Severe injury-induced osteoporosis and skeletal muscle mineralization: Are these related complications?

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1. Background

Due to advances in critical care medicine, patients now survive severe injuries, such as burn, blast, or polytraumatic injuries, that were once considered fatal. While being fortunate to survive, many patient experience complications during convalescence, including poor tissue repair, and hemostasis. Heterotopic ossification (HO), dystrophic calcification, and bio-mineralization (Muschiitz et al., 2017; Klein, 2006; Vanden Bossche and Vanderstraeten, 2005; Dey et al., 2017). Specifically, patients can experience a loss of mineralization from the skeleton, referred to as severe injury-induced osteoporosis, that diminishes skeletal integrity, and increased risk of fragility fractures; yet they also accrue mineralization in soft tissues, resulting in complications such as heterotopic ossification (HO). The pathophysiology leading to dysregulated biomineralization in severely injured patients is not well defined. It has been postulated that these pathologies are linked, such that mineralization is “transferred” from the bone to soft tissue compartments. The goal of this study was to determine if severe injury-induced osteoporosis and soft tissue calcification are temporally coincident following injury. Using a murine model of combined burn and skeletal muscle injury to model severe injury, it was determined that mice developed significant progressive bone loss, detectable as early as 3 days post injury, and marked soft tissue mineralization by 7 days after injury. The observed temporal concordance between the development of severe injury-induced osteoporosis and soft tissue mineralization indicates the plausibility that these complications share a common pathophysiology, though further experiments are required.

Abstract

Severely injured patients are beleaguered by complications during convalescence, such as dysregulated biomineralization. Paradoxically, severely injured patients experience the loss of bone (osteoporosis), resulting in diminished skeletal integrity and increased risk of fragility fractures; yet they also accrue mineralization in soft tissues, resulting in complications such as heterotopic ossification (HO). The pathophysiology leading to dysregulated biomineralization in severely injured patients is not well defined. It has been postulated that these pathologies are linked, such that mineralization is “transferred” from the bone to soft tissue compartments. The goal of this study was to determine if severe injury-induced osteoporosis and soft tissue calcification are temporally coincident following injury. Using a murine model of combined burn and skeletal muscle injury to model severe injury, it was determined that mice developed significant progressive bone loss, detectable as early as 3 days post injury, and marked soft tissue mineralization by 7 days after injury. The observed temporal concordance between the development of severe injury-induced osteoporosis and soft tissue mineralization indicates the plausibility that these complications share a common pathophysiology, though further experiments are required.

Keywords: Severe injury, Burn, Trauma, Soft tissue mineralization, Heterotopic ossification, Osteoporosis, Severe injury-induced osteoporosis, Biomineralization, Dystrophic calcification.
delays rehabilitation, increases length of hospital stays, and increases the risk of fragility fractures (Kaewboonchoo et al., 2019; Fontaine and Herrmann, 1933). Furthermore, severely injured patients can also experience the accrual of mineralization in soft tissues such as skeletal muscle, resulting in complications such as dystrophic calcification (DC) and heterotopic ossification (HO). A recent report in CTE demonstrated that DC, if persistent within damaged skeletal muscle, is sufficient to promote HO formation (Moore-Lotridge et al., 2019), which clinically is associated with pain, joint dysfunction, and, in severe cases, may necessitate amputation of the affected limb (Dey et al., 2017; Kornhauser et al., 2017). Although estimates vary, it has been reported that up to 80% of patients develop HO after a severe injury. Furthermore, severe injury-induced HO has also complicated more than 60% of the severe orthopaedic extremity injuries during the Afghanistan and Iraq conflicts (Forsberg et al., 2009; Potter et al., 2006).

Following an injury, the acute phase response (APR) is activated in proportion to the injury severity (Gibson et al., 2020; Benvenuti et al., 2017; Baker et al., 2018). To ensure survival, the APR first activates coagulation and an acute inflammatory response to stop bleeding and prevent infection. Proportional to the amount of tissue injury, an exuberant survival inflammatory response following a severe injury can drive systemic changes in the musculoskeletal system (Gibson et al., 2020; Baker et al., 2018; Klein, 2019). Both pre-clinical and clinical studies have examined the molecular connections between traumatic injuries and hypermetabolism, skeletal muscle wasting, and rapid bone turnover (Klein, 2019; Pereira et al., 2005; Klein, 2015; Auger et al., 2017; Hew et al., 2020; Stanojcic et al., 2018). However, few studies have examined the relationship between severe injury-induced bone turnover, resulting in osteoporosis, and soft tissue calcification of injured tissues.

