

Heterotopic Ossification of the Elbow: A Literature Review

ABSTRACT

Background: Aberrant ectopic bone formation of the elbow is a common clinical presentation after neurologic, burn, and traumatic injuries to the joint. This represents a significant source of patient burden, delayed recovery times and increased medical costs. Although there is an abundance of literature on heterotopic ossification (HO) of the hip, there is little literature on HO of the elbow in comparison. **Aims:** This literature review seeks to summarize consensus regarding the appropriate system of classification, pathophysiology, clinical presentation, risk factors, and prophylactic treatment options associated with HO formation of the elbow. Clinicians may utilize this information to identify high risk patient populations for potential prophylactic therapy to prevent the occurrence/complications of HO at the elbow. **Methods:** A PubMed literature review was conducted using combinations of the key words “heterotopic ossification,” “elbow,” and “fracture/dislocation.” All study types were considered and relevant articles were utilized for this review. **Results:** Higher levels of injury, severe neurologic and burn injuries, delay to surgery, delay in fixation/stabilization of the elbow, multiple surgical treatments, and genetics were correlated with ectopic bone formation. Single dose pre/postoperative radiotherapy with 700cGy or preoperative NSAID regimens were found to be the main prophylactic treatments. **Conclusion:** Clinicians must consider the HO risk profile of their patients as well as the risk factors of treatment before deciding on prophylactic options. Surgical resection is reserved for the most severe cases.

Keywords: [heterotopic, ossification, elbow, fracture, dislocation, injury]

1. INTRODUCTION

Heterotopic ossification (HO) is the abnormal formation of mature and metabolically active lamellar bone in soft tissue[1]. HO most commonly presents after traumatic injury and/or surgery, significant burns and neurological injuries. HO is a significant cause of discomfort, leading to impaired ability to complete daily tasks, complications, and dissatisfaction for patients postoperatively. Additional surgical treatment is often required when joint spaces and/or impinged neuro-vasculature is involved. In one study of 142 patients with elbow fractures and fracture-dislocations, as many as 37% developed HO, with 20% of patients presenting with clinically relevant symptoms and up to 10% requiring additional surgical intervention[2]. The prominence of HO in traumatic and other forms of injury requires a better understanding of factors contributing to ectopic bone formation. Understanding the common clinical presentation and risk factors of HO formation is important in identifying at risk populations for prevention and treatment strategies, as well as minimizing patient burden.

31 There is a paucity of literature on the development and prophylactic treatment of HO
 32 of the elbow. The high incidence of elbow HO formation, combined with patient burden and
 33 high costs associated with additional medical intervention, warrants an in-depth
 34 understanding of HO pathophysiology and understanding of current preventative treatment
 35 modalities other than surgery. This literature review evaluates current research to establish a
 36 consensus on the pathophysiology, presentation, risk factors, and prophylactic treatments
 37 associated with elbow HO.

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 39 **2. Classification**

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 41 In order to systematically categorize HO severity and progression, many different
 42 classification methods have been created. The Brooker classification is popularly referenced
 43 in the literature, but like many other classification systems it was originally used for HO about
 44 the hip. We recommend clinicians instead utilize the Hastings and Graham classification[3]
 45 system which is specific for HO of the elbow and forearm. This offers a standardized
 46 approach to describing HO severity and functional limitation in the clinical setting. Class I is
 47 formation of HO without functional limitation. Class II is HO formation with functional
 48 limitation. Class III is HO formation with associated joint ankyloses. Classes II and III can be
 49 further subdivided into A, B, & C, subcategories that are utilized to describe the plane in
 50 which range of motion is compromised. These classifications may serve useful to identify the
 51 progression of elbow HO in patients, and quickly identify deficits in function. The
 52 classification is summarized in Table 1.

