bone diseases. Here, we show that inositol hexakisphosphate kinase 1 (Ip6k1) deletion in mice increases MSC yields from marrow and enhances cell growth and survival ex vivo. In response to the appropriate stimuli, Ip6k1<sup>−/−</sup> versus Ip6k1<sup>+/+</sup> MSCs also exhibit enhanced osteogenesis and hematopoiesis-supporting activity and reduced adipogenic differentiation. Mechanistic-based studies revealed that Ip6k1<sup>−/−</sup> MSCs express higher MDM2 and lower p53 protein levels resulting in lower intrinsic mitochondrial reactive oxygen species (ROS) levels as compared to Ip6k1<sup>+/+</sup> MSCs, but both populations upregulate mitochondrial ROS to similar extents in response to oxygen-induced stress. Finally, we show that mice fed a high fat diet exhibit reduced trabecular bone volume, and that pharmacological inhibition of IP6K1 using a pan-IP6K inhibitor largely reversed this phenotype while increasing MSC yields from bone marrow. Together, these findings reveal an important role for IP6K1 in regulating MSC fitness and differentiation fate. Unlike therapeutic interventions that target peroxisome proliferator-activated receptor gamma and leptin receptor activity, which yield detrimental side effects including increased fracture risk and altered feeding behavior, respectively, inhibition of IP6K1 maintains insulin sensitivity and prevents obesity while preserving bone integrity. Therefore, IP6K1 inhibitors may represent more effective insulin sensitizers due to their bone sparing properties. Stem Cells 2017; 00:000–000

Significance Statement

This study is the first to identify a role for inositol hexakisphosphate kinase 1 (IP6K1) in regulating primary mouse mesenchymal stem/stromal cell (MSC) fitness. Our data show that genetic deletion of Ip6k1 results in increased MSC yields from bone marrow, and enhances the growth, survival, hematopoiesis-supporting activity, and osteogenic potential of culture-expanded cells. Increased fitness correlated with lower intrinsic p53 protein expression and mitochondrial reactive oxygen species levels, consistent with the known sensitivity of MSCs to oxidative stress. Furthermore, pharmacological inhibition of IP6K1 in mice reversed high fat diet (HFD)-induced decreases in bone volume. Together, these studies indicate that IP6K1 influences MSC frequency and function in bone marrow by modulating responsiveness to systemic changes induced by HFD feeding, and identify IP6K1 as a therapeutic target to prevent skeletal involution in response to these conditions.

Introduction

Mesenchymal stem cells (MSCs) are multipotent stem/progenitors first identified in bone marrow based on their ability to generate heterotopic osseous tissue with an architecture capable of supporting hematopoiesis when serially transplanted into mice [reviewed in Ref. 1]. Subsequently, MSCs have been shown to localize adjacent to blood vessels in vivo, which has led to their characterization as vascular pericytes or perivascular stromal cells [2]. Most recently, several groups have identified specific MSC subsets within bone marrow that comprise the hematopoietic stem cell niche and regulate hematopoietic stem cell self-renewal and differentiation [3–6]. Included among these...
latter is a population of leptin receptor (LEPR) expressing perivascular cells, which are the principle source of chemokines that regulate hematopoietic stem cell self-renewal and retention in bone marrow [7]. Importantly, LEPR-MSCs also represent the main source of osteoblasts and adipocytes that form bone and fat within adult bone marrow [8], and recent studies indicate that the LEPR acts as a systemic energy sensor in MSCs promoting adipogenesis and inhibiting osteogenesis in response to high fat diet (HFD) feeding [9]. Therefore, MSC fate determination in marrow is modulated by nutritional status as a consequence of LEP/LEPR signaling. However, LEP/LEPR signaling is also involved in satiety. Therefore, targeting LER/LEPR signaling to reverse HFD-induced trabecular bone loss and increased marrow adiposity is not feasible. Similarly, peroxisome proliferator-activated receptor gamma (PPARG) agonists like rosiglitazone used to treat type II diabetes inhibit insulin resistance but induce osteoporosis [10]. Therefore, a novel drug target that can reverse HFD-induced weight gain and insulin resistance while preserving bone integrity remains to be identified. Recent studies indicate that mice with a germline deletion of inositol hexakisphosphate kinase 1 (Ip6k1), the major isoform of three known inositol hexakisphosphate kinases [11], are protected against HFD-induced weight gain and insulin resistance [12–14]. A family of three mammalian IP6 kinases (IP6Ks) primarily generates the inositol pyrophosphate 5-IP7 from IP6 [15–20]. However, at a lower ATP/ADP ratio, IP6Ks dephosphorylate IP6 to a distinct form of IPS [21]. IP6 and 5-IP7 bind protein targets to modulate their functions [15–17]. In addition, inositol pyrophosphate-mediated pyrophosphorylation alters functions of target proteins [15, 16]. HFD-fed Ip6k1−/− mice maintain insulin sensitivity by sustaining activity of the insulin sensitizing protein kinase Akt [12]. Moreover, IP6K1 reduces adipose tissue browning mediated thermogenesis by inhibiting AMPK (AMP stimulated protein kinase) activity [13]. Accordingly, the pan IP6K inhibitor TNP [N2-(m-Trifluorobenzyl), N6-([28] with the following exceptions. Adipogenic induction media consisted of Reduced Serum Medium (1:5). MSCs (10,000 cells per square centimeter) were plated onto the transfection reaction, cultures were fed with fresh media 2 days later, and every 2–3 days thereafter for up to 7 days. Cultures were photographed using a Leica DMi3000 B upright fluorescent microscope attached to a DFC295 digital camera (Micro Optics of Florida, Inc., Davie, FL, www.microopticsfl.com).