The overarching hypothesis of this study was that a shared pathophysiology exists, such that a “transfer” of mineralization from the bone to soft tissue compartments can occur. To determine if severe injury-induced osteoporosis and pathologic soft tissue calcification are mechanistically linked, we utilized a combined burn and skeletal muscle injury to model of severe injury. Unlike other preclinical severe injury models, burn injuries have a low risk of hemorrhage, thereby allowing for consistent investigation of the late complications of convalescence. This study aimed to examine if severe injury-induced osteoporosis and pathologic soft tissue calcification are temporally coincident. If found true, this temporal relationship may indicate a unifying pathophysiology for consistent investigation of the late complications of convalescence.

2. Materials and methods

2.1. Experimental overview

The overarching goal of this study was to examine, in a murine model of severe injury, if severe injury-induced osteoporosis and soft tissue calcification were temporally coincident, and therefore potentially pathologically connected. To examine this hypothesis, following a burn and/or skeletal muscle injury, bone quality and the presence of soft tissue calcification were assessed using both longitudinal and endpoint measures (Fig. 1).

2.2. Animal husbandry

All animal procedures were approved by Vanderbilt University IACUC and performed in accordance with ethical standards of the institution (M/15/024, M1800154). All studies were conducted in 6-week-old male C57BL/6J mice, weighing 20–25 g. Mice were individually housed in a regulated facility with 12-h light-dark cycle and provided food (5LOD chow) and water ad libitum.

2.3. Cardiotoxin-induced skeletal muscle injury mouse model

Skeletal muscle injury was induced by an intramuscular injection of 40 μL of 10μM Cardiotoxin (CTX, Accurate Chemical and Scientific Corp; Westbury, NY) in the posterior compartments of the left and right lower limbs, as previously described (Moore-Lotridge et al., 2019; Mignemi et al., 2017; Moore et al., 2016).

2.4. Burn injury mouse model

This model was adapted from a separate burn wound model (Patil et al., 2017) and modified here to be conducted in conjunction with or without a skeletal muscle injury to the lower extremity to model severe injury. Following a subcutaneous injection of buprenorphine (0.5-mg/kg) 30 min prior to the burn procedure, mice were anesthetized with isoflurane and the dorsal hair was removed (Fig. 1A). Following hair removal, if the mouse was designated to receive a focal skeletal muscle injury, the gastrocnemius and soleus muscles were injected with CTX at this time (Fig. 1B). 1 ml of saline was injected subcutaneously posterior to the spine to protect the underlying spinal column from thermal injury (Fig. 1C). The mouse was then placed in a heat-resistant template (Fig. 1D & E) and partially submerged in a 100-C water bath for 10 s to create a full thickness cutaneous burn covering approximately 30% of the total body surface area (Fig. 1F). Immediately following the burn, the mouse was dried and injected with 2 ml of intraperitoneal fluid resuscitation with lactated Ringer’s solution.

2.5. Radiographs

At 7, 14, 21, and 28 days post injury (DPI), mice were analyzed by digital radiography for the development of mineralization within the injured lower extremity as previously described (Fig. 1G) (Moore-Lotridge et al., 2019; Mignemi et al., 2017; Moore et al., 2016). Soft tissue mineralization was quantified using the previously validated soft tissue calcification scoring system (STiCSS) (Moore et al., 2016). Briefly, the operational definitions of each score are based on the percentage area of soft tissue calcification observed in the posterior compartment of the lower extremity: 0 (0%), 1 (1–25%), 2 (25–49%), 3 (50–75%) and 4 (>75%).

2.6. Longitudinal dual energy X-ray absorptiometry (DXA) imaging

Immediately prior to injury and at 1, 2, 5, 7, 14, 21, and 28 DPI, DXA imaging (Faxitron, LX-60) was obtained with an average of 3 images (Fig. 1G). Mice were laid in the prone position with their muzzle placed in a nose cone to maintain anesthetization. Legs of the mouse were placed laterally away from the body of the animal to obtain a clear image of the femurs. Scans were checked for quality, as movement of the whole femur was selected utilizing the arbitrary ROI analysis tool, and bone mineral density (BMD) values were obtained in g/cm2. Measures from the left and right femurs from a single mouse were averaged together and reported as a single value within. As mice were not fasted as part of this study, vertebral measures were found to be unreliable due to the presence of food pockets (Shi et al., 2016).