53
 54 **Table 1: Hastings and Graham Classification**

Class I	HO without functional limitation		
Class II	HO with functional limitation (limited ROM)	Class IIA	flexion/extension limitation
		Class IIB	pronation/supination limitation
		Class IIC	Both A and B
Class III	HO with ankylosis	Class IIIA	flexion/extension limitation
		Class IIIB	pronation/supination limitation
		Class IIIC	Both A and B

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 57 **3. Pathophysiology**

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 59 Several mechanisms have been suggested for the multifactorial process of HO bone
 60 formation. Ectopic bone is thought to be the result of mesenchymal stem cells that migrate to

61 areas of insult and are prompted to differentiate into osteocytes[4,5]. This newly formed
62 bone resembles normal bone, but is metabolically hyperactive and lacks a true periosteal
63 layer[1,4]. Studies suggest that many other body processes including the immune system,
64 inflammatory response, and the CNS are involved in bone formation[6]. The impairment of
65 these processes during severe neurologic injury in trauma cases may play a role in the
66 development of ectopic bone formation. However, the exact mechanism of HO formation due
67 to nervous system dysfunction remains unknown.

68 Several authors suggest the role of tissue expression of increased levels of Bone
69 Morphogenetic Protein (BMP), an impaired BMP pathway, and elevated alkaline phosphatase
70 levels (ALP) in the pathogenesis of HO[4,7,8]. BMP is thought to contribute by stimulating
71 the differentiation of pluripotential cells into osteoblast[9]. One of the many roles of ALP is to
72 remove factors that prevent mineralization of bone. One study found a significantly elevated
73 difference in ALP levels in patients that developed HO versus patients who did not,
74 suggesting a possible correlation[10]. Inflammation is also thought to play a pivotal role in
75 the formation of HO. An exact pathway has yet to be identified, but many factors are
76 potentially implicated. Leukotrienes and PGE2 released during the inflammatory process are
77 responsible for increased periosteal lamellar bone formation, and PGE2 specifically is
78 thought to stimulate mesenchymal cells to osteoblasts[11]. Despite the close connection with
79 the inflammatory process, there is a gap in evidence in the current literature on whether
80 elevated inflammatory markers such as c-reactive protein (CRP), creatine kinase (CK), and
81 erythrocyte sedimentation rate (ESR) may be useful in identifying high risk patients and
82 monitoring HO progression. These markers are non-specific for HO. Nevertheless, the
83 majority of cases of HO seem to most commonly be triggered by acute traumatic injury and
84 resultant hyperactive growth and inflammatory conditions. Due to the close relationship of
85 HO and inflammation, prophylactic therapy often focuses on NSAID (Indomethacin)
86 treatment[12,13].

87 There are also rare cases where patients have a genetic predisposition towards the
88 formation of ectopic bone in soft tissue. This could include genetic mutations anywhere
89 along the implicated BMP pathway[7]. Patients with known genetic mutations in the BMP
90 pathway, or family history of conditions such as fibrodysplasia ossificans progressiva [FOP]
91 should be considered prime candidates for prophylactic therapy.

92 **4. Clinical Presentation**

94 Not all cases of HO are clinically significant. Symptoms may range from mild to
95 severe depending on a case to case basis. After surgery or other traumatic event, it can take
96 up to 3-4 weeks for HO formation to occur. Upon the onset of bone formation, patients may
97 typically present with warmth, redness, swelling, and varying degrees of pain (from none to
98 severe)[7]. More often, patients present to the clinic when faced with severe symptoms such
99 as elbow stiffness or contractures, compromised range of motion (ROM), neurovascular
100 compression, pain/discomfort, and in rare cases, bony elbow ankylosis[7,14]. Elbow
101 ankylosis is a more severe clinical finding but can reduce elbow ROM by up to 90%,
102 debilitating the patient[15]. Such symptoms can severely compromise patients' ability to
103 complete even the simplest of daily tasks, interfering with quality of life and impinging on
104 patient independence. Furthermore, these symptoms may be severe enough to warrant
105 surgery (recurrent in some cases), which contributes to increased costs of management.

