SKELETAL MUSCLE WEAKNESS AND MITOCHONDRIAL DYSFUNCTION IN THE OSTEOGENESIS IMPERFECTA MURINE (OIM) MODEL

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Doctor of Philosophy

by

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The undersigned, appointed by the dean of the Graduate School, have examined the dissertation entitled

**SKELETAL MUSCLE WEAKNESS AND MITOCHONDRIAL DYSFUNCTION IN THE OSTEOGENESIS IMPERFECTA MURINE (OIM) MODEL**

presented by Victoria L Gremminger

a candidate for the degree of Doctor of Philosophy of Biochemistry,

and hereby certify that, in their opinion, it is worthy of acceptance.

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Dr. Antje Heese

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Dr. Michael Petris
DEDICATION

I would like to dedicate this dissertation to my wonderful husband, Thomas Gremminger. I am beyond grateful for your continued love and support. You have been such a rock to me, providing constant encouragement throughout the progression of my dissertation. I could think of no one else I would have rather shared this journey with and cannot wait to see where our next adventure takes us. I would also like to dedicate this dissertation to my sweet baby, Gus, who has been patient with me as I have learned to navigate motherhood while finishing my degree and has brought so much joy into my life, during the sometimes challenging and difficult moments of finishing my dissertation research.
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SKELETAL MUSCLE WEAKNESS AND MITOCHONDRIAL DYSFUNCTION IN THE OSTEOMELIOGENESIS IMPERFECTA MURINE (OIM) MODEL

Victoria L Gremminger
Dr. Charlotte L. Phillips, Dissertation Advisor

ABSTRACT

Osteogenesis imperfecta (OI), also commonly referred to as brittle bone disease, is a heritable connective tissue disorder occurring in roughly 1:15,000 births. OI arises as a result of mutations in the type I collagen genes, COL1A1 and COL1A2, approximately 85% of the time with the remaining 15% of cases arising from mutations in genes involved in posttranslational modification of type I collagen, osteoblast maturation or mineralization. OI is a heterogeneous disorder that can be classified into four major types with severity ranging from perinatal lethality to premature osteoporosis. As the name suggests, osteogenesis imperfecta, literally translating to imperfect bone formation, results in bone fragility with patients often experiencing many fractures throughout their lifetime. While bone fragility is the most prominent manifestation of OI, skeletal muscle weakness, cardiopulmonary complications, short stature, and craniofacial abnormalities are also common. There is currently no cure for OI and therapeutic options rely on mitigating symptoms, primarily through the use of bone anti-resorptive agents referred to as bisphosphonates. Although, current treatment options focus solely on bone health, skeletal muscle weakness is a common manifestation in OI, where 80% of patients with mild OI experience muscle force deficits, and with even higher percentages in patients with more clinically severe OI. Historically, OI muscle weakness was largely attributed to inactivity with recent studies highlighting its inherent nature in both patients and
mouse models. Studies investigating the mechanisms by which skeletal muscle weakness arises in OI are limited, despite the large prevalence.

My research sought to better understand OI muscle weakness primarily through the investigation of mitochondrial health in a mouse modeling a severe human type III OI (oim/oim), as mitochondria are important regulators of energy metabolism and overall cell health. We hypothesized that oim/oim mice, exhibiting severe skeletal muscle weakness, would exhibit mitochondrial dysfunction suggesting a correlation between skeletal muscle and mitochondrial function. To test this hypothesis, we assessed mitochondrial function and content in the oim/oim mouse. One of our major findings was the observation that oim/oim mice exhibit >50% reductions in gastrocnemius mitochondrial respiration rates relative to wildtype littermates. Additionally, we found that citrate synthase activity in oim/oim isolated gastrocnemius mitochondria was reduced relative to wildtype littermates. Furthermore, to determine if skeletal muscle mitochondrial function correlated with skeletal muscle severity, we evaluated mitochondrial respiration in a mouse model of mild OI (+/oim). We did not find differences between +/oim and WT gastrocnemius mitochondrial respiration suggesting that mitochondrial function does correlate with skeletal muscle function. Moreover, we did not observe changes in mitochondrial respiration in oim/oim liver and heart suggesting the mitochondrial dysfunction is not global in the oim/oim mouse.

Additionally, we sought to investigate whole body metabolic alterations, as skeletal muscle comprises roughly 50% of body mass and is a significant contributor to the body’s resting metabolic rate. We hypothesized that skeletal muscle mitochondrial dysfunction in the oim/oim mouse would lead to changes in metabolic parameters.
including altered substrate utilization, altered body composition, and changes in energy expenditure. Interestingly, we did not observe changes in substrate utilization, although we did note increased energy expenditure and subtle changes in body composition with oim/oim animals exhibiting reduced percentages of fat mass and increased percentages of lean mass relative to wildtype littermates.

Overall, my research was the first to implicate mitochondrial dysfunction in the pathophysiology of OI using a mouse model of severe OI. This work has led to numerous studies in other mouse models evaluating mitochondrial function and energy metabolism. While there is more work to be done to further understand the mechanisms and correlation between mitochondrial dysfunction and skeletal muscle weakness in OI, this novel finding has initiated a new area of research in OI and has contributed to the overall understanding of OI muscle weakness.
CHAPTER I

INTRODUCTION AND REVIEW OF THE LITERATURE
Osteogenesis imperfecta

Osteogenesis imperfecta (OI), also commonly referred to as “brittle bone disease,” is a genetically and clinically heterogeneous heritable connective tissue disorder affecting roughly 1:15,000 births[1,2]. While the most striking feature of the disease is bone fragility and skeletal dysplasia, more recent research has implicated inherent muscle weakness as part of the pathophysiology of the disease[3–6] and a small number of studies have highlighted the presence of a metabolic phenotype in OI [7,8]. Other manifestations include short stature, cardiopulmonary abnormalities, kyphoscoliosis, and the presence blue-gray sclera[1,2].

The heterogeneity of OI is exemplified by the more than 1500 recognized disease-causing mutations[9,10]. Roughly 85% of the time OI is the result of qualitative (dominant negative) or quantitative (haplo-insufficient) defects in type I collagen due to mutations in the type I collagen genes: COLIA1 and COLIA2[1,2,9,11]. The remaining 15% of cases are the result of mutations in genes implicated in bone mineralization, posttranslational modifications, folding, and secretion of type I collagen, as well as those involved in osteoblast maturation and function[1,2,12].

As might be expected from the vast number of disease-causing mutations affecting multiple genes exhibiting unique functions, the clinical severity of the disease ranges considerably, from asymptomatic to perinatal lethality[1,2,12,13]. Classically, there are four subtypes of OI (Sillence classification; I-IV) with type I being the most mild form of the disease, type II being the most severe (resulting in perinatal lethality), type III being the most severe viable form (often resulting in the non-ambulation of patients), and type
IV being of moderate severity [13]. As the genetic understanding and the number of identified novel OI-causing mutations has expanded, there has been a shift in the classification of OI, with 20 types currently recognized in the Online Mendelian Inheritance in Man (OMIM) database [12,14]. A summary of known genetic loci responsible for human OI with corresponding mouse models can be seen in Table I-1.
Table I-1: Summary of known osteogenesis imperfecta genetic loci and representative mouse models.

<table>
<thead>
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<th>Gene Symbol</th>
<th>Silence Type</th>
<th>Protein</th>
<th>Mouse Models (OI type)</th>
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<td><strong>Group A: Autosomal Dominant and defects in collagen synthesis, structure, or processing</strong></td>
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<td></td>
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<tr>
<td>COL1A1 or COL1A2</td>
<td>I, II, III, and IV</td>
<td>Collagen type I, alpha1 or alpha2</td>
<td><strong>COL1A1 Mutations:</strong> BrlIV/+ (type IV), Jrt/+ (type IV/III) <strong>COL1A2 Mutations:</strong> oim/oim (type III), +/oim (type I), +/G610C (type IV)</td>
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<tr>
<td><strong>Group B: Defects in collagen post translational modification</strong></td>
<td></td>
<td></td>
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<tr>
<td>CRTAP</td>
<td>VII</td>
<td>Cartilage-associated protein</td>
<td>Crtap&lt;sup&gt;−−&lt;/sup&gt; (type VII)</td>
</tr>
<tr>
<td>LEPRE1/P3H1</td>
<td>VIII</td>
<td>Leucine proline-enriched proteoglycan1/prolyl 3-hydroxylase-1</td>
<td>P3H1&lt;sup&gt;−−&lt;/sup&gt; (type VIII)</td>
</tr>
<tr>
<td>PPIB</td>
<td>IX</td>
<td>Peptidylprolyl isomerase B/cyclophilin B</td>
<td>Ppib&lt;sup&gt;−−&lt;/sup&gt; (type IX)</td>
</tr>
<tr>
<td>TMEM38B</td>
<td>XIV</td>
<td>Transmembrane protein 38B</td>
<td>Tric-b (type XIV)</td>
</tr>
<tr>
<td><strong>Group C: Defects in collagen folding and cross linking process</strong></td>
<td></td>
<td></td>
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<tr>
<td>SERPINH1</td>
<td>X</td>
<td>Serpin Peptidase inhibitor, clade H, member1/heat shock protein 47</td>
<td>Hsp47&lt;sup&gt;−−&lt;/sup&gt; (type X)</td>
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<tr>
<td>FKBP10</td>
<td>XI</td>
<td>FK506 binding protein 65</td>
<td>Fkbp10&lt;sup&gt;−−&lt;/sup&gt; (type XI)</td>
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<td><strong>Group D: Defects in bone mineralization</strong></td>
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<td></td>
<td></td>
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<tr>
<td>IFITM5</td>
<td>V</td>
<td>Interferon-induced transmembrane protein 5</td>
<td>Ifitm5&lt;sup&gt;−−&lt;/sup&gt; (type V)</td>
</tr>
<tr>
<td>SERPINF1</td>
<td>VI</td>
<td>Pigment epithelium-derived factor</td>
<td>Pedf&lt;sup&gt;−−&lt;/sup&gt; (type VI)</td>
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<tr>
<td><strong>Group E: Defects in osteoblast development and function</strong></td>
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<td></td>
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<tr>
<td>SP7</td>
<td>XII</td>
<td>Transcription factor 7/osterix</td>
<td>SP7&lt;sup&gt;−−&lt;/sup&gt; (Osx) (type XIII)</td>
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<tr>
<td>CREB3L1</td>
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<td>cAMP responsive element binding protein 3 like 1</td>
<td>OASIS&lt;sup&gt;−−&lt;/sup&gt; (type XVI)</td>
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The genetic and clinically heterogeneous nature of OI presents a challenge when it comes to treatment strategies. The diversity in OI mutations and severities prevents use of a “one size fits all” approach; instead physicians focus on mitigating symptoms, as there is currently no cure for OI [2,12]. Current treatments for OI are predominately focused on mitigating the bone fragility, with little to no consideration of muscle weakness. These include the use of bone antiresorptive and osteoanabolic pharmaceutical compounds, physical therapy, and surgical interventions such as intermedullary rodding in pediatric patients[2,12,15–18].

The current standard of treatment in OI includes the administration of intravenous or oral bone antiresorptive bisphosphonate therapies, which improve bone mineral density (BMD) although their efficacy in fracture reduction remains debated and the safety and efficacy in pediatric patients remains unclear [11,12,16,19]. Additionally, long term use of bisphosphonates may have an overall detrimental effect on bone due to decreased bone remodeling, increased accumulation of microfractures thus compromising the bone integrity, and potential association with osteonecrosis of the jaw[11,15,19–21]. The use of bone anabolic therapies, including parathyroid hormone, anti-sclerostin, and anti-TGFβ, are currently being evaluated for use in OI[15,22–24]. It is important to note that while all these compounds increase bone formation, they do not alter the compromised integrity of bone material due to type I collagen defects.

**Type I collagen structure and synthesis**

There are 28 members of the collagen superfamily, making collagen the most abundant protein in the body, comprising roughly 30% of total protein mass [25]. Type I collagen,
found throughout the body in various connective tissues and accounting for approximately 95% of bone collagen content, is a fibrillar type of collagen organized as a heterotrimer composed of $\alpha 1(1)$, $\alpha 2(1)$ chains [26]. Each of these alpha chains exists as a polyproline II-like helix, which supercoils to form a triple helical motif 300 nm in length and 1.5 nm in diameter [26,27]. This process is facilitated by the repeating primary amino acid sequence Gly-X-Y, where X and Y are often proline and hydroxyproline, respectively. The presence of glycine at every third amino acid position allows for the tight packing of the triple helix, as glycine lacks a side chain, and can be buried within the helix [28].

The biosynthesis of collagen (Figure I-1) is complex and requires numerous temporally and spatially coordinated processes including many post translational modifications (PTMs)[26,29–31]. The initial step of type I collagen biosynthesis is the transcription of COLIA1 and COLIA2 mRNA. Once transcribed, the mRNA enters the cytoplasm where it is translated on the ribosomes of rough endoplasmic reticulum (ER) in the N- to C-terminal direction. The newly translated molecule, referred to as a pre-pro-peptide, contains a signal on the N-terminus directing it to the ER where it is recognized by a signal recognition particle [26,29–31]. Once inside the ER lumen, three main events take place signifying the transition to procollagen: 1) the signal on the N-terminus is cleaved, 2) select lysine and proline amino acids are hydroxylated and 3) hydroxylysine is glycosylated via the addition of monosaccharides[26,29–31]. This hydroxylation and glycosylation contribute to the overall thermal and mechanical stability of type I collagen[26,29–31]. Following the PTM of the pro-$\alpha$ chains, the alignment of three pro-$\alpha$ chains, facilitated by disulfide bonding of cysteine residues in the C-telo-peptide, occurs
leading to triple helix formation in a zipper-like fashion from C- to N-terminus, a process that on average takes an astounding 14 minutes[26,29–31]. Once the triple helix is formed, the chaperone protein hsp47 binds to and increases stability of the triple helix while preventing premature procollagen aggregates as procollagen moves to the golgi[26,29–31]. From the golgi, procollagen is transported to the extracellular space, via secretory vesicles, where the globular N- and C- pro-peptides are cleaved [the rate limiting step of fibrillogenesis] [32]. Following cleavage of the pro-peptides, lysyl oxidase catalyzes the formation of reactive aldehydes on lysine and hydroxylysine residues which will spontaneously react to form intramolecular crosslinks[29].
Figure I-1 Collagen Synthesis. Originally published in Trends in Cell Biology (Makareeva et al., 2011)[33] 1) Cotranslational translocation of procollagen chains into the lumen of rough ER. 2) C-propeptide domains of two pro-α1(I) and one pro-α2(I) chains assemble into a trimer and fold. 3) The triple helix is nucleated at the C-terminal end. 4) The triple helix propagates towards the N-terminal end. 5) Procollagen is cotransported with HSP47 to the cis-Golgi network following N-propeptide folding. 6) Procollagen is transported through the Golgi stack. 7) Procollagen is secreted into the extracellular space. 8) N- and C-propeptides are cleaved via extracellular N- and C-proteinases. 9) Mature collagen assembles into fibers.
Bone derives its mechanical properties and strength in part from its mineral and organic components, hydroxyapatite and collagen fibrils, respectively [26,29,34]. The collagen matrix has been demonstrated to contribute mainly to bone toughness and strength whereas the mineral component contributes predominately to bone stiffness [35]. Type I collagen fibrils themselves exhibit great tensile strength that is “gram for gram” stronger than steel allowing them to withstand great forces [31], as evidenced by tendon, where type I collagen is abundant, connecting bone and muscle while enduring large forces generated by muscle pull [31].

That osteogenesis imperfecta is caused most often by mutations in the type I collagen genes exemplifies the significance of type I collagen strength and organization for overall bone biomechanics, demonstrating that qualitative and/or quantitative defects in type I collagen severely compromise bone [1,26]. For example, the osteogenesis imperfecta murine (oim) model, containing a mutation in the Col1a2 gene, has been used to analyze the role of intrinsic bone properties on bone. Bone biomechanical properties of oim mice demonstrate reduced biomechanical strength, evidenced by a 60% reduction in post-yield deformation, a parameter often associated with accumulated microdamage, viscoelastic response, and plastic deformation, ultimately indicating a weaker bone as the result of the Col1a2 mutation [26,36–38].

**oim mouse model**

The osteogenesis imperfecta murine (oim) mouse model contains a spontaneous deletion of the 3983rd nucleotide in the Col1a2 gene[39]. This mutation results in the production of homotrimeric type I collagen[α1(I)3] rather than the normal heterotrimeric type I
collagen $[\alpha 1(\text{I})_2 \alpha 2(\text{I})]$[39]. The homozygous $oim/oim$ mouse models a patient with severe type III OI [40,41], exhibiting severe skeletal fragility, whereas the heterozygous $+/oim$ mouse models more mild type I OI, exhibiting less severe skeletal fragility [39].

Bone fragility in the $oim$ mouse can be at least partially explained by the reduced ability of homotrimeric collagen to resist tensile stress in the $oim$ mouse as compared to wildtype collagen[42]. Another contribution to the skeletal fragility noted in the $oim/oim$ mouse is likely the reduced ability to form “normal” mineral crystals relative to wildtype animals, a result of homotrimeric collagen altering the scaffold on which this mineral is laid[43]. Numerous studies have been conducted on the $oim$ mouse, leading to a better understanding of the contribution of type I collagen to overall bone biomechanical properties including biomechanical strength[36,37,42,44,45].

In addition to the severe bone fragility, the $oim/oim$ mice exhibit severe muscle weakness and studies in the $oim$ mouse were the first to implicate inherent skeletal muscle weakness as part of the disease pathology of OI[3]. Gentry et al. demonstrated that $oim/oim$ skeletal muscles had reduced specific contractile generating force, [contractile generating force normalized to myofiber cross sectional area] in the gastrocnemius, tibialis anterior, and plantaris muscles in addition to reduced body weights, muscle weights, and relative muscle weights[3]. This study launched the field of muscle research in osteogenesis imperfecta.
Skeletal muscle weakness and energy metabolism in OI

Historically, muscle weakness in osteogenesis imperfecta had been attributed to hypoactivity, with patients reporting fatigue and reduced exercise capacity in addition to muscle weakness[6,46]. Studies in the *oim/oim* mouse were the first to demonstrate an inherent muscle pathology as part of the pathophysiology of OI[3]. Following this novel discovery, clinicians and basic scientists began to consider and investigate muscle function in OI patients and other mouse models of OI. Of these studies, one of the most significant findings was that 80% of patients with a type I OI phenotype caused by a mutation in either *COL1A1* or *COL1A2* experienced muscle force deficits when muscle force was assessed via mechanography[6,47]. Muscle function correlates with OI severity, with muscle force deficits often more severe in patients with moderate and severe types IV and III OI, respectively [48]. Another interesting finding by Pouliot-Laforté et al., was that children with OI who experienced muscle force deficits were just as active as their healthy peers, although neither group of children met the daily recommended amount of activity, suggesting that reduced physical activity is not the leading cause of muscle weakness in children with OI[47]. Over the past decade, efforts to better characterize and understand muscle weakness in patients with OI have drastically increased. However, the cause of muscle weakness in OI remains unknown, although numerous mechanisms continue to be investigated including the role of altered collagen in muscle connective tissue layers and tendon, reduced mobility in patients with OI, and altered signal transduction and mechanotransduction between muscle and bone[2,14,48].
In addition to patient studies, skeletal muscle function has also been assessed in several mouse models of OI, providing useful insights to potential mechanisms of muscle weakness, as well as for evaluation of novel therapeutic options targeting muscle weakness in OI[3,14,49]. These studies have demonstrated a positive correlation between OI severity and skeletal muscle weakness. Skeletal muscle weakness is absent in the +/-G610C or “Amish” mouse model of mild OI, containing a glycine to cysteine substitution in the triple helical domain; a mutation also found in an old Amish kindred affected by OI[49,50]. In contrast, in models of severe OI such as the oim/oim and Colla1Jrt/+ , reduced activity and skeletal muscle weakness are observed[3,8,51,52]. The idea that skeletal muscle weakness may be mutation- and severity- specific in OI warrants further investigation to better understand the mechanisms of this muscle weakness.

The Colla1Jrt/+ is the first combined model of both severe type IV OI and Ehlers Danlos Syndrome (EDS) due to a COL1A1 mutation leading to skipping of exon 9 and subsequent 18 amino acid deletion in the triple helical domain[51]. EDS is a genetic connective tissue disorder arising primarily due to mutations in type V collagen, resulting in skin hyper-elasticity, joint hypermobility, skin fragility, and blood vessel fragility, although a subset arise from mutations near the amino terminal end of type I collagen [53,54]. In addition to reduced physical activity and skeletal muscle weakness, recent evidence in the Colla1Jrt/+ mouse suggests a metabolic phenotype exists and contributes to the pathophysiology of the disease[8]. While there are limited studies evaluating energy metabolism in OI, Cropp and Myers, in 1972, described the presence of a hypermetabolic state in adolescent male OI patients[7]. Taken together, the report of a
metabolic phenotype in both the mentioned patient cohort and in the Colla1Jrt/+ model suggest energy metabolism as an important contributor to the pathophysiology of OI. The idea of altered energy metabolism in OI may not be that surprising due to the wide prevalence of skeletal muscle weakness in patients with OI and the fact that skeletal muscles account for roughly 20% of the basal metabolic rate[15,48,55,56].

