Minimally Invasive Surgery for Intracerebral and Intraventricular Hemorrhage

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Spontaneous intracerebral hemorrhage (ICH), especially related to intraventricular hemorrhage (IVH), is the most devastating type of stroke and is associated with high mortality and morbidity. Optimal management of ICH remains one of the most controversial areas of neurosurgery and no effective treatment exists for ICH. Studies comparing conventional surgical interventions with optimal medical management failed to show significant benefit. Recent exploration of minimally invasive surgery for ICH and IVH including catheter- and mechanical-based approaches has shown great promise. Early phase clinical trials have confirmed the safety and preliminary treatment effect of minimally invasive surgery for ICH and IVH. Pending efficacy data from phase III trials dealing with diverse minimally invasive techniques are likely to shape the treatment of ICH.

Keywords: intracerebral hemorrhage, intraventricular hemorrhage, minimally invasive surgery, mechanical-based approach, pharmacological catheter-based approach

INTRODUCTION

Spontaneous intracerebral hemorrhage (ICH) is the most serious public health issue and impinges on 2 million people in the world each year (1–3). The incidence of ICH has been increasing because of an aging population and increased use of anticoagulation and antiplatelet agents for thromboembolic diseases. Although ICH accounts for 6.5–19.6% of strokes, it is related to the high rate of mortality and morbidity; the 30-day mortality rate is in the range of 35–52% and only 20% of survivors live independently (4). When ICH involves intraventricular hemorrhage (IVH), the outcome is even worse, with an estimated mortality rate of between 50 and 80% (3). Since most cases occur in working adults in a large number of people in low- and middle-income countries, ICH has a huge social and economic impact due to the loss of productive life years (5).

Risk factors for ICH include hypertension, anticoagulation, and amyloid angiopathy. Most patients (50–70%) with ICH suffer from hypertension (6). Hypertensive hemorrhage is inclined to occur in the external capsule (42%), pons (16%), thalamus (15%), cerebellum (12%), and white matter (10%) (7). Evidence from studies indicates that the incidence of ICH is reduced due to improved control of hypertension (8, 9). Anticoagulation-related ICH is more severe and associated with more extensive hemorrhage and a higher mortality rate in comparison with non-anticoagulation-related ICH and has risen in incidence over the past decade (10, 11). Cerebral amyloid angiopathy (CAA) is associated with 12–15% lobar ICH, particularly in the elderly (12, 13).
CAA-related ICH usually occurs in people aged > 60 years. Compared with other causes of ICH, CAA-related ICH has a lower mortality rate, but an increased rate of recurrence (14).

Unfortunately, there has been no beneficial medical treatment for ICH and surgery lacks definitive evidence and is still in debate. Theoretically, hematoma evacuation could remove the neurotoxic blood products, reduce the mass effect and intracerebral pressure (ICP), and control cerebral perfusion pressure, helping to prevent brain edema and secondary brain injury. Open surgery, however, may make damage to the normal brain tissue and affect functional outcomes, especially for patients with deep hemorrhage. The International Surgical Trial in ICH (STICH) and STICH II were the two largest prospective randomized clinical studies to evaluate the efficacy of open surgery for ICH. These clinical trials did not find a significant benefit in functional outcomes between patients who underwent surgery and patients who were medically treated (15, 16). The failure of traditional surgical treatment to improve the prognosis of patients with supratentorial ICH is because, in many patients, the damage of surrounding brain tissues caused by surgical methods neutralizes the benefits of hematoma clearance. This makes minimally invasive surgery (MIS) the most promising surgical strategy for patients with ICH. Because of the shorter operation time, the possibility of bedside treatment, and less damage to the brain tissue, minimally invasive treatment is more attractive than conventional craniotomy in the treatment of ICH (17).

In this review, we discuss different minimally invasive techniques for ICH and IVH, emphasizing clinical trials (Table 1) for this condition.