2.7. Micro-computed tomography (μCT)

To more sensitively assess changes in bone quality, μCT imaging was performed on the distal femur, distinct from the site of skeletal muscle mineralization at 1, 3, 7, and 28 DPI (Fig. 1G), as previously described (Moore-Lotridge et al., 2019; Mignemi et al., 2017; Moore et al., 2016). The bone volume/tissue volume (BV/TV) ratio, trabecular thickness, trabecular space, and trabecular number were assessed within the metaphyseal region. Average cortical thickness, total cross sectional area, and tissue mineral density were assessed within the diaphyseal...
region (O’Neill et al., 2012). To visualize skeletal muscle mineralization, μCT imaging of the gastrocnemius and soleus muscles were conducted at 7 DPI as previously described (Moore-Lotridge et al., 2019; Mignemi et al., 2017; Moore et al., 2016).

### 2.8. Histologic analysis of skeletal muscle mineralization

Injured skeletal muscle was isolated at 7 and 28 DPI (Fig. 1G) and routinely fixed in 10% neutral-buffered formalin, processed, and embedded (Moore-Lotridge et al., 2019). 6 μm sections were stained with hematoxylin and eosin (H&E) or von Kossa to visualize mineralization (Moore-Lotridge et al., 2019; Mignemi et al., 2017). At least 5 mice were analyzed per cohort/timepoint with >3 sections per mouse. Images included in manuscript represent the average result per cohort.

### 2.9. Statistical analysis

Differences in BMD from DXA assessment were assessed utilizing a 2-way ANOVA corrected for multiple comparisons. Differences in endpoint μCT parameters were assessed individually at each time point between cohorts and between 1 and 28 DPI in the same cohort by a non-parametric Mann-Whitney U test given that they are discrete samples. Difference in STICSS Score was analyzed with a non-parametric 1-way ANOVA corrected for multiple comparisons. Statistical analyses were performed in GraphPad Prism (v6, GraphPad Software, La Jolla, CA) with α=0.05. Two-sided testing was applied throughout. N>3 for all timepoints assessed. Significance reported was relative to adjusted p values where appropriate.
3. Results

3.1. Severe injury-induced osteoporosis

In mice receiving a combined burn and CTX muscle injury (model of severe injury), significant bone loss was observed compared to mice receiving only a CTX muscle injury (lesser injury) (Fig. 2A). Longitudinal DXA imaging detected a significant reduction in average BMD of the left and right femurs by 7 DPI. Given that the mice examined are still undergoing bone development, a significant change in BMD between 5 and 28 DPI (p = 0.050) was observed in mice receiving a CTX injury, indicative of longitudinal bone growth. Alternatively, over the same time period, no change in BMD was observed in mice that received CTX + Burn injury (p = 0.627), leading to the greatest magnitude change in BMD between cohorts at 28 DPI (Fig. 2A). To more sensitively measure changes in bone quality and bone architecture, samples were collected and analyzed by μCT. By 3 DPI, a significant loss of percent metaphyseal bone volume/tissue volume, reduced trabecular thickness and trabecular number, and increase in trabecular space was detected in mice receiving CTX + Burn injury (Fig. 2B & C, Table 1). Assessment of cortical bone volume at 28 DPI between cohorts revealed a significant loss in the average cortical bone thickness in mice receiving CTX + Burn injury compared to unburned controls (CTX-0.157 ± 0.013; Burn + CTX-0.135 ± 0.006; p = 0.001), with no marked changes in total cross-sectional area of the diaphysis (p = 0.279) or tissue mineral density (p = 0.493).

3.2. Skeletal muscle mineralization

By 7 DPI, marked skeletal muscle mineralization was observed in the gastrocnemius and soleus muscles by radiographic analysis and μCT reconstruction in mice that received CTX + Burn injury. Importantly, no significant mineralization was observed at any timepoint in mice that received either a focal skeletal muscle injury (CTX) or a burn injury alone (Fig. 3A). Histological analysis confirmed that the skeletal muscle mineralization observed at 7 and 28 DPI was morphologically indicative of DC (Fig. 3B) (Moore-Lotridge et al., 2019). Longitudinal radiographic analysis demonstrated that the deposited DC, while significantly greater in mice receiving CTX + Burn injury at 7DPI, progressively regressed from injured muscle between 7 and 28 DPI resulting in comparable levels of mineralization to CTX controls by 28 DPI (Fig. 3C).

4. Discussion

This study identified that a temporal concordance exists in which both severe injury-induced osteoporosis and soft tissue mineralization occur. Specifically, progressive bone loss was observed as early as 3 DPI with μCT assessment and a subsequent gain in DC that was radiographically evident by 7 DPI. These results demonstrate the plausibility that a unifying pathophysiology may exist between these complications in which a “transfer” of mineralization from the bone to soft tissue compartments may occur, similar to what has been described previously in cardiac literature (Duke et al., 2016).

