Management of intracerebral hemorrhage

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Abstract: Currently, intracerebral hemorrhage (ICH) has the highest mortality rate of all stroke subtypes (Counsell et al 1995; Qureshi et al 2005). Hematoma growth is a principal cause of early neurological deterioration. Prospective and retrospective studies indicate that up to 38% hematoma expansion is noted within three hours of ICH onset and that hematoma volume is an important predictor of 30-day mortality (Brott et al 1997; Qureshi et al 2005). This article will review current standard of care measures for ICH patients and new research directed at early hemostatic therapy and minimally invasive surgery.

Keywords: ICH, hemostatic therapy, recombinant factor VII, surgical management

Intracerebral hemorrhage

An intracerebral hemorrhage (ICH) account for only 15% of all strokes but it is one of the most disabling forms of stroke (Counsell et al 1995; Qureshi et al 2005). Greater than one third of patients with intracerebral hemorrhage (ICH) will not survive and only twenty percent of patients will regain functional independence (Counsell et al 1995). This high rate of morbidity and mortality has prompted investigations for new medical and surgical therapies for intracerebral hemorrhage.

Primary ICH develops in the absence of any underlying vascular malformation or coagulopathy. Primary intracerebral hemorrhage is more common than secondary intracerebral hemorrhage. Hypertensive arteriosclerosis and cerebral amyloid angiopathy (CAA) are responsible for 80% of primary hemorrhages (Sutherland and Auer 2006). At times it may be difficult to identify the underlying etiology because poorly controlled hypertension is often identified in most ICH patients. Patients with CAA-related ICH are more likely to be older and the volume of hemorrhage is usually > 30 cc (Ritter et al 2005). Hypertension related ICH is frequently seen in younger patients, involving the basal ganglia, and the volume of blood is usually < 30 cc (Lang et al 2001). However these characteristics are nonspecific and histopathological studies are needed to confirm a definitive diagnosis of CAA or hypertension related ICH. Hypertension causes high pressure within the Circle of Willis resulting in smooth cell proliferation followed by smooth muscle cell death. This may explain why hypertension related ICH are frequently located deep within the basal ganglia, thalamus (Figure 1), cerebellum, pons and rarely the neocortex (Campbell and Toach 1981; Sutherland and Auer 2006). In contrast, preferential amyloid deposition within leptomeningeal and intraparenchymal cortical vessels may explain the reason for large superficial lobar hemorrhages with amyloid angiopathy (Auer and Sutherland 2005). It is important to identify those afflicted with cerebral amyloid angiopathy because of the high risk of recurrent lobar hemorrhage and predisposition for symptomatic hemorrhage with anticoagulants and thrombolytics (Rosand and Greenberg 2000).

Secondary ICH is due to underlying vascular malformation, hemorrhagic conversion of an ischemic stroke, coagulopathy, intracranial tumor, etc. Arteriovenous malformations and cavernous malformations account for majority of underlying vascular malformations (Sutherland and Auer 2006). An AVM (Figure 2) is usually a singular lesion composed...
of an abnormal direct connection between distal arteries and veins. AVMs account for only 2% of all ICH but are associated with an 18% annual rebleed risk (Al-Shahi and Warlow 2001). Cavernous malformations are composed of sinusoidal vessels and are typically located in within the supratentorial white matter. The annual risk of recurrent hemorrhage is only 4.5% (Konziolka and Bernstein 1987). Intracranial aneurysms usually present with subarachnoid hemorrhage but anterior communicating artery and middle cerebral artery may also have a parenchymal hemorrhagic component near the interhemispheric fissure and perisylvian region respectively (Wintermark and Chaalalron 2003). Embolic ischemic strokes can often demonstrate hemorrhagic conversion without significant mass effect (Ott and Zamani 1986). Sinus thrombosis should be suspected in patients with signs and symptoms suggestive of increased intracranial pressure and radiographic evidence of superficial cortical or bilateral symmetric hemorrhages (Canhoo and Ferro 2005). An underlying congenital or acquired coagulopathy causing platelet or coagulation cascade dysfunction can result in ICH. Cogential disorders account for Hemophilia A, Hemophilia B, and other rare diseases. Acquired coagulopathy may be attributed to longstanding liver disease, renal disease, malignancy, or medication. Particular attention has been directed towards oral anticoagulant (OAT) associated hemorrhage due to greater risk for hematoma expansion as well as increased 30 day morbidity and mortality rates (Flibotte et al 2004; Roquer et al 2005; Toyoda et al 2005; Steiner and Rosand 2006). Metastatic tumors account for less than ten percent of ICH located near the grey white junction with significant mass effect. The primary malignancy is usually melanoma, chorocarinomina, renal carcinoma, or thyroid carcinoma (Konziolka and Berstein 1987).