MINIMALLY INVASIVE SURGERY FOR ICH

The use of MIS for ICH evacuation started in the 1960s (32). The endoscope was used to evacuate the clot after ICH (33). Since then, several improvements have been made to the method to improve the efficiency of hematoma removal (34). In the late 1980s, mechanically-assisted thrombolysis by using ultrasound techniques was explored and favorable outcomes were described (35). MIS continues to evolve with the advancement of technology. Generally, MIS approaches can be divided into two categories: primarily pharmacological catheter based and primarily mechanically based. The pharmacological catheter-based approach involves the placement of a drainage catheter under the guidance of images and gentle aspiration for hematoma, followed by the infusion of the catheter with a thrombolytic agent to prevent clogging, facilitating passive drainage of the hematoma over several days. This requires intermittent imaging to confirm the catheter position and to monitor the progress of hematoma drainage. Mechanical methods for hemorrhage evacuation involve the surgical removal of the hematoma in a single procedure without leaving a drainage catheter and by using a thrombolytic agent. The evacuation of the hematoma is performed with either an endoscope or other devices. These two approaches have their own advantages and disadvantages, which may or may not contribute to functional outcomes in patients with ICH.

Pharmacological Catheter-Based MIS for ICH

Matsumoto and Hondo reported that they performed CT-guided stereotactic evacuation of hypertensive intracerebral hematoma in 51 patients. First, they inserted one silicon tube into the center of the hematoma and then, the hematoma was aspirated with a syringe, followed by the administration of urokinase until the hematoma was completely evacuated. Of 51 patients, 38 patients had functional outcomes (36). This method was also used in the posterior fossa hematoma. In one study, the aspiration and administration of urokinase were performed in 11 patients (9 patients with cerebellar hematoma and 2 patients with pontine hematoma) and 7 patients had functional outcomes (37). Thereafter, the recombinant tissue-type plasminogen activator (rt-PA) was used for clot thrombolysis and it was well tolerated and more effective in evacuating hematoma (26, 38). In addition, sonothrombolysis, in combination with pharmacological thrombolysis, has been used for clot evacuation. Newell et al. reported that rt-PA and 24-h of continuous ultrasound were delivered to the clot in 33 patients with ICH and they found that the combination of sonothrombolysis and rt-PA was more effective than rt-PA alone in hemorrhage lysis (39). Recently, Sun and Liu developed a new catheter-based MIS technology called cubic oriented stereotactic aspiration (40) and invented associated devices. This technology combines brain anatomy with solid geometry (Figure 1). Luo et al. confirmed the safety and efficacy of the technology (41) (Figure 2). At present, this technology has been used for supratentorial and infratentorial ICH in China (42–44).

These preliminary studies have shown that pharmacological catheter-based MIS can effectively remove hematomas. However, there is no unified standard for the selection of patients, the timing of MIS, and the proportion of hematoma evacuated. More importantly, the clinical benefit is still unclear. Therefore, it is essential to conduct randomized controlled trials (RCTs) to evaluate the effect of the pharmacological approach of MIS.