While the bone phenotype in OI has been well characterized and thoroughly evaluated in both patients and countless animal models, skeletal muscle weakness in OI is a relatively recent finding, and a thorough understanding of the mechanisms leading to this phenomenon remain unknown. However, understandings of these mechanisms should be pursued to identify novel therapeutic targets that will not only improve the observed muscle phenotype, but also to ultimately improve bone strength as bones and muscles are highly synergistic tissues that communicate extensively with one another[57–61].

**Bone Biology**

Bone is a highly metabolic and dynamic endocrine tissue that has major roles in locomotion, calcium and phosphate storage, and energy metabolism[62–64]. Bone can be divided into a mineral phase and an organic phase consisting of primarily hydroxyapatite and type I collagen, respectively and is composed of four major cell types: osteoblasts, osteoclasts, osteocytes, and bone lining cells [62,63]. These four cell types facilitate the extensive remodeling necessary for the skeletal adaptation to mechanical strain, fracture repair, and calcium homeostasis[63,64].

Osteoclasts control the first step of the bone remodeling process; reabsorption of bone. Osteoclasts are multinucleated cells coming from a macrophage/hematopoietic lineage,
For bone resorption to occur, there must be proliferation of the osteoclast precursor cells and differentiation of these cells into mature osteoclasts, a process referred to as osteoclastogenesis [65,66]. Osteoclasts then adhere to the cell surface where they will break down the mineral hydroxyapatite component of bone followed by the degradation of the organic collagen matrix, a process facilitated by the secretion of multiple factors including tartrate resistant alkaline phosphatase or TRAP[65,66]. Osteoclast differentiation and activity are regulated by two primary essential factors: macrophage colony stimulating factor (M-CSF) and receptor for activation of nuclear factor kappa B ligand (RANKL)[66]. The process of bone resorption is also highly regulated by the osteoblast and cells from the osteoblast lineage including pre-osteoblasts, osteocytes, and bone-lining cells[67].

Osteoblasts, deriving from the mesenchymal stem cell lineage, facilitate the formation of bone in the remodeling process[63,64,67]. Following bone resorption, osteoblasts are recruited to the resorption site where they will deposit new osteoid onto the bone matrix[63]. This process is largely enabled by expression of Runx2, a master regulator of osteoblast differentiation, and canonical WNT signaling[63]. One gene upregulated by Runx2 is COL1A1, resulting in production of type I collagen, which will be secreted by the osteoblast[63,68]. Expression of COL1A1 in addition to Runx2 signifies the transition of pre-osteoblast to mature osteoblast [63]. In addition to their significant role in bone formation, osteoblasts are also essential regulators of osteoclasts and thus, bone resorption[63,64]. The best characterization of this regulation is observed via osteoblasts production of osteoprotegerin (OPG), a decoy receptor for RANKL, which results in
inhibition of osteoclast differentiation[67]. Other cells of the osteoblastic lineage also play important roles regulating the tightly coordinated bone remodeling process.

Osteocytes and bone lining cells both arise from mature osteoblasts and are key players in bone remodeling[63]. The function of bone lining cells has not been well characterized and little is known about these quiescent cells other than they tend to physically separate the osteoclast from the bone matrix[63]. Upon initiation of bone remodeling, these cells retract, allowing for bone resorption to begin[68]. Osteocytes also arise from osteoblasts, comprise up to 95% of all bone cells, and are known for their roles in orchestrating bone remodeling via control of osteoblasts and osteoclast, as well as in mechano-sensation[69]. Osteocytes are embedded in the bone matrix and have an average half-life of 25 years. Osteocytes possess dendritic processes which aid in communication with other osteocytes, osteoblasts, and osteoclasts[69,70].

Bone remodeling is a highly coordinated and tightly regulated process facilitated by the different bone cells. Due to the extensive remodeling that occurs in bone, it may not come as a surprise that alterations in the molecular pathways underlying these processes can result in bone disease such as osteogenesis imperfecta. As previously stated, ~85% of OI cases arise due to mutations in type I collagen with the remaining cases arising due to mutations in collagen modification and folding, osteoblast differentiation and function, and bone mineralization; all of which are essential to bone remodeling[15] (Table I-1). These mutations lead to changes in several important signaling pathways regulating bone remodeling including the WNT and RANK/RANKL pathways, ultimately resulting in uncoordinated bone remodeling[15].
Skeletal Muscle Biology

With the presence of over 600 muscles, skeletal muscle is the largest organ in the human body, comprising roughly 40-50% of body mass[71–74]. Skeletal muscle participates in many roles including body mechanics, energy metabolism and serves as a secretory organ[71–73]. Here, important aspects of skeletal muscle biology, including development and growth, structure and organization, and mechanical function, will be discussed.

Embryonic Development

Skeletal muscle arises from the paraxial mesoderm that gives rise to somites that eventually diverge into sclerotome and the dermomyotome, giving rise to bone/cartilage and skin/skeletal muscle, respectively[74,75]. Skeletal muscle formation progresses in five major steps including: delamination, migration, proliferation, determination, and differentiation[75]. Cells that give rise to skeletal muscle delaminate from the dermomyotome and migrate to the limb bud, processes facilitated by the factors Pax3, c-met, hepatocyte growth factor (HGF), and Lbx1[75,76]. Once myogenic precursors reach the limb bud, they begin expressing MyoD and Myf5, signifying the commitment of the cells to the myoblast lineage[77]. Evidence suggests that proliferation of myoblast precursor cells occurs before and after determination and is likely facilitated by Pax3 and c-met[77]. Differentiation occurs via the expression of MRF4 and myogenin, which together with MyoD and Myf5 make up the myogenic regulatory transcription factor family (MRFs)[76,77].

Following the final step of differentiation, myoblasts begin to express myosin heavy chains as part of terminal differentiation, at which point myoblasts will fuse to form
multinucleated myotubes[76]. These myotubes will then assemble to form muscle fibers leading to the formation of specific muscle groups[76,77].

Skeletal muscle consists of multinucleated muscle fibers that can be broadly classified as either fast/white/glycolytic or slow/red/oxidative[78]. The expression of different myosin heavy chains is what distinguishes muscle fiber types and their cellular energy metabolism, although skeletal muscles are highly plastic allowing fibers to adapt to the muscles’ activity level and needs[79]. There are four primary adult fiber types (MyHC-β/slow, MyHC-2A, MyHC-2B, and MyHC-2X) and two developmental types (MyHC-emb and MyHC-neo) in skeletal muscle[78]. Type 1, or slow, fibers are red in appearance, have a high oxidative capacity, are slower to fatigue, and have relatively high mitochondrial volume density (2-3 fold higher compared to fast fibers)[80]. Type 2 or fast fibers are white in appearance, have a lower oxidative capacity, fatigue quickly, and have relatively lower mitochondrial volume density[80]. Type 2 fibers can be further broken down to the types 2A, 2B, and 2X based on their oxidative capacities as fast oxidative, fast glycolytic, and fast intermediate; respectively[78].

In the mouse, all skeletal muscles express the embryonic (MyHC-emb) and slow (MyHC-β/slow) myosin heavy chains prior to embryonic day 16 (E16), at which point some fibers will begin to lose their MyHC-β/slow expression, which is replaced by MyHC-neo, giving rise to two distinct populations of fibers expressing either MyHC-emb/MyHC-neo or MyHC-emb/MyHC-β/slow[78]. The fibers expressing MyHC-emb/MyHC-neo or MyHC-emb/MyHC-β/slow will develop into fast or slow fiber types, respectively[78].
Muscle growth and regeneration

While myoblasts arise from the mesenchymal stem cells during development, post-natal muscle growth and repair is facilitated by a population of stem cells referred to as satellite cells. These quiescent cells appear in limbs at embryonic day 17.5 arising from cells of the dermomyotome, similar to myoblasts[81,82]. Satellite cells lie between the basal lamina and the sarcolemma of each myofiber, and are heterogeneous in terms of cell number and Pax7 expression [83–85]. Although satellite cells are able to self-renew, the total number of satellite cells has been shown to decrease overtime with age as the number of myonuclei increase[81,82,86].

Due to the postmitotic nature of muscle fibers, they are not able to divide. They rely on satellite cells to provide the myonuclei required for maintenance of muscle by normal regeneration, hypertrophy, and repair[84]. Satellite cells are identified by their expression of Pax7, an important transcription factor in myogenesis and the canonical satellite cell biomarker [82,84]. Once satellite cells are activated, in response to an environmental signal or damage, they begin to express muscle regulatory factors (MRFs) which are also present during development, including MyoD1, Myf5, and myogenin [81,84]. The activation, differentiation, and migration of these satellite cells ultimately lead to the cells fusing with one another to generate new myofibers or with a damaged muscle fiber for repair [81]. While there are many signals and factors regulating satellite cell function, three stand out that maybe relevant in OI, including WNT/β-catenin signaling, NOTCH signaling, and extracellular matrix composition and stiffness[81]. However, the relationship of these factors to OI satellite cell function has not yet been characterized.
**Structure and organization**

The basic functional unit of skeletal muscle, a sarcomere, consists of the region from one Z-disc to another, as shown in figure I-2. The sarcomere is composed of thin actin and thick myosin filaments that facilitate muscle contraction. The thin actin filaments (F-actin) are composed of actin monomers (G-actin) that exhibit polarity within the sarcomere. Additionally, thin actin filaments contain the proteins troponin and tropomyosin, important regulators of muscle contraction [87]. The thick myosin filament is composed of myosin II, a hexameric protein composed of four light chains and two heavy chains containing a rod and head region[87]. The two globular heads also serve as important facilitators of contraction[87].

Upon visualization of the sarcomere via electron micrograph (Figure I-2), several conserved and obvious structures stand out. The dark banded Z disc or Z band can be seen at either end of the sarcomere adjacent to the light isotropic or I band consisting of thin actin filaments[88]. Between the I-bands lies another dark banded region referred to as the anisotropic or A-band consisting of the thick myosin filaments[88]. In the very center of the sarcomere exists the M-band composed of proteins of the myomesin family serving as cross linkers between the thick myosin filaments and contributing to the hexagonal lattice within the A-band [88–90]. On either the side of the M-band a faint light line can be seen, a result of the absence of myosin heads on this region of the thick filament[89]. The area between these light bands within the A-band is referred to as the H-zone which shortens during contraction[34,88].
Figure I-2 Sarcomere organization. Originally published in Comprehensive Physiology (Henderson et al., 2017)[91]. A) Schematic of sarcomere organization. B) Electron micrograph image of a sarcomere. C) Electron micrograph of the M zone of the sarcomere.
As previously described, the basic structural and functional unit of a muscle is referred to as a sarcomere and is composed of thick myosin and thin actin filaments. Together these myofilaments make up the myofibril. Bundles of myofibrils are surrounded by a plasma membrane denoted as the sarcolemma and make up a muscle fiber. Skeletal muscles possess a highly organized extracellular matrix composed of a hierarchical arrangement of endo-, peri-, and epimysium (Figure I-3)[92]. The endomysium surrounds bundles of muscle fibers referred to as fascicles and bundles of fascicles surrounded by the perimysium make up skeletal muscle which is covered by the epimysium[92]. At the base of a skeletal muscle the epimysium connective tissue converges to the tendon which connects the muscle to bone[93]. The composition of collagen within these various extracellular components differ greatly. Type I collagen is most abundant in both the tendon and perimysium of the skeletal muscle while types I and III collagens are equally abundant in both the epi- and endomysium[92]. Interestingly, the organization of type I collagen differs from tendon and perimysium with the type I collagen in tendon being much more organized, aligning with the muscle axis, than in the perimysium which is likely a result from the different functions of each component [92]. This highly organized hierarchical extracellular matrix surrounding the muscle fibers is believed to help facilitate force transduction with all of the different levels believed to be interconnected mechanically[92,94]. In addition to the collagens found in these extracellular matrix components, type IV collagen networks are highly present in the basement membranes of skeletal muscles with Types VI, XV, and XVIII also present in smaller amounts[92]. Furthermore, while the endomysium and basement membrane are separate entities, they are interconnected with the basement membrane likely playing an important role in force
transduction[92]. In addition to collagen which comprises roughly 1-10% of dry muscle weight [92,95,96], the extracellular matrix consists largely of proteoglycans (such as decorin), glycosaminoglycans, and glycoproteins (including integrins, fibronectin, and laminins) all interacting with one another to facilitate force transduction throughout the ECM of the skeletal muscle[92].
Figure I-3 Skeletal muscle and extracellular matrix organization. Originally published in *Muscle Nerve* (Gillies and Lieber, 2012)[92]. A) Schematic of skeletal muscle extracellular matrix organization. B) Cross section of skeletal muscle to show the continuous nature of the perimysium with the tendon.
**Mechanical function**

As for its role in body mechanics, skeletal muscle contraction provides the power and force necessary for body movement, for maintenance of skeletal structure, body posture, and offers protection of internal organs [97]. Early work done by Huxley and Niedergerke and Huxley and Hanson in two, consecutive, 1954 Nature articles laid the foundation for what would become known as the sliding-filament model, observing that as contraction occurs, actin and myosin filaments do not change size despite the overall shortening of the sarcomere and deducing instead that the actin and myosin filaments were overlapping in an ATP-dependent manor[88,98]. This work laid the foundation leading to our current understanding of muscle contraction.

Muscle contraction is facilitated largely by the thin actin and thick myosin filaments overlapping one another in an ATP-dependent manner in response to a calcium signal[87,99–101]. The process by which muscle contractions occurs in response to this signal is referred to as excitation-contraction coupling[87,97,100]. Muscle contraction begins with the muscle fiber receiving a signal from a motor neuron at the neuromuscular junction which “excites” the muscle via propagation of an action potential terminating at the neuromuscular junction at which point acetylcholine is released into the synaptic cleft where it will bind to its receptor leading to a local depolarization of the muscle fiber and the opening of voltage gated sodium channels. This leads to further/ depolarization of the transverse tubules within the sarcolemma, resulting in the release of calcium from the sarcoplasmic reticulum[97,100]. Skeletal muscle can exist in an off (resting) or on (actively contracting) state. In the resting state, intracellular calcium concentration is less than 100 nM, and when calcium is released from the sarcoplasmic reticulum, the
concentration increases to ~20 uM[100]. This influx of calcium ultimately leads to contraction by exposing the myosin binding sites on the thin actin filaments. Actin contains the proteins troponin and tropomyosin, and in a resting muscle tropomyosin covers myosin binding sites. Calcium will bind the troponin causing a conformational change uncovering the myosin binding sites[87,100,102].

The uncovering of these binding sites leads to muscle contraction via a process referred to as cross-bridge cycling wherein the myosin heads actually appear to walk along the actin filaments via ATP hydrolysis[87,102]. This process exists in four main steps: 1) formation of the cross bridge is facilitated by the binding of myosin to its binding sites on actin, 2) the power stroke of the myosin head occurs releasing ADP and Pi, 3) a new molecule of ATP binds, and 4) ATP is hydrolyzed and the myosin head is cocked and ready to re-form the cross bridge with actin[97,100,102]. Muscle relaxation is facilitated by the dissociation of the myosin heads from actin filaments in addition to the reuptake of calcium by the sarcoplasmic reticulum, both relatively slow processes[100].

**Bone muscle crosstalk**

Osteogenesis imperfecta translates to imperfect bone formation and, as the name literally suggests, is a bone disease. However, as mentioned, 80% of patients with mild OI experience muscle force deficits, suggesting the wide prevalence of muscle weakness in addition to the bone fragility[6,47,48]. It is important to consider the impact that muscle weakness may have on further compromising the bone of OI patients. Until recently, the understanding of the relationship between these two organs has been limited to the context of physical forces and biomechanics[60,61,103]. In this instance, the muscle acts
as a pulley exerting force on the bone, acting as a lever to move the skeleton[103]. The understanding of this relationship has evolved over the past decade, with the current understanding incorporating the idea that both bones and muscles act as secretory organs communicating via biochemical signaling in addition to the historical understanding of mechanotransduction[60,61,103].

**Biomechanical**

The earliest understanding of the relationship between bone and muscle was that of their physical interactions, with early studies suggesting a correlation between muscle weight and bone mass[104]. This idea, initially beginning as Wolf’s law, eventually leading to Frost’s mechanostat theory, postulates that bone strength is tightly coupled to skeletal muscle action[57,105,106]. Significant evidence in the field of musculoskeletal health has corroborated this idea since its initial conceptualization in 1892[106,107]. Countless studies evaluating the consequences of disuse or unloading on these two organs support the notion that muscle force and bone strength are tightly coupled[5,14,103,108–112]. A summary of selected studies highlighting the mechanical relationship of bone and muscle can be seen in Table I-2.

The ability of bone to remodel its geometry and size in response to the mechanical strain from muscle is primarily facilitated by the osteocyte, a mechanosensory cell[69]. For mechanotransduction to occur, the osteocyte must perceive a mechanical signal then convert it to a biochemical signal to elicit a response by osteoblasts and/or osteoclasts to facilitate bone remodeling[69,113–115]. The exact mechanisms by which the osteocytes act as a mechanosensing cells as well as the components of the osteocyte that facilitate
mechanosensing, remain unknown, although there is evidence that the dendritic processes, cell body, and cilium contribute to the osteocytes’ ability to function as a mechanosensory cell[69,114]. It is hypothesized that the osteocyte can respond to several factors in response to mechanical strain, including the physical deformation of the bone matrix, fluid flow shear stress, and streaming potentials[69,114,116].
<table>
<thead>
<tr>
<th>Reference (Author)</th>
<th>Title</th>
<th>Area of Research</th>
<th>Significant Findings</th>
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<tbody>
<tr>
<td>Scheld K. et al., 2001</td>
<td>Nitrogen metabolism and bone metabolism markers in healthy adults during 16 weeks of bed rest</td>
<td>Bed Rest</td>
<td>Within 2 weeks of bed rest in healthy adults, bone metabolic marker, osteocalcin was decreased suggesting reduced bone formation. Additionally, body nitrogen was decreased within the initial 2 weeks of bed rest which was associated with a rapid decline in muscle mass[117].</td>
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<td>Lang T. et al., 2004</td>
<td>Cortical and trabecular bone mineral loss from the spine and hip in long-duration spaceflight</td>
<td>Microgravity</td>
<td>Areal bone mineral density was reduced in both spine and hip at rates of approximately 0.9% and 1.4%, respectively in 14 members aboard the International Space Station that were followed during their 4 to 6 month flights[118].</td>
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<tr>
<td>Novotny SA. et al., 2011</td>
<td>Bone is functionally impaired in dystrophic mice but less so than skeletal muscle</td>
<td>Musculoskeletal Disease</td>
<td>Mdx (dystrophin deficient) and dko (utrophin deficient) mice exhibit altered bone geometry and reduced bone material properties including 50% reduction in bone strength compared to WT littermates[119].</td>
</tr>
<tr>
<td>Tagliaferri C. et al, 2015</td>
<td>Muscle and bone, two interconnected tissues</td>
<td>Bed Rest, Aging, Microgravity</td>
<td>A comprehensive review discussing the consequences of reduced mechanical loading providing a case that mechanical loading links bone and muscle functions[120].</td>
</tr>
<tr>
<td>Yuan Y. et al., 2016</td>
<td>The roles of exercise in bone remodeling and in prevention and treatment of osteoporosis</td>
<td>Aging and Exercise</td>
<td>This thorough review summarizes the effects that exercise can have on different types of bone cells in terms of their mechanical and signaling responses to exercise as well as how this translates prevention of age-related effects of disuse such as osteoporosis[121].</td>
</tr>
<tr>
<td>Veilleux LN. &amp; Rauch F., 2017</td>
<td>Muscle-Bone Interactions in Pediatric Bone Diseases</td>
<td>Musculoskeletal Disease</td>
<td>This review highlights the clinical perspectives in regards to pediatric patients on the roles of bone in muscle disease and the role of muscle in bone disease[5].</td>
</tr>
<tr>
<td>Bettis T., Kim B-J., and Hamrick MW., 2018</td>
<td>Impact of muscle atrophy on bone metabolism and bone strength: implications for muscle-bone crosstalk with aging and disuse</td>
<td>Aging, Bed Rest, and Microgravity</td>
<td>This review offers a comprehensive overview of bone muscle crosstalk including both biomechanical and biochemical examples in clinical cases and animal model studies focused on the effects of inactivity or muscle disuse specifically in instances of aging, bed rest, and microgravity[111].</td>
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In response to a mechanical stimulus, intracellular calcium levels increase and the osteocyte begins to regulate and integrate osteoblasts and osteoclasts to control bone remodeling[114]. To respond to mechanical stress, the osteocyte releases signals including nitrous oxide, prostaglandins, and ATP, which are believed to have direct effects on osteoblasts and osteoclasts, in response to this increased intracellular calcium[114]. One of the most important signaling pathways modulated by the osteocyte’s response to mechanical stress is the Wnt/β-catenin pathway[122,123].