106 Diagnosis of HO is primarily via clinical findings and confirmed via radiography of the
107 affected area. Ultrasound is a rapid, cost efficient modality that may be utilized to detect
108 early HO, but its efficacy is user dependent and requires a trained operator and experienced
109 radiologist[16]. Triple phase bone scans remain the most sensitive method of detecting early
110 HO and assessing maturity of HO bone formation[17]. MRI and CT scans can be utilized
111 when neurovasculature is at risk of being compromised by HO, and can aid in planning for
112 surgical resection approaches. MRI is useful for identifying well-developed HO, but recent

113 research indicates that CT joint imaging may help in distinguishing early vs late HO in soft
114 tissue[18,19]. The addition of CT scanning allows the operator to recognize early HO foci
115 and differentiate them from other soft tissue lesions. Earlier recognition could identify
116 patients ideal for prophylactic treatment.

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118 **5. Risk Factors**

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120 **5.1 Trauma**

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122 Since HO is a multifactorial disease process, it is difficult to ascertain direct risk factors.
123 The results are often mixed depending on the type of study, the patient population, and the
124 statistical analysis utilized. However, a great majority of the literature agrees that HO
125 formation is generally greater in patients who have previously had HO[20], as well as those
126 who have been exposed to acute traumatic injury, thermal burns, or neurogenic
127 insult[4,7,18,21]. The incidence and severity of HO correlates with the extent of injury and
128 degree of surgical trauma[4]. In acute injury, the presence of fracture and dislocation of the
129 elbow, as well as joint instability is linked to increased risks of HO formation[2,4,12,13].
130 Severe elbow injuries such as open fractures and a delay in fracture fixation were found to
131 be risk factors for HO[2,12,13,22,23]. One study found the surgical approach used, total
132 operating time, formation of a hematoma, extensive dissection and disseminated bone dust
133 to be potentially implicated[4]. The research on this is not conclusive. Multiple studies
134 emphasized delay to surgical treatment of elbow trauma to be a risk factor for
135 HO[2,12,13,23]. This may be the result of longer periods of joint immobilization, which can
136 increase the risk of developing HO[12,20]. Additionally, Wiggers, et al. found that the number
137 of surgeries (within the first 4 weeks) was also an independent predictor based on their 417
138 adult elbow fracture patient sample[23]. They suggested this is due to high muscle
139 manipulation and retraction during operative procedures. Waiting over a week before
140 surgery for fracture fixation was found to result in 10 times the odds of radiographic HO
141 formation, and 7 times the odds of clinically relevant HO formation[12]. Studies further
142 suggest that fixation of unstable fractures within 48 hours of injury may reduce the chances
143 of ectopic bone formation[14,24]. For these reasons, it is important for surgeons to weigh the
144 risks of delayed ORIF and consider early definitive fixation when treating elbow
145 fracture/dislocation injuries.

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147 **5.2 Neurogenic Injury**

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149 The high incidence of HO formation related to neurogenic injuries represents significant
150 risk factors as well. In patients with combined neurological and elbow injuries, one study
151 found the incidence of HO to be up to 70%[4]. Perhaps this is due in part to the high
152 incidence of elbow fracture injuries, accounting for up to 30% of upper limb injuries[12]. A
153 systematic review of clinical reports on 626 patients undergoing HO excision of the elbow
154 found that 55% of cases were in patients with trauma, 28% in burn patients, and 17% in
155 patients with traumatic brain injury[7,25]. In many cases, these injuries may not even directly
156 involve the elbow, yet HO of the elbow is still commonly found[7]. The mechanism behind
157 CNS dysfunction and HO formation remains unclear, but several authors suggest theoretical
158 mechanisms. In patients with head and spinal cord injury, the healing response can often be
159 found to be accelerated[4]. Dysfunction of this pathway is thought to lead to new bone
160 formation in abnormal locations such as joint spaces and soft tissue. Interestingly enough,
161 Bidner et al. found that the serum of patients with head injuries contained increased growth
162 factor activity of cells of the osteoblast phenotype[26]. This suggests a central humoral
163 and/or neurological mechanism involved in enhanced osteogenesis following head/CNS
164 injury[26]. In one study, paroxysmal sympathetic hyperactivity and dysregulation of the CNS
165 as a result of brain injury was found to be associated strongly with HO formation[27]. The

166 authors identified sympathetic hyperactivity as paroxysmal increase in heart rate, respiratory
167 rate, diaphoresis, motor hyperactivity with or without increased blood pressure and/or
168 hyperthermia. Although a strong association was found, a causal role remains to be
169 identified.