YL-1 Craniopuncture

Wang et al. randomized 377 patients with 25–40 ml basal ganglion hematoma in a multicenter controlled study into conservative medical treatment or the stereotactic craniopuncture and aspiration with YL-1 puncture needle followed by infusion of urokinase. Although there were significantly more complications in the craniopuncture group, the case fatality in these two groups was not significantly different and MIS led to a significant reduction of dependence at the end of the third month, compared with the conservative treatment (40.88 vs. 63.03%) (23). Moreover, other two RCTs were performed to evaluate and compare the effect of YL-1 craniopuncture plus urokinase with craniotomy with a small bone flap or conventional surgery in patients with ICH. Compared with craniotomy with a small bone flap and conventional surgery, the MIS had fewer complications and had a trend to improve long-term outcomes (25, 27). These
| References          | Treatment | Included patients (MG/OG) | Age* (years) (MG/OG) | Hematoma location                  | Volume* (ml) (MG/OG) | Onset to surgery (hours) | Outcomes | Follow-up (months) |
|---------------------|-----------|---------------------------|---------------------|-----------------------------------|----------------------|--------------------------|----------|-------------------|
| Auer et al. (18)    | ES        | CMT                       | 100 (50/50)         | 30–80                             | ≥10                  | Within 48                | ①②③    | 6                 |
| Naff et al. (19)    | EVD with urokinase | EVD with saline            | 12 (7/5)            | 49.6/55.2                          | ICH: 5.3/13.2        | NA                       | ①②③    | 1                 |
| Teemstra et al. (20) | SA with urokinase | CMT                       | 71 (36/35)          | 67/69                             | Supratentorial       | 66/52                    | Within 72 | ①②    | 6                 |
| Zhang et al. (21)   | ES with EVD | EVD with urokinase        | 42 (20/22)          | 31-75                             | ICH < 30             | Within 48                | ①②    | 2                 |
| Kim and Kim (22)    | SA        | CMT                       | 387 (204/183)       | 64.3/67.1                         | Basal ganglia and thalamus | 24.3/21.0               | NA       | ①②③    | 6                 |
| Wang et al. (23)    | CP with urokinase | CMT                       | 377 (195/182)       | 56.6/56.9                         | Basal ganglion       | 33.9/31.3               | 4–72     | ①②③    | 3                 |
| Chen et al. (24)    | ES with EVD | EVD                       | 48 (24/24)          | 65.54/62.17                       | ICH with thalamic hemorrhage | ICH:10.5/11.5           | NA       | ①②③    | 3                 |
| Sun et al. (25)     | CP with urokinase | CC with small bone flap  | 304 (159/145)       | 56.9/55.2                         | Basal ganglion       | 52.3/51.7               | Within 72 | ①②③    | 3                 |
| Naff et al. (26)    | EVD with rt-PA | EVD with placebo            | 48 (26/22)         | 54.1/56.6                         | ICH: 7.2/7.9         | ≤12                     | ①②③    | 1                 |
| Zhou et al. (27)    | CP with urokinase | CC with large bone flap    | 168 (90/78)         | 57.6/59.2                         | Basal ganglia or brain lobe | 30–100 ml              | 6–24     | ①②③    | 12                |
| Hanley et al. (29)  | SA with rt-PA | CMT                       | 96 (54/42)          | 60.7/61.1                         | Lobar or deep        | 48.2/43.1               | NA       | ①②③    | 12                |
| Vespa et al. (29)   | ES        | CMT                       | 24                  | 59/62                             | Supratentorial       | 36.4/41.4               | Within 48 | ①②③    | 12                |
| Hanley et al. (30)  | EVD with rt-PA | EVD with saline            | 500 (249/251)       | 59/59                             | ICH with or without supratentorial ICH | ICH:8.3/7.2 | IVH:21.2/22.4 | ≤12 | ①②③    | 6                 |
| Hanley et al. (31)  | SA with rt-PA | CMT                       | 506 (255/251)       | 62/62                             | Basal ganglia or lobar region | 42.7/41.5              | NA       | ①②③    | 12                |

CC, conventional craniotomy; CMT, conservative medical treatment; CP, craniopuncture; ES, endoscopic surgery; EVD, external ventricular drainage; ICH, intracerebral hemorrhage; IVH, intraventricular hemorrhage; MG, minimally invasive surgery group; NA, not available; OG, other treatment options group; SA, stereotactic aspiration. ① good functional outcome; ② death; ③ rehemorrhage; ④ ventriculitis; ⑤ ventriculoperitoneal shunt.

*Age and volume are usually expressed as mean or median. If there was no such information in literatures, a range would be given.
studies suggest that minimally invasive craniosurgery combined with urokinase is safe and might improve independent survival in patients with ICH. Nonetheless, the YL-1 needle is sharp and rigid, which may induce damage to brain tissues and rebleeding. Therefore, large-scale RCTs are needed to evaluate the safety and efficacy of YL-1 craniopuncture with thrombolytic agents.

