Heterotopic Ossification in Wartime Wounds

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Heterotopic ossification (HO) refers to the formation of mature lamellar bone in nonosseous tissue. In the setting of high-energy wartime extremity wounds, HO is expected to complicate up to 64% of patients, has a predilection for the residual limbs of amputees, and remains a significant source of disability. Although the inciting events and the definitive cell(s) of origin continue to remain elusive, animal models and human histology samples suggest that HO formation follows a predictable sequence of events culminating in endochondral ossification. Primary prophylaxis is not medically or logistically practical in most cases because patients have generally sustained massive wounds and are undergoing serial debridements during an intercontinental aeromedical evacuation. Surgical excision of symptomatic lesions is warranted only after an appropriate trial of conservative measures and is associated with low recurrence rates in appropriately selected patients. Future research regarding prognostication and defining the early molecular biology of actobone may permit individualized prophylaxis and development of novel targeted therapies. (Journal of Surgical Orthopaedic Advances 19(1):54–61, 2010)

Key words: heterotopic ossification, trauma, war wounds

The term heterotopic ossification (HO) refers to the formation of mature lamellar bone in nonosseous tissue. In moderate and severe cases, this disorder can lead to significant disability, though most cases are mild and asymptomatic. Classically, HO is associated with severe systemic insults including spinal cord injury, traumatic brain injury, and neoplasm (1–8). Also, HO forms as sequelae to hip arthroplasty and fractures of the acetabulum or elbow, particularly those requiring operative fixation (9–12). These associations imply a relationship between HO and muscle traumatized due to injury and/or surgical dissection (9, 12–20). Less common causes of heterotopic bone formation include the genetic disorders fibrodysplasia ossificans progressiva and progressive osseous heteroplasia (21–23). Although both proven risk factors and genetic predispositions exist, the underlying cause(s) of HO, the initiating molecular biology, and the cellular origin remain largely unknown.

The Combat Wounded Population

Recently, HO has been observed to be more common than previously reported in patients sustaining high-energy wartime extremity wounds (24–26). Blasts and high-velocity projectiles inflict a high percentage of modern war wounds and predominately affect the extremities (27–38). This injury mechanism results in a unique injury pattern — one comprised of severely traumatized limbs, open fractures, and extensive zones of injury with frequent bone and soft tissue loss, often in association with both gross foreign body and bacterial contamination. Serial debridement procedures are performed every 24–72 hours prior to definitive wound closure or coverage in an effort to remove devitalized tissue and gross contamination. Antibiotic-impregnated polymethylmethacrylate beads are routinely used to reduce the bacterial bioburden, as are negative pressure wound dressings. Despite the severity of these injury patterns, patient survival approaches 90%, due in part to improved body armor, the judicious use of tourniquets, and a robust casualty treatment and evacuation system (39).
**4. TITLE AND SUBTITLE**

Heterotopic Ossification in Wartime Wounds

**9. SPONSORING/MONITORING AGENCY NAME(S) AND ADDRESS(ES)**

Naval Medical Research Center, Regenerative Medicine Department, 503 Robert Grant Avenue, Silver Spring, MD, 20910

**12. DISTRIBUTION/AVAILABILITY STATEMENT**

Approved for public release; distribution unlimited

**16. SECURITY CLASSIFICATION OF:**

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Prepared by ANSI Std Z39-18
The incidence of HO in combat-wounded service members has consistently been reported as 63%-64.6%, far greater than that described in civilian trauma centers. Formation of HO in this patient population is associated with blast injuries, a combat-related amputation within the zone of injury, and injury severity scores greater than 16 (24, 26). In contrast, the largest civilian series examining fracture care and HO found that ectopic bone complicated the extremities in 11% of severe traumatic brain-injured patients and 20% of spinal cord injuries (40). Earlier work in civilian patients reported baseline rates of ectopic bone growth in various long-bone fractures, including forearm fractures (20%) (16), femoral shaft fractures (52%) (41), and tibial shaft fractures (0%) (42), all in the setting of significant head injury. Nevertheless, the incidence of clinically relevant or symptomatic HO in this setting is generally considered to be low (7, 43–45).