Adipogenic and osteogenic potential of MSCs (1 × 10^4 cells per square centimeter) was quantified as described previously [28] with the following exceptions. Adipogenic induction media consisted of CCM supplemented with 0.5 μM dexamethasone, 0.5 mM isobutylxanthine, and 50 μM indomethacin, and lipid accumulation was quantified by staining with adipogenic RNAiMAX Transfection Reagent (Thermo Fisher Scientific). Briefly, transfection reagent was prepared by mixing Lipofectamine RNAiMAX (1:2,000) and the appropriate RNAi duplex (10nM, Thermo Fisher Scientific) in Opti-MEM™ Reduced Serum Medium (1:5). MSCs (10,000 cells per square centimeter) were plated onto the transfection reagent, cultures were fed with fresh media 2 days later, and every 2–3 days thereafter for up to 7 days. Cultures were photographed using a Leica DMi3000 B upright fluorescent microscope attached to a DFC295 digital camera (Micro Optics of Florida, Inc., Davie, FL, www.microopticsfl.com).

**Materials and Methods**

**MSC Isolation, Transfection, and Differentiation**

MSCs were isolated from Ip6k1−/− and congenic wild type mice via immunodepletion as described previously [27]. Briefly, whole bone marrow obtained from the long bones of mice was suspended in complete culture media (CCM; α minimal essential media containing l-glutamine and supplemented with 10% fetal bovine serum, 100 U/ml penicillin, and 100 U/ml streptomycin), plated at a density of 1.4 × 10^6 cells per square centimeter, and cultured for 72 hours after which the non-adherent cells were removed by aspiration. Adherent cells were cultured an additional 5–7 days with a single media change and then harvested by gentle scraping after a 5-minutes incubation in 0.25% trypsin with 1 mM EDTA. The cell pellet was dispersed by gentle agitation, suspended in 20 ml of Hank’s balanced salt solution (HBBS), and filtered through a 70-μm filter. Cells were suspended in HBBS at a maximum density of 40 × 10^6 cells per milliliter and incubated on a rotator for 1 hour at 4°C, then successively for 50 minutes intervals with streptavidin-conjugated Dynabeads (five beads per cell) (ThermoFisher Scientific, Waltham, MA, www.thermofisher.com) conjugated to anti-CD11b, anti-CD34, and anti-CD45 antibodies (10 μg/mg beads) (BD Biosciences, San Jose, CA, www.bdbiosciences.com). All cell isolation and culture manipulations were performed in a modular airtight chamber (BioSpherix Ltd., Lacona, NY, www.biospherix.com/) flushed with 5% O_2 balanced with N_2. MSCs were cultured at 37°C in a humidified chamber with 5% CO_2 and 5% O_2 unless stated otherwise. The total yield of MSCs per mouse was calculated based on the total number of MSCs obtained following immunodepletion. Population doubling (PD) times were calculated as PD = t log2/(log NT – log No) where t is time period, NT is the number of cells at time t, and No is the initial number of cells plated. Cumulative cell numbers and total PDs for each passage were determined from initial plating densities post-immuno-depletion. Delivery of siRNA into MSCs was accomplished using the Lipofectamine RNAiMAX Transfection Reagent (Thermo Fisher Scientific). Briefly, transfection reagent was prepared by mixing Lipofectamine RNAiMAX (1:2,000) and the appropriate RNAi duplex (10nM, Thermo Fisher Scientific) in Opti-MEM™ Reduced Serum Medium (1:5). MSCs (10,000 cells per square centimeter) were plated onto the transfection reaction, cultures were fed with fresh media 2 days later, and every 2–3 days thereafter for up to 7 days. Cultures were photographed using a Leica DMi3000 B upright fluorescent microscope attached to a DFC295 digital camera (Micro Optics of Florida, Inc., Davie, FL, www.microopticsfl.com).