**Stereotactic Treatment of Intracerebral Hematoma by Means of a Plasminogen Activator (SICHPA)**

The SICHPA trial was a multicenter RCT to investigate the efficacy of stereotactic treatment of ICH with the use of a plasminogen activator. This trial enrolled 71 patients with ICH volume $\geq 10$ ml and randomly assigned them to surgery with catheter placement and urokinase infusion or medical treatment. The SICHPA significantly reduced the ICH volume. However, the rebleeding rate and mortality at 180 days were similar in these two groups (20). This study suggests that stereotactic aspiration is safe and effective for reducing the ICH volume. Nevertheless, the number of patients enrolled in this trial is small and, thus, the results should be interpreted carefully.

**Minimally Invasive Surgery Plus Recombinant Tissue Plasminogen Activator in Intracerebral Hemorrhage Evacuation (MISTIE)**

The MISTIE was an international, randomized, open-label, phase II study that enrolled 96 patients with ICH from 26 hospitals and randomized them into medical care or MIS with alteplase. According to the MISTIE protocol, a rigid cannula was inserted into the hematoma under the image guidance and then one soft drainage cannula was placed. The administration of rt-PA in a dose of 0.3 or 1.0 mg was carried out every 8 h until 9 doses were given or until the volume of remaining hematoma was $\leq 15$ ml or until a clinically significant rebleeding occurred. MIS plus alteplase decreased the number of patients with the modified Rankin Scale (mRS) $\geq 3$ in 1-year follow-up and significantly decreased the perihematoma edema volume. In addition, 30-day mortality,
infection, and symptomatic bleed were not significantly different between MIS plus alteplase and the conservative groups. Moreover, the volume of hematoma remaining was statistically significantly correlated with functional outcomes when other variables were controlled. These results suggest that MIS plus alteplase is safe to effectively evacuate hematoma and may improve functional outcomes through reducing clot volume (28). However, the sample size is relatively small and the objective of this study is to detect the threshold of safety not efficacy. In addition, the wide range of benefits estimated in intention-to-treat analyses makes it difficult to draw conclusions about benefits.

To test the conclusion of the MISTIE II and ensure its generalizability, the MISTIE III was performed in 78 centers. Appropriate patients were ≥ 18 years with supratentorial hemorrhage more than 30 ml plus the Glasgow Coma Scale (GCS) ≤ 14 or the National Institutes of Health Stroke Scale (NIHSS) score ≥ 6 and the hematoma remained stable (growth < 5 ml) for at least 6 h after diagnostic CT. A total of 506 patients were enrolled in this study. The process for cannula placement and rt-PA injection was the same as described in the MISTIE II, except that the dose of rt-PA was different and limited to 1.0 mg. The mean reduction in hematoma size was 69% in the MISTIE group vs. 3% in the standard medical care group. Nonetheless,
only 58% of patients met the preset goal that the hematoma was <15 ml after surgical treatment and the functional outcome between these two groups did not significantly differ. In addition, the safety events rates were not significantly different between the two groups (31). The main conclusion of this clinical study is that the MISTIE is safe and effective for ICH evacuation, but it cannot improve the prognoses of patients. Therefore, it cannot be recommended as an intervention to improve the functional outcome of all the patients with ICH. However, the MISTIE may improve functional outcomes when the protocol-defined surgical aim can be achieved (residual hematoma smaller than 15 ml) (31). In an exploratory analysis, reduction of hematoma volume to ≤15 ml associated with a good functional outcome (mRS 0–3) and reducing the hematoma volume by 70% or more increased the chance of achieving a good outcome, i.e., if the reduction exceeded the 15 ml threshold, the probability of good results would increase by 10% for every additional 1 ml of hematoma removed. The failure to achieve ≤15 ml goal evacuation was significantly associated with initial hematoma volume, history of hypertension, irregular-shaped hematoma, number of alteplase doses given, surgical protocol deviations, and catheter manipulation problems. Additionally, greater surgeon/site experience was associated with avoiding poor hematoma clearance. Taken together, the results indicate that surgical performance may determine the outcomes of patients with ICH. Enhanced surgical performance may improve the prognosis of patients.