**Clinical presentation**

The classic presentation of ICH is sudden onset of a focal neurological deficit that progresses over minutes to hours with accompanying headache, nausea, vomiting, decreased consciousness, and elevated blood pressure. Rarely patients present with symptoms upon awakening from sleep. Neurologic deficits are related to the site of parenchymal hemorrhage. Thus, ataxia is the initial deficit noted in cerebellar hemorrhage, whereas weakness may be the initial symptom with a basal ganglia hemorrhage. Early progression of neurologic deficits and decreased level of consciousness can be expected in 50%
of patients with ICH. The progression of neurological deficits in many patients with an ICH is frequently due to ongoing bleeding and enlargement of the hematoma during the first few hours (Kazui et al 1996; Brott et al 1997; Fujii et al 1998). Compared with patients with ischemic stroke, headache and vomiting at onset of symptoms is observed three times more often in patients with ICH (Gorlick et al 1986; Rathore et al 2002). Despite the differences in clinical presentation between hemorrhagic and ischemic strokes, brain imaging is required to definitively diagnose intracerebral hemorrhage.

### Diagnosis

Computed tomography (CT) is more widely available so CT of the brain has become the initial diagnostic test of choice for ICH. However, recent studies suggest MRI and CT are equally efficacious in diagnosing hyperacute ICH (<6 hours) (Fiebach et al 2004; Kidwell et al 2004). In addition, in 2004, Fiebach and colleagues conducted a multicenter study and concluded that visual identification of ICH is not difficult with MRI with mean sensitivities = 95% with expert readers as well as final-year medical students (Fiebach et al 2004). MRI and magnetic resonance angiography (MRA) can also help elucidate any underlying cause of the hemorrhage. Sometimes the pattern and topography of bleeding can give important clues about a secondary cause of ICH. For example, subarachnoid blood should raise suspicion for a ruptured aneurysm, multiple inferior frontal and temporal hemorrhages may be seen after head trauma, and fluid levels within the hematoma suggest an underlying coagulopathy (Figure 3). Active contrast extravasation...
into the hematoma seen with CT angiography may predict hematoma expansion and is predictive of poor outcome (Becker et al 1999; Murai et al 1999). Angiography is not required for older hypertensive patients with hemorrhages in deep subcortical structures with no findings suggestive of an underlying structural lesion. Secondary intracerebral hemorrhage should be suspected in patients <45 years of age, no risks for hypertensive hemorrhage, presence of subarachnoid hemorrhage, prominent vascular structures, perisylvian or interhemispheric hemorrhage and angiography should be pursued. Angiography should always be considered in young non-hypertensive patients with ICH who have no obvious explanation for their hemorrhage, or when the only risk factor is cocaine or sympathomimetic-drug use (Halpin et al 1994; Griffiths et al 1997; Zhu et al 1997; Broderick and Adams 1999).