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**Colony Assays**

To quantify CFU-Fs, MSCs (10,000 cells) were plated in 100 mm dishes and were cultured for 7–10 days with media changes every 2–3 days. On day 7, the monolayers were washed with phosphate-buffered saline, stained with 0.5% crystal violet (Sigma-Aldrich, St. Louis, MO, www.sigmaloadrich.com) in 25% methanol for 10 minutes at room temperature, and colonies ≥ 0.01 square centimeter in area were counted and their total area quantified using ImageJ software. To perform standard colony assays for hematopoiesis, conditioned media (serum
free) was collected over a 24-hour period from confluent monolayers of wild type and \(^{l p 6 k 1^{-/-}}\) MSCs, filtered, then used to suspend whole bone marrow cells isolated from wild type mice. The cells were then mixed with growth factor supplemented MethoCult media (StemCell Technologies, Vancouver, BC, www.stemcell.com) and plated at 60 mm dishes at 100,000 cells per dish. The plates were incubated 37°C in a humidified chamber in 5% \(\text{CO}_2\) and 5% \(\text{O}_2\) balanced with \(\text{N}_2\) for 8–10 days after which colonies were enumerated based on morphology.

**Flow Cytometry**

MSCs (2.5 × 10\(^5\) cells per milliliter) were suspended in HBSS and incubated with antibodies against CD11b, CD19, CD29, CD31, CD34, CD44, CD45, CD73, CD105, CD106, CD146, CD166, SCA1, F4/80, SSEA-1, and SSEA-4 (R&D Systems Inc., Minneapolis, MN, www.rndsystems.com) for 30 minutes at 4°C. Cells were washed twice with HBSS, incubated with the appropriate fluorochrome-conjugated secondary antibody to 30 minutes at 4°C, and analyzed using an Epics FC 500 FACS scanner equipped with CXP software (Beckman Coulter, Inc., Indianapolis, IN, www.beckmancoulter.com). Alternatively, MSCs were stained with MitoSOX Red (Thermo Fisher Scientific) to quantify mitochondrial superoxide production. For microscopic analysis, 10 \(\mu\)M MitoSOX Red was used while 100 nM was used for fluorescent quantification. To confirm mitochondrial localization of the dye, cells were counterstained with 5 \(\mu\)M Mito-Tracker Green (Thermo Fisher Scientific) for 10 minutes. Normalization of fluorescent data was carried out using counterstaining with Hoechst 33342 (1 \(\mu\)M) for 5 minutes. Fluorescent wavelength pairs for the individual dyes were 510 nm/580 nm for MitoSOX Red, and 490 nm/516 nm for Mito-Tracker Green.

**RNA Sequencing**

Total RNA was quantified on a Qubit 2.0 Fluorimeter (Thermo Fisher Scientific) and run on an Agilent 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA, www.agilent.com) for quality assessment. Samples with an RNA integrity number > 8.0 were further analyzed for further analysis. Briefly, mRNA was isolated from total RNA (100–200 ng input) using poly-dT oligos attached to magnetic beads, and the purified mRNA processed using the TrueSeq Stranded mRNA sample prep kit (Illumina, San Diego, CA, www.illumina.com). RNA samples were chemically fragmented in buffer containing divalent cations at 94°C for 8 minutes, primed with random hexamers, and reverse transcribed to generate the first strand cDNA. The second strand was synthesized after removing the RNA template and incorporating dUTP in place of dTTP. The ds cDNA was then repaired and adenylated at its 3′-ends. Adaptor-ligated DNA was purified using magnetic Ampure XP beads and polymerase chain reaction (PCR) amplified using 13 cycles to generate the final libraries, which were size selected and purified using 1.0× Ampure XP beads to remove any primer dimers. The final library size was typically 200–600 bp with insert sizes ranging from 80 to 450 bp. Libraries were validated using bioanalyzer DNA chips and quantitative PCR (qPCR) using primers that recognize the adaptors, pooled at equimolar ratios, loaded onto a NextSeq 500 flow cell at 1.8 pM final concentration, and sequenced using 2 × 75 bp paired-end chemistry. On average 18–20 million reads are generated (base quality score > Q30 suggesting less than 1 error in 1,000 bp). Data (fastq) generated from the NextSeq 500 were trimmed using Flexbar 2.4 and aligned to the Human reference database (Hg19) using TopHat version 2.0.9 [29]. HTseq-count version 0.6.1 was used to generate gene counts and differential gene expression analysis was performed using Deseq2 [30].

**Quantitative Polymerase Chain Reaction**

Total RNA was isolated using the RNaseasy Kit (Qiagen, Valencia, CA, www.qiagen.com), converted to cDNA using the Transcriptor First Strand cDNA Synthesis Kit (F. Hoffmann-La Roche, Basel, Switzerland, www.roche.com), and amplified by PCR using the TaqMan® EZ RT-PCR kit (Applied Biosystems, Carlsbad, CA, www.appliedbiosystems.com) according to the manufacturer’s instructions. Reactions were performed on a 7900 HT sequence detector (Applied Biosystems) using the FastStart Universal SYBR Green Master Mix (F. Hoffmann-La Roche) and transcript levels quantified using the relative Ct method with \(\text{Gapdh}\) as an internal control using the following primer pairs: \(lp6k2\), 5′-ggcacagtggaagctgttatag-3′ and 5′-ccagctcttgaggtctttc-3′; \(lp6k3\), 5′-cgctgtggaagataaagat-3′ and 5′-ctcctgtctctgtgagct-3′; and \(\text{Gapdh}\), 5′-tcacagacagcccccatcttca-3′ and 5′-acctgcttgtagctgccattca-3′.