Mechanically-Based Approaches for ICH Endoport-Mediated Evacuation

The endoport system for clearing ICH originates from the computer-aided stereotactic method for resection of deep and central intracranial tumors. The system aims to effectively remove the hematoma and minimize damage to adjacent tissues. The endoport-mediated evacuation involves a small craniotomy with dura opening, stereotactic placement of an endoport to the hematoma (typically along the longest axis of the hematoma), and removal of hematoma with traditional surgical tools or special devices. Compared with the pharmacological catheter-based approach, this technique allows direct access to the hematoma cavity with irrigation, suction, and cautery instruments and clears more hematoma at once. It, however, needs craniotomy and may increase brain damage along the endoport path.

Currently, the most commonly used device is the NICO endoport system (Indianapolis, Indiana, USA). The NICO endoport system, consisting of the Myriad handpiece and the BrainPath sheath (Figure 3), is cleared by the United States Food and Drug Administration for the removal of tissue and visualization of the surgical field during neurosurgery (45). Ding et al. presented a case report about the use of NICO endoport system to evacuate a large basal ganglia hematoma (46). And then, they performed a retrospective study of 11 patients with ICH (9 with supratentorial hemorrhage and 2 with infratentorial hemorrhage and the volume of ICH from 8 to 168 ml) and used the same technique to evacuate the hematoma. They found that the ICH volume was reduced by 87% and 36% of patients were functionally independent (mRS 0–2) at 90 days (47).

Minimally Invasive Subcortical Parafascicular Access for Clot Evacuation (MISPACE)

A multicenter study to evaluate the safety and feasibility of the MISPACE by using the NICO endoport system was conducted in 11 centers in the United States and Canada. A total of 39 patients with ICH (ICH volume from 27 to 65 ml) were enrolled. The result of this study showed that hematoma volume was reduced by more than 90% in 72% of patients and 52% of patients had good functional outcomes with the mRS ≤ 2 at 90 days (48). The authors concluded that this method was safe and feasible to remove the clot. However, the sample size of this study is small and it is a retrospective, uncontrolled study. RCTs are still needed to assess the safety and effectiveness of the MISPACE.

Early Minimally-Invasive Removal of Intracerebral Hemorrhage (ENRICH)

The ENRICH is a multicenter randomized controlled study comparing early (<24 h) surgical hematoma evacuation by using minimally invasive parafascicular surgery (MIPS) with the NICO endoport system with standard medical management in the treatment of acute spontaneous supratentorial ICH. This study is based on the solid foundation supplied by the results from the multicenter study for the MISPACE and other clinical studies by using the NICO system. The ENRICH is designed to enroll 300 patients with ICH and was started in 2016. It is still in progress.

Endoscope Evacuation

The ICH evacuation by an endoscope with the use of a single burr hole was first reported by Auer et al. (49). Thereafter, the same group performed a single-center randomized controlled study that compared endoscopic evacuation vs. medical treatment in patients with ICH. A total of 100 patients with supratentorial hematoma more than 10 ml and neurological or consciousness impairment were enrolled and assigned randomly. The clearance rate of hematoma in most patients was 50–70%. The mortality rate in the surgical group was significantly lower than the medical treatment group 6 months after hemorrhage (42 vs. 70%). The rate of better outcomes was higher in the surgical group than in the medically treated group (40 vs. 25%). However, these results were restricted in patients under 60 years old and with hematoma smaller than 50 ml (18). Since these promising results were obtained by the first generation of endoscopy, it is difficult to transfer the findings to the clinical situation of today. A more recent retrospective analysis of 68 patients with supratentorial ICH treated with endoscope-assisted ICH evacuation showed that rebleeding, morbidity, and mortality rates were low and long functional outcome was favorable when compared with the traditional craniotomy (50). Moreover, another retrospective analysis of 43 patients with putamen hematoma (volume > 31 ml), cerebellar ICH (hematoma > 3 cm in diameter), or thalamic ICH (hematoma volume > 20 ml) and acute hydrocephalus treated with endoscopic surgery and craniotomy procedure demonstrated higher evacuation rate and the higher GCS score at day 7 in the group of
endoscopic surgery (51). However, the safety and efficacy of endoscopic surgery for ICH evacuation should be evaluated in RCTs.