**Amputees**

The predilection of heterotopic bone for growth within the residual limbs of amputees is an important recent observation (24, 26). Definitive amputations are often performed within or near the zone of injury (which is extensive in blast injuries) in an effort to preserve residual limb length, joint levels, and subsequent function. As a result, there exists a strong association between these injuries and the subsequent development of both radiographic and symptomatic HO (26).

Several grading classification systems exist to classify its formation about the hip, knee, and elbow (5, 9, 10, 20, 46–48). These were later extrapolated to other joints, but none apply or adapt directly to the residual limbs of amputees. For these patients, a classification system, originally described by one of the authors (BKP) has been adopted. The severity of HO is graded using the single radiographic projection (anteroposterior, lateral, or oblique) that maximizes the extent of the ectopic bone within the soft tissues of the residual limb. For example, ectopic bone formation is considered to be mild if it occupies less than 25%, moderate if it occupies 25%–50%, and severe if it occupies >50% of the soft tissues on a single radiographic projection (Fig. 1).

**Basic Science Efforts**

Recent HO research by Gannon and others (49) has successfully identified genetic mutations that localize to chromosome 4q (27–31). Although the BMP4 gene itself does not harbor a genetic mutation, overexpression of BMP4 and its receptor BMPRIA coupled with underexpression of its antagonists is thought to be required for HO formation (49–52). This phenomenon, first identified in patients with fibro dysplasia ossificans progressiva, firmly establishes a link between some forms of HO and traditional osteoblastic signaling. Davis, in association with Gannon (53), further defined the microenvironment by identifying the presence of brown (hypoxic) adipocytes in the early stages of HO development. The hypoxic environment induces both chondrogenesis and neovascularization. The result is an increase in oxygen tension enabling endochondral ossification to occur. Nesti and coauthors (54) isolated a population of mesenchymal progenitor cells present in traumatized muscle. The authors concluded, based on their ability to demonstrate pluripotency, that these cells may play a central role in the pathologic osteogenic response. The team also noted that the progenitor cells derived from traumatized muscle had a certain propensity to become osteoprogenitor cells, more so than those derived from non-age- or sex-matched geriatric bone marrow donors (55). They further concluded that
muscle-derived progenitor cells are the "putative osteo-
progenitor cells that initiate ectopic bone formation in
HO," but provided no suitable justification for this conclud-
ion and thus the matter requires further study. In another
study, Lounev and others (56) implicate progenitor cells
of a vascular lineage. It is therefore plausible that more
than one source of progenitor cells plays a role in the ini-
itation of ectopic bone formation, either as the cells of
origin or the source of the sentinel cellular signals, but
the precise inciting event(s) and cellular origin(s) remain
elusive.

Ongoing studies from our own institutions examine
sera, tissue, and wound effluent from high-energy wartime
extremity wounds. We are developing predictive bio-
marker and gene-based profiles for HO formation in these
patients. These profiles will permit the early identification
of patients most at risk for HO via computer-based algo-

rithms, potentially allowing aggressive primary prophyl-
xsis. We are characterizing the differentiation propensity
and genetic expression of muscle-derived progenitor
cells isolated from high-energy wounds, compared to age- and sex-matched healthy controls. Finally, we
have successfully induced stem-cell production of bone
in vitro utilizing patient sera and wound effluent, with
the composite goal of identifying molecular triggers of HO
production, evaluating therapeutic targets, and developing
and testing novel preventative treatments.

Factors Associated With HO Formation

The Injury Severity Score (ISS) is associated with
the development of HO (24, 57). Critics of ISS utility
as a prognostic factor for HO growth argue that head-
injured patients score higher and therefore are inherently
more likely to develop heterotopic bone. However, Stein-
berg and coauthors (43) reported that the ISS, independent
of a head injury, remained an important predictor of
the development of HO in a civilian trauma popu-
lation after intramedullary nailing of femoral fractures.
These findings add to the growing body of evidence
suggesting that systemic factors, arguably related to the
degree of systemic inflammation, initiate or contribute to
an exaggerated osteogenic response that may ultimately
be responsible for the development of heterotopic bone.