**Western Blot**

Protein lysates were prepared using the Qproteome Mammalian Protein Prep Kit (Qiagen) and protein concentrations were determined using the Bradford assay (Bio-Rad, Hercules, CA, www.bio-rad.com). Protein samples (20 \(\mu\)g) were prepared in Laemmli sample buffer (Bio-Rad) containing 5% β-mercaptoethanol, denatured at 95°C for 5 minutes, electrophoresed on NuPAGE 10% Bis-Tris gels using 1× NuPAGE MES SDS Running Buffer (Thermo Fisher Scientific), and then transferred to 0.45 \(\mu\)m nitrocellulose membranes in 1× NuPAGE transfer buffer containing 10% methanol. Membranes were washed with Tris-buffered saline (TBS) for 5 minutes, incubated in TBS with 0.1% Tween-20 (TBST) and Odyssey® blocking buffer (LI-COR Biosciences, Lincoln, NE, www.li-cor.com) overnight at 4°C, washed an additional 3× in TBST, and then incubated with anti-\(\text{IP6K1}\) (1:500, GeneTex, Irvine, CA, www.genetex.com), anti-p53, anti-MDM2, anti-p16, and anti-GAPDH (1:200, Santa Cruz Biotechnology Inc., Dallas TX, www.scbt.com) antibodies in Odyssey® blocking buffer for 2 hours at room temperature with gentle agitation. Membranes were washed 5× in TBST and probed with an Infrared (IR) dye-labeled secondary antibody at a 1:10,000 dilution in Odyssey® blocking buffer for 1 hour. Blots were scanned using Odyssey® infrared image system (LI-COR Biosciences).

**Chemokine Quantification**

Confluent monolayer cultures of \(lp6k1^{-/-}\) and wild type congenic MSCs were maintained in serum-free media for 24 hours, the conditioned media collected, and concentrated 50-fold using Amicon Ultra-15 centrifugal filters (Millipore, Sigma, Billerica, MA, www.endmillipore.com). Media was then analyzed using the Mouse Pre-Mixed Multi-Analyte Kit (R&D Systems) for simultaneous detection of the indicated analytes. Quantification of analyte abundance was performed using a Lumineo 200 dual-laser, flow-based detection platform (Lumineo Corp., Austin, TX, www.lumineox.com). Alternatively, media was analyzed using the mouse SCF ELISA Kit per the manufacturer’s instructions (Thermo Fisher Scientific).
**Results**

**Ip6k1−/−** MSCs Exhibit Enhanced Growth and Survival Resulting in Increased Yields from Bone Marrow

To predict whether deletion of *Ip6k1* impacts MSC function, we first demonstrated by RNA sequencing that *Ip6k1* is the major isoform of three known family members (*Ip6k1–3*) detected in primary MSCs isolated from the bone marrow of wild type mice (Fig. 1A). Moreover, we confirmed by qPCR that genetic deletion of *Ip6k1* did not significantly alter expressed levels of *Ip6k2* and *Ip6k3* in wild type MSCs (Fig. 1B). Subsequently, we found that marrow from the long bones of *Ip6k1−/−* mice yielded significantly more MSCs on a per mouse basis following enrichment by immunodepletion as compared with age-matched, wild type littermates (Fig. 1C), and that yields of *Ip6k1−/−* versus *Ip6k1+/−* MSCs were also significantly higher after short term (1 week) culture expansion based on cell counting (Fig. 1D). Consistent with these results, *Ip6k1−/−* versus *Ip6k1+/−* MSCs underwent significantly more PDs (Fig. 1E) following long-term (1 month) culture resulting in a ~40-fold increase in total cell yield over three passages (Fig. 1F). Two-way ANOVA revealed a significant effect of genotype [F(1,8)=160.63, p < 0.0001] and passage number [F(3,8)=42.89, p < 0.0001] on the number of PDs. Although MSC yields following immunodepletion were lower in aged (18 months old) versus young (2 months old) *Ip6k1+/−* mice (Fig. 1C, 1G), overall MSC yields were significantly higher from aged *Ip6k1−/−* mice as compared with aged-matched *Ip6k1+/−* mice.
littermates (Fig. 1G). MSCs from Ip6k1−/− versus Ip6k1+/+ aged mice also exhibited enhanced growth in short-term culture (Fig. 1H). Despite differences in growth and survival, staining with a panel of established cell surface markers failed to detect significant differences in the phenotype of Ip6k1+/+ versus Ip6k1−/− MSCs (Supporting Information Fig. S1). Together these data suggest that Ip6k1 deletion enhances the overall fitness of primary MSCs from bone marrow.