**Intraoperative Stereotactic CT-Guided Endoscopic Surgery (ICES) Trial**
The ICES trial was a randomized arm of the MISTIE trial. In this study, 24 patients with hematoma volume more than 20 ml from different hospitals were randomized 3:1 to endoscopic surgery or standard medical management. The result showed that endoscopic surgery led to a 71.2% reduction of ICH with an associated 12% increase in good functional outcomes defined as the mRS score from 0 to 3 at 1 year (29). Although the number of patients involved in this preliminary study was small, the result of the study suggests that the endoscopic technique can be performed by different surgeons in a different location and can produce valuable results worthy of further evaluation in the phase III study.

**Apollo System and the Minimally Invasive Endoscopic Surgical Treatment With Apollo in Patients With Brain Hemorrhage (INVEST)**
There are some promising tools used for endoscopic ICH evacuation and the Apollo System (Penumbra Incorporation, Alameda, California, USA) is one of them. The Apollo System is a nonclogging aspiration–irrigation system designed to fit down the working channel of a three-port neuroendoscope (Figure 4). The aspiration–irrigation system can be connected with the Apollo wand through a flexible tube. The wand is equipped with an internal stirrer line that vibrates at ultrasonic frequency and impregnates the clot material to keep the system unobstructed during the suction process. The aspiration–irrigation system provides the ability of suction and continuous saline flushing and transmits vibration energy to the internal agitator elements in the wand. The advantage of the Apollo device in increasing the emptying of the endoscope is that it allows more directional suction and can extend beyond the end of the endoscope in a controlled way. Spiotta et al. performed a retrospective analysis of 29 patients with ICH who underwent ICH evacuation with the Apollo system at 4 centers. The mean ICH volume decreased from 45.4 ± 30.8 to 21.8 ± 23.6 ml after evacuation (52). Fiorella et al. presented case reports of 3 patients with supratentorial ICH who underwent hematoma evacuation by using the Apollo System with combined neuronavigation, neuroendoscopy, and cone-beam CT. The volume of hematoma was reduced from 93.4 to 15.6 ml and 11.2 to 0.9 ml after evacuation. Moreover, no complications occurred for the procedure (53). More clinical data are needed to determine the extent of hematoma removal to improve clinical symptoms and whether it can achieve more complete blood product emptying, thereby offsetting the disadvantages of large channels.

The INVEST trial is a multicenter single-arm phase II trial, which is evaluating the safety and efficacy of the minimally invasive hemorrhage evacuation with Apollo device compared with medical management. The study has initiated enrollment in 2016.

**Other Mechanically-Based MIS Approach**
Kim et al. performed a prospective study, which enrolled 387 patients with ICH volume ≤ 30 ml limited to the basal ganglia and thalamus and randomized them into either MIS by using an Archimedes aspirator or to medical management. If the remaining hematoma exceeded 10 ml after surgical evacuation, the instillation of urokinase in the hematoma cavity was considered. In the MIS group, patients had better functional outcomes with higher mean of the Barthel Index scores (90.9 vs. 62.4) and the lower mRS scores (1.2 vs. 3.0) at 180 days.
The Apollo System consists of a suction-irrigation system that can be connected to the Apollo wand through a soft tube. The wand (A) can be placed through the working channel of a neuroendoscope. The wand is equipped with an internal stirrer line that vibrates at ultrasonic frequency and impregnates the clot material to keep the system unobstructed during suction. The wand is connected to a stand-alone suction-irrigation system (B) [from Fiorella et al. (45) with permission].
Meanwhile, the mortality was not significantly different between the MIS and medical treatment groups (22).

