PEDF: An essential stem cell regulator in models of Osteogenesis Imperfecta Type VI

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ABSTRACT

The rare bone disease Osteogenesis Imperfecta (OI) type VI is caused by mutations in the gene coding for PEDF, Serpinf1. Individuals with OI type VI have an accumulation of unmineralized bone matrix and multiple fractures. Our lab group has previously shown that PEDF restoration in the mouse model of OI type VI increases bone mass and mineralization. Further, we demonstrated that PEDF directs mesenchymal stem cells to the osteoblast lineage. One mechanism appears to involve modulation of canonical Wnt/β-catenin signaling in a temporally-defined manner. We have also induced pluripotent stem cells from a patient with OI type VI and differentiated the cells into osteoblasts to investigate how PEDF regulates the expression of various bone matrix proteins such as IBSP. In this brief review, we provide an overview of PEDF biology and highlight how PEDF’s role as a stem cell regulator lends support to its causative role in OI type VI.

PEDF Null Mutations Result in Osteogenesis Imperfecta Type VI

Osteogenesis Imperfecta (OI) type VI is an autosomal recessive bone disease characterized by decreased bone mineralization and multiple early fractures1,2. The histological markers of OI type VI are accumulations of unmineralized bone matrix (osteoid) and the appearance of a fish like pattern under polarized light1.

Unlike classic forms of OI that involve collagen mutations, OI type VI occurs in the setting of normal collagen synthesis, post-translations modification, and secretion1. Exome sequencing has identified null mutations in the gene coding for pigment epithelium-derived factor (PEDF), Serpinf1, as the causative genetic defect in OI type VI3. This results in complete absence of circulating PEDF4-6. The PEDF knockout (KO) mouse recapitulates key features of human OI type VI.7 Another OI subtype, Type V, is caused by mutations in the gene coding for the bone enriched protein BRIL, IFITM58. In some instances, OI Type V presents with a phenotype consistent with the PEDF null state. Farber et al. have identified an IFITM5 mutation at codon 40 as the cause of this form of OI Type V. This mutation results in a S40L substitution and results in the functional absence of secreted PEDF9. How BRIL influences PEDF secretion remains unknown.

In addition to null mutations, homozygous in-frame insertion and deletion mutations in Serpinf1 have been identified and result
in retention or degradation of PEDF within intracellular compartments. In these cases, PEDF levels are >10 fold lower compared to normal individuals and present clinically with a phenotype similar to PEDF null patients. Recently, a point mutation in intron 6 of *Serpinf1* (c.787-10C>G) has been identified as another genetic defect leading to OI type VI. This point mutation results in an altered splice site, addition of three amino acids (p.Lys262_Ile263InsLeuSerGln), and an inability to secrete PEDF. The absence of PEDF through null mutations or those that lead to its inability to be secreted result in OI type VI and provides insight into PEDF's role as a key regulator of bone development and stem cell biology.

**PEDF Biology**

PEDF is a 50kDa secreted glycoprotein first identified and isolated from the conditioned medium of cultured human fetal retinal pigment epithelium cells. PEDF is a member of the serine protease inhibitory (SERPIN) family, and lacks protease inhibitory function due to differences within the reactive center loop (RCL). Expression levels are highest in the liver and then adipose tissue. PEDF is approximately 100nM in normal individuals, increased up to five-fold with obesity and the metabolic syndrome, and is undetectable or dramatically reduced in OI type VI and provides insight into PEDF's role as a key regulator of bone development and stem cell biology.

PEDF appeared to stimulate the Wnt co-receptor LRP6 in early undifferentiated MSCs, while in later stages of terminal differentiation, PEDF appeared to block LRP6 activation. These effects on osteoblast lineage specification were accompanied by blockade of MSC to adipocyte differentiation that was restricted at an early time point. Once adipocyte lineage specification beyond 48 hours was achieved, PEDF could not block adipogenesis. In cells without LRP6 PEDF-mediated effects were not observed. Further, our group found that PEDF KO mice displayed reduced bone volume and high fat content consistent with this effect on early MSC lineage specification. These data provided evidence of PEDF's role in modulating MSC differentiation to the osteoblast lineage while concurrently blocking adipocyte maturation.

**PEDF's Role in an Important Developmental Pathway**

Other groups have shown that PEDF modulates MSC differentiation to the osteoblast lineage. Li et al. have shown that media supplemented with PEDF induces expression of osteoblast associated genes while also facilitating mineral deposition. In addition, Niyibizi and colleagues demonstrated decreased osteoblast differentiation and mineralization in PEDF knockdown hMSCs that were rescued with exogenous PEDF. Together, these data provide evidence for PEDF's direct role in osteoblast differentiation from MSC precursors and help explain why the absence of PEDF is the cause of a debilitating bone disease.

In our most recent work, we used the murine model of OI type VI (PEDF null state) to determine whether *in vivo* PEDF reconstitution would recover bone loss. In mature mice (6 months old) PEDF injection had no effect in wild-type (WT) mice, but increased trabecular bone volume/total volume (BV/TV) by 52% in PEDF KO mice. Similarly, PEDF reconstitution recovered BV/TV by 35% in young PEDF KO mice (19 days old). Functional studies demonstrated that PEDF treatment improved various biomechanical parameters in PEDF KO femurs such as bone stiffness and plasticity. Notably, young mice had relatively less bone recovery compared to older mice, suggesting the possibility that older mice may have accumulated more unmineralized matrix characteristic of OI type VI.

