Cerebral Fat Embolism: A Neuropathological Study of a Microembolic State

ELIZABETH KAMENAR, M.D., AND PETER C. BURGER, M.D.

SUMMARY Multiple cerebral petechiae associated with intravascular globules of neutral fat and localized primarily within the white matter are distinctive lesions which secure the pathologic diagnosis of cerebral fat embolism. The abundance of these lesions in an unknown, but presumably small, percentage of cases of fat embolism, along with the even more widespread distribution of embolic fat droplets throughout both white and gray matter, suggest that these lesions and emboli must have a profound effect on neurologic function. Nevertheless, respiratory insufficiency is by far a more common clinical manifestation of the fat embolism syndrome and the neurologic involvement of such patients is often attributed to the secondary effects of generalized hypoxia. The following patient with overt respiratory and neurologic symptoms re-emphasizes the direct primary effect of fat emboli within the central nervous system as a cause of white matter hemorrhages and neurologic deterioration. Explanations for the selectivity of the lesions for the cerebral white matter are explored.

FOR OVER A CENTURY, fat embolization has been recognized as a potentially serious, but poorly understood, sequela of skeletal trauma. Relative to the frequent subclinical embolization of fat droplets following fractures, the "fat embolism syndrome" is less common and is unpredictable in the severity with which it affects the major target organs, the lungs and the brain. Because of its greater incidence, the pulmonary, rather than the cerebral, component has received primary attention in clinical and experimental studies which attempt to define the relationship between embolic intravascular fat and clinical disease. In contrast to the lung, however, the brain may present more distinctive, if not diagnostic, pathologic lesions in this syndrome. With the aid of an illustrative patient, this discussion reviews cerebral fat embolism from the perspective of the pathologist and surveys our incomplete understanding of its pathogenesis.

Patient Presentation
The patient was a 45-year-old man who sustained in an automobile accident multiple right-sided rib fractures and a comminuted fracture of the right femur. He did not lose consciousness. After initial treatment for shock at an outside hospital, he was transferred to Duke University Medical Center where his blood pressure on admission was 120/80 mm Hg, pulse 120/min, and respiratory rate 28/min. He was dyspneic, but neurologically intact except for anisocoria (right 4 mm, left 2 mm). Arterial blood gases revealed a Po2 of 51 mm Hg, O2 saturation of 83%, pH of 7.31, HCO3 of 14 mEq/l, and a Pco2 of 28 mm Hg. Bilateral pneumothoraces were noted; he was intubated and chest tubes were placed. Alveolar densities were also seen throughout both lung fields. His right leg was splinted in traction.

Four hours after arrival (12 hours after the accident), he became lethargic and responded to pain with semipurposeful movements of all extremities. His pupils, which were unchanged in size, reacted slowly to light; the fundi were normal. Deep tendon reflexes were 3+ bilaterally except for the right leg which could not be tested. Toes were upgoing and there was sustained ankle clonus on the left. Skull and cervical spine films were normal; a CAT scan revealed a slight compression of the right lateral ventricle which was attributed to cerebral edema. There was no midline shift. Blood gases were Po2 100 mm Hg, pH 7.28, Pco2 34 mm Hg, HCO3 15 mEq/l, and O2 saturation 95%.

Because of abdominal tenderness and a peritoneal lavage which returned bright red blood, an exploratory laparotomy was performed, but no lesions were encountered. A chest x-ray in the recovery room showed bilateral fluffy densities. The patient remained obtunded.

Twenty-two hours after the accident, he developed myoclonic jerks. An EEG was diffusely abnormal with greater amplitudes on the left. Mild hypotension responded to vasopressors. His blood gases deteriorated but with the use of a volume respirator with a 60% FiO2, the Po2 rose to 190 mm Hg. The Pco2, however, was difficult to control and gradually rose to high of 104 mm Hg. The Pco2, however, was difficult to control and gradually rose to high of 104 mm Hg. The patient's vital signs and blood gases were stabilized over the 5 hours preceding his death with the Pco2 in the 50–60 mm Hg range. He then developed bradycardia, asystole, and expired 42 hours after the accident.