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171 **5.3 Burn Injury**

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173 Burn injury is another complex risk factor for HO that also consists of multiple pathways.
174 In a study of nearly 3000 patients, there were 11.5 times higher odds of developing HO if the
175 patient had suffered more than 30% total body surface area burns[28]. A literature review of
176 51 studies on HO and bony ankyloses formation in post burn injuries found incidences
177 ranging anywhere from 0.1 to 35.3%[29]. Similar to neurologic injury, burn injuries activate
178 multiple pathways that induce hyperactive inflammatory and resultant growth responses.
179 Inflammation sets in motion pathways that prepare healthy cells to proliferate and replace
180 dead cells and injured/necrotic tissue and matrix[18]. It may be relevant to note that even in
181 patients without HO formation, severe burns can lead to post burn contractures that limit the
182 effected joint mobility quite significantly, thereby producing similarly debilitating symptoms.
183 This highlights how the elbow is especially susceptible to becoming stiff after injuries. Early
184 mobilization is important in prophylaxis, and active range of motion (AROM) or passive
185 range of motion (PROM) can help prevent stiffness of the elbow joint after injury or
186 surgery[1].

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188 **5.3 Additional Risk Factors**

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190 Other risk factors found to be significant by some studies include male gender[12,20,22],
191 and excessive stretching of affected joints[4]. Demographic data such as age and sex also
192 remain a source of debate in the literature, as some studies report no age[12] or other
193 patient related demographic factors to be significantly related to formation of symptomatic
194 HO[20,23].

195 Genetic risk factors include a statistically significant association amongst three SNP
196 variants (beta2-adrenergic receptor, toll-like receptor 4, complement factor H) to the
197 development of HO or lack of protection against it[6]. Other genetic risk factors may include
198 mutations along the BMP pathway such as those seen in Fibrodysplasia Ossificans
199 Progressiva (FOP) where patients have disseminated HO formation of ligaments and soft
200 tissues[4,8].

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202 **6. Prophylaxis/Treatment**

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204 Physicians can take three overarching approaches to HO management and
205 treatment. One is prophylaxis in high risk patients who have not developed HO but may be
206 likely too. Second, to opt for no treatment in patients whose HO formation is minimal, not
207 interfering with daily activity, or causing pain and/or discomfort. The third and most invasive
208 approach would be surgical treatment and resection of HO in patients with advanced bone
209 formation. This approach should be reserved to patients with significantly limited range of
210 motion, neurovascular impediment, and/or pain and discomfort.

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212 **6.1 Radiotherapy**

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214 Prophylactic treatment can be either radiotherapy or pharmacologic treatment. The
215 accepted approach for radiotherapy currently seems to be 700cGy single-dose radiologic
216 treatment 24 hours preoperatively or within 24-48 hours post operatively[7,14,3-32]. Single
217 dose peri-operative radiation therapy (700cGy) has been reported to reduce HO formation
218 after surgical treatment for elbow fractures[12,30,31,33]. Despite the effective results, these

219 patients are exposed to higher risks of nonunion. Post-operative single radiation therapy was
220 found to potentially play a role in increasing the rate of nonunion at fracture sites[5,34].
221 Hamid, et al. had to terminate their study prematurely due to the significantly higher rate of
222 nonunion in the radiotherapy group[30]. Other potential risks of radiation at the elbow are
223 adverse skin effects such as ulceration and infection[5]. Physicians that choose to utilize
224 radiotherapy for their high risk HO patients must follow up closely due to these potential
225 adverse effects.