**Management**

**Emergency management**

ICH is a neurological emergency and initial management should be focused on assessing the patient’s airway, breathing capability, blood pressure and signs of increased intracranial pressure. The patient should be intubated based on risk of aspiration, impending ventilatory failure (\(\text{PaO}_2 < 60 \text{ mmHg}\) or \(\text{pCO}_2 > 50 \text{ mmHg}\)), and signs of increased intracranial pressure. (Broderick et al 1999) Emergency measures for ICP control are appropriate for stuporous or comatose patients, or those who present acutely with clinical signs of brainstem herniation. The head should be elevated to 30 degrees, 1.0–1.5 g/kg of 20% mannitol should be given by a rapid infusion, and the patient should be hyperventilated to a \(\text{pCO}_2\) of 30–35 mmHg. (Allen and Ward 1998) These measures are designed to lower ICP as quickly as possible prior to a definitive neurosurgical procedure (craniotomy, ventriculostomy, or placement of an ICP monitor) can be done. A number of these patients will present after a fall so particular attention should be directed to lacerations, skeletal fractures, stabilization of the cervical spine.

**Blood pressure**

Elevated blood pressure is seen in 46%–56% of patients with ICH. (Dandapani et al 1995) It remains unclear if elevated blood pressure directly causes hematoma expansion but studies have shown elevated systolic, diastolic, and mean arterial pressure are associated with a poor outcome in ICH (Terrayama et al 1997; Leonardi-Bee et al 2002; Vemmos

![CT scan showing large left parietal lobe lobar hemorrhage with a fluid level (arrow) after the patient received r-TPA.](image-url)
et al 2004). However, physicians have been reluctant to treat hypertension in ICH patients because the fear of overaggressive treatment of blood pressure may decrease cerebral perfusion pressure and theoretically worsen brain injury, particularly in the setting of increased intracranial pressure. In 1999, a special group consisting of healthcare professionals from the American Heart Association Stroke Council addressed these 2 rational theoretical concerns while attempting to write guidelines for the management of intracerebral hemorrhage. The task force recommended maintaining a mean arterial pressure below 130 mmHg in patients with a history of hypertension (level of evidence V, grade C recommendation). In patients with elevated ICP who have an ICP monitor, cerebral perfusion pressure (MAP–ICP) should be kept >70 mmHg (level of evidence V, grade C recommendation) (Broderick et al 1999).

Early hemostatic therapy

In the past, early neurologic deterioration in ICH was attributed to edema and mass effect around the hematoma. Pathological, CT, and SPECT studies suggest that continuous rebleeding into congested damaged tissue is associated with poor clinical outcome and is now an exciting new target of treatment (Fisher 1971; Kazui et al 1996; Fujii et al 1998; Becker et al 1999; Mayer 2005). Recent interest in hemostatic therapy is based on early hematoma growth often seen within six hours of onset of ICH in 14%–38% of patients (Kazui et al 1996; Brott et al 1997; Fujii et al 1998; Flibotte et al 2004; Roquer et al 2005). Initial efforts should be directed towards identifying thrombolytic, antiplatelet or anticoagulant use and reversing their effects. The biologic half-life of recombinant tissue plasminogen activator (rt-PA) at the site of the thrombus is limited to 45 minutes and accordingly hemorrhagic complications from rt-PA occur within the first few hours of use. Information is scarce to guide recommendations about treatment of hemorrhagic complications of thrombolytic therapy (levels of evidence III through V). According to guidelines devised by the American Heart Association Stroke Council, if bleeding is suspected the following measures should be taken: (1) blood should be drawn to measure the patient’s hematocrit, hemoglobin, partial thromboplastin time, prothrombin time/INR, platelet count, and fibrinogen (2) blood should be typed and cross-matched if transfusions are needed (at least 4 U of packed red blood cells, 4–6 U of cryoprecipitate or fresh frozen plasma, and 1 U of single donor platelets) (Adams et al 1996). These therapies should be made available for urgent administration. The risk of intracerebral hemorrhage with heparin is related to the level of anticoagulation. Heparin can be inactivated by 1 mg of protamine sulfate for every 100 IU of heparin administered (Wakefield and Stanley 1996). FFP should not be used to correct heparin related coagulopathy because FFP contains heparin binding antithrombin III (AT-III) which may prolong the anticoagulated status (Badjatia and Rosand 2005). Warfarin prevents recycling of vitamin K and indirectly inhibits synthesis of vitamin K dependent coagulation factors. Replenishing vitamin K via the oral or intravenous route helps reverse the effect of warfarin but an effective response may be delayed over 24 hours. Concomitant use of vitamin K with FFP, cryoprecipitate, or clotting factor concentrates are recommended to hasten reversal of warfarin induced coagulopathy. Considering the short half-life of coagulation factors at least 5–20 mg of vitamin K is required to sustain reversal of anticoagulation. Intravenous administration of vitamin K should be limited due to concerns of allergic and anaphylactic reactions. In an acute setting, vitamin K should not be administered subcutaneously because reversal of anticoagulation is neither rapid nor reliable (Steiner et al 2006). However, the variable content of vitamin K-dependent clotting factors in FFP and the effects of dilution have raised concerns that a coagulopathic state may persist despite correction of the international normalized ratio (INR) (Makris et al 1997). It is not clear at this time whether prothrombin complex concentrate is more reliable than FFP in repleting coagulation factors but it has proven to correct the INR faster than FFP which reduces the incidence and extent of hematoma expansion (Fredriksson et al 1992; Huttner et al 2006). Use of antithrombotic agents prior to ICH is a risk factor for continuous bleeding and poor outcome so it is reasonable to treat these patients with platelet infusions and desmopressin (Janssen and van der Meulen 1996; Saloheimo et al 2006).