The association between heterotopic bone growth and
the number and method of surgical debridement proce-
dures, including the use of intermediate-pressure pulsatile
lavage irrigation devices and negative pressure wound
therapy, is not well understood. Two recent studies
reported trends toward an association between HO forma-
tion and the number of debridement procedures as well
as the duration of negative pressure dressing therapy
(24, 26). However, these results should be interpreted
with caution because the increases in both the number
of debridement procedures and the duration of nega-
tive pressure dressing therapy are ostensibly also indica-
tors of greater local injury severity; therefore, establish-
ment of a causal linkage between local ectopic bone and
these wound care modalities is difficult and fraught with
confounding factors.

The type of definitive fracture treatment (internal fixa-
tion, external fixation, or amputation) appears unrelated
to the formation of HO in extremity trauma, despite an
historic association with certain surgical approaches to the
hip and acetabulum (9, 11, 15, 20, 58–62). This theoretical
concern has not been borne out in clinical studies of
extremity trauma (24).

Prophylaxis

Several randomized studies have documented the effi-
cacy of primary prophylaxis for the prevention of HO.
This type of prophylaxis is given following high-risk
index procedures, such as revision total hip arthroplasty
or operative fixation of acetabular fractures (63–73).
Typically, 5–10 Gy of local radiation therapy is dosed
in a single fraction, with or without nonsteroidal anti-
inflammatory medication. Nonsteroidal anti-inflammatory
medications alone can be expected to provide a cost-
effective, dose-related decrease in heterotopic bone forma-
tion, though the risk of treatment-related complications
(i.e., gastrointestinal, renal, or hemorrhagic), as well as
patient noncompliance, appears higher (64, 74). Although
some randomized series have demonstrated no difference
in ectopic bone formation between nonsteroidal treatment
and radiation therapy (63, 69, 72), the bulk of the litera-
ture, including two meta-analyses, modestly favors radia-
tion therapy, arguably related to compliance issues with
medical treatment (67, 73, 75, 76). Two randomized series
found no difference between preoperative and postoper-
ative radiation when dosing single fraction of 7–10 Gy,
provided it is given less than 4 hours prior or 48 hours
after surgery (65, 71).

Evidence supporting secondary prophylaxis following
excision of symptomatic HO is lacking. The authors are
aware of no randomized trials of any secondary preven-
tion modality. Nevertheless, the rate of recurrence in the
appropriate surgical candidate is generally accepted to be
low, and the theoretical benefit of secondary prophylaxis
outweighs the risks of symptomatic recurrence for most
patients.

Pitfalls of Prophylaxis

The use of the aforementioned methods of primary and
secondary HO prophylaxis is not without consequence.
Following radiation therapy, wound- and implant-related
complications have been reported (60, 73). Considering the relatively high prevalence of wound and fracture-related complications in patients with high-energy penetrating extremity wounds, external beam radiation is theorized to result in an unacceptably high wound complication rate as well as potential untoward effects on fracture healing. As such, radiation as primary prophylaxis for HO remains highly controversial and is not currently recommended by the authors for use in this patient population.

Nonsteroidal anti-inflammatory drugs (NSAIDs) may also be problematic in certain patient populations. Cyclooxygenase-2 is required for endochondral bone formation, a mechanism critical to the development of heterotopic ossification, as well as early fracture healing (53). Concerns about NSAIDs in an orthopaedic population stem from this blunting of "helpful" inflammation necessary for endochondral ossification (77–81), leading to increased time to union and increase in the number of delayed unions in several studies (77, 78, 80–83). NSAIDs are also contraindicated in patients with intracranial vascular trauma that is common in severe traumatic brain and penetrating head injuries. The potential benefit of NSAIDs for HO prophylaxis must be weighed heavily against potential fracture-related complications. The authors, nevertheless, emphasize the importance of individualizing primary prophylaxis and that the concerns regarding fracture healing are somewhat moot in patients without long-bone fractures, including many amputees.