Under a homeostatic state, plasma contains near-saturating levels of calcium and phosphate (Moore et al., 2015). These levels are essential to allow for proper cellular signaling and to maintain musculoskeletal integrity. Yet, soft tissues are therefore, likewise exposed to these near-saturating levels and can experience pathologic mineralization if a nidus, such as tissue injury, occurs. Therefore, throughout life the body employs protective mechanisms, such as plasmin, pyrophosphate, and osteopontin, to block the mineralization within soft tissues (Kornhaber et al., 2017; Mignemi et al., 2017; Moore et al., 2015). As we age, in addition to accruing microtissue injuries, the body likewise experiences a diminished capacity to regulate biomineralization, leading to conditions such as calcific artery disease and osteoporosis (Sell and Scully, 1965; Giachelli, 1999; Mitsuyama et al., 2007; TEALE et al., 1989; Ettinger, 1999).
Given these temporal changes under homeostatic conditions, it is also possible that age may influence the body’s capacity to regulate biomineralization following a severe injury. This is observed in both burn patients and those with traumatic injuries, where the incidence of HO is estimated to be lower in pediatric patients compared to adults with injuries of similar severity (Kluger et al., 2006; Hurvitz et al., 1992; Orchard et al., 2015; Gaur et al., 2003; Bossche and Vanderstraeten, 2005). Interestingly, reports from veterans suffering polytraumatic injuries indicate a lower age, specifically <30 years of age, was independently predictive of heterotropic ossification development on multivariate analysis (Forberg et al., 2009). While many mechanisms can influence this change in incidence, further studies characterizing the capacity for protection mechanisms to regulate biomineralization throughout life may provide critical insight and pharmacologic direction.

Severe injuries such as burn or polytrauma can lead to the systemic derangement of multiple body systems including coagulation, inflammation, and the immune system (Muschitz et al., 2017; Klein, 2019; Hew et al., 2020; Smolle et al., 2017; Peterson et al., 2015). Given the quick onset of bone loss following severe injuries, prior studies have attributed bone loss to a variety of mechanisms including endogenous glucocorticoids, production of pro-inflammatory cytokines, progressive vitamin D deficiency, and disruption of calcium homeostasis (Klein, 2019; Muschitz et al., 2016). Independent of the mechanism, numerous studies have illustrated that release of calcium, phosphate, and magnesium from the bone can have systemic effects on muscle physiology, regulation of inflammation, and glucose handling. To date, few investigations have examined how severe injuries can predispose soft tissues to pathologic mineralization. Given the temporal concordance between the development of osteoporosis and soft tissue calcification, it is plausible that a severe injury can drive bone turnover, thereby increasing the availability of calcium and phosphate in the plasma (Klein, 2019; Lisiecki et al., 2018) and 2) simultaneously diminish soft tissue protection mechanisms against calcification (Gibson et al., 2017; Caprini et al., 1977; García-Avello et al., 1998; Ball et al., 2020); thereby establishing an environment in damaged skeletal muscle where DC formation is still functioning in our model. However, if the severity of the injury was increased or comorbidities were added to the model, both of which are linked to reduced macrophage function, DC regression may be impaired, leading to the possible formation of HO.

Table 1

| Analysis between 1 and 28 DPI | CTX ± Burn | CTX | CTX ± Burn | CTX | CTX ± Burn | CTX | CTX ± Burn | CTX | CTX ± Burn |
|-------------------------------|-----------|-----|-----------|-----|-----------|-----|-----------|-----|-----------|
| %BV/TV                        | 13.3 ±    | 2.8 | 16.8 ±    | 0.024 | 11.9 ±    | 2.9 | 15.3 ±    | 0.41 | 5.69 ±    |
| (N=5)                         | 16.6 ± 2.2 |     | 9.5 ± 2.1** |     | 6.3 ± 1.2* |     | 5.8 ± 1.0*** |     | 0.43*** |
| Trabecular thickness (µm)     | 27 ± 3    | 31 ± 10 | 23 ± 3* | 27 ± 3 | 21 ± 2 | 30 ± 4 | 23 ± 1* | 203 ± 31 | 319 ± 47* |
| (N=5)                         | 26 ± 3    |     |     | 21 ± 3 |     |     |     | 170 ± 34 | 376 ± 72*** |
| Trabecular space (µm)         | 181 ± 37  | 150 ± 26 | 225 ± 29** | 203 ± 31 | 319 ± 47* | 170 ± 34 | 376 ± 72*** |     | 0.43*** |
| (N=5)                         | 197 ± 22  |     |     |     |     |     |     |     | 0.998 |
| Trabecular number (1/mm)      | 4.92 ± 0.72 | 5.69 ± 0.46 | 4.06 ± 0.43* | 4.40 ± 0.57 | 2.98 ± 0.38* | 5.14 ± 0.88 | 2.57 ± 0.43*** |     | 0.988 |
| (N=5)                         | 4.54 ± 0.41 | 0.99 |     |     |     |     |     |     | p=0.002 |