A summary of osteocyte signaling on osteoblasts and osteoclasts in response to mechanical stimuli can be seen in Figure I-4[114]. Briefly, prostaglandins, nitrous oxide, and ATP have a direct positive effect on bone formation, while nitrous oxide has a negative effect on osteoclasts, resulting in a net gain of bone[114]. Furthermore, members of the Wnt/β-catenin signaling pathway, and thus bone formation, are inhibited via sclerostin, dickkopf-1 (DKK1), and selected frizzled-related protein 1 (sFRP1)[114,124]. Receptor activator of nuclear kappa-B ligand (RANKL), another important regulator in bone remodeling, activates the osteoclast resulting in increased bone resorption whereas osteoprotegerin (OPG) inhibits RANKL resulting in increased bone formation[114,124]. Finally, macrophage colony stimulating factor (M-CSF) regulates differentiation of osteoclasts from hematopoietic stem cells[124]. Importantly, several of these signaling molecules serve as targets of novel therapies being investigated in OI including the use of anti-sclerostin and RANKL inhibitor antibodies[15].
Figure I-4 Osteocyte regulation of bone remodeling. Originally published in *Endocrine Reviews* (Dallas et al., 2013)[114]. The osteocytes secrete factors that act on osteoblast and osteoclast precursors to modulate bone remodeling.

Fifty percent of peak adult bone mass is accrued during adolescence and there is strong evidence that 20-40% of adult peak bone mass is determined by modifiable factors including exercise, further reiterating the importance of exercise in children with OI[125,126]. Studies have demonstrated that children with OI have activity levels comparable to those of their healthy peers, and there is evidence that increased physical activity may also improve aerobic capacity and muscle strength[47,115,127,128]. The use of exercise and whole body vibration therapy in OI as modalities of treatment to exploit the biomechanical relationship between bone and muscle have also been investigated[14].

Whole body vibration therapy (WBV) has also been implemented as treatment in OI and is an especially attractive target for children with severe OI and lower ambulation and ability to participate in traditional physical activities [129–131]. Studies in the *oit* mouse evaluating the use of WBV demonstrated improvement of several bone parameters including increased cortical bone area, cortical thickness, and trabecular bone volume while no significant changes in bone mechanical properties were observed [130]. Limited
studies on the use of WBV in OI exist and the results from these studies are conflicting with one study by Hoyer-Kuhn et al. demonstrating improved bone mineral density and motor function in patients with OI while another study by Högler et al. demonstrated increased lean mass, but no changes in bone mass or muscle function[129,132]. This discrepancy and inability of OI patients to respond to WBV may be the result of genetic mutations specificity or differing levels of OI severity or may reflect a reduced biomechanical responsiveness of OI bone[132]. Regardless, the use of exercise and WBV as therapeutic approaches in OI are attractive targets that are minimally invasive and cost-effective ways to exploit the biomechanical relationship of bone and muscle.

Biochemical

Until recently, the nature of bone-muscle crosstalk was thought to be limited to the biomechanical relationship. However, more recent research has highlighted the presence of extensive biochemical crosstalk that occurs with both bone and muscle acting as secretory and endocrine organs[60,61,133]. One of the earliest reports suggesting biochemical crosstalk between muscle and bone, specifically a positive effect of muscle tissue on osteogenesis independent of contraction, demonstrated improved fracture healing (increased speed and quality) when an open tibial fracture was wrapped with a muscle flap relative to non-wrapped tibias in a canine model[134]. This has been reproduced in rodent models by showing improved tibia open fracture healing in the presence of a muscle flap compared to a fasciocutaneous flaps[135]. Since these findings, the field of musculoskeletal interaction research has grown exponentially, with roughly 600 articles published in 2019 [136].
Proposed musculoskeletal biochemical interactions can be summarized into three main mechanisms[136]. The first two mechanisms suggest that both bone and muscle secrete soluble factors into the blood stream due to their highly vascularized nature[136]. The third mechanism consists of direct cell-to-cell contact; due to the close proximity of bone and muscle, it is likely that small molecules are able to be directly signaled between the two organs via extracellular vesicles or passive diffusion [136]. These factors secreted by muscle and bone are referred to as myokines and osteokines, respectively.

Myostatin, discovered in 1997, was the first and remains the best characterized myokine, although the term myokine was not coined until 2003, by Pedersen et al[133,136–138]. Since then, over 3,000 myokines have been identified[139]. Additionally, there is evidence that different myokines may be preferentially secreted by certain types of muscle fibers (i.e. glycolytic vs. oxidative)[139,140]. Some of the most prominent myokines, especially in the context of muscle-bone crosstalk include myostatin (mstn), interleukin-6 (IL-6), irisin, and β-aminoisobutyric acid (BAIBA) [14,61,136,140]. Just as muscle acts as a secretory organ releasing myokines that will act on bone, bone reciprocates this action, releasing osteokines that act on muscle including osteoclacin, Wnt3a, TGFβ, and sclerostin[14,136]. The role of these myokines and osteokines are summarized in Tables I-3 and 1-4, respectively. Several examples of altered biochemical signaling pathways in OI muscle and bone have been described and novel therapeutics have emerged acting on these altered pathways.
### Table I-3: Myokines and their roles on bone

<table>
<thead>
<tr>
<th>Myokine</th>
<th>Role in bone</th>
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<tr>
<td>Myostatin (mstn)</td>
<td>A member of the TGF-β superfamily and negative regulator of muscle mass. In addition to its inhibitory effect on muscle growth, mstn has been repeatedly shown to have a negative impact on bone formation via increased osteoclastogenesis, increased osteocyte expression of negative regulators of bone including sclerostin, and reduced expression of osteoblast differentiation markers[133,136,138,141–143].</td>
</tr>
<tr>
<td>Interleukin-6 (IL-6)</td>
<td>Although secreted by multiple tissues, large amounts of IL-6 are secreted by muscle in response to exercise leading to its characterization as a myokine[137]. Interestingly, IL-6 has been shown to upregulate both osteoclast and osteoblast formation[136,144,145].</td>
</tr>
<tr>
<td>Irisin</td>
<td>Irisin has been shown to be positively correlated with BMD and negatively correlated with serum sclerostin[146,147]. Additionally, in mice, treatment with recombinant irisin led to improved bone geometry and mechanical properties[148].</td>
</tr>
<tr>
<td>β-aminoisobutyric acid (BAIBA)</td>
<td>BAIBA, a myokine secreted in response to exercise was originally identified for its function in the browning of white adipose tissue though more recent studies have demonstrated a role in osteocytes as a protective agent against reactive oxygen species[149,150].</td>
</tr>
</tbody>
</table>

### Table I-4: Osteokines and their roles on muscle

<table>
<thead>
<tr>
<th>Osteokine</th>
<th>Role in muscle</th>
</tr>
</thead>
<tbody>
<tr>
<td>Osteocalcin (OCN)</td>
<td>OCN, secreted by the osteoblast plays an important role in glucose and energy homeostasis and is believed to promote nutrient catabolism and uptake in skeletal muscle as well as improve exercise capacity[151–153].</td>
</tr>
<tr>
<td>Wnt3a</td>
<td>Wnt3a, secreted by osteocytes, was shown to promote myogenesis via increased myogenin and myoD expression in a cultured osteocyte cell line (C2C12) [154,155].</td>
</tr>
<tr>
<td>Transforming growth factor-β (TGF-β)</td>
<td>TGF-β, an osteokine secreted by osteoblasts has been shown to negatively impact skeletal muscle function via calcium leakage and increased oxidative stress[156,157].</td>
</tr>
<tr>
<td>Sclerostin</td>
<td>Although there is not evidence of sclerostin, secreted by the osteocyte, directly effecting muscle; it has been shown to inhibit Wnt-3a action in skeletal muscle, thus indirectly negatively impacting muscle[154,158,159].</td>
</tr>
</tbody>
</table>
The role of these osteokines in osteogenesis imperfecta have been well characterized, while less information exists regarding the role of different myokines in the pathology of OI. Of the myokines, myostatin has been the most studied in the context of OI. Myostatin is a negative regulator of muscle mass that signals through the activin receptor type IIB to inhibit muscle growth[160]. Several studies in the oim and +/G610C mouse models have highlighted the potential for myostatin inhibition to have a beneficial outcome on bone properties. The first report highlighting the benefit of myostatin (mstn) inhibition demonstrated that +/oim mice, modeling a mild to moderate patient OI, when also deficient in mstn (+/oim +/-mstn), exhibited improved femoral bone biomechanical properties compared to their +/oim littermates[161]. Other data suggest that the responsiveness to myostatin inhibition is OI mutation-specific [162,163]. Studies using the activin receptor IIB decoy molecule resulted in improved skeletal properties in the +/G610C mouse, modeling a mild to moderate human OI, and improved skeletal muscle contractile properties in the oim/oim mouse[162,163]. These studies suggest that myostatin may serve as a novel therapeutic target in the treatment of OI, although responses may be based on specific mutations.

As indicated above, relative to the myokines, more data exists on osteokines in OI. Osteocalcin levels are elevated in both patients and the Col1a1+/Jrt OI mouse model [8,164,165]. Additionally, dysregulated TGF-β signaling has been observed in OI mouse models with increased signaling observed in both recessive (Crtatp-/-) and autosomal dominant (+/G610C and Col1a1+/Jrt) variants of the disease[15,22,166]. However, treatment with a TGF-β neutralizing antibody only improved bone properties in two of these models (Crtap-/- and Col1a1+/Jrt) suggesting the importance of mutation specific
treatment approaches[15,22,166]. Clinical trials in OI patients are also ongoing using TGF-β antibody[2]. Finally, there is not a clear consensus on whether or not the levels of sclerostin are altered in OI patients, although there is evidence of improved bone properties in several mouse models of OI (oim/oim[24], +/-G610C, Brlt-/-[167], Col1a1+/+m[168], and Crtap-/-[169]) and patient clinical trials utilizing sclerostin antibody are ongoing, with initial results showing increased markers of bone formation and reduced markers of bone resorption [2,14,15,170,171].

The relationship between bone and muscle is highly synergistic, exhibiting both biomechanical and biochemical crosstalk. Biomechanical and biochemical crosstalk are not mutually exclusive. The role of myostatin in bone muscle crosstalk highlights this point. Inhibition of the myokine, myostatin, leads to altered mechanotransduction via increased muscle size; in addition this inhibition has a direct effect on biochemical signaling leading to a downregulation of osteoclasts. Understanding the complexity of bone muscle crosstalk in OI is crucial to determining novel therapies that can exploit this relationship by improving muscle function to ultimately improve bone strength.

**Skeletal muscle pathology in disease**

Skeletal muscle weakness arises as part of the pathology of cancer cachexia, numerous congenital myopathies and neuromuscular diseases, and with aging[172–175]. Much research has been compiled on each of these individual instances, but there is not a single mechanism that leads to muscle weakness. Furthermore, the need for research in OI skeletal muscle weakness is apparent due to the lack of studies aiming to understand skeletal muscle weakness mechanisms in OI. Thus, it is necessary to review the
mechanisms of muscle weakness in other diseases to get an understanding of how muscle weakness might arise in OI.

One field with extensive research is that of the muscular dystrophies including Duchenne and Becker Muscular dystrophies, both arising from mutations in the dystrophin gene, as well as other congenital muscular dystrophies which include Ullrich syndrome and Bethlem myopathy, arising from mutations in the type VI collagen genes[175]. In Duchenne and Becker muscular dystrophies, the absent or highly reduced amount of dystrophin disrupts skeletal muscle organization, sarcolemma fragility, and abnormal calcium homeostasis [176–178]. Excessive calcium and damaged mitochondria have been observed in the dystrophic muscle with damaged mitochondria further perpetuating calcium dysregulation, ultimately resulting in increased damage of mitochondria in a feed-forward fashion[176]. Similarly, the mutations in type VI collagen genes can lead to disruptions in the extracellular matrix (specifically the perimysium and endomysium [179]) and increased apoptosis via mitochondrial dysfunction[180,181]. Mitochondrial dysfunction has also been implicated in aging and neurodegenerative disorders[182,183], so while there is no single mechanism that leads to skeletal muscle weakness, mitochondrial dysfunction appears a common theme amongst these different instances of skeletal muscle weakness.

**Mitochondria: the powerhouse of the cell?**

Mitochondria are commonly and simply referred to as the powerhouse of the cell, and although they were originally known for their role in oxidative phosphorylation and ATP production, mitochondria are also crucial in other cellular functions including calcium
homeostasis, cell signaling, ER stress, autophagy, apoptosis, and others[184].

Mitochondria are double membrane-bound organelles that possess their own circular, double stranded DNA encoding members of the electron transport chain necessary for oxidative phosphorylation[185,186]. Several parameters dictate mitochondrial function and health including mitochondrial dynamics and morphology as well as mitochondrial biogenesis and mitophagy[184,186,187].

A textbook image of a mitochondrion often highlights its spherical shape with a well-defined outer and inner membrane, as well as the presence of membrane invaginations known as cristae. However, a mitochondrion’s morphology is more complex and is regulated by the dynamic nature facilitated by mitochondrial fusion and fission in response to a cell’s unique needs[188]. Because mitochondrial morphology and function are tightly coordinated, mitochondria can exhibit unique morphology from cell to cell with mitochondria in skeletal muscle consisting of a network of interconnected mitochondria, although recent studies suggest mitochondrial morphology in skeletal muscle may be more complex than this, with the network and interconnectedness varying across the muscle[184,186,189].

Mitochondrial fusion is the union of two mitochondria, often allowing for the combination of two smaller organelles to ultimately increase the size of the mitochondrial network, and is facilitated by the proteins mitofusin 1 and 2, as well as optical atrophy 1 and 2[184,186]. Mitochondrial fission is the fragmentation of mitochondria often in response to damage and is facilitated by the proteins dynamin-related protein 1, mitochondrial fission factor, and fission protein 1[184,186].

Mitochondrial biogenesis and mitophagy are coordinated with the mitochondrial dynamics and also play critical roles in overall mitochondrial health by controlling
production of new mitochondria and mitochondrial clearance, respectively[186,187]. The process of mitochondrial biogenesis and mitophagy involve complex regulation via different signaling pathways. Increased intracellular calcium is one of the most common activators of mitochondrial biogenesis, leading to activation of AMP-activated protein kinase and ultimately PGC1-α whom is considered the “master regulator” of mitochondrial biogenesis[187]. Briefly, activation of PGC1-α along with its co-activators NRF-1 and NRF-2 lead to activation of TFAM, a mitochondrial transcription factor leading to mitochondrial DNA transcription and replication[190]. Mitophagy is a form of selective autophagy often initiated in response to damaged or dysfunctional mitochondria[187,191]. Two major pathways of mitophagy are the PINK-1/Parkin ubiquitin pathway and the BNIP3/NIX pathway[187,191]. In the PINK-1/Parkin dependent pathway, PINK-1 is recruited to a depolarized mitochondrial membrane at which point Parkin, an E3 ubiquitin ligase, is recruited ultimately leading to the mitophagy[191]. BNIP3 and NIX are both atypical members of the BCL2 family, a group of proteins regulating cell death[191]. These proteins through there interaction with LC3, an important component of autophagosome formation, lead to degradation of the mitochondria[191].

As previously discussed, the roles of mitochondria in cell health and maintenance are diverse. Briefly, their well-known roles in ATP production and calcium homeostasis will be discussed. The mitochondrial membrane is the site where the respiratory chain carries out oxidative phosphorylation, the process of electron transport ultimately leading to ATP production[192]. The process of oxidative phosphorylation includes the transfer of electrons through complexes I-IV of the electron transport chain with O₂ as the terminal
electron acceptor. During this process protons are pumped across the membrane generating an electro-chemical gradient which is used to drive ATP synthesis via complex V (ATP synthetase) (Figure I-5) [192]. Calcium signaling is an important regulator of many different cellular processes and the mitochondria aid in regulating calcium homeostasis by serving as a reservoir for taking up extracellular calcium, although excessive calcium influx can lead to mitochondrial damage and dysfunction[193].

In skeletal muscle, three distinct populations of mitochondria exist: the intermyofibrillar mitochondria, subsarcolemmal mitochondria, and a third less characterized yet distinct population extending from the subsarcolemmal mitochondria[194]. These distinct populations differ in terms of location, morphology, and function and have been shown to be differentially affected in disease states[186,194]. The subsarcolemmal mitochondria lie beneath the mitochondrial membrane and are believed to play roles in ATP generation necessary for membrane transport whereas the intermyofibrillar mitochondria are located near Z-lines in skeletal muscle and are believed to generate ATP necessary for contraction and regulate calcium signaling[186]. Due to their diverse cellular roles, regulation of their distinct mitochondrial content and function is essential for skeletal muscle health.
Figure I-5 Oxidative Phosphorylation. Originally published in Genetics (van der Bliek et al., 2017)[192]. Electrons are shuttled through electron transport chain complexes I-IV to oxygen, the final electron acceptor. This process generates a proton gradient that drives ATP synthesis via protonmotive force.
Links between collagen and mitochondrial function

While there is no clear link between type I collagen and mitochondrial function, a link between type VI collagen and muscle mitochondrial function can be seen in cases of Bethlem myopathy and Ullrich Congenital Muscular dystrophy[180,181,195]. In these cases, type VI collagen defects result in a mitochondrial myopathy believed to be the result of induction of mitochondrial permeability transition (the rapid increase of inner membrane permeability to solutes[196]) due to the opening of the mitochondrial permeability transition pore (mPTP), a high conductance channel whose components and structure remain to be elucidated[180,197]. Under normal circumstances, the mPTP functions in the “direction” of ATP synthesis with occasional transient opening facilitating regulated apoptosis[198]. However, prolonged opening, the result of excessive calcium stress, reactive oxygen species or a combination of the two, can cause dissipation of the proton gradient and uncoupling of oxidative phosphorylation leading to mitochondrial dysfunction[198]. In animal studies and clinical trials, cyclosporine A, an inhibitor cyclophilin D (an activator of the mPTP) has shown promising results improving the mitochondrial dysfunction associated with type VI collagen defects by inhibiting mitochondrial permeability transition [181,199–202].

While there is no clear link between type I collagen and mitochondrial function, there is limited evidence of interactions between type I and type VI collagen, which may imply an indirect relationship between type I collagen and mitochondrial function. An early report demonstrates the binding of type I collagen to the α3 chain of type VI collagen in chicken connective tissue[203]. Furthermore, an in silico analysis of the mouse α3 chain of type VI collagen (Col6a3) on the STRING: functional protein association network
database reveals that the type I collagen genes, \textit{Col1a1} and \textit{Col1a2}, are amongst the top five predicted interaction partners (https://string-db.org). Type VI collagen is an important extracellular matrix protein found in many diverse tissues including skeletal muscle[195]. In skeletal muscle, the extracellular matrix plays important role in maintaining structure and lateral force transmission throughout the muscle[204]. The collagen network of the perimysium is composed of mainly type I collagen with types III, VI, and XII present in lesser amounts[205]. Defects in type I collagen may lead to destabilization of the perimysial collagen bundles and ultimately the perimysial junction plate, a region important for lateral force transmission where the perimysial collagen bundles interact with the endomysium[205]. Moreover a recent study of teriparatide (a bone anabolic) treatment in OI patients and post-menopausal women revealed increased presence of collagen biomarkers including types II, III, IV, V, and VI in OI patients, but not in post-menopausal women[206]. This further suggests the possibility of dysregulated extracellular matrix interactions as part of the muscle pathology in OI.

\textbf{Research Goals and Objectives}

Skeletal muscle weakness is a prominent manifestation in OI with over 80% of mild patients exhibiting muscle force deficits and the prevalence of muscle weakness correlating with severity of disease. The wide occurrence of skeletal muscle weakness in OI, limited studies aimed at understanding this weakness, and lack of therapeutic options targeting skeletal muscle weakness in OI demonstrate the need for further research evaluating muscle function and mechanisms of skeletal muscle weakness in OI. The goal of this work was to begin to develop an understanding of the skeletal muscle weakness observed in OI using the \textit{oim/oim} mouse, modeling severe human type III OI, to evaluate
mechanisms underlying skeletal muscle weakness. We focused our studies on the role of mitochondrial health, hypothesizing that abnormal mitochondria in the oim/oim skeletal muscle would adversely impact skeletal muscle function.

