History of Chronic Subdural Hematoma

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Trephination or trepanation is an intentional surgical procedure performed from the Stone Age. It looks like escaping a black evil from the head. This technique is still used for treatment of chronic subdural hematoma (SDH). Now, we know the origin, pathogenesis and natural history of this lesion. The author try to explore the history of trephination and modern discovery of chronic SDH. The author performed a detailed electronic search of PubMed. By the key word of chronic SDH, 2,593 articles were found without language restriction in May 2015. The author reviewed the fact and way, discovering the present knowledge on the chronic SDH. The first authentic report of chronic SDH was that of Wepfer in 1657. Chronic SDH was regarded as a stroke in 17th century. It was changed as an inflammatory disease in 19th century by Virchow, and became a traumatic lesion in 20th century. However, trauma is not necessary in many cases of chronic SDHs. The more important prerequisite is sufficient potential subdural space, degeneration of the brain. Modifying Virchow’s description, chronic SDH is sometimes traumatic, but most often caused by severe degeneration of the brain. From Wepfer’s first description, nearly 350 years passed to explore the origin, pathogenesis, and fate of chronic SDH. The nature of the black evil in the head of the Stone Age is uncovering by many authors riding the giant’s shoulder. Chronic SDH should be categorized as a degenerative lesion instead of a traumatic lesion.

Key Words: Hematoma, subdural, chronic · Trephination · History · Craniocerebral trauma · Aging.

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lesions.\textsuperscript{26,56} Differentiation is not always possible, however, evidence of healing from the edge of holes clearly implies surgery in living human skull.\textsuperscript{26,56,64} The shapes of the holes depended on the technique and tools.\textsuperscript{6} Previous authors thought trephined skulls were relatively rare in the Far East and China, more trephined skulls have been found in this region than in the rest of the world combined.\textsuperscript{26,39}

Why they trephined the skull from the Neolithic time to Middle age? Since trephined skulls were found in all around the world, it is hard to develop in a certain area, then spread over. It is also hard to think that all trephination had the same purpose. The reasons for trephination and the instruments used for the procedure differ with time and from culture to culture.\textsuperscript{4,26,34} Trephination was performed as part of tribal or superstitious rituals. It was also used as a treatment for a head injury.\textsuperscript{71} Obvious fracture lines were found on many specimens, often coinciding with, or near, the site of the trephine defect.\textsuperscript{2,26} It may be used to treat disorders such as headaches, epilepsy, hydrocephalus and mental disorders. Hippocrates recommend trephination for patients with fissured fractures, bone contusion with or without associated fracture, and head injuries with associated bone contusion with or without fracture. He also described surgical technique of trephination with caution.\textsuperscript{1,26,50}

Surprisingly, Inca surgeons achieved an average survival rate of 50 to 70% of their craniectomy patients, with little incidence of infection or other complications.\textsuperscript{72,74} The outcomes of trepanation in Papua New Guinea were good, in that 70% of patients were thought to survive in 19th century, contrasting with a 75% mortality for cranial surgery in London in the 1870s.\textsuperscript{2,26,72} Surgeons of the ancient Peru executed postmortem trepanations on corpses as a means of better understanding cranial anatomy and improving techniques.\textsuperscript{30} Postmortem trepanations was used as an ancient hands-on workshop.

How they know the trepanations could save the life? It is impossible to prove the fact happen before prehistoric age without any writings, we can guess only. The Stone Age was a clan or a tribal society. The leader of those societies would be a man who was the strongest and the cleverest. The young leader became the aged with an aged brain. Brain atrophy representing the aged brain is an important pre-requisite for development of chronic subdural hematoma (SDH). Asymptomatic chronic SDH can be developed after a trivial injury. If the hematoma became enlarged, typical symptoms of increased intracranial pressure would appear. Shifting of the brain or compression of the motor cortex will brought hemiplegia with loss of consciousness, which might fall down over a sharp stone, which made a natural trephination. The liquefied hematoma would drain out, which resolve the shifting and compression. The old leader recovered from coma and hemiplegia. It looks like a black evil leak out from the head of the leader.\textsuperscript{29} People of the Stone Age might think that the black evil in the head made the leader confused. They saw the evil escaping from the head through a hole made by trepanation (Figure 1).