MINIMALLY INVASIVE SURGERY FOR IVH

Usually, IVH is a complication of spontaneous ICH. However, it sometimes occurs without an identifiable cause. The extension of ICH into the ventricular is an independent risk factor for poor prognosis (54). In addition, the volume of IVH is related to the clinical outcomes (54, 55).

The application of MIS in the treatment of IVH is based on the hypothesis that the reduction of hematoma volume and the restoration of cerebrospinal fluid circulation can reduce secondary brain tissue damage. Like ICH, MIS for IVH includes pharmacological catheter-based and mechanically-based approaches. The pharmacological catheter-based method involves the injection of fibrinolytic drugs (urokinase or rt-PA) through external ventricular drainage (EVD) tube. The mechanically-based approach involves the direct movement of intraventricular hematoma with the endoscopy. These methods have their disadvantages. The disadvantages of the pharmacological catheter-based method include the risk of infection from the repeated injection of a lytic agent through EVD, potential hematoma expansion induced by the lytic agent, a relatively long period for adequately clearing IVH, and risk of failure to reduce the volume of intraventricular hematoma (56). The mechanically-based approach is more invasive and needs craniotomy with a small bone flap.

Pharmacological Catheter-Based Treatment for IVH

Injection of fibrinolytic drugs through EVD has been proved to be safe and effective in animal studies (57–59). And then, this result has been confirmed in small clinical case series (60–62). Nieuwkamp et al. conducted one systemic review of 343 patients with severe IVH caused by extension from subarachnoid hemorrhage or ICH from 18 observational studies and these patients received conservative treatment, EVD or EVD with fibrinolytic agents. They found that the case fatality rate was significantly lower in patients receiving EVD combined with fibrinolytic agents. The poor outcome rate was also lower for EVD with fibrinolytic agents compared with conservative treatment or EVD. Treatment with EVD combined with fibrinolytic agents may improve the outcome of patients with severe IVH (63). Furthermore, Khan et al. conducted a meta-analysis to investigate the impact of intraventricular fibrinolytic therapy on mortality, functional outcome, ventriculitis, shunt dependence, and rehemorrhage. The meta-analysis enrolled 24 studies and demonstrated the benefit of intraventricular fibrinolytic on mortality and good functional outcome. In addition, intraventricular fibrinolytic decreased the rate of shunt dependence and was not associated with increased rates of ventriculitis or rehemorrhage (64). This meta-analysis concluded that intraventricular fibrinolytic was safe and could be an effective strategy for the treatment of IVH. Nonetheless, those results are from observational or small randomized studies and randomized clinical trials are needed. Then, a randomized, double-blind, controlled, multicenter, pilot study was performed to assess the safety and efficacy of intraventricular thrombolysis. A total of 12 patients with IVH associated with supratentorial ICH of <30 ml were enrolled in this study and randomized to receive intraventricular injections of normal saline solution or urokinase at 12-h intervals. Compared with treatment with EVD alone, intraventricular thrombolysis with urokinase speeded the resolution of intraventricular blood clots and the frequency of adverse events did not differ significantly between the two groups (19). The number of patients enrolled in this study, however, is small and the findings need to be validated in a large study.

Clot Lysis: Evaluating Accelerated Resolution of IVH (CLEAR-IVH)

The pilot study led to the CLEAR-IVH trial which was a phase II trial and was to evaluate the safety and efficacy of using multiple injections of low-dose rt-PA to accelerate lysis and evacuation of IVH. In this study, 48 patients with IVH were randomized into treatment with rt-PA and placebo. The administration of rt-PA could accelerate the clearance of IVH clots. Although the symptomatic bleeding rate in patients treated with rt-PA was higher than that in patients receiving placebo, a trend to improve clinical outcomes at 30 days was observed. Moreover, the rate of IVH clot resolution was significantly correlated with early clinical improvement (26). This phase II trial optimized the dose effect of rt-PA and demonstrated a trend toward improved clinical outcome at 30 days in patients treated with rt-PA, promoting a more definitive phase III trial.