The first aim of my study, described in chapter II, was to characterize mitochondrial content and function in oim/oim mouse skeletal muscle. Prior to this work, citrate synthase activity was determined to be increased in oim/oim whole gastrocnemius muscle relative to WT littermates suggesting an increase in mitochondrial content. Here, we demonstrate that mitochondrial function is significantly reduced in isolated oim/oim gastrocnemius mitochondria evidenced by significantly reduced mitochondrial respiration rates and significantly reduced mitochondrial citrate synthase activity. Furthermore, we found glycogen content to be equivalent between oim/oim and WT gastrocnemius muscles, further indicating that reduced mitochondrial function rather than another mechanism, such as changes in substrate preferences, was likely contributing to skeletal muscle weakness. Additionally, we observed reduced mitochondrial DNA to nuclear DNA ratios, increased protein content of markers of mitochondrial biogenesis (PGC1-α and TFAM), and reduced protein content of LC3, an important marker of mitophagy in the skeletal muscle of oim/oim mice relative to WT littermates. This study was the first to characterize mitochondrial function in a mouse model of OI and will hopefully stimulate new research in the field regarding the role of mitochondria in OI skeletal muscle weakness.

In chapter III, I describe the second aim, the further characterization of mitochondrial function in liver and cardiac muscle in addition to the evaluation of several parameters of whole body and skeletal muscle metabolism in the oim/oim mouse. One of the most
significant findings of this study was that mitochondrial dysfunction in the oim/oim mouse is not global, as the mitochondrial dysfunction was present in oim/oim gastrocnemius mitochondria, but not in liver or heart mitochondria. We also observed reduced complete and incomplete fatty acid oxidation in oim/oim gastrocnemius mitochondria, but not in oim/oim whole quadriceps muscle, further confirming the presence of a mitochondrial dysfunction and suggesting a compensatory effect in the whole muscle. Moreover, we noted increased whole-body energy expenditures consistent with previous reports in the Col1a1^RT/+ mouse and OI patients. We also saw reduced production of H₂O₂ in oim/oim gastrocnemius mitochondria and a change in soleus muscle fiber type distributions with a decrease in the percentage of slow twitch type I fibers and an increase in the percentage of fast twitch type IIa fibers in oim/oim mice relative to WT. Completion of this study further characterized skeletal muscle specific mitochondrial dysfunction in the oim/om mouse and demonstrated the potential of a metabolic phenotype in the oim/oim mouse.

Finally, chapter IV provides a brief summary of the work presented in this dissertation followed by three primary objectives that are necessary to move the field forward.

The aims of this research were to 1) characterize mitochondrial function in the hindlimb muscles of a mouse modeling a severe human type III OI (oim/oim), 2) determine if mitochondrial dysfunction is a global phenomenon in the oim/oim mouse, and 3) determine if whole body and skeletal muscle metabolic parameters are altered in the oim/oim mouse. To this end, we discovered that 1) oim/oim mice exhibit severe skeletal muscle mitochondrial dysfunction evidenced by drastic reductions in mitochondrial respiration and mitochondrial citrate synthase activity, 2) this mitochondrial dysfunction
is not global as reduced mitochondrial respiration is not observed in heart or liver tissue, and finally, 3) a subtle metabolic phenotype is present in the oim/oim mice including the presence of altered body composition and increased energy expenditure.

This work contributes to the understanding of muscle weakness in OI by beginning to investigate the role of mitochondrial function in the oim/oim mouse. These results should stimulate further research into mechanisms of muscle weakness in patients and mouse models of OI highlighting the important need for future research into mitochondrial health and metabolism in OI. Elucidation of these mechanisms could potentiate novel therapeutic targets aimed at improving skeletal muscle function which, due to the highly synergistic nature of muscle and bone, would ultimately lead to improved bone strength.
CHAPTER II

COMPROMISED EXERCISE CAPACITY AND MITOCHONDRIAL DYSFUNCTION IN THE OSTEOGENESIS IMPERFECTA MURINE (OIM) MOUSE MODEL

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Abstract

Osteogenesis imperfecta (OI) is a heritable connective tissue disorder that most often arises from type I collagen, COL1A1 and COL1A2, gene defects leading to skeletal fragility, short stature, blue-gray sclera, and muscle weakness. Relative to the skeletal fragility, muscle weakness is much less understood. Recent investigations into OI muscle weakness in both patients and mouse models have revealed the presence of an inherent muscle pathology. Understanding the mechanisms responsible for OI muscle weakness is critical, particularly in light of the extensive cross-talk between muscle and bone via mecanostransduction and biochemical signaling. In the following study we initially subjected wild-type (WT) and oim/oim mice, modeling severe human OI type III, to either weight-bearing (voluntary wheel running) or non-weight bearing (swimming) exercise regimens as a modality to improve muscle strength and ultimately bone strength. The oim/oim mice ran only 35-42% of the distance ran by age and sex-matched WT mice and exhibited little improvement with either exercise regimen. Upon further investigation, we determined that oim/oim gastrocnemius muscle exhibited severe mitochondrial dysfunction as characterized by 52-65% decrease in mitochondrial respiration rates, alterations in markers of mitochondrial biogenesis, mitophagy, and the electron transport chain components, as well as decreased mitochondrial citrate synthase activity, relative to age and sex-matched WT gastrocnemius muscle. Thus, mitochondrial dysfunction in the oim/oim mouse likely contributes to the compromised muscle function and reduced physical activity levels.
Introduction

Osteogenesis imperfecta (OI) is a heritable connective tissue disorder characterized most often by bone fragility, short stature, blue-gray sclera, and muscle weakness[1,17]. OI is a heterogeneous disease, both in phenotype and genotype that ranges in severity and can be classified into four main types: I-IV, and at least nine subtypes[1,13]. Type II is perinatal lethal; type III is the most severe viable form with patients often non-ambulatory, while patients with Types I and IV experience more mild and moderate manifestations, respectively. Roughly 85% of OI cases are the result of autosomal dominant mutations in the genes that encode the alpha chains of type I collagen (COL1A1 and COL1A2), while the remaining cases are due to rare mutations of genes primarily involved in type I collagen post-translational modifications and folding, bone mineralization, and osteoblast differentiation[1,17].

In addition to the skeletal fragility, patients with OI often experience muscle weakness and exhibit reduced cardiopulmonary fitness[46,48,207]. Both muscle weakness and reduced cardiopulmonary fitness in OI are less characterized compared to other manifestations, and whether they reflect an inherent pathology or are the result of inactivity is only beginning to be elucidated[115,208]. Recent investigations have begun to unveil the nature of the OI muscle weakness in both patients and mouse models, including recognition that approximately 80% of patients with type I OI due to mutations in the type I collagen genes exhibit muscle force deficits[4,6].

In humans, peak bone mass, the amount of bone accrued at the end of skeletal maturity, is achieved at or around the age of 30 with dramatic accumulation of bone during the pre-
pubertal growth period and then gradual decline initiating around 50 years of age[209]. Bone mass and size are largely attributed to heritable factors, as well as environmental factors, such as activity levels and hormonal and nutritional status. Currently, there is no cure for OI, and treatment options for OI patients are limited primarily to anti-resorptive bisphosphonate therapy and surgical rodding[17,210]. Although the contribution of environmental factors to bone strength are not as great as genetic factors, they are postnatally modifiable. Specifically, physical exercise has been suggested as one of the safest therapeutic options to enhance bone strength due to the mechanosensitive characteristics of bone. Several studies have demonstrated increases in bone mineral density (BMD) and bone mineral content (BMC) in more physically active children or children who underwent an exercise regimen[211,212]. Increased BMD has been attributed partly to increases in muscle mass and strength where muscle and bone mass are positively associated. Van Brussel et al. demonstrated improved aerobic capacity and muscle force in OI patients following a physical training regimen which suggests that exercise may be a viable treatment option for patients with mild and moderate OI[115].

In young and mature rodents, treadmill/voluntary wheel running exercise improved BMD, bone microarchitecture and biomechanical strength[213–218], and was protective against estrogen deficiency induced bone loss[216,217]. Non-weight bearing swimming exercise also prevented bone loss associated with ovariectomy or hindlimb suspension[219–221]. In the following study we investigated the effect of weight bearing (voluntary wheel running) and non-weight bearing (swimming) physical activity on musculoskeletal mass and strength in wildtype (WT) and homozygous osteogenesis
imperfecta murine (oim/oim) mice. Gentry et al. was the first to identify an intrinsic muscle pathology in OI by demonstrating that oim/oim mice have reduced specific muscle contractile function (described as the peak tetanic force normalized to myofiber cross sectional area) as compared to WT littermates[3]. The oim/oim mice model moderately severe human OI type III due to a nucleotide deletion in the Col1a2 gene leading to a frameshift resulting in non-functional α2(I) chains and the formation of homotrimeric type I collagen, α1(I)3, rather than the normal heterotrimeric type I collagen, [α1(I)2 α2(I)][39].

We hypothesized that wildtype and oim/oim mice that underwent 1) voluntary wheel running would exhibit increased skeletal mass and strength (ultimate force) as previously seen in C57BL mice[214], and 2) a swimming regimen would exhibit increased skeletal stiffness and sheer modulus of elasticity, with less of an impact on ultimate force[222,223]. We demonstrate that the present exercise regimens were unable to improve either muscle or bone strength in the oim/oim mice compared to their in-cage control oim/oim mice, even though WT mice demonstrated a positive correlation between distances ran and bone strength.

In addition to the severe bone phenotype, oim/oim mice exhibit decreased activity levels, muscle weights, reduced contractile force and elevated skeletal muscle citrate synthase activity compared to age and sex matched WT littermates[3,163]. The presence of altered citrate synthase activity and the recent report of altered energy metabolism in another moderately-severe OI mouse model, Col1a1<sup>Ind/Ind</sup>, which in younger mice exhibited reduced physical activity levels with increased oxygen consumption and CO<sub>2</sub> production concomitant with increased whole body energy expenditure, and altered energy homeostasis[8], led us to investigate skeletal muscle mitochondrial bioenergetics in the
In the second half of this study, we demonstrate that oim/oim mouse skeletal muscle has poor mitochondrial respiration, decreased mitochondrial citrate synthase activity, and decreased mitochondrial DNA content in addition to alterations in mitochondrial electron transport chain (ETC) complex IV, markers of mitochondrial biogenesis, and markers of autophagy and mitophagy, suggesting a potential and critical role of mitochondria in the muscle weakness associated with OI.

**Methods**

**Animals**

Col1a2<sup>oim</sup> mice (stock #001815) are publicly available from Jackson Laboratory (Bar Harbor, ME, USA). All mice were maintained on the C57BL/6J background and genotyped as previously described[39,224]. Mice were housed in an AAALAC-accredited facility at the University of Missouri, and all experimental manipulations were performed under an approved University of Missouri Animal Care and Use protocol.

**Exercise regimen**

Four week old male and female WT and oim/oim mice were randomly divided into swimming exercise, volunteer wheel running, or control (non-exercise in-cage activity) groups. At six weeks of age, mice in the swimming exercise group were acclimated to the swimming exercise protocol in the first 3 weeks as described in Table II-S1, and then full 60 minutes (3 sessions of 20 minutes of swimming exercise with 3 minutes of rest in between each session) of swimming exercise was maintained. Four to five week old mice in the volunteer wheel running group were housed individually in cages equipped with a running wheel [Kaytee Silent Spinner, 4.5inch diameter,136g (wheel weight alone; 75g),
Kaytee Products, Inc, Walnut Creek, CA] assembled with cyclocomputer (SpeedZone® Comp, Specialized Bicycle Components Inc., Morgan Hill, CA) for recording max speed (km/h), average speed (km/h), distance (km), running time (hr), and odometer reading (km) daily. Mice were monitored daily for evidence of adverse effects of exercise [injury, lameness, weight loss, poor grooming, and change in behavior (major change in time on the wheel)]. At 4 months of age, mice were euthanized and hindlimb muscles and femurs evaluated.

**Contractile Properties**

Contractile properties of the soleus (sol), plantaris (plant), gastrocnemius (gast), and tibialis anterior (TA) muscles in male and female WT and oim/oim mice were evaluated as previously described. Briefly, mice were anesthetized and the left sol, plant, gast, and TA muscle surgically exposed at their distal insertions. The distal tendon of each muscle was attached to the Grass force transducer and sequentially tested, sol→plant→gast→TA. The distal tendon was adjusted in length so that the passive tension was zero grams. The sciatic nerve was isolated and placed on a stimulating electrode and a twitch was obtained as previously described[3]. At optimal length, a peak tetanic contraction (P_o) was elicited by pulses delivered at 150Hz, 300-ms duration, and an intensity of 6V for each type muscle[225]. Previous studies of force curves generated at 15, 50, 75, 100 and 125Hz (6V, 300ms) demonstrated that all the muscles were maximally recruited by the time 100Hz was reached[3]. All data were collected using PowerLab® (ADinstruments, Colorado Springs, CO).
Femoral Geometry and Torsional Loading to Failure

Right femora were evaluated by μCT scan analysis [Siemens Inveon μCT equipped with Siemens Inveon Acquisition Workplace Software Version 1.5 (Siemens Preclinical Solutions, Knoxville) with an X-ray peak of 80 kVp and an exposure time of 140ms] prior to ex vivo torsional loading to failure analyses as previously described[224]. μCT image slices were analyzed using the Amira 5.3.3 software package (Mercury Computer Systems/TGS, Chelmsford, MA) to give a cubic voxel dimension of 0.083 mm$^3$. The mid-shaft slice located and modeled as a hollow elliptical cross-section. The marrow cavity diameter (MCD) reported as the average of the two endosteal diameters, the cortical bone width (CBW) reported as the average of the four thicknesses on the major and minor axes of the cortical mid-slice, and the polar moment of area (K: mm$^4$) determined as previously described. Following μCT analyses, the femora were potted into individualized cylindrical holders and the torsional loading to failure evaluated using the TA-HDi testing machine (Stable Micro Systems, Surrey, UK)[224]. Applied torque T (Nmm) was calculated and plotted as a function of relative angular displacement θ (degrees). The whole bone parameters of strength [torsional ultimate strength (T$_{\text{max}}$, Nmm) and strain energy to failure (U, Nmm)] and stiffness [torsional stiffness (Ks, Nmm/rad), which take into account bone geometry and material properties, were determined as previously described[224]. The bone material properties, tensile strength (Su, N/mm$^2$) and stiffness [shear modulus of elasticity, (N/mm$^2$)] were also determined according to Carleton et al[224].
Western Blot Analysis

Western blots were performed using a modified version of a previously described protocol[226]. Briefly, right gastrocnemius muscle, previously stored at -80°C, was homogenized in lysis buffer. Protein was mixed with 6x Laemmli buffer, boiled, and loaded on a Criterion TGX Pre-cast gel (Bio-Rad, Hercules, CA). After size separation via gel electrophoresis, the proteins were transferred to PDVF membrane. 5% milk in tris-buffered saline with Tween 20 (TBST) was used for blocking. The blots were labeled with primary antibodies against the total OXPHOS Antibody Cocktail (Abcam, Cambridge, MA, ab10413) containing 5 mouse monoclonal antibodies [CI subunit NDUFB8 (ab110242), CII-30kDa (ab14714), CIII-Core protein 2 (ab14745) CIV subunit I (ab14705) and CV alpha subunit (ab14748)], TFAM (Abcam, Cambridge, MA, ab89818), PGC-1α (MilliporeSigma, Burlington, MA, 516557), BNIP3 (Cell Signaling, Danvers, MA, 3769), PINK1 (Cell Signaling, Danvers, MA, 6946), Parkin (Cell Signaling, Danvers, MA, 4211S), P62 (Cell Signaling, Danvers, MA, 5114), and LC3 A/B (Cell Signaling, Danvers, MA, 12741). Primary antibodies were diluted to 1:1,000 in 5% BSA in TBST. Secondary horseradish peroxidase-conjugated antibodies were diluted to 1:5,000 in 5% milk in TBST. Blots were imaged and analyzed using Image Lab™ Software (Bio-Rad, Hercules, CA); protein content was normalized using amido black total protein stain (Figure II-S4).

Mitochondria Isolation and Respiration

Mitochondrial isolation and high resolution respirometry were carried out using modified versions of previously described methods[227–229]. For mitochondrial isolation, both
right and left gastrocnemius were removed, cleaned of extraneous fat and tissue, and placed in mitochondrial isolation buffer (100 mM KCl, 50 mM MOPS, 5 mM MgSO₄, 1 mM EGTA, and 1 mM ATP), on ice. Gastrocnemius muscles were homogenized using a rotor type homogenizer, the homogenate was centrifuged (800xg, 4°C, 10 min), and the supernatant filtered through gauze. The pellet was re-suspended in 5 mL mitochondrial isolation buffer, homogenized using a Potter-Elvehjem homogenizer (Teflon on glass), and centrifuged (800xg, 4°C, 10 min). The supernatant was filtered, added to the previously filtered supernatant, and centrifuged (12,000xg, 4°C, 10 min). The pellet was re-suspended in 3 mL isolation buffer with glass on glass homogenization, centrifuged (8,000xg, 4°C, 10 min), and the process repeated with 2 mL isolation buffer. 0.2% Bovine Serum Albumin (BSA) was added to 2 mL of the isolation buffer and used to re-suspend the pellet a third time via glass on glass homogenization followed by centrifugation (8,000xg, 4°C, 10 min). Finally, the pellet was re-suspended in 200 µL MiPO₃ buffer (0.5 mM EGTA, 3 mM MgCl₂-6H₂O, 60 mM K-lactobionate, 20 mM Taurine, 10 mM KH₂PO₄, 20 mM HEPES, 110 mM Sucrose, 1g/l BSA, 20 mM Histidine, 20 µM vitamin E succinate, 3 mM glutathione, 1 µM leupeptine, 2 mM glutamate, 2 mM malate, 2 mM Mg-ATP)[227] and allowed to equilibrate on ice for 30 min.

High resolution respirometry was used to measure mitochondrial respiration by the isolated mitochondria using the Orosboros Oxygraph-2k (Orosboros Instruments; Innsbruck, Austria). 30-45 µl of isolated mitochondria were added to the respiration chambers to measure basal respiration. Steady-state oxygen flux was measured by the addition of 2 mM malate and 5 mM glutamate. Oxygen flux through complex I was
measured by the titration of 125-375 µM of ADP; through complex I+II by the titration of 1-5 mM succinate, and maximal uncoupled respiration was measured by the addition of 0.125 µM FCCP. Finally, 10 µM cytochrome C was added to assess the quality of mitochondrial isolation. All measurements were normalized to the mitochondrial protein content.

**Transmission Electron Microscopy and Analysis**

Soleus muscle was excised, cleaned of any extraneous tissue or fat and placed in glutaraldehyde fixative at room temperature for 1 hour before being transferred to 4°C. Tissue was processed and prepared by the University of Missouri Electron Microscopy Core. Samples were imaged using a JEOL JEM-1400 120kV TEM. Images were captured at a magnification of 3,000X. Analysis was done using classical stereological methods as previously described to calculate the following, intermyofibrillar mitochondria: mitochondrial volume density, cross sectional area, and numerical density using the low magnification images[230,231].

**Mitochondrial DNA Content**

Mitochondrial DNA content was measured as a ratio of mitochondrial DNA to nuclear DNA using quantitative PCR (qPCR) methods and primers described by Malik et al[232]. Briefly, DNA was isolated from gastrocnemius muscle using a DNeasy Blood and Tissue Kit (Qiagen, Germantown, MD). qPCR was carried out in 10 µl reaction volume containing 5 µl PowerUp SYBR green master mix, 0.5 µl forward primer, 0.5 µl reverse primer, and 4 µl of diluted DNA. The mitochondrial DNA forward and reverse primers used were mMitoF1 (5’-CTAGAAACCCCGAAACAA-3’) and mMitoR1 (5’-
CCAGCTATCACCAGCTCGT-3’), respectively, and the nuclear DNA forward and reverse primers used were mB2MF1 (5’-ATGGGAAGCCGAACATACTG-3’) and mB2MR1 (5’-CAGTCTCAGTGGGGTGAATA-3’), respectively. A standard curve with copy numbers ranging from $10^2 – 10^9$ was amplified simultaneously with the isolated DNA used to quantify the ratio of mtDNA to nuclear DNA.

**Citrate synthase activity**

Citrate synthase activity was measured in mitochondria isolated from mixed gastrocnemius muscle using methods by Srere et al.[233], as previously described[226].