Autopsy Findings
Evidence of recent trauma to the face, trunk, and extremities was obvious externally. Multiple posterior rib fractures were present on the right and there was a fracture of the right femoral shaft with soft-tissue swelling and hemorrhage. The 650 gm right lung was crepitant throughout except for a 7.0 \( \times \) 3.0 \( \times \) 1.5 cm firm, hemorrhagic area of the right lower lobe consistent with a contusion. From the remaining parenchyma, pink frothy material could be expressed and there were multiple, widely scattered 3–10 mm hemorrhagic foci within all lobes. The 750 gm left lung...
was similar to the right with multiple small hemorrhages and a 4 × 4 × 3 cm hemorrhagic area anteriorly in the upper lobe. Microscopically, both lungs had intra-alveolar edema and blood in the grossly hemorrhagic areas and mild edema or unremarkable parenchyma elsewhere. Oil red O and osmium stains revealed numerous bilateral, widely distributed intravascular globules of fat within arterioles, capillaries, and venules. The emboli were not concentrated in the hemorrhagic areas. The heart was not dilated and there were no septal defects.

The 1,560 gm brain gave no external evidence of asymmetry or herniation. Innumerable petechiae were dispersed over the pial surface of the cerebellum (fig. 1). The cerebrum was externally unremarkable.

Internally, petechiae up to 2–3 mm in diameter diffusely stippled the white matter of the cerebral hemispheres but conspicuously avoided the cerebral cortex and deep nuclei (fig. 2). A few were subependymal. In the cerebellum, in contrast to the cerebrum, petechiae involved both the white matter and gray matter (fig. 3). They were also randomly scattered throughout the brain stem and rapidly decreased in number through the medulla and spinal cord. Where there was intertwining of nuclei and fiber tracts within the brain stem, it was difficult to assign petechiae to either the white matter or gray matter by gross examination (fig. 4).

Microscopically, the petechiae in any site consisted of ball, ring, or perivascular hemorrhages with their long axes usually parallel to the myelinated fibers. Ball hemorrhages, which varied from 2.0 mm to less than 1.0 mm, were solid round-to-oval collections of erythrocytes occasionally associated with one or more vessels. Macrophages and a few polymorphonuclear leukocytes were noted centrally within many of these hemorrhages. The ring hemorrhages, usually a millimeter or less in size, were also infiltrated by inflammatory cells, but, in contrast, contained a central amorphous area of necrosis in which the remnants of a small vessel were sometimes apparent (fig. 5). Perivascular hemorrhages consisted of a sleeve of red cells about a centrally placed vessel. The distinction between these 3 types of hemorrhages or hemorrhagic microinfarcts was sometimes arbitrary, and it is likely that some ball hemorrhages were tangential sections of the ring type.

All 3 types of hemorrhagic lesions were individually scattered throughout the cerebral and cerebellar white matter, cerebellar cortex, and brain stem with a minor tendency to cluster in a few areas. In the cerebellar cortex, the hemorrhages were slightly more variable in shape as they flattened out along anatomical lines of
Figure 4. Petechiae in the brainstem and spinal cord decrease in number with progression caudally.

The molecular, Purkinje, and granular cell layers. They also involved the cerebellar subarachnoid space. Although the hemorrhages in the brainstem often extended into adjacent gray matter, they seemed to be centered within the white matter. The few hemorrhages noted in the spinal cord, cerebral cortex, and deep nuclear masses did not include those of ring type.