**Modern History of Chronic SDH**

D’Errico and German\textsuperscript{10} described that the first authentic report of chronic SDH was that of Johannes Wepfer in 1657. He found a large blood-filled cyst beneath the dura on a patient who died after an “apoplectic stroke”. Ninety years later, Morgagni reported a similar finding on a patient who died of an “apoplectic attack”. At that time, this condition was regarded as an apoplectic stroke. In 1817, Houssard described the nature of this condition as the clot and its enveloping membranes. Bayle ascribed the pathophysiology of chronic SDH to ‘chronic rebleeding’ in 1826. However, until 1857, histology and etiology of this lesion remained obscure. Two hundred years after the Wepfer’s report, Virchow described the histology of the membranes and explained their formation. He recognized that hematoma durae matris sometimes was traumatic, but he believed that this lesion was most often caused by chronic inflammation (pachymeningitis chronica hemorrhagica) of the dura with extravasation of blood into the subdural space and formation of a film of fibrin over the inner surface of the dura.\textsuperscript{10,37} Virchow’s hypothesis became widely accepted notion over 50 years.\textsuperscript{75} Successful neurosurgical treatment of chronic subdural hematoma was first reported by Hulke in 1883,\textsuperscript{75} however, this lesion was regarded as an inflammatory disease in a textbook published in 1911.\textsuperscript{10} Trauma got attention as a possible cause of this lesion from late 19th and early 20th century.\textsuperscript{75} In 1914, Trotter\textsuperscript{49} empha-
sized the traumatic etiology. Following Putnam and Cushing’s report in 1925, this lesion has generally been called chronic SDH instead of pachymeningitis hemorrhagica interna. SDH was regarded as a stroke in 17th century, changed as an inflammatory disease in 19th century, and became a traumatic lesion in 20th century. Although the cause of chronic SDH was revealed as trauma, this lesion has still many unrevealed secrets. At first, doctors thought that missed or asymptomatic acute SDHs will be the source of chronic SDH. There was no clear explanation for the latent interval between head injury and onset of symptoms. In 1826, Bayle suggested the chronic rebleeding as a pathogenesis. In 1925, Putnam and Cushing thought that recurrent hemorrhage caused progressive enlargement of the hematoma. However in 1932, Gardner proposed that expansion of an original subdural clot occurred through osmotic attraction of cerebrospinal fluid (CSF) by blood within the semipermeable hematoma neomembranes. Although oncotic pressure theory and effusion theory were proposed in 1934 and 1955 respectively, oncotic pressure theory was a general opinion for about 40 years. Weir discarded osmotic and oncotic pressure theories comparing the osmolality and oncotic pressure of SDH fluid, venous blood, and CSF in 1971 and 1980. However, Weir could not explain the mechanism of hematoma enlargement by himself.

Attempts to produce chronic SDH in experimental animals were usually failed. Injecting blood into the subdural space of mice or dogs was a reasonable method. However, small amount of blood would absorbed, while too much blood killed the animal. In 1972, Watanabe et al could produce a clinical form of chronic SDH by inoculating a clot of blood mixed with CSF. However, Apfelbaum et al failed to prove that CSF was essential to produce chronic SDH. Till the end of 20th century, we could not develop any experimental model demonstrating progressive enlargement of chronic SDH, except a similar pathology of liquefied hematoma enveloped with neomembrane.

Meanwhile, attempts to explore the nature of chronic SDH was continued by studying the structure of the neomembrane and content of the hematoma. In 1975, Sato and Suzuki found repeated microhemorrhage from the capillaries of the outer membrane. They reported that repeated microhemorrhage was responsible to the enlargement of chronic SDH. Markwalder supported that the mechanism of hematoma enlargement was repeated microhemorrhage from the membrane of chronic SDH by reviewing the literature in 1981. Markwalder’s review of rebleeding theory was widely accepted. However, asymptomatic acute SDHs were suspected as the origin of chronic SDH till 1985. Till the end of 20th century, there were controversies on the origin and natural history of this lesion.