Antifibrinolytic agents such as e-aminocaproic, tranexamic acid, aprotinin, and activated recombinant Factor VII (rFVIIa) have been receiving attention for early hemostatic therapy in patients with no underlying coagulopathy. However, rFVIIa is the only agent whose role in treating primary ICH has been evaluated in the randomized placebo control trial. The Novoseven Phase II trial was an international, multicenter, double-blinded trial that clearly demonstrated a reduction in early hematoma expansion in patients administered rFVIIa within 4 hours of symptom onset compared with placebo. In fact, the hemostatic effect was more pronounced with incremental doses of rFVIIa (Mayer et al 2005). Despite these promising results, early results from the Phase III Fast trial showed use of rFVIIa did not alter severe disability or...
mortality rates at 90 days (Forbes 2007). Complete results from the Phase III FAST trial are expected later this year.

Management of ICP
Elevated ICP is defined as intracranial pressure >20 mmHg for over 5 minutes. Large volume ICH is commonly associated with high ICP and brain tissue shifts related to ICP gradients. This problem can be exacerbated by intraventricular hemorrhage, which leads to acute obstructive hydrocephalus. The therapeutic goal of treating elevated ICP is to maintain ICP < 20 mmHg while maintaining cerebral perfusion pressure >70 mmHg. When ICP is monitored, use of a standard management algorithm results in better control, fewer interventions, and shorter duration of therapy. Initially, acute and sustained increase in ICP should prompt a repeat CT to assess the need for a definitive neurosurgical procedure. An intravenous sedative such as propofol (0.6–6.0 mg/kg/h) or fentanyl (0.5–3.0 µg/kg/h) should be given to the agitated patient to attain a motionless state. Thereafter, therapy should be directed at controlling blood pressure with vasopressors such as dopamine and phenylephrine if the CPP is < 70 mmHg or with antihypertensive agents if the CPP is > 70 mmHg. If ICP does not respond to sedation and cerebral perfusion management, osmotic agents and hyperventilation should be considered (Mckinley et al 1999). Of the 3 osmotic agents frequently used (mannitol, glycerol, and sorbitol), each has characteristic advantages and disadvantages. Sorbitol and glycerol are metabolized by the liver and interfere with glucose metabolism. However, sorbitol is infrequently used due to a short half life and poor penetration into the cerebrospinal fluid (CSF). Glycerol has a half-life less than one hour but it penetrates into the cerebrospinal fluid the best. Mannitol is commonly used because it is renally metabolized, has a half-life up to 4 hours, and achieves intermediate concentrations within the CSF (Nau 2000). Large ICH associated with elevated intracranial pressure refractory to these measures is fatal in most patients but a barbiturate coma may considered as a last resort to try to reduce intracranial pressure (Broderick et al 1999; Mckinley et al 1999). Corticosteroids are not recommended in the management of ICH because they have been proven to offer no benefit in randomized trials (Tellez and Bauer 1973; Poungvarin et al 1987).