Much less frequent than the hemorrhages were the anemic lesions which were almost as common in the cerebral gray matter as in the cerebral white matter. Elsewhere, their distribution was similar to that of the hemorrhages. In the white matter, they were irregular, often spongy, areas containing eosinophilic swollen axis cylinders surrounded by ballooned myelin sheaths (fig. 6). These varied from 4 mm to less than 1 mm. A few of the larger areas of coagulative necrosis followed the path of, and surrounded, larger penetrating arteries. In the gray matter, the anemic lesions were small, ill-defined, spongy, pale areas which contained hyper eosinophilic, "ischemic" neurons. These anemic microinfarcts were randomly distributed within the cerebral cortex (fig. 7).

Multiple sections of cerebral and cerebellar cortices and of the hippocampi disclosed no laminar necrosis, diffuse Purkinje cell loss, or Sommer sector necrosis suggestive of a systemic hypoxic/ischemic insult.

Figure 5. The erythrocytes of ring hemorrhages surround central zones of necrosis. A vessel cut longitudinally enters the center of the larger lesion (top right) which contains a discrete hole (arrow). This space was previously occupied by fat which has been dissolved out in processing. The smaller lesion is similar. (Hematoxylin and eosin/Luxol fast blue, X 130.)

Figure 6. A focal white matter anemic microinfarct (arrows) is illustrated in the compact fiber tracts of the brainstem. The injured axons are glassy, pink, large, round and surrounded by ballooned myelin sheaths. (Hematoxylin and eosin/Luxol fast blue, X 200.)
FIGURE 7. The anemic microinfarcts of the cortical gray matter (arrows) of fat embolism are characterized by the focal sponginess of the neuropil. Hypereosinophilic neurons may be found in these lesions. (Hematoxylin and eosin/Luxol fast blue, × 80.)

In all paraffin-embedded sections, the presence of embolic fat within lesions or vessels was inferred from the appearance of empty circular or oval spaces which displaced and compressed the surrounding cells and tissue (fig. 5). Oil red O and osmium stains confirmed the presence of discrete fat globules within the hemorrhagic lesions (fig. 8) as well as in small vessels throughout the brain (fig. 9). There was no consistent relationship between fat globules and the anemic microinfarcts, although for some, globules were concentrated in the capillaries at the perimeter. The irregular distribution of fat droplets within a given anatomic region made comparisons somewhat difficult, but they were clearly more numerous in the cerebral cortex than in the underlying deep white matter by a ratio of almost 4:1. This ratio was determined from the total number of fat emboli counted in 20 separate 1 mm² areas at 100× magnification in both the cortical gray and underlying white matter from each lobe of both cerebral hemispheres (table). In the cerebellum, they were present in numbers roughly proportional to the capillary density and were, therefore, most plentiful in the Purkinje and granular cell layers. In any cerebellar layer, most petechiae contained globules of fat. Some of the petechiae in the brain stem were also associated with local fat emboli.

Fat emboli within the microvasculature were best appreciated in a flat retinal preparation stained with oil red O where the emboli in arterioles sat as globules or long cylinders branching and filling successively smaller vessels (fig. 10). Fat droplets were abundant in the renal glomeruli, but were also seen in the myocardium, liver, pancreas, and gastric mucosa. In contrast to the brain, the emboli in these viscera or the retina were not associated with structural alterations of the vessels or parenchyma.

**Discussion**

Since the clinical diagnosis of cerebral fat embolism must usually be made in the presence of hypoxemia, hypotension, and/or possible cranial trauma, it is rarely made with certainty and should be considered only after other causes of post-traumatic neurologic deficit are remedied or excluded. The diagnosis is

<table>
<thead>
<tr>
<th>Table Number of Cerebral Fat Emboli/80 mm³</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td><strong>Cortex</strong></td>
</tr>
<tr>
<td>Right Frontal</td>
</tr>
<tr>
<td>Left Frontal</td>
</tr>
<tr>
<td>Right Parietal</td>
</tr>
<tr>
<td>Left Parietal</td>
</tr>
<tr>
<td>Right Occipital</td>
</tr>
<tr>
<td>Left Occipital</td>
</tr>
<tr>
<td>Right Temporal</td>
</tr>
<tr>
<td>Left Temporal</td>
</tr>
<tr>
<td><strong>Total</strong></td>
</tr>
</tbody>
</table>