Etidronate is the only drug FDA approved for the primary prophylaxis of HO and thus warrants discussion. The FDA label states that etidronate is indicated following total hip replacement or spinal cord injury, though the drug has been evaluated off-label in other settings such as civilian orthopaedic extremity trauma and in burns. Etidronate blocks the aggregation, growth, and mineralization of hydroxyapatite crystals, necessary for the formation of heterotopic bone. Early randomized and pseudo-randomized trials demonstrated efficacy (84–89), but only as long as the drug was administered. "Rebound" formation of HO following cessation of therapy was common (84–87, 89), and follow-on studies failed to corroborate earlier results (90–92). In fact, a recent Cochrane database review did not demonstrate pharmacologic efficacy and could not recommend etidronate treatment for the primary prophylaxis of HO (93). Additionally, etidronate is relatively nonselective and inhibits osteoblasts as well as osteoclasts, prompting concerns similar to those applicable to NSAIDs, which are known to delay fracture healing in orthopaedic trauma patients. For these reasons, etidronate is infrequently utilized for primary HO prophylaxis in our patient population.

Treatment

The treatment of heterotopic ossification is individualized. Numerous series in many different patient populations report that most cases are mild and result in little or no functional impairment (10, 11, 14, 15, 17, 46–48, 57, 58, 62, 66–68, 70, 71, 74, 94–102). Moderate to severe cases can be highly debilitating, particularly in perarticular locations or in the residual limbs of amputees (26, 96, 103). Once heterotopic ossification has been identified by plain radiographs, one must assess the impact on the patient’s level of function and activities of daily living. In amputees, it is imperative that other likely sources of residual limb pain, such as painful bursae, myodesis failure, and neuromata, are identified and treated, prior to considering surgical management (104, 105).

Conservative management including rest, local and systemic medications, activity modification, and prosthetic socket/suspension modifications requires a multidisciplinary approach. Close consultation with skilled prosthetists, physical therapists, and physiatrists is critical. Likewise, in nonamputees, alternative causes of pain and functional limitations, including infection, fracture nonunion, and neuropathic pain syndromes, must be evaluated and treated. Surgical excision is reserved for pain, ulceration, or joint stiffness attributable to HO that remains refractory to exhaustive conservative measures.

Timing and Results of Excision

The timing of excision for symptomatic lesions remains controversial. Historically, excision was advocated only after prolonged observation ensuring that the ectopic bone was "mature," as evidenced by quiescent three-phase bone scans and the relative normalization of the serum alkaline phosphatase (106–108). This practice has long been questioned because these measures do not accurately predict recurrence (5). Numerous other studies support earlier excision based on the roentgenographic appearance of the lesion(s) (26, 109–119). This approach has been shown to allow earlier range of motion and return of functional mobility, with recurrence rates similar to that of late excision (110). Garland (5) identified other prognostic factors for HO excision in patients with head injuries, using a classification system based on the patient’s cognitive and physical disability. In his series, motion-related outcomes and recurrence rates were excellent in classes I and II and uniformly poor, with a 100% recurrence rate, in class V. He theorized that the latter group of patients possessed a systemic osteogenic stimulus, possibly the result of a prolonged systemic inflammation, which may persist for years after the initial injury. Knowledge of this can help set patient and family expectations, particularly in cases involving severe traumatic brain injury.
After appropriate patient selection and preoperative counseling, we advocate surgical excision as soon as symptoms warrant following appropriate efforts at conservative management. Regarding the amputee with variable disability due to pain and joint stiffness. Primary prophylaxis via radiation therapy is neither practical nor recommended in patients with high-energy penetrating extremity wounds, though nonsteroidal anti-inflammatory drugs may be effective in carefully selected patients. After an appropriate trial of conservative measures, operative excision of symptomatic heterotopic bone provides generally good results with low recurrence rates in appropriately selected patients treated with secondary prophylaxis. Future research regarding biomarker-based prognostication and identification of initiating chemokines, genes, and cellular origin of ectopic bone may permit individualized prophylaxis and development of novel targeted therapies.

**References**


