PATHOGENESIS AND PREVENTION OF DYSBARIC OSTEONECROSIS

Abstract  Japanese diving fishermen are known to have a high prevalence of dysbaric osteonecrosis. Their typical dive practices are characterized by long exposure to high pressure and by repetitive diving.

The etiology of dysbaric osteonecrosis still remains controversial. Because the development of dysbaric osteonecrosis may occur silently, the diver may be unaware of the actual ischemic event. Moreover, key etiologic evidence is often lost before the diver is examined.

The early stage of dysbaric osteonecrosis was evaluated in four autopsy cases of divers who died of acute decompression sickness. Hypercoagulability of blood was present in these cases.

Hypercoagulability of blood has also been observed in experimental rats after decompression.

Dysbaric osteonecrosis has been experimentally induced in six sheep at the University of Wisconsin – Madison. These findings are reviewed by comparing dysbaric osteonecrosis in both divers and experimental animals.

In conclusion, a bone compartment syndrome and hypercoagulability appear to be pathogenetic factors in dysbaric osteonecrosis.

Key Words: Dysbaric osteonecrosis, Pathogenesis, Bone compartment syndrome, Hypercoagulability, Decompression sickness, Bends

Introduction  The first case of dysbaric osteonecrosis in divers was reported by Grutzmacher in 1941. The prevalence of dysbaric osteonecrosis in Japanese diving fishermen is the highest. In our recent surveys of the Kyushu area, radiological investigation revealed 467 cases (51.6%) osteonecrosis among 905 divers. As we already reported to the past meetings of the United States – Japan Cooperative Program in Natural Resources (UJNR), a characteristic of their diving profile is prolonged hyperbaric exposure and a rapid decompression.

As Elliot and Kindwall (1982) described, prolonged hyperbaric exposure and rapid decompression can provoke a high incidence of limb bends. As we already reported, there appears to be a strong association between limb bends and dysbaric osteonecrosis.

The University of Wisconsin findings of bone necrosis in sheep long bones after hyperbaric exposure provides the basis for a proposed etiological and pathogenic model of dysbaric osteonecrosis.

Our hematological experiments in rats and autopsies in four divers also provides a basis for an etiological and pathogenic model of dysbaric osteonecrosis.

Histopathology of Early Stage Osteonecrosis
This investigation was based on the histopathological examination of femoral heads taken from four divers who died of decompression sickness.

Case 1 presented the most acute clinical course, with fatal chokes. Multiple air bubbles were found in the femoral sinusoids. A characteristic finding was platelet aggregation in the vicinity of the femoral bubbles.

The clinical course of case 2 was also acute with chokes and spinal cord paralysis. Fat emboli were seen in the lung vessels. Air bubbles, platelet aggregation, and an accumulation of fat were seen in the dilated sinusoids.

The diver of case 3 died 5 days after the onset of spinal cord paralysis. An extensive necrosis was noted around the dilated sinusoids. Platelet aggregation and thrombosis were seen in the sinusoids.

The diver of case 4 died 14 days after the onset of spinal cord paralysis. Slight hemorrhage, migration of phagocytes, and fibrosis were found around the femoral sinusoids.

**Experimental Hypercoagulability in DCS**

In 1978 Kitano & Kawashima succeeded in producing DCS in experimental rats. Ten male rats were subjected to a hyperbaric exposure of two hours at 6 atm abs and were quickly decompressed to ambient pressure in five minutes. Platelets were counted before compression and after decompression. Thrombelastographical examination was performed in three rabbits which was subjected to a hyperbaric exposure of one hour at 6 atm abs and decompressed in five minutes to ambient pressure.

Before compression, the average number of platelets was 569,700/mm$^3$. After decompression, the average number of platelets was 491,300/mm$^3$. A statistically significant reduction of the number of platelets was seen.

**Dysbaric Osteonecrosis in Sheep**

Since 1985 when Lehner and Lanphier at the University of Wisconsin (U.S.A.) had succeeded in producing dysbaric osteonecrosis in sheep long bone, Kitano and others performed a histopathological evaluation of the dysbaric osteonecrosis in sheep as the part of cooperative research in UJNR.

Six, 2-year-old, crossbred female sheep underwent a series of 29-30 hyperbaric exposures of 0.5 to 4 hours duration over a 2.5 year period. The maximum pressure of the hyperbaric exposures reached 2.68 atm abs in the 4 hour exposures and 4.31 atm abs in the 0.5 hour exposures.

Macrosopic Findings: Sectioned bone specimens from the sheep femoral contained fatty marrow necrosis which is a common finding in dysbaric osteonecrosis. Fatty marrow necrosis appeared as opaque, shiny, yellow-white inclusions in the marrow cavity.