Ventricular drains should be used in patients with or at risk for hydrocephalus. Drainage can be initiated and terminated according to clinical performance and ICP values. The volume of IVH strongly affects morbidity and mortality at 30-days (Tuhrim et al 1988). Preliminary studies with urokinase have suggested use of intraventricular thrombolysis within 72 hours of IVH may help drain the blood filled ventricles, speed clot resolution and decrease 30-day mortality rate (Naff et al 2000; Naff et al 2004). Patients are currently being recruited for Phase III trials assessing thrombolytic use in intraparenchymal and intraventricular hemorrhage.

Anticonvulsant therapy
The 30-day risk of seizures after ICH is about 8%. Seizures most commonly occur at the onset of hemorrhage and may even be the presenting symptom. Loban location is an independent predictor of early seizures (Passero et al 2003). Although, no randomized trial has addressed the efficacy of prophylactic antiepileptic in ICH patients, the Stroke Council of the American Heart Association suggest prophylactic antiepileptic treatment may be considered for 1 month in patients with intracerebral hemorrhage and discontinued if no seizures are noted (Broderick et al 1999; Temkin 2001). Acute management of seizures entail administering intravenous lorazepam (0.05–0.10 mg/kg) followed by an intravenous loading dose of phenytoin or fosphenytoin (15–20 mg/kg), valproic acid (15–45 mg/kg), or phenobarbital (15–20 mg/kg).

Fever control
Fever after ICH is common and should be treated aggressively because it is independently associated with a poor outcome (Schwarcz et al 2001). Sustained fever in excess of 38.3 °C (101.0 °F) should be treated with acetaminophen and cooling blankets. Patients should be physically examined and should undergo laboratory testing or imaging to determine the source of infection. Fever of neurologic origin is diagnosis of exclusion and may be seen when blood extends into the subarachnoid or intraventricular (Commichau and Scarmeas 2003). Intracerebral hemorrhage patients with persistent fever that is refractory to acetaminophen and without infectious cause may require cooling devices to become normothermic. Adhesive surface-cooling systems and endovascular heat-exchange catheters are better at maintaining normothermia than conventional treatment. However, it is still unclear whether maintaining normothermia will improve clinical outcome (Dringer 2004).

Deep venous thrombosis prophylaxis
Immobilized state due to limb paresis predisposes ICH patients for deep vein thrombosis and pulmonary embolism. Intermittent pneumatic compression devices and elastic stockings should be placed on admission (Lacut et al 2005). A small prospective trial by Boeer and colleagues using...
low-dose heparin on hospital day 2 to prevent thromboembolic complications in ICH patients significantly lowered the incidence of pulmonary embolism and no increase in rebleeding was observed (Boeer et al 1991).

**Surgical management**

Numerous surgical trials since the 1960s offered conflicting results and until recently no firm conclusions could be reached regarding the operative management of intracerebral hemorrhage. In 1995 randomization for the landmark Surgical Trial in Intracerebral Hemorrhage (STICH) had commenced. This trial was an international, multicenter trial that randomized 1033 patients with spontaneous supratentorial intracerebral hemorrhage within twenty-four hours to early surgery or conservative best medical therapy. Size, location, and volume of hemorrhage were similar in both treatment groups. Patients randomized to early surgery had their hematoma evacuated within twenty-four hours of randomization by the method of choice of the designated neurosurgeon. In 77% of cases, craniotomy was the surgical procedure and the remainder of cases had hematoma removal by burr hole, endoscopy, or stereotaxy in similar numbers. Thus, the STICH Trial is primarily a trial of craniotomy for ICH removal and left the role of less invasive surgery to remove ICH unanswered. Structured postal questionnaires were used to assess outcomes with the Glasgow Coma Scale, modified Rankin Scale, Barthel index, and mortality at 6-months. Overall, the STICH trial revealed no benefit from early craniotomy in supratentorial intracerebral hemorrhage when compared to initial conservative management. Of the prespecified subgroups that were examined, patients with an ICH within a centimeter of the cortical surface showed a benefit for early surgery. However, the statistical testing of this subgroup was not adjusted for in the multiple subgroup comparisons in this trial. In addition, early surgery was delayed with median time from onset to treatment for early surgery group was 30 hours and that may have affected the outcome (Broderick 2005; Mendelow et al 2005).