Ip6k1−/− MSCs Exhibit Enhanced Osteogenic Differentiation at the Expense of Adipogenesis, and Enhanced Hematopoiesis-Supporting Activity

MSCs are best characterized by their differentiation potential and capacity to support hematopoiesis. Therefore, we evaluated how Ip6k1 deletion impacted these functions, which serve as metrics to assess overall MSC fitness. As shown in Figure 2, Ip6k1−/− MSCs isolated from both young (Fig. 2A) and old (Fig. 2C) mice exhibited a significantly reduced capacity for stimulus driven adipogenic differentiation as compared with MSCs obtained from wild type, age-matched littermates. This decline in adipogenic potential was offset by a modest but significant increase in stimulus-driven osteogenic differentiation of Ip6k1−/− MSCs (Fig. 2B, 2D). Next, we assessed the hematopoiesis-supporting activity of Ip6k1−/− versus Ip6k1+/− MSCs using standard colony forming assays. This analysis revealed a significant difference [F(12,35) = 148.9, p = 2.6 × 10−12] in the ability of conditioned media from Ip6k1+/− versus Ip6k1−/− MSCs to support growth of hematopoietic progenitors from the bone marrow of wild type mice in vitro (Fig. 3A). Post hoc analyses of these data confirmed that conditioned media from Ip6k1−/− versus Ip6k1+/+ MSCs yielded significantly more CFU-GM, CFU-E, CFU-G, and CFU-M colonies. Consistent with these findings, Luminex-based multiplex arrays further revealed that conditioned media from Ip6k1−/− versus Ip6k1+/− MSCs contained significantly higher levels of SDF1 (Fig. 3B) and much lower levels of interleukin 6 (Fig. 3C) although the latter difference failed to reach statistical significance (Fig. 3C). Enzyme linked immunosorbent assay analysis further showed that Ip6k1−/− versus Ip6k1+/− MSC conditioned media contained significantly higher levels of SCF (Fig. 3D). As perivascular MSCs are the principle source in marrow of the HSC maintenance factors SDF1 and SCF [6], these findings further illustrate how IP6K1 inactivation enhances MSC fitness.

Pharmacological Inhibition of IP6K1 in Mice Reversed Adipogenesis in Bone Marrow

To explore whether IP6K1 inactivation protects bone and marrow from the effects of HFD feeding, we placed wild type mice on a HFD for 8 weeks and treated them with or without daily injections of TNP (10 mg/kg, IP), a pan-inhibitor of IP6K activity. As shown in Figure 4A, the yield of MSCs from the bone marrow of HFD-fed mice treated with TNP was significantly higher than compared with vehicle treated mice at the end of the 8 week feeding period. Additionally, bone marrow cells from TNP treated mice exhibited an enhanced capacity to generate CFU-Fs (Fig. 4B), a measure of progenitor frequency within populations, as compared to cells from vehicle treated mice. CFU-Fs generated from the marrow of TNP treated mice were also significantly larger in surface area as compared to those obtained from the marrow of vehicle treated mice (Fig. 4C). Histological analysis of bone sections further revealed that TNP treatment resulted in conservation of trabecular bone (Fig. 4D). Morphometric analysis confirmed that the overall area of bone tissue was increased in TNP versus vehicle treated tissue sections (57.94 ± 9.8 vs. 48.89 ± 13.0) although differences between groups failed to reach statistical significance owing to wide variations in the two-dimensional cross sections of the three-dimensional trabecula (Fig. 4E).

Furthermore, while tissue processing precluded a quantitative analysis of marrow adiposity, bone marrow parenchyma of TNP-treated mice also contained fewer vacuoles as compared to vehicle treated mice based on visual inspection (Fig. 4D; Supporting Information Fig. S2), which was indicative of decreased marrow adiposity due to IP6K1 inhibition. Together, these data demonstrate that HFD feeding inhibits osteogenesis and promotes adipogenesis in adult bone marrow and that TNP treatment protects mice from these detrimental effects by inhibiting IP6K1 activity.