Histological Findings: Focal liquefaction of fat occurred in the necrotic lesion. Endosteal new bone formation was present in the cortical bone. Necrotic fatty marrow is surrounded by a fibro – osseous layer. Individual necrotic fat cells were enveloped by basophilic material of calcified tissue.

Articular cartilage appeared intact. The histological findings were also essentially identical to those found in Japanese diver’s dysbaric osteonecrosis which was reported by Kawashima (1976) Kitano (1976). However in divers, the prevalence of necrotic foci appeared to be almost as high in the femoral heads as in the distal femoral shafts.

**Discussion**

The etiology of dysbaric osteonecrosis still remains controversial. Osteonecrosis associated with fat embolism of bone was first demonstrated clinically by Jones in 1965 and first confirmed experimentally by Jones in 1966. In our Case 2, and accumulation of fat and aggregation of platelets were noted in the vicinity of air
bubbles, and fat emboli were seen in the lung vessels. Jones proposed three mechanisms which would be potentially capable of producing intraosseous fat embolism and of triggering a process leading to focal intravascular coagulation and osteonecrosis (Fig. 1).

**MECHANISMS OF LIPID METABOLISM RESULTING IN FAT EMBOLISM AND OSTEONECROSIS**

<table>
<thead>
<tr>
<th>Hyperlipemia</th>
<th>Destabilization / Coalescence</th>
<th>Disruption Of</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fatty Liver A</td>
<td>Of Endogenous Plasma Lipoproteina</td>
<td>Depot Or Marrow Fat</td>
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**FAT EMBOLISM Stage O**

**PHASE I**  
Mechanical Intraosseous Vascular Occlusion  
Lipase

**PHASE II**  
Chemikal Free Fatty Acids Increased  
Prostaglandins Increased  
Focal Intravascular Coagulation

**PHASE III**  
Thrombotic Platelet Aggregation  
Fibrin Thrombosis

**OSTEONECROSIS Stage I**  

*Fig. 1. Schematic representation of three mechanisms potentially capable of producing intraosseous fat embolism and triggering a process leading to focal intravascular coagulation and osteonecrosis. (Jones, 1985).*

*Long Hyperbaric Exposure & Rapid Decompression*

*Nitrogen Bubbles*

*Bone Compartment*
Syndrome
Fat
Embolism
Propagation
Microthrombi
Intravascular
Coagulation
acids
Ischemia / hemostasis
Platelet
Activation
Endothelial
Damage:
Free fatty

Osteonecrosis

Fig. 2. Pathogenesis of dysbaric osteonecrosis.

In our experimental DCS in rats, a marked reduction of the number of circulating platelets after decompression was observed. Intraosseous hypercoagulation and fat emboli appeared to be factors also important in the pathogenesis of dysbaric osteonecrosis.

Lehner and Lanphier have proposed a bone compartment syndrome as the primary mechanism in the early pathogenesis of dysbaric osteonecrosis. Nitrogen is about five times more soluble in fat than in water. As Behnke observed, the cortical bone surrounding fatty marrow establishes a semi-closed compartment with a complex vasculature potentially vulnerable to vascular obstruction due to bubble formation.

Hypercoagulation and thrombosis in long bone sinusoids would increase bone marrow pressure and promote a bone compartment syndrome (Fig. 2).

Lehner has proposed prompt recompression therapy to prevent or block the development of dysbaric osteonecrosis. Increased ambient pressure in recompression will reduce the volume of any bubbles within the marrow cavity and thereby decrease tissue pressure within the marrow cavity and thereby decrease tissue pressure within the long bones.

Another proposed therapy to prevent or block the development of dysbaric osteonecrosis involves anticoagulants. Anticoagulants might prevent dysbaric osteonecrosis if it is administered at the time of prompt recompression therapy for limb bends.

Conclusions
1) Prolonged hyperbaric exposures permit the substantial accumulation of dissolved nitrogen in the fatty marrow of the long bones.
2) Decompression may form bubbles in the semi-closed compartment of the long bone. If significant numbers of bubbles form, tissue pressures can rise and ischemia may result. Persistent limb bends may reflect elevated bone marrow pressure and ischemia in the long bones. If the ischemia is sustained, then dysbaric osteonecrosis may result.
3) Persistent ischemia may occur by the proposed mechanism of a bone compartment syndrome in the development of dysbaric osteonecrosis.
4) A bone compartment syndrome may be promoted by intraosseous hypercoagulation and fat embolism. These factors appear to be the pathogenetic mechanisms most likely responsible for dysbaric osteonecrosis.
5) Early recompression therapy together with anticoagulant therapy may prevent the development of dysbaric osteonecrosis.
References

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