**Glycogen Content**

Snap frozen gastrocnemius muscle was powdered using a mortar and pestle on liquid nitrogen and approximately 15-30 mg of powdered muscle was added to 0.5 mL of 1 N HCL. The muscle was placed in a boiling water bath for 2.5 hours. After boiling, tubes were vortexed, and 1.5 mL of 0.67 M NaOH was added. The acid hydrolysis converts glycogen to free glucose which was then measured using Infinity Glucose Hexokinase Liquid Stable Reagent (Thermo Fisher Scientific, Waltham, MA) according to the manufacturer protocol using a sample to reagent ratio of 22.5:1[234].
Statistical Analysis

Statistical analyses were performed using SAS (SAS Institute Inc., Cary, NC). Swimming and voluntary wheel running exercise regimens were analyzed separately as a 2X2X2 factorial [2 genotypes (WT and oim/oim) X 2 sexes X 2 treatments (control and exercise)] and mitochondrial analyses were analyzed as 2X2 factorials [2 genotypes (WT and oim/oim) X 2 sexes] using Fisher’s Protected Least Significant Difference[235,236]. If heterogeneous variations were present, a log transformation was used to stabilize the variation. If the log transformation failed to stabilize the variation, a nonparametric ranked analysis was performed according to Conover et al[237]. Differences were considered significant at $p \leq 0.05$.

Results

Effect of volunteer wheel running and swimming exercise on muscle weight and function

As was demonstrated in our previous study[3], oim/oim mice exhibited reduced hindlimb muscle weights and absolute contractile forces as compared to WT littermates (Figure II-1), with minimal impact on relative contractile forces. Previously the decreased absolute contractile forces in male oim/oim hindlimb muscles was directly associated with decreased specific contractile force in male oim/oim plantaris (plant), gastrocnemius (gast), and tibialis anterior (TA) muscles, even though relative contractile forces did not always reach significance[3,163]. Volunteer wheel running and swimming exercised male oim/oim mice exhibited decreases in plant, gast, and TA muscle wet weight when compared to in-cage control counterparts (Figure II-1 B-D). Peak tetanic force of plant,
gast, and TA muscle in volunteer wheel running exercised and plant and TA muscle in swimming exercised male oim/oim mice were reduced as compared to their in-cage control counterparts (Figure II-1F-H). These decreases were still evident in the relative contractile force \([\text{peak tetanic force (P}_{o};\text{g})/\text{muscle weight (mg)}]\) of volunteer wheel running exercised male oim/oim plant and gast muscles, and swimming exercised male oim/oim plant muscle (Figure II-1J&K). However, interestingly, female mice did not exhibit any changes in muscle weight or function with both type of exercises regardless of genotypes (Figure II-S1). Both male and female oim/oim mice ran shorter total distances, 42% and 35% of the total distances ran by WT littermates, respectively (Figure II-S2). The average speed (km/hr) ran per week was also reduced in male and female oim/oim mice being 57% and 63% of WT speeds, respectively. In addition to reduced distance and average speed, the average time on the wheel (minutes) per week spent by oim/oim was 37% and 51% less that WT mice for both males (p=0.122) and females, respectively (Figure II-S2).
Figure II-1: Volunteer wheel running and swimming exercised male oim/oim mice had reduced hindlimb skeletal muscle weight and peak tetanic force as compared to in-cage control counterparts. (A) Soleus, (B), Plantaris, (C) Gastrocnemius, and (D) Tibialis Anterior muscle wet weights. (E) Soleus peak tetanic force ($P_o$), (F) Plantaris Peak $P_o$, (G) Gastrocnemius Peak $P_o$, (H) Tibialis Anterior Peak $P_o$, (I) Soleus relative $P_o$, (J) Plantaris relative $P_o$, (K) Gastrocnemius relative $P_o$, and (L) Tibialis Anterior relative $P_o$ of 4-month-old male WT and oim/oim in-cage control, volunteer wheel running, and swimming exercised mice. Graphs are presented as MEAN±STDEV. *$p$$\leq$0.05 vs. WT; †$p$$\leq$0.05 vs. in-cage control. (n=6-13 per group). Red and blue symbols represent WT and oim/oim, respectively. Circles, squares, and triangles represent control, wheel running, and swimming, respectively.
Effect of volunteer wheel running and swimming exercise on skeletal properties

Similar to previous investigations, oim/oim mice exhibit compromised bone microarchitecture and biomechanical properties as compared to their WT littermates regardless of sex and exercise regimen (Tables II-S2&S3, Figure II-2 A-D). Male WT mice that underwent volunteer wheel running exercise demonstrated decreases in several geometric and biomechanical parameters, including cortical bone width (CBW), polar moment of area (K), torsional ultimate strength ($T_{\text{max}}$), torsional stiffness (Ks), and energy to failure (U), while shear modulus of elasticity increased as compared to in-cage control counterparts (Figure II-2 A & C and Table II-S2). Swimming exercised male WT mice also showed decreases in CBW and K and increase in marrow cavity diameter (Table II-S2). Male oim/oim and female WT and oim/oim mice did not exhibit any changes in bone geometric or biomechanical properties with either of exercise regimen; except that female WT mice had decreases in $T_{\text{max}}$ and tensile strength (Su) and female oim/oim mice had decreased in $T_{\text{max}}$ with swimming exercise (Figure II-2 B and D and Table II-S3). Pearson’s correlation coefficients were evaluated between total distances run and bone biomechanical parameters in WT and oim/oim mice. The total distances run by male and female WT mice exhibited positive correlations with $T_{\text{max}}$ (+0.508; $p=0.022$) and U (+0.500; $p=0.025$), while oim/oim mice did not (Figure II-2E-H). The lack of positive correlations in oim/oim mice suggest that oim/oim mice are not able to respond as WT mice to weight bearing exercise, leading us to further question the musculoskeletal function of oim/oim.
**Figure II-2:** *oim/oim* mice exhibited compromised femoral biomechanical properties as compared to WT littermates. Volunteer wheel running exercised WT mice showed positive correlation between biomechanical properties and the total distance ran (km), but *oim/oim* mice did not. Torsional ultimate strength ($T_{\text{max}}$) of (A) male and (B) female mice; energy to failure ($U$) of (C) male and (D) female mice. *Red* and *blue* symbols represent WT and *oim/oim*, respectively. *Circles*, *squares*, and *triangles* represent control, wheel running, and swimming, respectively.

Linear regression analysis of the correlation between $T_{\text{max}}$ and total distance run by (E) WT and (F) *oim/oim* mice, linear regression analysis of the correlation between $U$ and total distance run of (G) WT and (H) *oim/oim* mice. Graphs are presented as MEAN±STDEV. *p*≤0.05 vs. WT; †p≤0.05 vs. in-cage control. (n=7-13 per group). *White* and *black* squares represent male and female animals, respectively.
Protein levels of electron transport chain (ETC) complexes

Previously, our group determined that citrate synthase activity was increased in oim/oim mixed gastrocnemius muscle homogenates relative to WT mixed gastrocnemius muscle homogenates[163], which in light of the response of oim/oim mice to the exercise regimens, prompted the evaluation of the ETC complexes in the current study. Oim/oim mice exhibited reduced levels of ETC complex IV, mitochondria encoded cytochrome oxidase I [51% decrease in females and 20% decrease (which did not reach significance) in males] as compared to WT counterparts (Figure II-3), while the protein levels of other complexes did not differ between genotypes.
Figure II-3: Female oim/oim mixed gastrocnemius muscle homogenate had reduced ETC complex IV mitochondrial encoded cytochrome c oxidase I (c4). Western blot analysis of electron transport chain (ETC) complexes measured in 4-month-old mixed gastrocnemius muscle of (A) male and (B) female WT (red diamonds) and oim/oim (blue squares) mice. (C) Representative blot images. Graphs are presented as MEANS±SE. *p≤0.05 vs. WT. (n=5-6 per group).
Skeletal Muscle Mitochondrial Function

To investigate skeletal muscle mitochondrial function more directly, mitochondria were isolated from 4-month-old male WT and oim/oim mice mixed gastrocnemius muscles, and oxygen consumption was measured using high resolution respirometry. Oim/oim skeletal muscle mitochondria demonstrated reductions in respiration during basal (-65%), state 2 (-52%), state 3-Complex I (-61%), state 3-Complex I+II (-53%), and maximal uncoupled respiration (-52%) as compared to WT skeletal muscle mitochondria (Figure II-4A). These decreases clearly demonstrate inefficient mitochondrial function with compromised energy production in oim/oim gastrocnemius muscle.

In addition, the isolated oim/oim gastrocnemius muscle mitochondria exhibited a 33% decrease in citrate synthase activity as compared to WT gastrocnemius muscle mitochondria (Figure II-4B). While citrate synthase activity in whole muscle homogenate is often an indirect marker of mitochondrial content/mass, in isolated mitochondria, citrate synthase activity can provide insight into mitochondrial function[238–240].
**Figure II-4:** (A) Male oim/oim mice exhibit reduced mitochondrial respiration and (B) mitochondrial citrate synthase activity compared to WT littermates. (A) Basal, state 2, state 3, and maximal uncoupled mitochondrial respiration was significantly decreased in 4-month-old oim/oim (blue squares) male mice compared to WT (red diamonds) littermates. Oxygen consumption was measured with the addition of substrates to stimulate different respiration states: basal (mitochondria in MiPO₃ buffer), state 2 (Glutamate and Malate (GM)), state 3 respiration with electron flux through complex I (GM and ADP), state 3 respiration with electron flux through complex I and II (succinate), and maximal uncoupled respiration (FCCP). (B) Citrate synthase activity was decreased in isolated mitochondria from 4-month-old male oim/oim mice compared to WT littermates. Graphs are presented as Means±SE of the genotype values. *p≤0.05 vs. WT. (n=5-8 per group).
Mitochondrial Biogenesis and Mitophagy

To determine if \textit{oim/oim} mice have altered mitochondrial turnover, markers of mitochondrial biogenesis and autophagy/mitophagy were evaluated. Mitochondrial biogenesis markers, PGC1-\(\alpha\) (a nuclear co-activator) and TFAM (a mitochondrial transcription factor)[187] were increased in male \textit{oim/oim} mixed gastrocnemius muscle while TFAM was increased in female \textit{oim/oim} mixed gastrocnemius muscle compared to their WT counterparts (Figure II-5A, Supplemental Figure II-3A). Several markers of autophagy and mitophagy were also evaluated including LC3, a known component of autophagosome formation, as well as the LC3-II to LC3-I ratio, a method to assess autophagy; with increases in the LC3-II/I ratio being suggestive of elevated autophagy[186,187,241,242] and warranting direct studies of autophagy flux. In the skeletal muscle of male \textit{oim/oim} mixed gastrocnemius, the LC3-II/I ratio was reduced by greater than 50\% compared to WT counterparts; a decrease largely driven by increases in LC3-I (117\%) (Figure II-5B & C). The levels of other autophagy/mitophagy markers including PINK1, Parkin, BNIP3, and p62 did not differ between genotypes.
Figure II-5: Male oim/oim mice exhibit increased markers of mitochondrial biogenesis and decreased markers of autophagy and mitophagy. Western blot analysis of markers of (A) mitochondrial biogenesis (PGC-1α and TFAM) and (B) autophagy and mitophagy (PINK1, Parkin, BNIP3, LC3 II/I, and p62) in protein homogenate isolated from 4-month-old male oim/oim mixed gastrocnemius muscle. (A) TFAM and PGC-1α (p=0.07), a marker of mitochondrial biogenesis was increased in oim/oim (blue squares) mice compared to WT (red diamonds) mice and (B) LC3 II/I, a marker of mitophagy, was decreased in oim/oim mice compared to WT mice driven by an increase in (C) LC3-I. Graphs are presented as MEANS±SE. *p≤0.05 vs. WT; †p=0.07 vs WT. (n=5-6 per group).
To determine if the decrease in the LC3-II/I ratio seen in the whole gastrocnemius muscle homogenate was indicative of decreased mitochondrial degradation rather than decreases in general autophagy, we measured LC3-II and BNIP3 levels in isolated mitochondria of mixed gastrocnemius muscle and found that LC3-II levels in the 4-month-old male oim/oim mice were dramatically decreased compared to their WT counterparts while BNIP3 levels appeared reduced but did not reach significance, further suggesting that mitophagy is decreased in the skeletal muscle of oim/oim mice (Figure II-6).

Figure II-6: BNIP3 and LC3-II levels were measured in isolated mitochondria of 4-month-old male WT (red diamonds) and oim/oim (blue squares) mixed gastrocnemius muscle. LC3-II levels were significantly reduced in oim/oim mitochondrial compared to WT mitochondria. Graphs are presented as MEANS±SE *p≤0.05 vs WT. (n=6-8 per group).
Mitochondrial Content

Electron microscopic analyses of 4-month-old male oim/oim and WT soleus muscle found no evidence of gross abnormal morphological changes in oim/oim soleus muscle. In addition no differences were found in intermyofibrillar mitochondrial volume density, cross-sectional area, and numerical density (Figure II-7). To investigate further, the relative mitochondrial DNA content was evaluated by the ratio of mitochondrial DNA to nuclear DNA in 4-month-old male and female oim/oim and WT gastrocnemius muscles. The male and female oim/oim gastrocnemius mitochondrial/nuclear DNA ratios were decreased by 40% and 34% relative to the mitochondrial/nuclear DNA ratios of sex matched WT gastrocnemius muscles (although not reaching significance), respectively (Figure II-8).
Figure II-7: Ten representative images of 4-month-old WT (red diamonds) and oim/oim (blue squares) intermyofibrillar mitochondria from soleus muscle were assessed using Transmission Electron Microscopy and classical stereological methods. The intermyofibrillar (A) mitochondria volume density, (B) cross sectional area, and (C) mitochondrial numerical density as well as gross morphology were not different between (D) WT and (E) oim/oim soleus muscle. Graphs are presented as MEANS±SE. (n=3 per group).
Figure II-8: *Oim/oim* mice exhibit a reduced mitochondrial DNA to nuclear DNA ratio. The ratio of mitochondrial DNA copy number to nuclear DNA copy number was decreased by 40% in (A) male and by 34% in (B) female 4-month-old oim/oim (blue squares) mixed gastrocnemius muscle as compared to WT (red diamonds) mixed gastrocnemius muscle. Graphs are presented as MEANS±SE. †p=0.08 vs. WT. (n=8-11 per group)
**Skeletal Muscle Glycogen Content**

Effective glycogen metabolism provides glucose to muscle cells to induce muscle contraction, and alterations could result in reduced muscle contractile forces. To investigate glycogen metabolism, skeletal muscle glycogen content was evaluated in WT and *oim/oim* gastrocnemius muscles. No differences between WT and *oim/oim* mice were found, further suggesting that the mitochondrial dysfunction itself is likely the major contributor to the reduced contractile force in *oim/oim* mice (Figure II-9).

**Figure II-9:** Glycogen content is equivalent in *oim/oim* (blue squares) relative to WT (red diamonds) mixed gastrocnemius muscle. Glycogen content was measured in gastrocnemius muscle of 4-month-old (A) male and (B) female *oim/oim* and WT. Graphs are presented as MEANS±SE. (n=4–5 per group).
Discussion

The beneficial impact of exercise on overall health is well established, and particularly its positive impact on musculoskeletal health during the pre-pubertal/pubertal growth period[115,126,129,211,212]. We chose to test the impact of two different exercise modalities, voluntary wheel running (weight-bearing) and swimming (non-weight bearing)[243] on WT and oim/oim hindlimb muscle and bone. Voluntary wheel running provides unlimited access without aversive stimuli; but it does not control for duration and intensity. Swimming provides minimal ground reaction forces and aqua therapy is often a physiotherapeutic recommendation for individuals with osteogenesis imperfecta[244], however it may induce physiological stress responses, at least initially, in mice[243]. Unlike previous reports of the positive influence of physical exercise on muscle and bone health[115,212], we found male WT and oim/oim mice exhibited decreases in muscle weight and function and slightly compromised bone microarchitecture and biomechanical properties, while female mice were minimally affected. Sex differences in bone structure and properties in the oim mouse model have previously been reported[245]. One caveat between our exercise study and the previous reports of the positive impact of physical exercises is that our study focused only on aerobic exercise (swimming and voluntary wheel running) while exercise studies in OI patients incorporated a variety of activity including aerobic and strength training exercises[115,212]. In addition, our exercise study focused on mice modeling severe OI while previous reports were in children with mild and moderate OI[115]. This suggests that both the type of exercise and the type of OI are important factors when considering exercise as a potential treatment option. WT mice ran greater than average distances
compared to their age-matched oim/oim counterparts regardless of sex. This is consistent with previous studies of reduced physical activity levels in oim/oim mice as well as in Col1a1Jrt/+ mice, a model of moderately severe OI as a result of a Col1a1 gene defect[8,163]. Growing evidence suggests bone is most responsive to dynamic moderate to high magnitude loads, which are quick, short in duration, and non-repetitive in direction[246,247]. Interestingly, WT mice showed positive correlations between total distances run and bone strength, while oim/oim mice did not, which may reflect impaired musculoskeletal response to the weight bearing exercise or that the reduced distances ran, average speeds and time spent on the wheel by the oim/oim mice did not reach the minimal threshold for musculoskeletal responsiveness. The skeleton is striving for the optimum geometry and mass to be able to bear loads while remaining light enough for efficient mobility[126]. Therefore, one should consider that the lack of an overall positive response to the extended running and swimming regimens may also reflect desensitization of the osteocytes to repetitive low magnitude loads[58,248,249]. In separate studies with rats extended durations of running exercise have been shown to not change or to decrease BMD and serum alkaline phosphatase levels, a bone formation biomarker[213,216].

The lack of positive musculoskeletal effects in both WT and oim/oim animals that underwent training regimens suggests that perhaps a more controlled exercise regimen consisting of both endurance and strength training exercises with increasing loads is required to improve muscle and bone properties.

In a previous study, whole muscle homogenates from oim/oim gastrocnemius muscle had increased citrate synthase activity as compared to WT gastrocnemius muscle
homogenates[163], which led us to hypothesize that oim/oim mice have either 1) increased mitochondrial numbers (mitochondrial mass) or 2) increased mitochondrial function as compared to WT mice. However, our current study demonstrates that oim/oim mice have dramatic reductions in skeletal muscle mitochondrial respiration rates, markers of autophagy and mitophagy, ETC complex IV mitochondrial encoded cytochrome oxidase I, and mitochondrial citrate synthase activity with more mild reductions in mitochondrial DNA /nuclear DNA ratios compared to WT littermates. In addition, markers of mitochondrial biogenesis are increased in oim/oim mice, which may reflect compensatory increases in an attempt to overcome the severely reduced mitochondrial respiration or mitochondrial damage.

The decreases in mitochondrial respiration, ETC complex IV protein content, and mitochondrial DNA copy number may be reflective of the functional oim/oim muscle phenotype of decreased muscle contractile force in oim/oim mice[3]. The mitochondria are responsible for generating the energy necessary for muscle contraction[250,251], and are the primary source of ATP production; ATP is generated by complex V of the ETC chain, ATP synthase, via a proton gradient which is primarily generated and sustained as electrons are transferred through complex IV to oxygen while four protons are simultaneously pumped across the membrane[250,252]. The decreased mitochondrial oxygen consumption in skeletal muscle of oim/oim mice suggests that the oim/oim mitochondrial function is not as efficient as WT mitochondria. Decreased ETC complex IV mitochondrial encoded cytochrome oxidase I in oim/oim gastrocnemius muscle is consistent with the decreased potential to transfer electrons to oxygen which may lead to the inability to maintain an adequate proton gradient.
Decreased mitochondrial DNA / nuclear DNA ratios suggests decreased mitochondria number[232], although it is controversial as to whether mitochondrial DNA/nuclear DNA ratio is an adequate marker for mitochondrial content especially due to the large variation in DNA copy number per mitochondria[239]. However, increased protein markers of mitochondrial biogenesis, PGC-1α and TFAM, and decreased LC3-II in isolated mitochondria favor increased mitochondrial number[187,239]. Although, mitochondrial DNA appears reduced in oim/oim gastrocnemius, increased PGC-1α and TFAM support the hypothesis that oim/oim mice are trying to modulate the signaling pathways in favor of increasing mitochondria, likely to compensate for reduced function. This would be consistent with the indirect measure of mitochondrial content, whole muscle citrate synthase activity, previously observed to be increased in oim/oim whole muscle gastrocnemius protein homogenates[163].

This is the first report of severe muscle weakness in an OI mouse model being attributed, at least in part, to compromised mitochondrial function. Another moderately-severe OI mouse model, Col1a1Jrt/+ , which has an autosomal dominant 18 amino acid deletion in the triple helical domain of the procα1(I)collagen chain[51], exhibits reduced physical activity levels with increased oxygen consumption and CO2 production concomitant with increased whole body energy expenditure and altered energy homeostasis[8]. Interestingly, we did not observe any changes in skeletal muscle glycogen content of oim/oim mice compared to WT littermates, further suggesting that the muscle defects, specifically reduced contractile generating force, may be associated with mitochondrial dysfunction, but future studies warrant the inclusion of fatiguing protocols to decipher the role of mitochondria in oim/oim muscle weakness more clearly. Further investigation into
metabolic homeostasis and energy utilization and expenditure in the oim/oim mouse are critical to understanding the altered mitochondrial function.