Ip6k1−/− MSCs Are Resistant to Oxidative Stress

We previously reported that primary MSCs from mouse bone marrow are highly sensitive to oxidative stress, exposure to which induces growth arrest and cellular apoptosis via a p53 dependent mechanism [31]. Therefore, we questioned if Ip6k1 deletion improved the fitness of MSCs by affecting the cellular redox balance. Flow cytometric analysis of unstained populations revealed that a higher proportion (58% vs. 46%) of Ip6k1−/− versus Ip6k1+/+ MSCs were characterized by low forward and side scatter (Fig. 5A; Supporting Information Fig. S3A), physical properties associated with low intrinsic mitochondrial reactive oxygen species (ROS) levels and resistance to oxygen-induced stress [31]. Consistent with this observation, flow cytometric analysis of MitosOX Red stained cells revealed that Ip6k1−/− MSCs exhibited lower intrinsic mitochondrial ROS levels when cultured in 5% oxygen saturation as compared to wild type MSCs (Fig. 5B; Supporting Information Fig. S3B), while we did not detect a significant difference in oxygen consumption rates between the two populations using Seahorse technology (data not shown). ImmunobLOTS of whole cell extracts revealed that p53 expression levels were markedly lower in Ip6k1−/− versus Ip6k1+/+ MSCs cultured in 5% oxygen (Fig. 5C). Expression of p16INK4A was also downregulated whereas expression of the E3 ubiquitin ligase MDM2, which mediates p53 degradation, was upregulated. Owing to the fact that oxygen-induced stress responses in primary MSCs are p53 dependent [31], lower intrinsic p53 levels likely account for lower overall mitochondrial ROS levels in Ip6k1−/− MSCs. Flow cytometric analysis further revealed that Ip6k1−/− versus Ip6k1+/+ MSC populations contain fewer pre-apoptotic and apoptotic cells based on Annexin V and propidium iodide (PI) staining (Fig. 5D; Supporting Information Fig. 3C) and this difference in apoptosis persisted in populations through passage 4 (not shown). To evaluate whether Ip6k1 deletion conferred onto cells resistance to oxygen-induced stress, Ip6k1−/− and Ip6k1+/− MSCs cultured in 5% oxygen were switched to atmospheric (~21%) oxygen for 1 week then stained with MitosOX Red. Flow cytometric analysis demonstrated that both populations exhibited a marked increase in mitochondrial ROS levels when exposed to 21% oxygen such that there was no significant difference in the overall abundance of MitosOX Red high cells under these conditions (Fig. 5E). Together, these findings demonstrate that Ip6k1−/− MSCs express intrinsically lower levels of mitochondrial ROS and p53 protein, which likely accounts for their enhanced growth and survival under physiological (~5%) oxygen levels but still retain sensitivity to oxygen-induced stress.
DISCUSSION

Skeletal homeostasis in adult organisms is controlled by a balance between bone formation and resorption, which is mediated by marrow resident osteoblasts and osteoclasts, respectively. During skeletal aging changes in the bone marrow microenvironment, such as increased oxidative stress and accumulation of advanced glycation end products, impairs osteogenesis resulting in reduced bone mineral density and increased marrow adiposity [32–35]. These age-associated changes have been linked, in part, to impaired osteogenesis and enhanced adipogenic differentiation of marrow resident MSCs [26, 36]. A number of disease states are also known to disrupt bone metabolism leading to defects in skeletal function. For example, patients with type 1 and type 2 diabetes have an increased risk of fracture due to diminished bone mineral density and bone fragility, and while the mechanisms responsible for these changes are varied [34, 37], these clinical observations demonstrate a strong link between nutritional status and skeletal homeostasis. Similarly, obesity induced by chronic steroid use is accompanied by rapid bone loss, indicating that obesity and bone metabolism are interrelated [38]. Finally, genetic studies in rodents have shown that HFD feeding induces adipogenesis and reduces osteogenesis in bone marrow, and that HFD-induced increases in leptin expression mediate these effects by acting on marrow-resident, LEPR-expressing MSCs.

Figure 2. Enhanced osteogenic and reduced adipogenic differentiation of inositol hexakisphosphate kinase 1 (Ip6k1) null mesenchymal stem/stromal cells (MSCs). (A, C): Quantification of adipogenic differentiation by MSCs isolated from young (2 months, A) and old mice (18 months, C). Photomicrographs (top panels) are cell monolayers stained with AdipoRed at 3 days post-induction. (B, D): Quantification of osteogenic differentiation by MSCs isolated from young (C) and old (D) mice. Photomicrographs (right panels) are cell monolayers stained with Alizarin Red at 10 days post-induction. All data represent mean ± SD (n = 3) from experiments repeated three times. Student’s t test (*, p < .05; **, p < .01; ***, p < .001). Magnification of photomicrographs is ×50. Abbreviations: DAPI, 4',6-diamidino-2-phenylindole; Ip6k1, inositol hexakisphosphate kinase 1.
These findings are consistent with other studies showing that mouse models of type 2 diabetes mellitus exhibit delayed fracture healing [39]. Nevertheless, as LEP/LEPR signaling exerts anorectic effects, its inhibition to prevent diabetes-induced bone loss is contraindicated. Importantly, certain anti-diabetic medications that have been developed that function as insulin sensitizers, such as the thiazolidinediones [40], have been shown to decrease bone mineral density by suppressing bone formation [41] and also increase marrow adiposity [42]. Therefore, while new generation drugs that function as PPARG partial inverse agonists may reduce unwanted effects on bone metabolism [10], identification of proteins/pathways that restore insulin sensitivity while preserving bone integrity are needed.