226 **6.2 NSAIDs**

227 NSAIDs offer a cheaper alternative to prophylactic care. This is also a better option
228 for patients who do not want to be exposed to radiation therapy. By reducing inflammation
229 and interfering with BMP pathways, NSAID administration has the potential to interfere with
230 the environment conducive to ectopic bone formation[7]. There are a number of
231 recommendations as to the type and dosing of NSAID therapy. Indomethacin is the most
232 commonly used NSAID that can be prophylactic for complex elbow fracture cases[13]. It is
233 typically administered as an oral dose of 75mg two times per day or 25 mg 3 times per day
234 for 3-6 weeks preoperatively. Indomethacin however can be toxic with cardiac risk, GI
235 bleeding, and reduced fracture healing[7,12,35]. Factors to consider before use are patient's
236 hemodynamic stability and cardiac risk status. These patients may be better candidates for
237 radiotherapy. Other options include COX-2 inhibitors, which have less GI risks. In a
238 retrospective review of 152 patients treated prophylactically with COX-2 inhibitor celecoxib,
239 Sun, et al. found more common and severe cases of HO in the untreated group[36]. Their
240 regimen included celecoxib (200mg) administration daily for 28 days and produced a
241 significant difference.

242 Surgical treatment of HO should be reserved for the most severe cases since it is in
243 itself a form of soft tissue trauma. Of the various surgical approaches and fixation options,
244 the least invasive and traumatic resection approach should be selected to optimize recovery
245 and decrease recurrence of ectopic bone formation.

246 **7. Conclusion**

247 Heterogenic ossification is a relatively common clinical finding and can lead to
248 significant patient burden. The highest incidence of HO seems to be related to degree of
249 severity of acute trauma to the elbow and severity of burn or neurological injuries. The
250 pathological mechanism thought to be implicated is an overactive inflammatory response
251 due to injury, leading to hyperactive growth and resultant ectopic bone formation. The
252 overarching trend seems to follow the higher the level of injury and aggravation to soft
253 tissue, the higher the chance of ectopic bone formation. These patients should be screened
254 for prophylactic therapy to prevent HO. Other than traumatic injury, the literature supports
255 delay to surgery, delay in fixation or stabilization of the elbow, multiple surgical interventions
256 and genetics as significant risk factors for HO bone formation. Physicians are recommended
257 to minimize delay to surgical treatment/stability over 48 hours after elbow trauma to avoid
258 increased risks of HO formation. Furthermore, the least invasive surgical approach that will
259 minimize soft tissue manipulation is also recommended. Imaging modalities such as triple
260 phase bone scans, ultrasound and CT can help detect early HO in high risk patients that are
261 candidates for prophylactic treatment, and measure HO severity before considering
262 prophylaxis and/or surgical treatment.

263 There seemed to be mixed or very little to no support for other patient demographics
264 such as age and gender. Despite HO being closely related to the inflammatory response,
265 there is little research showing the utility of monitoring serum inflammatory molecules such
266 as ALP, CRP, CK and ERP to predict risks of HO formation.

271 In regard to prophylaxis, 700cGy seems to be the one of the mainstay prophylactic
272 treatment but has been cited in the literature to be associated with many potential adverse
273 outcomes. NSAIDs are a cheaper alternative. Both therapies however are related to
274 potential increases in fracture healing and present with their own side effect profiles that
275 must be considered on a case by case basis. In high bleed risk patients, radiotherapy may
276 be a better alternative. In hemodynamically stable patients with low cardiac risks and whom
277 may be averse to radiotherapy, NSAIDs offer an effective option.

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284 **COMPETING INTERESTS**

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286 Authors have declared that no competing interests exist.
287 manuscript.

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289 **CONSENT (WHERE EVER APPLICABLE)**

291 No patient consent was needed for this literature review.

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294 **ETHICAL APPROVAL (WHERE EVER APPLICABLE)**

296 No ethical approval was required to conduct this study.

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298 **REFERENCES**

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