*Cortex/White Matter = 3.8*
nevertheless suspected in the classic case of the "fat embolism syndrome" in which long bone fracture and a 12- to 48-hour symptom-free interval are followed by respiratory distress, fever, tachycardia, a characteristic petechial rash, and neurologic dysfunction progressing to coma, and perhaps to death. Some cases may deviate from the classic evolution by the absence or abbreviation of the latent interval, or by a marked predominance of respiratory or, less commonly, neurologic symptomatology. The latter may consist of impairment of consciousness, seizures, pyramidal signs, decerebrate rigidity, and focal neurologic signs. Although there is no single pathognomonic clinical finding, systemic fat embolism may be best demonstrated by ophthalmoscopy or skin or renal biopsy. Detection of fat globules in urine, sputum, cerebrospinal fluid or blood is an unreliable diagnostic aid.

Considerable support for the diagnosis of clinically significant cerebral fat embolism is found at autopsy if gross examination of the brain reveals multiple petechiae in the white matter. Characteristically, these minute hemorrhages are most abundant in the centrum semiovale, digitate white matter, internal capsule, and cerebellar white matter, but may also be noted in the brain stem and spinal cord. Generally, the gray matter is notably spared or only minimally involved but, occasionally, it may be more conspicuously affected. This is particularly true of the cerebellar cortex as seen in the above case. White matter petechiae, however, are not definitive evidence of cerebral fat embolism as they may occur in other conditions including hypoxic-ischemic leukoencephalopathy, acute hemorrhagic leukoencephalitis, white matter watershed infarcts, some cases of head trauma, malaria, air embolism, and intoxication with organic arsenicals, carbon monoxide, and morphine. The firm diagnosis requires confirmation by light microscopy.

As illustrated above, the classic histologic lesions of cerebral fat embolism are ball, ring, and perivascular hemorrhages which are usually apparent to the naked eye as petechiae. The less obvious and usually less frequent anemic lesions become conspicuous on microscopic examination, especially with the application of a myelin stain. Occasionally they may predominate numerically. With time, the hemorrhages also may become less obvious grossly as they undergo resolution. Microscopically, the latter may have a cellular reaction within 2 days consisting mostly of mononuclear cells. This is followed later by the appearance of gitter cells, reactive astrocytes, and hemosiderin. Based on long-term survivals (3 months and 7 years) following presumed cerebral fat embolism, more advanced lesions may be sufficiently contractile in aggregate to produce gross white matter atrophy. Although any of these histologic findings, especially ball and ring hemorrhages, support the possibility of cerebral fat embolism, they are nonspecific and an absolute diagnosis rests on the demonstration of intravascular fat.

The presence of this substance may be inferred from
paraffin-embedded sections by empty round spaces which displace and compress the surrounding cells (fig. 5), but fat stains are needed for confirmation. With oil red O or osmium stains, globules can be visualized within arterioles or capillary-sized vessels throughout the brain and spinal cord. These globules are often located in the center of the ring and ball hemorrhages, but are usually more plentiful within vessels of the gray matter where lesions are less common. The significance of diffuse intravascular fat in the absence of associated hemorrhages or microinfarcts is unclear, although some studies suggest that such involvement could produce neurologic deficit or death.5,6,7 Timing is important in the identification of the intracerebral fat for, in contrast to the lung, hours must usually elapse following trauma before fat is detected in the cerebral vessels. In addition, embolic cerebral fat may largely disappear by the end of the subsequent week.4 The presence of the characteristic CNS petechiae without demonstrable CNS fat emboli should prompt a search for fat emboli in the lung and especially other systemic organs such as the kidney, posterior pituitary, or a flat preparation of the retina.