Our findings reveal that IP6K1 inactivation increases overall MSC fitness as evidenced by increased cell yields from bone marrow and enhanced growth and survival ex vivo under physiological (5%) oxygen saturation levels. Moreover, MSCs from \(lp6k1\) null mice exhibit augmented osteogenic potential and reduced adipogenic potential as compared to those isolated from age-matched congenic mice, consistent with an anti-aging phenotype, and also secreted higher levels of hematopoiesis-supporting cytokines. Furthermore, we found that \(lp6k1^{-/-}\)MSCs express lower basal p53 and mitochondrial ROS levels as compared to wild type MSCs. We previously reported that primary mouse MSCs isolated from bone marrow via immunodepletion are highly sensitive to oxygen-induced stress, that exposure to atmospheric oxygen rapidly induced mitochondrial ROS generation resulting in growth inhibition and increased cellular apoptosis, and that this stress response was p53-dependent [31]. Consistent with these results, we further demonstrated that exposure of \(p53^{-/-}\)MSCs to atmospheric oxygen failed to induce ROS generation, reduce cell viability or arrest growth. Therefore, considering the important role played by p53 in oxygen-induced growth inhibition, these findings

**Figure 3.** Inositol hexakisphosphate kinase 1 (\(Ip6k1\)) deletion enhances mesenchymal stem/stromal cell (MSC) hematopoiesis-support activity. (A): Methylcellulose colony forming assays using conditioned media from \(Ip6K1^{+/+}\) versus \(Ip6k1^{-/-}\)MSCs. Data are mean ± SD (\(n = 3\)). ***, \(p < .001\) by analysis of variance (ANOVA) and Tukey post hoc test. (B): Conditioned media from \(Ip6K1^{+/+}\) versus \(Ip6k1^{-/-}\)MSCs was concentrated 50-fold and analyzed using Luminex-based multiplex bead arrays. Data represent mean ± SD (\(n = 2\)). (A) is two-way ANOVA with Tukey post hoc test (**, \(p < .001\)) and (B–D) are Student’s \(t\) test (*, \(p < .05\); **, \(p < .01\)). Abbreviations: BFU, blast forming unit; CFU, colony forming unit; E, erythroid; G, granulocyte; GEMM, granulocyte, erythrocyte, monocyte, megakaryocyte; GM, granulocyte, monocyte; M, monocyte; IL-6, interleukin-6; IP6K1, inositol hexakisphosphate kinase 1; SCF, stem cell factor; SDF, stromal derived factor; VEGF, vascular endothelial growth factor.

[9]. These findings are consistent with other studies showing that mouse models of type 2 diabetes mellitus exhibit delayed fracture healing [39]. Nevertheless, as LEP/LEPR signaling exerts anorectic effects, its inhibition to prevent diabetes-induced bone loss is contraindicated. Importantly, certain anti-diabetic medications that have been developed that function as insulin sensitizers, such as the thiazolidinediones [40], have been shown to decrease bone mineral density by suppressing bone formation [41] and also increase marrow adiposity [42]. Therefore, while new generation drugs that function as PPARG partial inverse agonists may reduce unwanted effects on bone metabolism [10], identification of proteins/pathways that restore insulin sensitivity while preserving bone integrity are needed.

Our findings reveal that IP6K1 inactivation increases overall MSC fitness as evidenced by increased cell yields from bone marrow and enhanced growth and survival ex vivo under physiological (5%) oxygen saturation levels. Moreover, MSCs from \(lp6k1\) null mice exhibit augmented osteogenic potential and reduced adipogenic potential as compared to those isolated from age-matched congenic mice, consistent with an anti-aging phenotype, and also secreted higher levels of hematopoiesis-supporting cytokines. Furthermore, we found that \(lp6k1^{-/-}\)MSCs express lower basal p53 and mitochondrial ROS levels as compared to wild type MSCs. We previously reported that primary mouse MSCs isolated from bone marrow via immunodepletion are highly sensitive to oxygen-induced stress, that exposure to atmospheric oxygen rapidly induced mitochondrial ROS generation resulting in growth inhibition and increased cellular apoptosis, and that this stress response was p53-dependent [31]. Consistent with these results, we further demonstrated that exposure of \(p53^{-/-}\)MSCs to atmospheric oxygen failed to induce ROS generation, reduce cell viability or arrest growth. Therefore, considering the important role played by p53 in oxygen-induced growth inhibition, these findings
indicate that lower basal p53 levels in \( \text{Ip6k1}^{+/+} \) MSCs accounts in large part for their enhanced growth and survival at 5% oxygen saturation as compared to wild type cells. Rao et al. [43] showed that IP6K1 interacts with the Cullin-RING ubiquitin ligase 4 (CLR4), which is activated in response to DNA damage and targets substrates involved in cell cycle regulation and cell death. In these studies, expressed levels of p21 and p53 were diminished in \( \text{Ip6k1}^\text{-}\text{-}^\text{-} \) knockout MEFs due to de-repression of CLR4 activity. While we did not explore changes in this family of ubiquitin ligases in this study, IP6K1-mediated repression of CLR4 could account for reduced basal p53 expression. Importantly, both \( \text{Ip6k1}^{+/+} \) and \( \text{Ip6k1}^{+/+} \) MSCs exhibited robust mitochondrial ROS generation in response to oxygen-induced stress, indicating that \( \text{Ip6k1}^{+/+} \) MSCs retain oxygen sensitivity via p53 induction.