Origin of Chronic SDH and Relations of Traumatic Subdural Lesions

There are three kinds of traumatic subdural lesions; acute SDH, chronic SDH, and subdural hygroma (SDG). They
are related to trauma and take place in the subdural space. However, they have clearly different characteristics. Acute SDH is a clot without a membrane, while chronic SDH is a liquefied hematoma with neomembranes. SDG is an accumulation of clear or xanthochromic CSF in the subdural space (Figure 2). Endtz reported that the first description on the SDG was made by Schwencke in 1733. However, nobody concerned on this lesion over a hundred years. Oka et al. reported that Payr presented the first four cases of meningitis serosa traumatica in 1916, while Mayo and Richter reported subdural hydroma and duramater hygroma in 1894 and 1899, respectively. Naffziger proposed a ball valve mechanism for production of SDG in 1924. This flap hypothesis was hard to observe and impossible to explain the low incidence in the young people, while they were more frequently injured. SDG after head injuries became a common lesion after development of computed tomographic scan in 1971. Diagnosis of this lesion became easy and accurate. Although a lot of studies were reported on traumatic SDG after 1980s, the pathogenesis, symptoms, diagnostic criteria, and natural history of this lesion remained obscure. In 1994, we reported a study on the pathogenesis and fate of traumatic SDG (Figure 3). To develop SDG, there should be a sufficient potential subdural space and separation of the dural border cell (DBC) layer. The later alone is impossible to develop SDG, if there is no enough space. Separation of the DBC layer is usually made by trauma, however, it can occur after any surgery opening the skull, dehydration, artificial brain shrinkage, or excessive CSF drainage. SDG actually occurred at both ends of life, before 5 years or after 50 years, where the potential subdural space was enough. Age distribution of SDG is the same as that of chronic SDH, which implies prerequisite of these two lesions is identical. The subdural fluid of SDG is made from CSF by effusion. Most SDG resolve when the brain expansion or absorption exceeds effusion. However, when the brain remains shrunken or effusion exceeds absorption, it will be changed into chronic SDH by the following mechanism. Immediately after the separation, the DBC layer begins proliferation. Fibroblast appeared within 24 hours makes visible outer membrane within a week, and inner membrane around three weeks. These neomembrane envelops the subdural space. In-growth of new vessels will follow, especially along the outer membrane, then bleeding from these vessels occurs. These unresolved SDGs become chronic SDHs by repeated microhemorrhage from the neomembrane. Such a transformation from SDG to chronic SDH was first reported by Yamada et al. in 1979. In 1987, Ohno et al. reported nearly 50% of patients with a SDG developed a chronic SDH. They suggested that a chronic SDH usually developed as a consequence of a traumatic SDG. There were numerous reports observing such transformation.

![Figure 3](image-url)
lesion depends on the interaction of the pre-morbid status, the dynamics of absorption-expansion and maturation of the neomembrane. In 2000, we could confirm that the unresolved SDG is the precursor of chronic SDH. Patients with chronic SDH are prone to fall or slip down. If they slip, even though the injury itself is trivial, it may tear the cortical bridge veins or fragile vessels in the neomembrane. Repeated trauma may cause acute bleeding, which would make a lump or a layer of hyperdensity within hypo- or isodense hematoma. Like the repeated microhemorrhages from the outer membrane, repeated trauma may cause acute bleeding over the chronic SDH as a mechanism of hematoma enlargement. Sometimes repeated trivial trauma may cause a subdural hygroma, which became a chronic SDH. Although the ages of the SDHs were different, such a chronic-on-chronic SDH may produce the mixed density. Such multiple episodes of trivial trauma are hard to remember.

Although the acute SDH is the worst traumatic mass lesion, this lesion may become a chronic SDH if there is a sufficient potential subdural space or the amount of blood is small (Figure 4). The acute SDH was enveloped by a new membrane with time. The clot became liquefied by fibrinolytic activity. Neovascularization of the neomembrane results fragile and permeable vessels, which is easy to bleed. Repeated microhemorrhage is responsible to the enlargement of chronic SDH. Chronic SDH is not a static lesion, but an ever-changing lesion. Any forces to shrink the brain can be the precipitating factors, while the opposite forces to expand the brain will be the inhibiting factors. The initial thin neomembrane becomes matured, organized or even calcified with time. Chronic SDHs enlarge when rebleeding exceeds absorption and they become symptomatic. Symptoms of the chronic SDH depend on the pressure and the reserving capacity of the cranial cavity. A few chronic SDHs may resolve spontaneously. When the neomembrane is matured, the neocapillary becomes no longer fragile. If absorption or brain expansion exceeds rebleeding, the hematoma will disappear. Maturation of the neomembrane and stabilization of the neovasculature eventually bring about spontaneous resolution.

The origin of a chronic SDH is multiple. It can be developed spontaneously, or changed from a SDG or an acute SDH. Although trauma may separate the DBC layer, a significant number of chronic SDHs, often more than a half of cases, lacks any history of trauma. A more important prerequisite is sufficient potential subdural space that is aging, degeneration of the brain. At this time, we should answer whether this lesion is a traumatic lesion or a degenerative lesion? Modifying Virchow’s description,
chronic SDH is sometimes traumatic, but most often caused by severe degeneration of the brain. From Wepfer’s first description in 1657, nearly 350 years passed to explore the origin, pathogenesis, and fate of chronic SDH (Table 1). The cause of SDG was uncovered in 1994. Relations of traumatic subdural lesions and the origin of this lesion was uncovered in 1996 and 1998, respectively. The nature of the evil in the head of the Stone Age is uncovering by authors riding the giant’s shoulder.

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