To understand the mechanisms by which PEDF reconstitution recovers bone mass, we examined the canonical Wnt pathway in hMSCs undergoing osteoblast differentiation. A Wnt/β-catenin-GFP reporter demonstrated that Wnt activity was highest early in the differentiation time period but was absent by the end of the 21 day osteogenic differentiation protocol. Meanwhile, endogenously secreted PEDF increased over time, suggesting that it plays an important role in the terminal phase of osteoblast differentiation and directs MSCs to the osteoblast lineage in a temporally restricted manner. Continuous Wnt3a stimulation during the entire differentiation protocol resulted in decreased PEDF secretion and hMSC mineralization. However,
mineralization was recovered when PEDF was added during the terminal phase of osteoblast differentiation. Therefore, our data suggest that PEDF antagonizes Wnt activity during osteoblast differentiation, thereby allowing for osteoblast maturation and matrix mineralization (Figure 1). To confirm PEDF’s inhibitory role of LRP6, we also showed that PEDF suppresses Wnt activity to the same extent as the well-known Wnt inhibitor, DKK1. In fact, DKK1 has also been shown to impede terminal osteoblast differentiation\(^3\), demonstrating that multiple endogenous inhibitors are present to turn off Wnt signaling. These mechanisms highlight the importance of PEDF-directed Wnt inhibition for proper osteoblast maturation.

**Generation of PEDF null iPSC**

To assess the mechanisms leading to the OI type VI phenotype, we generated induced pluripotent stem cells (iPSCs) from a patient with OI type VI, differentiated them into MSCs, and assessed osteoblast-related gene expression\(^3\). Expression of bone sialoprotein, IBSP, was increased over 5-fold compared to control cells. IBSP is a major component of the extracellular matrix secreted by osteoblasts and its overproduction has been associated with reduced bone density and impaired bone homeostasis\(^34,35\). Interestingly, IBSP levels decreased in response to exogenous PEDF. We have also seen that the PEDF 34mer peptide decreased levels of type I collagen in PEDF null iPSC (unpublished data). Thus, PEDF possesses multiple functions during bone development, including directing MSC differentiation and regulating extracellular matrix proteins.

**PEDF in Alcohol-Induced Bone Loss**

Because the PEDF null state causes OI type VI, we questioned whether suppressed PEDF occurs in other disease states associated with bone loss. It is well established that alcohol abuse is a risk factor for osteoporosis and thus bone fragility\(^36\). In addition, alcoholic liver disease is associated with increased fractures\(^37\). The recent identification of secreted factors from the liver regulating bone mass support these clinical associations\(^38\). Given that PEDF is most highly expressed in the liver\(^14\), our most recent data is evaluating PEDF within the context of alcohol-induced bone loss. In wild-type and PEDF KO mice, ethanol feeding (5% Lieber-DeCarli diet) resulted in loss of trabecular bone volume in the femur and vertebral bodies. Bone mass was recovered following PEDF reconstitution in PEDF KO vertebral bodies (unpublished data). Moreover, we demonstrated that PEDF is suppressed in the bone and sera of wild-type mice fed an ethanol diet. This data is consistent

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**Figure 1: PEDF directed osteoblast differentiation**

PEDF blocks Wnt3a ligand binding, allowing for proper osteoblast terminal differentiation. In the absence of PEDF, either in the PEDF null state (OI Type VI) or in the PEDF deficient state (alcohol induced PEDF suppression), continuous Wnt3a exposure hinders terminal osteoblast differentiation and normal mineralization. PEDF is necessary for osteoblast precursors to differentiate into mature osteoblasts. MSC, mesenchymal stem cell.
with previous reports from our group demonstrating that loss of PEDF occurs in human and murine hepatic tissues with ethanol-induced steatosis. This model highlights the role of PEDF in bone homeostasis and suggests that alcohol abuse may share similar mechanisms to the pathogenesis of OI type VI.

Conclusion

Initial work characterizing PEDF suggest that it is a critical factor modulating developmental pathways and stem cells. First studied as a neuronal differentiation factor in retinoblastoma cells, in vivo studies demonstrated that PEDF induced neoblastoma cells into a terminally differentiated state. However, PEDF in vivo transplantation in retinoblastoma cells resulted in tumor growth, suggesting that PEDF effects are context specific and in stem cell populations can lead to either cell renewal or differentiation. PEDF in vitro strongly affected the differentiation of photoreceptors and counteracted gliotoxic damage, i.e. inflammation, of radial glial cells.

While this paradox is still incompletely understood, the discoveries of the PEDF null state in OI Type VI and PEDF’s role as a Wnt agonist in many tissue sites have explained its diverse functionality. As a fundamental developmental factor in retinoblastoma cells, Wnt signaling is crucial in development and adult homeostasis. As a result, aberrant Wnt signaling is associated with a wide array of abnormalities including various bone diseases, diabetes, and irregular angiogenesis in diabetic retinopathy. Thus, it is important to determine if PEDF-directed Wnt signaling occurs in disease states where Wnt signaling is disrupted. In addition, it would be useful to understand how Wnt signaling is altered in PEDF deficient disease states such as alcoholic osteodystrophy.

Grant support: This work was supported by Digestive Diseases Research Core Center–SP30DK034989 (CC); NIH/NIAAA R2123607; Veterans Affairs Merit Grant (CC).

References