Although the role of the extracellular matrix (ECM) as a potential contributing factor to the mitochondrial dysfunction observed in the oim/oim mouse remains to be investigated, the role of the ECM in reduced contractile force and the mechanotransduction of force also needs to be elucidated. The skeletal muscle ECM is hierarchical and can be divided into three structures: the endomysium, perimysium, and epimysium[92,253]. The endomysium is a thin continuous extracellular matrix sheath containing collagen (types III, IV, V, VI, XII, and very little type I collagen) covering the myofibers. The endomysium interacts with the perimysium, which is composed largely of type I collagen and surrounds the muscle fascicles. The perimysial junctional plates (PJP) are the focal site of contact between the perimysium and the myofiber[254]. Through the PJP the perimysium is responsible for the lateral transmission of the contractile forces to the epimysium (largely composed of type I collagen) and the myoteninous junction into the tendon and subsequent tendon-bone interface[92,205,253,254]. The presence of not only reduced amounts of type I collagen[3], but also abnormal homotrimeric type I collagen in the oim/oim muscle contributes to an altered ECM, which likely disrupts force transmission and altering muscle-bone crosstalk. Passerieux et al. demonstrated intracellularly at the sites of the PJP the presence of type I collagen bundles (perimysial) in association with the subsarcolemmal mitochondria and myonuclei clusters[205]. These findings suggest that although the intermyofibrillar mitochondria of the oim/oim soleus muscle did not appear quantitatively or morphologically abnormal by EM, it is important to evaluate the subsarcolemmal mitochondria. Abnormal muscle ECM has been
demonstrated by defects in type VI collagen, a component of the perimysium, seen in Bethlem myopathy and Ullrich congenital muscular dystrophy, leading to mitochondrial dysfunction, calcium deregulation, elevated reactive oxygen species, increased apoptosis, and muscle weakness[181,255]. Future investigation into the role of the ECM in oim/oim skeletal muscle weakness may provide insight into the mechanisms of mitochondrial dysfunction in the oim/oim skeletal muscle.

Understanding the mechanisms of muscle weakness in OI are an important consideration for OI treatment, as bone is responsive to the force it feels from muscle contraction. Physiotherapeutic studies that target the muscle weakness in OI are critically needed[256]. Severe muscle weakness or disuse hinder patient quality of life as well as negatively impacting bone health. Understanding the mechanism(s) of muscle weakness in OI will provide new therapeutic targets that will have the potential to improve both muscle and bone strength.

This study is one of the first to demonstrate that extended voluntary wheel running and swimming exercise regimens were well tolerated by the oim/oim mouse a model of moderately severe OI. Although wild-type mice exhibited positive correlations for total distance run and bone biomechanical strength, this was not true for the oim/oim mice, suggesting that the oim/oim mouse muscle-bone unit mechano-responsiveness is compromised. This study is the first to demonstrate reduced mitochondrial function in skeletal muscle as a potential contributing factor to muscle weakness in a mouse model of OI, as evidenced by alterations in mitochondrial content and respiratory function. Further studies including the evaluation of reactive oxygen species and calcium homeostasis are essential to characterize the mechanism of mitochondrial dysfunction in oim/oim mice.
and to understand the impact of the mitochondrial energy utilization and expenditure on muscle weakness. Exploring the mechanisms responsible for muscle weakness and whole-body metabolic alterations promises to provide important insight into the musculoskeletal pathogenesis of OI.

*Authors' roles:* Study design: VLG, YJ, CLP, RSR. Study conduct: muscle (YJ), mouse exercise (YJ), bone biomechanics (YJ), mitochondrial markers in whole muscle homogenate (VLG, RPC, and GMM), mitochondrial isolation experiments (VLG and GMM), and electron microscopy (VLG). Data analysis: VLG, YJ, RPC, GMM, RSR, and CLP. Drafting manuscript: VLG, YJ, RPC, GMM, RSR, and CLP. Revising manuscript content: VLG, YJ, RPC, GMM, RSR, and CLP. Approving final version of the manuscript: VLG, YJ, RPC, GMM, RSR, and CLP. CLP takes responsibility for the integrity of the data analysis.
Supplementary Material for Chapter II

**Table II-S1.** The swimming exercise regimen of wild-type and *oim/oim* mice was initiated at 6 weeks of age and completed at 4 months of age.

<table>
<thead>
<tr>
<th>Week 1</th>
<th>Swim (min)</th>
<th>Rest (min)</th>
<th>Swim (min)</th>
<th>Rest (min)</th>
<th>Swim (min)</th>
<th>Total Swim Time (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 1</td>
<td>2</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
<td>2</td>
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<tr>
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<td>N/A</td>
<td>N/A</td>
<td>N/A</td>
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<tr>
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<td>N/A</td>
<td>N/A</td>
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<tr>
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<td>6</td>
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<td>N/A</td>
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</tr>
<tr>
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<td>3</td>
<td>8</td>
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<td>N/A</td>
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</tr>
<tr>
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<td></td>
<td></td>
<td></td>
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<td></td>
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<td>3</td>
<td>10</td>
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<tr>
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<td>3</td>
<td>12</td>
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<td>N/A</td>
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Table II-S2. Femoral microarchitecture and biomechanical properties of male WT and oim/oim in-cage control, volunteer wheel running, and swimming exercised mice.

<table>
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<tr>
<th></th>
<th>WT</th>
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<th>oim/oim</th>
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<td></td>
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<td>Wheel</td>
<td>Swim</td>
<td>Control</td>
<td>Wheel</td>
<td>Swim</td>
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<td>(n = 9-11)</td>
<td>(n = 11)</td>
<td>(n = 8-10)</td>
<td>(n = 8-9)</td>
<td>(n = 9)</td>
<td>(n = 8-9)</td>
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<td><strong>Cortical Bone Microarchitecture a</strong></td>
<td></td>
<td></td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>FL</td>
<td>16.07±0.26</td>
<td>15.46±0.29†</td>
<td>15.86±0.24</td>
<td>16.09±0.34</td>
<td>15.47±0.34†</td>
<td>15.89±0.29</td>
</tr>
<tr>
<td>MCD</td>
<td>1.00±0.09</td>
<td>0.90±0.05†</td>
<td>1.09±0.08†</td>
<td>0.96±0.04</td>
<td>0.96±0.07</td>
<td>0.96±0.07</td>
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<td>CBW</td>
<td>0.40±0.04</td>
<td>0.35±0.02†</td>
<td>0.34±0.02†</td>
<td>0.33±0.01*</td>
<td>0.30±0.03*</td>
<td>0.32±0.03*</td>
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<tr>
<td>K</td>
<td>0.86±0.16</td>
<td>0.53±0.03†</td>
<td>0.72±0.09†</td>
<td>0.52±0.08*</td>
<td>0.43±0.06*</td>
<td>0.50±0.07*</td>
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<tr>
<td><strong>Torsional Loading to Failure b</strong></td>
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<td></td>
</tr>
<tr>
<td>T&lt;sub&gt;max&lt;/sub&gt;</td>
<td>43.26±6.49</td>
<td>29.37±3.61†</td>
<td>38.36±3.96</td>
<td>9.76±6.38*</td>
<td>9.27±4.00*</td>
<td>9.61±2.71*</td>
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<tr>
<td>Ks</td>
<td>55.86±11.12</td>
<td>47.74±9.21†</td>
<td>59.34±14.31</td>
<td>28.68±10.24*</td>
<td>29.18±6.74*</td>
<td>31.86±8.40*</td>
</tr>
<tr>
<td>Su</td>
<td>52.10±7.20</td>
<td>50.06±6.62</td>
<td>51.14±5.06</td>
<td>17.90±13.07*</td>
<td>18.15±7.54*</td>
<td>16.98±5.09*</td>
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<tr>
<td>G</td>
<td>489.16±130.</td>
<td>658.12±188.3†</td>
<td>558.72±117.37</td>
<td>417.24±189.</td>
<td>475.37±105.48</td>
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<tr>
<td>U</td>
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<td>6.45±1.25†</td>
<td>8.53±1.16</td>
<td>1.67±1.59*</td>
<td>1.54±1.12*</td>
<td>1.57±0.73*</td>
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</table>

*aFemur Length (FL); Marrow Cavity Diameter (MCD); Cortical Bone Width (CBW); Polar moment of area (K)

*bTorsional ultimate force (T<sub>max</sub>); Torsional stiffness (Ks); Tensile strength (Su); Shear modulus of elasticity (G); Energy to failure (U)

Values are presented as MEAN±STDEV. *<i>p</i>≤0.05 vs. WT; †<i>p</i>≤0.05 vs. in-cage control
Table II-S3. Femoral microarchitecture and biomechanical properties of female WT and oim/oim in-cage control, volunteer wheel running, and swimming exercised mice.

<table>
<thead>
<tr>
<th></th>
<th>WT</th>
<th></th>
<th>oim/oim</th>
<th></th>
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</thead>
<tbody>
<tr>
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<td>Control (n = 8)</td>
<td>Wheel (n = 9-11)</td>
<td>Swim (n = 11-12)</td>
<td>Control (n = 7-9)</td>
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<td><strong>Cortical Bone Microarchitecture a</strong></td>
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<tr>
<td>FL</td>
<td>15.77±0.37</td>
<td>15.71±0.22</td>
<td>15.58±0.44</td>
<td>15.44±0.49</td>
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<td>0.89±0.05</td>
<td>0.92±0.05</td>
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<td>CBW</td>
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<td>0.35±0.02</td>
<td>0.34±0.02</td>
<td>0.29±0.03*</td>
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<tr>
<td>K</td>
<td>0.53±0.06</td>
<td>0.50±0.05</td>
<td>0.53±0.09</td>
<td>0.37±0.05*</td>
</tr>
<tr>
<td><strong>Torsional Loading to Failure b</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T&lt;sub&gt;max&lt;/sub&gt;</td>
<td>33.14±3.49</td>
<td>31.38±1.46</td>
<td>26.71±3.56†</td>
<td>13.54±1.43*</td>
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<tr>
<td>Ks</td>
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<td>42.20±2.88</td>
<td>47.07±15.28</td>
<td>36.29±8.22*</td>
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<tr>
<td>Su</td>
<td>55.37±3.13</td>
<td>54.71±3.82</td>
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<td>682.95±196.2</td>
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<td>U</td>
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<td>7.48±0.91</td>
<td>6.50±1.80</td>
<td>2.44±0.71*</td>
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a Femur Length (FL); Marrow Cavity Diameter (MCD); Cortical Bone Width (CBW); Polar moment of area (K)
b Torsional ultimate force (T<sub>max</sub>); Torsional stiffness (Ks); Tensile strength (Su); Shear modulus of elasticity (G); Energy to failure (U)

Values are presented as MEAN±STDEV. *p<0.05 vs. WT; †p<0.05 vs. in-cage control
Figure II-S1: Volunteer wheel running and swimming exercised female oim/oim mice had reduced hindlimb skeletal muscle weight and peak tetanic force as compared to in-cage control counterparts. (A) Soleus, (B) Plantaris, (C) Gastrocnemius, (D) Tibialis Anterior muscle wet weights, (E) Soleus peak tetanic force ($P_o$), (F) Plantaris Peak $P_o$, (G) Gastrocnemius Peak $P_o$, (H) Tibialis Anterior Peak $P_o$, (I) Soleus relative $P_o$, (J) Plantaris relative $P_o$, (K) Gastrocnemius relative $P_o$, (L) Tibialis Anterior relative $P_o$ of 4 months old female WT and oim/oim in-cage control, volunteer wheel running, and swimming exercised mice. Graphs are presented as MEAN±STDEV. *$p$≤0.05 vs. WT; (n=4-13 per groups). Red and blue symbols represent WT and oim/oim, respectively. Circles, squares, and triangles represent control, wheel running, and swimming, respectively.
Figure II-S2: (A) Average of the total distance (km) ran by male and female oim/oim (blue) mice was significantly less than wild-type (red) mice. Average of weeks 2-10 for (B) average distance ran (km) per day, (C) average speed (km/hr), and (D) average time spent on the wheel (mins) were reduced for oim/oim (blue) mice compared to wild-type (red) mice. Average body weight (g) per week of (E) male and (F) female WT and oim/oim mice. Average distance ran per day per week of (G) male and (H) female WT and oim/oim mice. Average speed (km/hr) ran per week of (I) male and (J) female WT and oim/oim mice. Average time on the wheel (minutes) per week of (K) male and (L) female WT and oim/oim mice. Graphs are presented as MEAN±STDEV. *p≤0.05 vs. WT. (n=9-12 per group)
Figure II-S3: (A) Female 4-month-old oim/oim mixed gastrocnemius muscle homogenates exhibit increased TFAM, a marker of mitochondrial biogenesis. Western blot analysis of markers of (B) autophagy and mitophagy in protein isolated from 4-month-old female oim/oim (blue symbols) mixed gastrocnemius muscle did not change compared to WT (red symbols). (C) LC3-I and LC3-II were quantified separately. Graphs are presented as MEANS ± SE. *p≤0.05 vs. WT. (n=5-6 per group).
Figure II-S4: A representative blots (for western blot analyses) following transfer and staining with the amido black stain for total protein content to show transfer quality and for normalization of the western blot quantification (Figures II-3, 5 & 6).
CHAPTER III

SKELETAL MUSCLE SPECIFIC MITOCHONDRIAL DYSFUNCTION AND ALTERED ENERGY METABOLISM IN A MURINE MODEL (OIM/OIM) OF SEVERE OSTEOGENESIS IMPERFECTA

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Abstract

Osteogenesis imperfecta (OI) is a heritable connective tissue disorder with patients exhibiting bone fragility and muscle weakness. The synergistic biochemical and biomechanical relationship between bone and muscle is a critical potential therapeutic target, such that muscle weakness should not be ignored. Previous studies demonstrated mitochondrial dysfunction in the skeletal muscle of oim/oim mice, which model a severe human type III OI. Here, we further characterize this mitochondrial dysfunction and evaluate several parameters of whole body and skeletal muscle metabolism. We demonstrate reduced mitochondrial respiration in female gastrocnemius muscle, but not in liver or heart mitochondria, suggesting that mitochondrial dysfunction is not global in the oim/oim mouse. Myosin heavy chain fiber type distributions were altered in the oim/oim soleus muscle with a decrease (-33 to 50%) in type I myofibers and an increase (+31%) in type IIa myofibers relative to their wildtype (WT) littermates. Additionally, altered body composition and increased energy expenditure were observed oim/oim mice relative to WT littermates. These results suggest that skeletal muscle mitochondrial dysfunction is linked to whole body metabolic alterations and to skeletal muscle weakness in the oim/oim mouse.
Introduction

Osteogenesis imperfecta (OI), also referred to as brittle bone disease, is a rare, heritable connective tissue disorder affecting approximately 1 in 15,000 births [1,17]. Roughly 85% of OI cases arise due to mutations in the type I collagen genes, *COL1A1* and *COL1A2*, while the remaining 15% result from mutations in genes involved in posttranslational modification of type I collagen, osteoblast maturation or mineralization [1]. Type I collagen, the most abundant of the collagens, is an important structural protein composing the majority of the organic component of bone. OI is an extremely heterogeneous disease in terms of severity with the most common and striking feature being bone fragility. There are four classical Sillence types of OI: type I is the least severe, type II results in perinatal lethality, type III being the most severe viable form, and type IV results in a moderate phenotype [13]. In addition to bone fragility, clinical manifestations include short stature and craniofacial abnormalities, with more recent investigations also highlighting the presence of an inherent muscle pathology in both mouse models and patients with OI [3,4,6,48]. However, current treatment strategies for patients with OI focus primarily on bone fragility, with emphasis on preventing bone resorption (bisphosphonates) and/or surgical rodding interventions[17]. Understanding the mechanisms of muscle weakness in OI could potentiate novel therapies to improve muscle function and ultimately bone strength.

Muscle weakness has been implicated as an inherent part of the pathophysiology of OI through investigation of the osteogenesis imperfecta murine (*oim*) model[3]. The *oim* mouse *Colla2* gene contains a nucleotide deletion, leading to a frameshift in the carboxy-terminal end resulting in a nonfunctional proα2(I) collagen. This ultimately leads to
production of homotrimeric type I collagen, \([\alpha_1(I)_3]\), rather than the normal heterotrimeric type I collagen, \([\alpha_1(I)\alpha_2(I)]\)[39]. The homozygous \((oim/oim)\) mice model severe human type III OI, exhibiting severe bone fragility, decreased bone mineral density, and small body size compared to wildtype (WT) littermates[39]. \(Oim/oim\) mice have significantly reduced muscle size as well as reduced specific contractile generating force [contractile force normalized to muscle cross sectional area][3]. Subsequently, numerous reports have revealed muscle weakness in OI patients, including that 80% of patients with mild/moderate OI experience muscle force deficits [4,6,47,48]. Because muscle and bone are highly synergistic tissues which “talk” to one another through both chemical signals and physical forces, it is important to understand this muscle weakness in OI in order to exploit this relationship to ultimately improve bone strength through the improvement of muscle function [14,57,257]. We recently demonstrated that \(oim/oim\) hindlimb skeletal muscle exhibits severe mitochondrial dysfunction and altered mitochondrial numbers, which likely contributes to their muscle weakness [162][258].

Evidence of metabolic alterations in OI, though limited, have been observed in both mouse models and the patient population. Previous evidence found that prepubescent OI patients exhibited higher body temperatures and a 45% increase in metabolic rate, but did not show altered respiratory quotients relative to children without OI[7]. Interestingly, they also reported that these changes subsided in patients after puberty, as do the overall fracture rates suggesting a link between hyper-metabolism, fractures, and sex-hormones[7]. More recently, evidence of metabolic alterations were demonstrated in a mouse model of severe OI; 4 week old \(Col1a1^{Jrv/+}\) mice (modeling human OI type IV/III) exhibited sex-dependent changes in metabolism that normalized by puberty (8 weeks of
These alterations included increased insulin levels and respiratory exchange rate, in males, as well as improved glucose tolerance in females. Additionally, a previous report found evidence that a patient with OI type III/IV, initially suspected to have malignant hyperthermia, was in a hypermetabolic state which was likely related to a primary type I collagen defect [259].

Based on these previous reports of metabolic alterations in both patients and the Coll1a1Jrt/+ mouse as well as our recent findings of mitochondrial dysfunction, we sought to investigate the potential of metabolic dysfunction in the oim/oim mouse. The current study provides further investigations of mitochondrial function and metabolism in the oim/oim mouse, by 1) addressing whether the skeletal muscle mitochondrial dysfunction is more global or tissue specific by evaluating cardiac muscle and liver mitochondrial function, 2) evaluating skeletal muscle fiber type distributions, and 3) evaluating several metabolic parameters including measures of whole body energy expenditure and skeletal muscle fatty acid oxidation relative to wildtype littermates. Better characterization and understanding of these alterations provide insight into the skeletal muscle weakness that occurs in OI and the nature of the relationship between mitochondrial function and type I collagen mutations and promise to pave the way for new therapies that will improve muscle function and ultimately bone strength.
Methods

Mice

Col1a2\textsuperscript{oim} mice (stock #001815 (\textit{oim})) were originally obtained from Jackson Laboratory (Bar Harbor, ME, USA). All mice were maintained on the C57BL/6 J background and genotyped as previously described [39,224]. Non-breeding male and female wild type (WT) and \textit{oim/oim} adult mice (13-19 weeks of age) were used for all experiments described, except for an additional subset of 8 week old WT and \textit{oim/oim} males whose gastrocnemius and quadriceps muscles were used to assess mitochondrial respiration. Age- and sex- matched WT littermates were used as control animals. All mice were given\textit{ ad libitum} access to food (stock chow, Purina Lab Diet 5053) and water. Mice were housed in an AAALAC-accredited facility at the University of Missouri; all experimental manipulations were performed under an approved University of Missouri Animal Care and Use Committee protocol. At the time of sacrifice, a mixture of xylazine/ketamine anesthesia was used followed by cardiac puncture (serum collected for future experiments), and cervical dislocation.

Mitochondria Isolation, Respiration, and \textit{H}_2\textit{O}_2 Analyses

Mitochondria were isolated from heart, liver, and mixed skeletal muscles including the gastrocnemius and quadriceps muscles (each muscle contains both white/fast-twitch/type II and red/slow-twitch/type I myofibers) following a series of homogenization and differential centrifugation procedures as previously described [227,229,260]. Mitochondria isolated from heart, liver and gastrocnemius and quadriceps muscles were used to assess mitochondrial respiration via high-resolution respirometry, while
mitochondria isolated from gastrocnemius muscle were also used to determine mitochondrial H$_2$O$_2$ production.