While we cannot confirm that IP6K1 inactivation results in expansion of the MSC progenitor pool in bone marrow, nor its effects specifically on LEPR\(^{+}\) MSCs, primary immunodepleted MSCs and LEPR\(^{+}\) MSCs both contain the majority of CFU-F activity within bone marrow [9, 31]. Therefore, while LEPR-MSCs may represent a fraction of the total MSC pool obtained after isolation, culture expansion and immunodepletion, increased MSC yields suggest that IP6K1 inactivation may result in expansion of the MSC pool in marrow. Furthermore, our data

Figure 4. Pharmacological inhibition of inositol hexakisphosphate kinase 1 (IP6K1) preserves mesenchymal stem/stromal cell (MSC) integrity and inhibits high fat diet (HFD)-induced adipogenesis and bone loss in adult bone marrow. (A): Total yield of immunodepleted MSCs from the bone marrow of mice fed a HFD for 2 months and treated with vehicle or N2-(m-trifluorobenzyl), N6-(p-nitrobenzyl)purine (10 mg/kg IP, \( n = 7 \) mice). Data represent mean ± SD (\( n = 3 \)). (B): CFU-F frequency measured in whole bone marrow from mice in (A). Data represent mean ± SD (\( n = 5 \)). (C): Total area occupied by CFU-Fs from (B). (D): Representative histological sections of long bones stained with H&E from mice in (A). Pink stained areas represent bone. Arrows demark lipid vacuoles. Magnification is \( \times 50 \). Scale bar = 180 \( \mu \)m. (E): Total area of bone determined by morphometric analysis of tissue sections (two sections per mouse, \( n = 6 \) mice per group) from mice in (A). Data represent mean ± SD (\( n = 10–11 \)). Student’s t test (**, \( p < .01 \); ***, \( p < .001 \)). Abbreviations: CFU, colony forming unit; F, fibroblast; MSC, mesenchymal stem/stromal cell; TNP, N2-(m-Trifluorobenzyl), N6-(p-nitrobenzyl)purine.
suggest that TNP treatment protects bone marrow from the detrimental effects of HFD feeding by preserving trabecular bone and preventing fat deposition within marrow.

Previous studies from one of us showed that genetic deletion and pharmacological inhibition of IP6K1 protects mice from HFD-induced weight gain and insulin resistance by enhancing insulin sensitivity via AKT activation, and decelerating weight gain by enhancing thermogenic energy expenditure in adipose tissue [12–14, 23]. Our current findings together with previous studies demonstrate that IP6K1 inactivation/inhibition restores glucose homeostasis and insulin sensitivity in HFD fed mice [12] without adversely affecting bone metabolism. Indeed, IP6K1 inhibition appears to expand the MSC pool in bone marrow and alter the cellular redox balance in cells in a manner that increases overall fitness and osteogenic potential. For this reason, IP6K1 promises to be a better and consistent target in diseases of aging including insulin resistance, obesity, and osteoporosis as opposed to the widely studied PPARG or LEP/LEPR pathways.

Figure 5. Inositol hexakisphosphate kinase 1 (Ip6k1) deletion enhances mesenchymal stem/stromal cell (MSC) survival and resistance to oxidative stress. (A): Flow cytometric analysis showing percentage of low forward scatter cells and side scatter cells in Ip6k1+/+ versus Ip6k1−/− MSC populations isolated from 2 months old mice and cultured in 5% oxygen saturation. (B): Left panel; representative histogram from flow cytometric analysis of MSC populations from (A) after staining with MitoSOX Red. Right panel: Percentage of MitoSOX Red high and low expressing cells in each MSC population. Data represent mean ± SD (n = 2). (C): Western blot of MSC cell extracts isolated from femurs and tibias of 8-week-old Ip6k1+/+ and Ip6k1−/− mice. (D): Flow cytometric analysis of Annexin V and PI stained MSCs from (A) showing percentage of preapoptotic (AnnexinV−/PI−) and apoptotic (AnnexinV+/PI+) cells. Data are mean ± SD (n = 2). (E): Left panel; representative histogram from flow cytometric analysis of MitoSOX Red stained MSCs cultured in 5% oxygen saturation or 21% oxygen saturation. Right panel: Percentage of MitoSOX Red high expressing cells in each MSC population. Data are mean ± SD (n = 2). Student’s t test (**, p < .01; ***, p < .001). Abbreviations: FSC, forward scatter cell; Ip6k1, inositol hexakisphosphate kinase 1; PE-A, phycoerythrin; SSC, side scatter cell.
MSCs play important roles in hematopoiesis and adult skeletal homeostasis, and both these processes are known to be impaired in response to systemic changes in energy metabolism, such as occurs in obesity and diabetes. The identification of LEPR expression by MSCs provided one mechanism by which systemic changes in energy metabolism may be sensed in bone marrow and impact hematopoiesis and skeletogenesis. Our findings that IP6K1 inactivation improves MSC fitness while promoting osteogenesis suggest a mechanism by which pharmacological inhibition of IP6K1 restores insulin sensitivity in HFD fed mice without promoting bone loss. Consequently, drugs that inhibit IP6K1 may be advantageous over other insulin sensitzizers, such as the thiazolidinediones, which negatively impact bone metabolism resulting in increased risk of fracture in diabetic patients.

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