2003). Given these temporal changes under homeostatic conditions, it is also possible that age may influence the body’s capacity to regulate biomineralization following a severe injury. This is observed in both burn patients and those with traumatic injuries, where the incidence of HO is estimated to be lower in pediatric patients compared to adults with injuries of similar severity (Kluger et al., 2006; Hurvitz et al., 1992; Orchard et al., 2015; Gaur et al., 2003; Bossche and Vanderstraeten, 2005). Interestingly, reports from veterans suffering polytraumatic injuries indicate a lower age, specifically <30 years of age, was independently predictive of heterotropic ossification development on multivariate analysis (Forberg et al., 2009). While many mechanisms can influence this change in incidence, further studies characterizing the capacity for protection mechanisms to regulate biomineralization throughout life may provide critical insight and pharmacologic direction.

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Injury is an essential component of skeletal muscle mineralization and the formation of HO. While clinical studies have detected the development of mineralization in muscle following a burn injury (Kornhaber et al., 2017), the location of these mineralized deposits is often unpredictable, making them difficult to reliably detect and track longitudinally. Unlike patients clinically, the ability to control essential variables is possible in a pre-clinical animal model. For example, the location of injury can be controlled in a pre-clinical animal model, allowing researchers to better predict the location of skeletal muscle mineralization. Yet, when utilizing an animal model of disease, one should consider the ability of the model to phenocopy the clinical condition and outcomes to support translation of associated findings.

5. Conclusion

By utilizing a murine model of combined burn and skeletal muscle injury as a model of severe injury, this study effectively examined late complications of convalescence, specifically dysregulated biomineralization. This study found that a temporal concordance exists in which both severe injury-induced osteoporosis and soft tissue mineralization occur. Going forward, future studies are required to investigate potential mechanism links between severe injury-induced osteoporosis and soft tissue mineralization, as well as potential therapeutic interventions for mitigating dysregulated biomineralization. The murine model presented within this manuscript can be utilized in such mechanistic studies, providing the foundation for future clinically translatable work.

CRediT authorship contribution statement

SNML contributed to study design, data collection, manuscript preparation, statistical analysis, and figure development.
RI assisted in study design, model development, and data collection.
BHYG and SLP contributed to data collection, manuscript revisions, and figure development. NAM assisted in study design and model development.
HAC assisted with critical manuscript revisions, data analysis, and figure development.
GDH and SU assisted with data collection and manuscript revisions.
JSN provided critical expertise for study design, data analysis, and manuscript revisions.
JGS oversaw the completion of this project, obtained funding, directed study design and data analysis.
All authors reviewed the paper critically for intellectual content and approved the final version. All authors agree to be accountable for the work and to ensure that any questions relating to the accuracy and integrity of the paper are investigated and properly resolved.
Transparency document

The Transparency document associated with this article can be found, in online version.

Declaration of competing interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

Dr. Moore-Lotridge has no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Dr. Ihejirka has no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Ms. Gibson has no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Dr. Posey is a member of the U.S. Air Force. The views expressed in this article are those of the author and do not reflect the official policy or position of the United States Air Force, Department of Defense, or the U.S. Government.

Dr. Mignemi has no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Dr. Cole has no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Mr. Hawley has no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Mr. Uppuganti has no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Dr. Nyman has no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Fig. 3. Assessment of skeletal muscle mineralization in a murine model of combined burn and skeletal muscle injury. A) By 7 DPI, marked skeletal muscle mineralization was radiographically evident by 3D μCT reconstruction. B) Representative histologic analysis is morphologically indicative of dystrophic calcification. N ≥ 3 for each cohort. C) Longitudinal radiographic assessment and STICSS scoring in mice undergoing CTX or CTX + Burn injury. Given that both the left and right lower limb is injury, each point upon the graph represents a single limb. STICSS: CTX: 7DPI - N=12 limbs assessed, CTX + Burn injury: 7DPI - N = 38 limbs assessed, 14 DPI - N = 38 limbs, 21 DPI - N=30 limbs, 28 DPI - N = 20 limbs. Error bars represent median + interquartile range. * notes significance (p) less than 0.05, *** p<0.001.

Transparency document

The Transparency document associated with this article can be found, in online version.
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