High-resolution respirometry was used to measure mitochondrial respiration by the isolated liver, heart, gastrocnemius (gast), and quadriceps (quad) muscle mitochondria using the Oroboros Oxygraph-2k (Oroboros Instruments, Innsbruck, Austria) following a modified protocol[227,258,261]. 20 µL of isolated mitochondria were added to the respiration chambers to measure basal respiration. Steady-state oxygen flux was measured by the addition of 2 mM malate and 5 mM glutamate. Oxygen flux through complex I was measured by the titration of ADP [50 to 125 µM (liver), 25-125 µM (heart), or 125-375 µM (gast and quad)]; through complex I + II by the titration of succinate [5 to 7.5 mM (liver), 10 mM (heart), or 1-5 mM (gast and quad)], and maximal uncoupled respiration was measured by the titration of carbonyl cyanide-4-(trifluoromethoxy) phenylhydrazone (FCCP) [0.125-0.25 µM (liver), 0.25 µM (heart), or 0.125 µM (gast and quad)]. Finally, cytochrome c was added [5 µM (liver and heart) and 10 µM (gast and quad)] to assess the quality of mitochondrial isolation. All measurements were normalized to the mitochondrial protein content.

An Oroboros Oxygraph-2k (Oroboros Instruments, Innsbruck, Austria) equipped with a fluorimeter was used to measure H$_2$O$_2$ production in mitochondria isolated from gastrocnemius muscles. Amplex Red (ThermoFisher Scientific; Waltham, MA) and horseradish peroxidase (HRP) were added to the respiration chamber followed by the addition of 20 uL of isolated mitochondria. 20-70 uM of palmitoyl-CoA was titrated into the respiration chambers and the amount of fluorescence assessed.
**Citrate Synthase Activity**

Citrate synthase activity was measured in gastrocnemius, liver, and heart mitochondria as previously described[233,258].

**Skeletal Muscle Mitochondrial Fatty Acid Oxidation**

Complete fatty acid oxidation leads to the production of carbon dioxide while incomplete fatty acid oxidation results in the generation of acid soluble metabolite (ASMs). Complete, incomplete, and total [Incomplete (ASM) + Complete (CO$_2$)] oxidation of (1-\textsuperscript{14}C) palmitate were measured in mitochondria isolated from gastrocnemius muscle as well as in whole quadriceps muscle homogenate as previously described [262].

**Myosin Heavy Chain Immunohistochemical Analysis**

Skeletal muscles can be characterized according to their oxidative capacity as type I slow twitch fibers and type II fast twitch fibers, also known as red and white muscle, respectively [79]. Furthermore, type II fibers can be further classified as type IIa, type IIb, and type IIx representing a fast oxidative, fast glycolytic and combination, respectively. Due to their increased endurance and oxidative properties, type I fibers tend to have more mitochondria relative to other fiber types[78]. Using myosin heavy chain antibodies against type I, IIa, and IIb fibers, oim/oim and WT soleus (sol) and extensor digitorum longus (EDL) muscles were visualized to determine fiber type composition.

Sol and EDL muscles previously stored in optimal cutting temperature (O.C.T.) compound at -80°C were cut to 12 µm sections. The primary antibodies mouse IgG2b BA.D5 (1:100), mouse IgG1 SC.71 (1:200), and mouse IgM BF.F3 (1:100) (DSHB,
University of Iowa, IA) against type I, type IIa, and type IIb fibers respectively were used. Additionally, rabbit IgG anti-laminin antibody (1:200) (ab11575; Abcam) was used to visualize fiber borders. The following Alexa Fluor secondary antibodies were used: 647 goat anti-mouse IgG2b (1:250), 488 goat anti-mouse IgG1 (1:500), 555 goat anti-mouse IgM (1:500), and 350 goat anti-rabbit IgG (1:400) (Invitrogen, IA, USA). Images were captured using the Zeiss Axiovert 200M with fluorescence and ORCA-ER camera and analysis was done using Image J software.

**Glucose Tolerance Testing**

Approximately two weeks prior to sacrifice, mice were fasted for six hours and then injected with 2 mg/kg glucose. Duplicate glucose measurements were taken and averaged using the ReliOn™ PRIME (Walmart) glucose monitoring system at the following time points: 0, 15, 30, 60 and 120 minutes.

**Metabolic Chambers and Body Composition Analysis**

Approximately one week prior to sacrifice, mice were placed in metabolic monitoring systems (PromethION; Sable Systems International). Mice were singly housed and given *ad libitum* access to food (stock chow, Purina Lab Diet 5053) and water. All metabolic chamber data are presented as day and night corresponding to the 12-hour diurnal and nocturnal blocks, respectively. Mice were placed in the chamber on the morning of day 1 and allowed to acclimate until the following morning; data from the subsequent 48 hours, 2 full day and night cycles (days 2 and 3), were analyzed and averaged for each animal. Mice were removed from the chambers on day 4. Data regarding oxygen consumption and carbon dioxide production were collected to determine energy expenditure.
(calculated based on the Weir equation [263]) and respiratory quotient (RQ; calculated as a ratio between oxygen consumption and carbon dioxide utilization [264]) via the PromethION software. RQ gives a sense of fuel utilization with a value of 1 indicative of predominately carbohydrate utilization and a value of 0.7 indicative of predominately fat utilization [264]. Food, water, and body weights were also monitored by the system in addition to activity being measured in three dimensions, in the x, y, and z planes, with the z indicating vertical standing on hindlimbs. All data are presented as absolute values normalized to body weight. Lean body mass and fat mass were determined via EchoMRI™ (Houston, USA) prior to mice being placed in the metabolic chamber.

**Fat Pad Analysis**

Following sacrifice, the inguinal white adipose tissue, gonadal white adipose tissue, and subscapular brown adipose tissue were removed and weighed as previously described [265].

**Serum Markers**

Blood collected via cardiac puncture sat at room temperature for approximately 15 minutes before being placed on ice. All blood was then centrifuged at room temperature in a tabletop centrifuge (15,000 rpm for 15 minutes) and the serum collected and stored at -80°C. Serum was then used to measure levels of total glycerol, free glycerol, triglycerides and non-esterified fatty acids (NEFA). Serum total glycerol, free glycerol, and triglycerides (Sigma-aldrich: St. Louis, MO; cat TR0100) as well as NEFA (Wako Diagnostics: 999-34691, 995-34791) levels were determined following the manufacturer instructions. The serum triglyceride kit measures the amount of free glycerol in the serum
prior to hydrolyzing triglycerides to glycerol and fatty acids at which point the total glycerol is measured again. Triglycerides are determined by subtracting the free glycerol from the total glycerol.

Statistical Analysis

Statistical analyses were performed using GraphPad Prism 8 program (GraphPad Software, Inc., La Jolla, CA, USA). If not mentioned otherwise, data are presented as mean ± SE. Male and female data were analyzed independently. For all experiments (minus glucose tolerance tests), an unpaired two-tailed Student’s t-test is used for comparisons between two groups if normal distribution of residual was confirmed. For non-normal distributions, Mann-Whitney tests are used for comparisons. For glucose tolerance tests (Figure III-7) we performed two-way repeated measures ANOVA with Geisser-Greenhouse correction for sphericity. Differences with p<0.05 were statistically significant.

Results

Mitochondrial respiration, citrate synthase activity and fatty acid oxidation

Previously we demonstrated severe mitochondrial dysfunction in 4-month old male oim/oim gastrocnemius muscle evidenced by reduced gastrocnemius mitochondrial respiration and citrate synthase activity [258]. To determine if this was a global phenomenon or isolated to specific tissues, we investigated mitochondrial respiratory function in the liver and heart as well as gastrocnemius of female oim/oim mice. We found that female oim/oim gastrocnemius mitochondrial respiration was decreased by 27-45% (p=0.05-0.07) with no significant differences in mitochondrial respiration of the
female oim/oim liver or heart (Figure III-1 A-C) when compared to WT littermates suggesting that mitochondrial dysfunction is tissue specific rather than a global phenomenon. Mitochondrial citrate synthase activity was reduced in female oim/oim gast relative to WT littermates, but did not reach significance. Female liver and heart mitochondrial citrate synthase activity was equivalent between oim/oim and WT littermates (Figure III-1D). To further evaluate the skeletal muscle specificity of mitochondrial dysfunction in the oim/oim mouse, we assessed mitochondrial respiration in 2-month old male mitochondria isolated from both gastrocnemius and quadriceps muscles and saw an overall trend of reduced mitochondrial respiration relative to WT counterparts (Figure III-S1). These findings are consistent with the previous findings in 4-month-old male oim/oim and WT gastrocnemius muscle mitochondria and further support compromised mitochondrial function is a generalized oim/oim skeletal muscle phenomenon that can be present as early as 8 weeks of age. Additionally, we evaluated mitochondrial respiration in heterozygous (+/oim) mice, which exhibit much milder muscle weakness as compared to their oim/oim littermates[3] and found no differences between +/oim gastrocnemius mitochondrial respiration relative to WT littermates (Figure III-S2).
Figure III-1: Isolated mitochondria from female *oim/oim* (blue) exhibit reduced (A) gastrocnemius (triangles) mitochondrial respiration, while (B) liver (diamonds) and (C) heart (hearts) mitochondrial respiration remains unchanged as compared to isolated mitochondria from WT (red) littermates. (D) There is no difference in isolated mitochondrial citrate synthase activity between female *oim/oim* and WT gast (triangles), liver (diamonds), or heart (hearts). Data are presented as Means±SE. *p*≤0.05 vs. WT. (n=3-5 per group).
Since skeletal muscle mitochondrial function is reduced in oim/oim mice, we assessed fatty acid oxidation (occurring in the mitochondria) of gast muscle mitochondria and whole quad muscle homogenate. The complete palmitate oxidation to CO$_2$ by gast muscle mitochondria tended to be reduced by 60% (p=0.07) in male oim/oim mice and was reduced by 85% (p=0.05) in female oim/oim mice relative to WT mice (Figure III-2 A), while incomplete and total palmitate oxidation were significantly reduced (p<0.05) in male oim/oim mice only (Figure III-2 B,C). Whole quad muscle homogenate palmitate oxidation was not different between oim/oim and WT littermates regardless of sex (Figure III-2 D-F).
Figure III-2: Isolated mitochondria from male and female oim/oim gastrocnemius (blue squares) exhibit reduced complete oxidation of [1-^{14}C]-palmitate to CO\(_2\) (A), while male oim/oim mitochondria exhibit reduced incomplete oxidation of [1-^{14}C]-palmitate to acid soluble metabolites (ASM) (B) and total oxidation of [1-^{14}C]-palmitate (ASM + CO\(_2\)) (C) relative to WT (red circles) littermates. Complete (D), incomplete (E), and total oxidation (F) of [1-^{14}C]-palmitate in oim/oim whole quadriceps muscle homogenate relative to WT littermates. Data are presented as Means±SE. *p≤0.05 vs. WT. (n=4-7 per group).
Myosin heavy chain analysis of soleus and extensor digitorum longus muscle

In WT C57BL/6J soleus (sol) muscle the distribution of myofibers is reported to be as follows: 43% type I, 46% type IIa, and 11% type IIx, while the distribution in WT C57BL/6J extensor digitorum longus (EDL) is: 0% Type I, 11% Type IIa, and 88% type IIb [266,267]. Since oim/oim mitochondria are severely dysfunctional [258], and oxidative capacity is largely dependent on mitochondria [268,269], a change in myofiber type composition of the oim/oim skeletal muscle was hypothesized. The size of oim/oim soleus muscles was reduced as was the total number of myofibers (p=0.10) when compared to WT littermates (Figure III-3C). Male and female oim/oim mice displayed a 33-50% decrease in the percentage of slow oxidative type I myofibers and a 31% increase in percentages of fast oxidative type IIa fibers relative to their WT littermates, respectively (Figure III-3 A-B, G-H). In the oim/oim male EDL muscle, type IIa myofibers were increased 129% (p=0.16) and type IIb myofibers were decreased 20% (p=0.4) relative to WT littermates, although not reaching significance (Figure III-3 D-E, I-J). Total myofiber count was equivalent between oim/oim and WT EDL muscles regardless of sex (Figure III-3F).
Type I (slow, oxidative), Type IIa (fast, oxidative), Type IIb (fast, glycolytic)
Figure III-3: Male and female oim/oim soleus muscles (blue squares) exhibit reduced percentages of type I myofibers (A) and type IIa myofibers (B) while the total number of myofibers are not altered (C) compared to WT (red circles) littermates. There were no significant changes between WT and oim/oim type IIa myofibers (D), type IIb myofibers (E), or myofiber count (F) in EDL muscles. Representative images of MyHC staining of male WT and oim/oim soleus (G&H) and EDL (I&J) muscles. Data are presented as Means±SE. *p≤0.05 vs. WT. (n=4-5 per group).

Mitochondrial H$_2$O$_2$ emission in isolated mitochondria

Due to the potential role of reactive oxygen species in mitochondrial dysfunction[186], we assessed gastrocnemius mitochondrial H$_2$O$_2$ emission. In male oim/oim gastrocnemius mitochondria (Figure III-4), we found H$_2$O$_2$ production was reduced by 32% (p=0.06) relative to WT littermates.

Figure III-4: Isolated male oim/oim (blue squares) gastrocnemius mitochondria produced reduced amounts of H$_2$O$_2$ when stimulated with palmitoyl-CoA compared to WT isolated gastrocnemius mitochondria (red circles) littermates. Data are presented as Means±SE. *p≤0.05 vs. WT. (n=3-5 per group).
**Body composition**

By EchoMRI, male *oim/oim* mice were determined to have a 3% (p=0.12) increase in lean mass and a 20% (p=0.07) decrease in fat mass relative to WT littermates (Figure III-5 A-B). Female *oim/oim* mice did not exhibit alterations fat or lean mass relative to female WT littermates. Both male and female *oim/oim* mice exhibited reduced relative gonadal white adipose tissue (although only reaching significance in female mice) while relative inguinal white adipose tissue was equivalent between *oim/oim* and WT littermates regardless of sex (Figure III-5 C-D). Subscapular brown adipose tissue (BAT) was not different between WT and *oim/oim* animals.
Figure III-5: Male oim/oim (blue squares) mice exhibit increased percentages of lean mass (A) and reduced percentages of fat mass (B) compared to WT (red circles) littermates. Additionally, relative gonadal fat pad weights were reduced in female oim/oim mice while brown adipose tissue and inguinal adipose depots remained unchanged in oim/oim mice compared to WT littermates (C-D). Data are presented as Means±SE. *p≤0.05 vs. WT. (n=11-17 per group).
**Indirect calorimetry**

WT and *oim/oim* mice were placed in metabolic chambers to determine their whole body energy metabolism; parameters measured included: energy expenditure, activity, and food/water consumption. When normalized to body weight, *oim/oim* mice displayed significantly increased energy expenditure in light and dark cycles (+9-19%), respectively (Figure III-6 A-D). This occurred despite a decrease in physical activity of female and male *oim/oim* relative to sex-matched WT mice during both light and dark cycles (Figure III-S3). Additionally, male *oim/oim* mice consumed ~36% more food per body weight during the 12 hour night cycle compared to their WT littermates (p=0.06; Figure S4).
Figure III-6: When normalized to body weight, both male and female oim/oim (blue squares) mice exhibit increased total (A,C) and average (B,D) energy expenditures for the night (A,B) and day (C,D) cycles when compared to WT (red circles) littermates. Data are presented as Means±SE. *p≤0.05 vs. WT. (n=5-6 per group).
During both day and night cycles, oim/oim animals consumed significantly greater amounts of oxygen per body weight (+9-20%) compared to their WT littermates (Figure III-S5 A&C), and oim/oim mice produced significantly greater amounts of carbon dioxide per body weight (+6-18%) relative to WT littermates during the night cycle (Figure III-S5 B&D). The respiratory quotient was not altered in oim/oim mice relative to WT littermates (Figure III-S5 E&F). Although, high variation and low sample number have likely limited the power of this analysis.
Glucose Tolerance

Despite reduced skeletal muscle mitochondrial respiration (Figure III-1 & S1) and fatty acid oxidation (Figure III-2), glucose tolerance was not different in oim/oim mice compared to WT littermates (Figure III-7).

Figure III-7: oim/oim (blue squares) mice do not exhibit changes in glucose tolerance compared to WT (red circles) littermates. Male (A) and female (B) mice were fasted for six hours, then injected with 2 mg glucose per gram of body weight, and glucose measurements taken at the following time points: 0, 15, 30, 60, and 120 minutes. Data are presented as Means ± SE (n=6 per group).
**Serum markers**

While serum triglycerides and NEFAs did not differ between oim/oim and WT littermates, serum free glycerol was drastically reduced in male oim/oim mice (~68%) relative to WT littermates (Figure III-8).

![Graphs A, B, C, D showing serum markers comparison between male and female WT and oim/oim mice.](image)

**Figure III-8:** Male oim/oim mice (blue squares) exhibit reduced total glycerol (A) and free glycerol levels in serum (B) compared to WT red circles) littermates while no differences were present in serum levels of triglycerides (C) and free fatty acids (NEFA) (D) between oim/oim and WT littermates. Data are presented as Means±SE of the genotype values. *p≤0.05 vs. WT. (n=4-10 per group).
Discussion

Skeletal muscle weakness is extremely prevalent in OI where roughly 80% of patients with mild to moderate types I and IV OI experience muscle force deficits and even higher levels in more severe patients as muscle weakness in OI correlates with the level of ambulation [4,6,48,115,207]. Bone and muscles are highly synergistic tissues that talk to one another via both biochemical signaling and mechanotransduction [57,59,257]. Thus, investigating the mechanisms by which skeletal muscle weakness arises is essential.

Previously, we found that skeletal muscle mitochondrial dysfunction was implicated as part of the pathophysiology of OI in the oim/oim mouse model [258]. Here, we further evaluated the impact and potential links between mitochondrial dysfunction, skeletal muscle weakness and whole-body metabolism. We demonstrate that the oim/oim mitochondrial dysfunction appears tissue specific manifesting in skeletal muscle (gastrocnemius and quadriceps) rather than globally as respiration in heart and liver mitochondria were not compromised in oim/oim mice. Additionally, we found alterations in skeletal muscle and whole body parameters related to energy metabolism.

Skeletal muscle, constituting roughly 50% of body mass, is a significant contributor to the body’s resting metabolic rate [55,270,271]. Yet, despite countless reports of skeletal muscle dysfunction in OI patients, there is a paucity of studies on the role of energy metabolism in OI [4,6,48]. Reports in prepubescent patients and the Col1a1Jrt/+ mouse suggest a metabolic phenotype in the pathophysiology of OI[7,8]. Reports in prepubescent patients suggest a hyper-metabolic phenotype consisting of increased body temperature, heart rate, and respiratory rates [7] while a more recent report by Boraschi-
Diaz et al. implicates whole-body metabolic changes in young mice (4 weeks), including reduced adiposity, increased energy expenditure, increased oxygen consumption, and increased carbon dioxide production, as part of the pathophysiology of OI in the $Col1a1^{Jrt/+}$ mouse model, containing an 18 amino acid deletion in the triple helical domain of the pro-$\alpha$1(I) collagen chain [8,51]. Similarly, our observations also implicate a metabolic phenotype in the $oim/oim$ mouse as evidenced by increased relative lean mass and reduced fat mass (Figure III-5 A-B), increased energy expenditure despite reduced activity (Figure III-6 and Figure III-S3), and increased oxygen consumption and carbon dioxide production (Figure III-S5). Although, the metabolic phenotype observed in the $Col1a1^{Jrt/+}$ mouse was diminished by 8 weeks of age, in the $oim$ mouse this was observed in adult $oim/oim$ mice aged 16-19 weeks.

Despite increased energy expenditures in the $oim/oim$ mouse, we did not observe changes in the respiratory quotient, glucose tolerance, or skeletal muscle fatty acid oxidation suggesting that the $oim/oim$ mice do not exhibit shifts in substrate utilization in response to altered energy metabolism. Additionally, we measured total glycerol, free glycerol, triglyceride, and NEFA levels in serum and found that while serum glycerol was the only marker significantly reduced, triglyceride and NEFA markers also appear slightly decreased in the $oim/oim$ male mouse compared to WT littermates (Figure III-8).

Glycerol, an important gluconeogenic substrate secreted by white adipose tissue [272,273], was shown to be increased in Swedish men with type II diabetes [274]. The decrease in serum glycerol seen in the $oim/oim$ mouse relative to WT littermates is likely associated with the reduced relative fat mass and may be indicative of upregulated
gluconeogenesis, although further studies to better understand the nature of this change are warranted.