With the appropriate clinical setting and distinctive gross and microscopic features as described above, cerebral fat embolism is seen by the pathologist as a well-defined clinicopathologic state. There is very little else about fat embolism, either clinically or pathologically, that can be considered as "well-defined." Since the "fat embolism syndrome" more commonly manifests itself as a "respiratory distress syndrome of shock and trauma," the clinical significance of cerebral fat embolism tends to be minimized. Some attribute any neurologic deficit and even the cerebral petechiae to secondary effects of generalized hypoxia resulting from the pulmonary fat embolization.

The pathogenesis of the cerebral lesions associated with the fat embolism syndrome has not been adequately explained; the source of the emboli themselves has even been a matter of debate. Most believe that these globules originate at the site of the fracture and are conveyed by the venous circulation to the lungs where, by passage through small pulmonary vessels, some gain entrance to the systemic circulation.1 However, cerebral fat emboli may produce white matter petechiae in some cases.

Whatever the genesis of fat emboli, the ischemic character of many cerebral lesions suggests that the latter are direct expressions of small vessel occlusion by the fat droplets. The association of the droplets with the lesions has been a consistent observation in most reported cases,4,7,8,46,47 and unless the lesions pre-exist and trap the circulating droplets, a cause and effect relationship seems logical. This hypothesis would receive support if the character and topography of the lesions of cerebral fat embolism were duplicated by some other human microembolic state, ideally one in which concurrent pulmonary emboli and hypoxia could be excluded. Unfortunately, in many cases the lesions associated with these other emboli are incompletely documented or differ in some critical detail from those of fat embolism. Thus, the recorded lesions associated with lipid contrast material48,49 and silicone50,51 are either too few, too anemic, or too indiscriminate in their gray/white involvement to be considered analogues of cerebral fat embolism. Cerebral air embolism varies in the descriptions of its pathology as well as in the apparent validity of the diagnosis.11,24,51,52 However, cerebral fat emboli may occur following cardiopulmonary bypass and produce white matter petechiae as seen in post-traumatic cases.52 Silicone emboli can also occur in this setting and, in our experience, can produce multiple anemic and hemorrhagic lesions in the white matter. Cerebral involvement with malaria, although not embolic, is a diffuse micro-occlusive state with ring and ball hemorrhages preferentially localized to the white matter similar to that of cerebral fat embolism.25,38 This, although there is need for more documentation, there is some evidence that other deformable microemboli may produce white matter petechiae in some cases.

If the cerebral lesions of fat embolism are expressions of ischemia, the preferential localization of the petechiae to the cerebral white matter remains a paradox since it is here that emboli are usually less frequent. Explanations have focused on the gray matter's rich vascularity which could have sufficient anastomotic potential to protect itself against the ischemic effects of microemboli, although such a protection may have an upper limit.38 Although this consideration of the anatomy of the vascular bed would offer the simplest explanation for the white matter predominance of the petechiae, it has also been suggested that the white matter lesions are produced indirectly by cortical emboli through venous sludging and white matter edema.

Experimental studies of fat embolization are numerous, but only a few have investigated the pathogenesis of the cerebral lesions. The latter have included injections of a variety of oily substances given by the carotid, intravenous, or intraarterial routes. Lipids used have included mayonnaise,46 olive oil,46,49 animal fat,41 Wesson oil,43,44 and cottonseed oil.45 Other experiments relevant to cerebral fat embolism are embolization of silicone,50,51 air,57 and non-deformable microspheres of plastic50 and paraffin.57 As a summary of these experiments, it can be said that microembolization to the brains of small laboratory animals may produce focal lesions which are both small and large, anemic and hemorrhagic, and located in both gray and white matter. The types of hemorrhages have not always been well-characterized although ring hemorrhages are seemingly rare. The small anemic type of lesions in both gray and white matter are most common. Thus, based on a limited body of evidence, experimental microembolization produces, in some instances, small focal lesions, sometimes with a hemorrhagic character, but not the white matter predominance of human cerebral fat embolism.