In contrast, infratentorial hemorrhages seem to benefit from early surgery. Most neurosurgeons believe cerebellar hemorrhages greater than 3 centimeters benefit from early surgical intervention because of the significant risk of brainstem compression and obstructive hydrocephalus within 24 hours (Ott et al 1974).

New areas of surgical research are focused on combination of minimally invasive surgery and clot lysis with rt-PA to remove intracerebral hemorrhage. Small preliminary trials have demonstrated that stereotactic aspiration and thrombolysis spontaneous intracerebral hemorrhage appears to be safe and effective in the reduction of ICH volume (Teermstra et al 2003; Barrett et al 2005; Vespa et al 2005). The National Institute of Health (NIH) has sponsored the Minimally Invasive Surgery Plus rtPA for Intracerebral Hemorrhage Evacuation (MISTIE) trial to determine the safety of using a combination of minimally invasive surgery and clot lysis with rt-PA to remove supratentorial primary ICH and compare efficacy to conventional medical management. The MISTIE trial is an open-label randomized treatment trial which is currently enrolling patients at multiple centers within the United States (NIH 2001).

**Conclusion**

Currently, no specific therapies improve the outcome after ICH. Although rFVIIa limits hematoma expansion, early Phase III results failed to show reduction in severe disability or mortality rates at 90 days. New trials evaluating the safety of the combination of minimally invasive surgery and clot lysis with rt-PA to remove intracerebral hemorrhage are currently underway.

**References**


Sahni and Weinberger


Abstract: Despite numerous studies on women's cardiac health throughout the past decade, the number of female deaths caused by cardiovascular disease still rises and remains the leading cause of death in women in most areas of the world. Novel studies have demonstrated that cardiovascular disease, and more specifically coronary artery disease presentations in women, are different than those in men. In addition, pathology and pathophysiology of the disease present significant gender differences, which leads to difficulties concerning diagnosis, treatment and outcome of the female population. The reason for this disparity is all steps for female cardiovascular disease evaluation, treatment and prevention are not well elucidated; and an area for future research. This review brings together the most recent studies published in the field of coronary artery disease in women and points out new directions for future investigation on some of the important issues.

Keywords: coronary artery disease, women, risk factors, prevention, diagnosis, treatment.

Introduction

The first female-specific recommendations for preventive cardiology were published in 1999 (Mosca et al 1999). Even though research in the treatment of cardiovascular disease (CVD) had advanced in many areas, it remains the leading cause of death in women in most parts of the world. Studies have shown that 500 thousand women die of CVD every year in the United States, somewhat near one death every minute (American Heart Association 2003). Such index exceeds not only the number of deaths in men, but also the next seven causes of death in women combined, and more importantly, coronary artery disease (CAD) is believed to be the major cause responsible for these deaths (American Heart Association 2003). Over a quarter of a million deaths per year are attributed to CAD alone in the United States (Merz et al 2004). Although already high, these figures are expected to rise even more during the next decades, due to an increase of diabetes and obesity, as well as the aging of the world population (Merz et al 2004).

Even though women have a higher frequency of chest pain/angina than men, the incidence of obstructive CAD in the female population is lower when compared with men with similar symptoms (Kenedy et al 1982; Diamond et al 1983; Merz et al 1999). In addition, it would appear that young women with obstructive CAD have a worse prognosis after acute myocardial infarction (AMI), whereas older women in similar circumstances often present with larger number of comorbidities that adversely influence the outcome, when compared to men (Coronado et al 1997). Women with acute coronary syndromes (ACS) are also less likely to receive rapid effective diagnosis and treatment than are men (Ayanian and Epstein 1991; Maynard et al 1996; Pope et al 2000).