Mitochondrial function and content exhibit tissue diversity due to differing cellular needs with cardiac tissue mitochondria having the highest electron transport chain activity relative to either liver or gastrocnemius mitochondria[275–277]. The tissue specific mitochondrial dysfunction observed in the oim/oim mouse may reflect differing tissue response to the type I collagen defect.

Skeletal muscle mitochondrial dysfunction may also reflect the observed change in fiber type distribution observed in the soleus muscle. The soleus muscle, containing the slow oxidative and fast oxidative fibers (type I and IIa, respectively) is considered mitochondria-rich, whereas the extensor digitorum longus (EDL) muscle containing types IIa and IIb fibers (representing fast oxidative and fast glycolytic, respectively), has more variable mitochondrial content[278]. Both male and female oim/oim mice exhibited decreased percentages of type I fibers and increased percentages of type IIa fibers in the soleus muscle relative to WT littermates (Figure III-3). Oim/oim male EDL muscles exhibited increased type IIa fibers and decreased IIb fibers relative to WT littermates, (although not reaching significance), while female fiber type distribution was equivalent between WT and oim/oim (Figure III-3). The decrease in type I fibers of the soleus muscle may serve as a compensatory mechanism for the dysfunctional mitochondria. Moreover, the contribution of reduced activity levels in oim/oim mice (Figure III-S3) to fiber type changes in addition to mitochondrial function cannot be discounted, as activity serves as important regulator of fiber type composition and mitochondrial bioenergetics[279,280].
Reduced activity levels in oim/oim mice have been previously reported[163], but the cause(s) of this reduced activity remain unknown. However, the increased prevalence of fractures in the oim/oim mice is potentially an important contributor to decreased activity levels. Specifically, increased long bone fracture rates of oim/oim mice [39] could potentially explain the reduced rearing activity in the z plane that was observed. It is important to note that skeletal muscle weakness in the oim/oim mouse is intrinsic, as demonstrated by the reduced specific contractile force (muscle contractile force normalized to muscle cross sectional area), suggesting that reduced activity alone is not solely responsible for the muscle weakness[3].

In our current study, we observed that skeletal muscle mitochondrial fatty acid oxidation in isolated oim/oim gastrocnemius mitochondria was reduced, but it was not different in whole quadriceps muscle homogenates relative to WT littermates (Figure III-2). This suggests that the reduction in mitochondrial fatty acid oxidation may be the result of dysfunctional mitochondria, as mitochondria are the primary site of fatty acid beta-oxidation [281]. The fact that fatty acid oxidation in the whole muscle was not altered suggest that there may be an increased number of mitochondria in the oim/oim skeletal muscle as proposed previously [258], which could lead to a compensation effect.

Additionally, we evaluated skeletal muscle mitochondrial ROS production. Although we hypothesized increased hydrogen peroxide as a contributor to mitochondrial dysfunction, we found that hydrogen peroxide emission per amount of mitochondria was reduced in male oim/oim gastrocnemius, which may again be the result of poorer functioning mitochondria (Figure III-4). Future studies are warranted to determine the amount of hydrogen peroxide produced within the whole muscle.
In this study, we provide further evidence that the oim/oim mouse exhibits tissue specific skeletal muscle mitochondrial dysfunction independent of heart and liver tissue as evidenced by changes in skeletal muscle mitochondrial respiration, mitochondrial fatty acid oxidation, and mitochondrial ROS production. Additionally, we suggest that a metabolic phenotype exists in the oim/oim mouse that likely contributes further to the pathophysiology of the disease as evidenced by increases in energy expenditure and altered body composition. Further studies are warranted to determine the nature of the relationship between a type I collagen mutation, mitochondrial defects, and metabolic alterations. Understanding these relationships could lead to novel therapeutics addressing the muscle weakness and bone fragility associated with osteogenesis imperfecta.

Limitations

The limitations of this study include the age of the mice, which were primarily from 13 weeks-19 weeks with a subset at 8 weeks. Narrower age range or an earlier prepubescent age may provide additional information concerning the relationship between metabolic alterations and metabolism during the most vulnerable fracture time period [282]. An additional limitation includes small animal number for certain analyses, including metabolic cages and fatty acid oxidations, which had to be balanced against time and fiscal costs.

Authors' roles: Study design: VLG, LCS, CLP, and RSR. Study conduct: mitochondrial isolation and mitochondrial experiments/respiration (VLG), fatty acid oxidation experiments (VLG), glucose tolerance testing (VLG, EH), metabolic chambers (VLG), fiber type experiments (VLG, EH, TKC), serum measurements (VLG). Data analysis:
VLG, EH, TKC, AO, LCS, RSR, and CLP. Drafting manuscript: VLG, EH, TKC, AO, LCS, RSR, and CLP. Revising manuscript content: VLG, EH, TKC, AO, LCS, RSR, and CLP. Approving final version of the manuscript: VLG, EH, TKC, AO, LCS, RSR, and CLP. CLP takes responsibility for the integrity of the data analysis.
Supplementary Material for Chapter III

**Figure III-S1:** Isolated gastrocnemius (A) and quadriceps (B) muscle mitochondria from 2 month old male *oim/oim* (*blue squares*) mice exhibit reduced mitochondrial respiration when compared to WT littermates (*red circles*). Mitochondrial respiration was measured using the Oroboros Oxygraph-2k (Oroboros Instruments, Innsbruck, Austria) as described in the methods. Data are presented as Means±SE. (n=3-5 per group).
Figure III-S2: Isolated gastrocnemius muscle mitochondria from female heterozygous +/oim (orange triangles) mice do not exhibit mitochondrial dysfunction when compared to WT littermates (purple circles). Gastrocnemius mitochondrial respiration was measured using the Oroboros Oxygraph-2k (Oroboros Instruments, Innsbruck, Austria) as described in the methods. Data are presented as Means±SE. (n=5-6 per group).
**Figure III-S3:** Male $oim/oim$ (blue squares) mice exhibit altered activity levels during the night (A) and day (C) cycles while female $oim/oim$ mice only exhibit changes in Z breaks (rearing) during the day (D) cycle and exhibit no changes during the night cycle (B) when compared to WT (red circles) littermates. Activity levels were assessed using metabolic monitoring systems (PromethION; Sable Systems International). Data are presented as Means±SE. *p≤0.05 vs. WT. (n=5-6 per group).

**Figure III-S4:** Male $oim/oim$ (blue squares) mice exhibit increased food consumption with no changes in water consumption while female $oim/oim$ mice do not exhibit changes
in their eating or drinking habits during either the night (A-B) or day (C-D) cycles compared to WT (red circles) littermates. Food and water consumption were assessed using metabolic monitoring systems (PromethION; Sable Systems International). Data are presented as Means±SE. *p≤0.05 vs. WT. (n=5-6 per group).
Figure III-S5: When normalized to body weight, both male and female oim/oim (blue squares) exhibit increased $V_O_2$ levels for both night and day cycles (A,C) while $V_{CO_2}$ levels increased during the night (B,D) cycle only, relative to WT littermates (red circles). The respiratory quotient, the ratio of carbon dioxide produced to oxygen consumed, remained unchanged during both the night and day cycles (E-F) compared to WT littermates. Oxygen consumption and carbon dioxide production were assessed using metabolic monitoring systems (PromethION; Sable Systems International). Data are presented as Means±SE. *p≤0.05 vs. WT. (n=5-6 per group).
CHAPTER IV

PERSPECTIVES AND FUTURE DIRECTIONS
The aims of this research were to 1) characterize mitochondrial function in the hindlimb muscles of a mouse modeling a severe patient type III OI \((oim/oim)\), 2) determine if mitochondrial dysfunction is a global phenomenon in the \(oim/oim\) mouse, and 3) determine if whole body and skeletal muscle metabolic parameters are altered in the \(oim/oim\) mouse. To this end, we discovered that 1) \(oim/oim\) mice exhibit severe skeletal muscle mitochondrial dysfunction evidenced by drastic reductions in mitochondrial respiration and mitochondrial citrate synthase activity, 2) this mitochondrial dysfunction is not global as reduced mitochondrial respiration is not observed in heart or liver tissue, and finally, 3) a subtle metabolic phenotype is present in the \(oim/oim\) mice including the presence of altered body composition and increased energy expenditure.

These studies were the first to demonstrate the presence of mitochondrial dysfunction in an OI mouse model. However, much more work is needed to fully understand the implications of these findings. The following three objectives outline the important future studies that are necessary to better understand these findings and to move the field forward.

**Objective 1**

Evaluate the role of the extracellular matrix and mitochondrial permeability transition pore induction on mitochondrial dysfunction and skeletal muscle weakness in the \(oim/oim\) mouse.
Hypothesis: The presence of abnormal type I collagen disrupts the interactions and organization within the extracellular matrix of skeletal muscle leading to mitochondrial dysfunction and skeletal muscle weakness in the oim/oim mouse.

Rationale: Type I collagen is the most abundant collagen in both the tendon and perimysium of the skeletal muscle while types I and III collagens are equally abundant in both the epi- and endomysium[92]. The organization of type I collagen differs from tendon and perimysium with the type I collagen in tendon being much more organized than in the perimysium. This likely reflects the different functions of each component[92]. This highly organized hierarchical extracellular matrix surrounding the muscle fibers is believed to help facilitate force transduction with all the different hierarchical levels believed to be interconnected mechanically[92,94]. Oim/oim mice exhibit abnormal type I collagen existing as a homotrimer composed of three alpha I chains, α1(I)₃, rather than normal heterotrimer [α1(I)₂ α2(I)][39]. Additionally, deficits of type VI collagen, an important and abundant extracellular matrix protein found in skeletal muscle, results in two classical myopathies, the milder Bethlem myopathy and the more severe Ullrich congenital muscular dystrophy, both characterized by muscle weakness and hypotonia, distal joint laxity, and proximal joint contractures [180]. Type VI collagen monomers are triple helixes composed of one α1 (VI), α2 (VI), and α3 (VI) chain (occasionally the α3 (VI) chain is replaced by an α4 (VI), α5 (VI), or α6 (VI) chains) [195]. These triple helical monomers form disulfide bonded dimers, and ultimately disulfide bonded tetramers giving type VI collagen a characteristic beaded microfibril pattern [195]. Collagen VI deficiencies in mouse models also lead to skeletal muscle weakness and mitochondrial dysfunction which is hypothesized to arise as a result
dysregulated calcium signaling and ultimately increased induction of the mitochondrial permeability transition pore [181]. The exact structure of the mitochondrial permeability transition pore has yet to be elucidated, however there is evidence implicating cyclophilin D, a prolyl isomerase, as a regulator of the mitochondrial permeability transition pore.

Furthermore, types I and VI collagens have been shown to interact with one another in vitro [283]. Moreover, there is evidence suggesting the interaction between types I, VI, III, and XII collagens in the organization of the perimysium[205]. This organization, especially at the sites where the perimysium converges with the endomysium, a region termed the perimysial junction plate, is believed to contribute significantly to lateral force transmission and ultimately to contractile forced [205]. This link between abnormal type VI collagen and mitochondrial dysfunction leads us to question the role of collagen interactions, such as those of type I and type VI collagen, within the extracellular matrix on skeletal muscle and mitochondrial function.

**Approach:** We will assess induction of the mitochondrial permeability transition pore, indirectly, in gastrocnemius muscle of the *oim/oim* mouse by measuring calcium retention capacity of isolated mitochondria, as excessive calcium is a known inducer of mitochondrial permeability transition. Furthermore, we will quantify cyclophilin D protein content, another known regulator of mitochondrial permeability transition, via western blot. Additionally, we will measure the gene expression and protein content of type VI collagen; specifically, we will measure these parameters in the α1, α2, and α3 chains of type VI collagen. We will evaluate the relationship of homotrimeric type I collagen and normal type VI collagen as compared to normal heterotrimeric type I collagen and type VI collagen by assessing binding strength via microscale
thermophoresis in conjunction with the University of Missouri Molecular Interactions Core.

**Objective 2**

Determine if skeletal muscle weakness and mitochondrial dysfunction are inherent to the oim/oim mouse or acquired as the mice age, as a consequence of inactivity or some other factor.

*Hypothesis:* Abnormal type I collagen in the oim/oim mouse leads to an inherent skeletal muscle weakness and mitochondrial dysfunction that is present at the neonate stage and independent of inactivity.

*Rationale:* Previous work demonstrates that oim/oim mice exhibit reduced specific contractile generating force (contractile generating force normalized to myofiber cross sectional area) suggesting that these mice exhibit skeletal muscle weakness that is independent of other factors that can lead to reduced function as a result of reduced muscle size, such as inactivity. Furthermore, oim/oim mice exhibit reduced activity levels as early as one month of age, and this inactivity may be the result of reduced skeletal muscle weakness.

*Approach:* Using primary myocytes cultured from 7 day old neonatal oim/oim and WT hindlimb muscles, we will assess myotube diameter and contractile function using electric pulse stimulation as previously described[284]. Additionally, mitochondrial function will be assessed using the Seahorse XF Analyzer to measure mitochondrial respiration in myocytes as previously described[285].
Objective 3

Evaluate the role of OI mutation in addition to the relationship between skeletal muscle and mitochondrial function using three distinct mouse models of OI (+/G610C, Col1a1Jrt/+ and CRTAP−/−).

Hypothesis: Mice modeling more severe forms of OI (Col1a1Jrt/+ and CRTAP−/−), similarly to the oim/oim mouse, will exhibit compromised skeletal muscle function and reduced mitochondrial function while mice with less impacted muscle function (+/G610C) will not exhibit as poor mitochondrial function.

Rationale: Recent evidence of mitochondrial dysfunction in the oim/oim mouse, including altered mitochondrial number and reduced mitochondrial function, in conjunction with reduced contractile generating force within skeletal muscle suggests mitochondrial dysfunction plays a role in the pathogenesis of OI muscle weakness and warrants the investigation of mitochondrial function in other mouse models of OI exhibiting different mutations and levels of severity[3,258]. +/G610C mice model a mild human type I/IV OI due to a COL1A2 gene variant (a glycine to cysteine substitution) identified in an extensive Old Order Amish kindred of 64 OI individuals in 37 nuclear families[50]. The Col1a1Jrt/+ mouse is the first combined model of both severe type IV OI and Ehlers Danlos Syndrome due to a COL1A1 mutation leading to skipping of exon 9 and subsequent 18 amino acid deletion in the triple helical domain[51]. The CRTAP−/− mice model a recessive type VII OI due to a loss-of-function mutation that results in the loss of 3-hydroxylation of the proline residue at position 986 (P986) in both α1(I) and α1(II) collagen chains[286]. Of these three models, skeletal muscle function has only
been partially evaluated in the $+/G610C$ and $Col1a1^{Jrt/+}$ mouse, with $+/G610C$ mice exhibiting normal muscle size and function, while $Col1a1^{Jrt/+}$ mice exhibit reduced muscle size and function. Furthermore, gastrocnemius mitochondrial respiration was equivalent to WT littermates in the $+/oim$ mouse (Figure III-S1), modeling a mild to moderate patient type I/IV OI, that does not exhibit muscle weakness[3]. This further supports the correlation between muscle weakness and mitochondrial function.

**Approach:** Hindlimb muscle (soleus, gastrocnemius, and tibialis anterior) contractile generating force will be assessed in six male and six female mice by measuring in vivo contractile generating force as previously described[3]. Parameters of mitochondrial function (mitochondrial respiration and citrate synthase activity in isolated mitochondria from gastrocnemius and quadriceps muscles) and mitochondrial content (citrate synthase activity in whole gastrocnemius and quadriceps muscles) will be evaluated in six male and six female mice of the three distinct OI ($+/G610C$, $Col1a1^{Jrt/+}$, and $CRTAP^{-/-}$) mouse models, as well as in their age and sex matched WT littermates. Mitochondria will be isolated from left and right fresh mixed gastrocnemius tissue, and mitochondrial respiration will be evaluated using the Orosboros Oxygraph-2k (Orosboros Instruments; Innsbruck, Austria) via the addition of malate and glutamate (steady-state oxygen flux), titration of ADP (oxygen flux through complex I), titration of succinate (oxygen flux through complex I + II), and addition of FCCP (uncoupled respiration). Finally, cytochrome C will be added to assess the quality of mitochondrial isolation. Citrate synthase activity will be determined in mixed quadriceps muscle and in mitochondria isolated from mixed gastrocnemius muscle as a measure of mitochondrial content and
function, respectively. This study will begin to elucidate if there is a severity and/or mutation specific relationship between muscle strength and mitochondrial function in OI.

**Significance and impact**

The most prominent manifestation of OI is bone fragility, with clinicians and researchers only recently beginning to consider skeletal muscle weakness as part of the pathophysiology of OI. Historically skeletal muscle weakness in OI has been attributed to inactivity, but recent research has demonstrated inherent muscle weakness in both mouse models and OI patients, including the finding that 80% of patients with mild OI exhibit muscle force deficits\[3,4,6,47,48\]. The findings presented in this dissertation were aimed to begin elucidation of the mechanisms responsible for skeletal muscle weakness in OI by characterizing mitochondrial health and function in a mouse modeling a severe human type III OI. These future proposed objectives will lead to a better and more comprehensive understanding of the relationship and mechanisms underlying mitochondrial function and skeletal muscle weakness in OI. Understanding these mechanisms are crucial to identifying novel therapeutic targets addressing muscle weakness in OI, which will ultimately lead to improved bone strength.
BIBLIOGRAPHY


[21] C.A. Miglierati, M.A. Siegel, L.S. Elting, Bisphosphonate-associated


[41] T. Pihlajaniemi, L.A. Dickson, F.M. Pope, V.R. Korhonen, A. Nicholls, D.J.


https://doi.org/https://doi.org/10.1016/j.bone.2015.08.001.

https://doi.org/10.1002/ajmg.c.31548.


https://doi.org/10.1007/BF00302070.


[88] B.D. Hugh Huxley, D. Jean Hanson, Changes in the Cross-Striations of Muscle during Contraction and Stretch and their Structural Interpretation, 1880.


Bone is functionally impaired in dystrophic mice but less so than skeletal muscle, Neuromuscul. Disord. 21 (2011) 183–193.

https://doi.org/10.1016/j.arr.2015.03.002.

https://doi.org/10.1016/j.pbiomolbio.2015.11.005.


https://doi.org/10.1016/S0070-2153(05)73002-7.


[125] K.L. Troy, M.E. Mancuso, T.A. Butler, J.E. Johnson, Exercise early and often:


https://doi.org/10.1038/nm.3917
https://www.nature.com/articles/nm.3917#supplementary-information.


https://doi.org/10.1155/2014/261545.

[147] V. Singhal, E.A. Lawson, K.E. Ackerman, P.K. Fazeli, H. Clarke, H. Lee, K. Eddy, D.A. Marengi, N.P. Derrico, M.L. Bouxsein, M. Misra, Irisin levels are lower in young amenorrheic athletes compared with eumenorrheic athletes and non-athletes and are associated with bone density and strength estimates, PLoS


[152] G. Karsenty, P. Mera, Molecular bases of the crosstalk between bone and muscle,


[169] I. Grafe, S. Alexander, T. Yang, C. Lietman, E.P. Homan, E. Munivez, Y. Chen,


https://doi.org/10.1038/ng1270
https://www.nature.com/articles/ng1270#supplementary-information.


osteogenesis imperfecta, Bone. (2020) 115703.


https://journals.lww.com/acsm-mss/Fulltext/1986/06000/Pathophysiological_factors_which_limit_the.4.aspx.


[283] P. Bonaldo, V. Russo, F. Buccioiti, R. Doliana, A. Colombatti, Structural and
Functional Features of the α3 Chain Indicate a Bridging Role for Chicken Collagen VI in Connective Tissues, Biochemistry. 29 (1990) 1245–1254. https://doi.org/10.1021/bi00457a021.


VITA

Victoria Lynn Gremminger was born January 4th, 1994 in St. Louis, MO and is the daughter of Patty Brooks and Brett Grill, step-daughter of Pam Grill and William Brooks, sister of Samantha and Alexandria Grill, wife of Thomas Gremminger and mother of Augustus Gremminger. Victoria first discovered her passion for science while taking biology and chemistry courses at Windsor High School. She decided to attend Missouri University of Science and Technology where she received a Bachelor of Science degree in Biology with a minor in Chemistry. Following her undergraduate studies, Victoria was accepted to the Biochemistry Graduate Program at the University of Missouri to further develop her interests and understanding of Biochemistry. Here, she was fortunate to join the laboratory of Dr. Charlotte L Phillips where she was able to study skeletal muscle weakness and mitochondrial dysfunction in mouse models of osteogenesis imperfecta. Under Dr. Phillips’ guidance, Victoria was able to blossom into an independent researcher while further developing her understanding of osteogenesis imperfecta and technical skills. Victoria intends to pursue a career in industry where she can put her knowledge of bone-muscle biology and crosstalk to use developing novel therapies.