The embolic/ischemic hypothesis of cerebral fat embolism has been modified by suggestions that there
is a synergism between cerebral fat emboli and some other local or systemic influence. Among the latter are increased venous pressure in the superior vena cava secondary to the pulmonary fat embolism, coagulopathy, hypotension, hypoxia, and toxic metabolites of lipids. These secondary factors have not been rigorously studied in the context of either human or experimental cerebral fat embolism, although some have been the subject of much interest with respect to the lung.

The role of free fatty acids has received particular attention with regard to pulmonary fat embolism and warrants consideration in the pathogenesis of the cerebral lesions. Briefly, it suggests that hydrolysis of embolic neutral fat by tissue or circulating lipases releases free fatty acids which exert a toxic effect on the alveolar-capillary membrane. The time required for the accumulation of these acids has been employed to explain the classic latent interval between the patient's injury and the onset of respiratory distress, while the hemorrhagic character of the pulmonary disease is attributed to the aforementioned toxic effect of the free fatty acids on the pulmonary vascularity. The hemorrhagic pulmonary lesions produced by intravenous administration of free fatty acids, and the observation that similar doses of neutral fat have considerably less effect, are proffered as supportive experimental evidence. It must be recognized, however, that hemorrhagic pulmonary lesions can also be produced by mineral oil which is metabolically inert. Furthermore, the lungs of patients with pulmonary fat embolism have no consistent or well-defined changes that distinguish it from other etiologies that are common in the post-traumatic setting.

In the present patient, for example, there was extensive intravascular fat, but relatively minimal intra-alveolar edema and hemorrhage. It seems, therefore, that an endpoint of hemorrhagic pulmonary edema may have been utilized too liberally as an experimental duplicate of the human pathology. In the brain, the fatty acid hypothesis should be accepted with even more reservation since hemorrhagic lesions are rare in the cortex where there are many emboli and common in the white matter where emboli are less numerous. Also, as noted above, silicone embolism may, in some instances, produce similar hemorrhagic white matter lesions. Since cerebral fat embolism is usually associated with, and often overshadowed by, coexistent respiratory insufficiency and other complications of trauma such as shock, investigation of the pathogenesis of the lesions of cerebral fat embolism should consider both the local effects of embolic fat as well as the influences on the brain of hypoxia and hypotension. Although neither this nor most other cases of cerebral fat embolism exhibit the classic features of hypoxia/hypotension such as necrosis in the basal ganglia, Sommers's sector, Purkinje cell layer, or specific cortical laminae, there are rare hypoxic/ischemic states which are characterized pathologically by white matter hemorrhagic infarcts which have similarities to those of cerebral fat embolism. The absence of embolic fat is critical in differentiating these conditions from cerebral fat embolism. These disorders do, however, suggest the need to consider a synergism between hypoxia/hypotension and embolic fat droplets to produce the characteristic topography of cerebral fat embolism, although a simple mechanical occlusive effect of fat droplets may be sufficient explanation.

References

Cerebral fat embolism: a neuropathological study of a microembolic state.

E Kamenar and P C Burger

*Stroke*. 1980;11:477-484
doi: 10.1161/01.STR.11.5.477

*Stroke* is published by the American Heart Association, 7272 Greenville Avenue, Dallas, TX 75231
Copyright © 1980 American Heart Association, Inc. All rights reserved.
Print ISSN: 0039-2499. Online ISSN: 1524-4628

The online version of this article, along with updated information and services, is located on the World Wide Web at:
http://stroke.ahajournals.org/content/11/5/477

Permissions: Requests for permissions to reproduce figures, tables, or portions of articles originally published in *Stroke* can be obtained via RightsLink, a service of the Copyright Clearance Center, not the Editorial Office. Once the online version of the published article for which permission is being requested is located, click Request Permissions in the middle column of the Web page under Services. Further information about this process is available in the Permissions and Rights Question and Answer document.

Reprints: Information about reprints can be found online at:
http://www.lww.com/reprints

Subscriptions: Information about subscribing to *Stroke* is online at:
http://stroke.ahajournals.org/subscriptions/