The CLEAR III was an investigator-initiated, phase III, randomized, multicenter, double-blind, placebo-controlled study comparing the use of EVD combined with rt-PA with EVD plus saline (placebo) for the treatment of IVH. A total of 500 patients with IVH and obstructive hydrocephalus in 73 centers were enrolled and assigned 1:1 to the alteplase group and the saline group. Alteplase (1 mg) or 0.9% saline was given up to 12 doses, 8 h apart through EVD. The administration of alteplase was stopped when 3rd and 4th ventricles open, intraventricular hemorrhage mass effect relieved, 80% of clot removed, or 12 doses were given. The study showed that intraventricular rt-PA did not affect the primary efficacy outcome, but there were less frequent adverse events and a lower case fatality at 180 days in the alteplase group. In addition, clot removal was significantly related with both the mRS ≤ 3 and case fatality (30). These results suggest that by using alteplase through EVD is safe, but it cannot improve functional outcomes at the mRS 3 cutoff compared with by using saline through EVD. However, the benefits of alteplase may be possible if greater clot removal can be achieved. Unfortunately, only 30% of patients treated with alteplase achieved the goal of 80% blood clot reduction, which may be one of the reasons that alteplase treatment cannot effectively improve functional outcomes.

Mechanically-Based MIS for IVH Treatment

Several case series and case reports showed the safety and efficacy of endoscopy to treat IVH. Longatti et al. reported 13 patients with spontaneous primary or secondary IVH who
underwent endoscopy treatment and they found that endoscopy management was safe and effective to remove the intraventricular blood clot. Compared with other more conventional treatments, endoscopy may improve the clinical outcome of patients with IVH (65). Basaldella et al. performed a comparative retrospective analysis of 96 patients with massive IVH (48 patients treated with endoscopy surgery and 48 patients treated with EVD alone) to assess the efficacy of endoscopy surgery on IVH. The analysis demonstrated that endoscopy treatment combined with EVD reduced the shunting rate by 34% compared with EVD alone. However, endoscopy treatment did not significantly affect the outcome at 1 year as determined by using the mRS (66). In addition, Rienzo et al. presented a retrospective analysis of the two groups of patients with cast IVH who received endoscopy-assisted surgery or EVD and found that endoscopy-assisted evacuation reduced ICU staying and cerebrospinal fluid (CSF) clearance times. Moreover, endoscopy seemed to improve neurological outcomes, but without affecting the need for permanent shunt (67). However, these are just experience from small case serials and RCTs are needed to evaluate the safety and efficacy of endoscopy treatment for IVH. Zhang et al. conducted a prospective and randomized study that enrolled 42 patients with IVH and these patients were randomly assigned to the endoscopic group and the EVD group. In the endoscopic group, EVDs were placed after the procedure and if the remaining IVH volumes were > 10 ml, injection of urokinase into the ventricle through EVD was performed. In patients allocated to receive an EVD alone, a lytic solution was also administered through the drain. The authors found that more patients in the endoscopic group got the high Glasgow Outcome Scores at 2 months. The mortality rate was not significantly different between these two groups (21). Chen et al. randomized 48 patients with IVH from thalamic hemorrhage (1:1) into the EVD group and endoscopic surgery followed by the EVD group. In the endoscopic surgery group, a standard suction tube was placed side by side with the endoscope through a hard sheath to evacuate blood from the ventricular system ipsilateral to the thalamus hemorrhage. The endoscopic surgery significantly reduced the length of the intensive care unit (ICU) stay and the rate of the shunt (47.62 vs. 90.48%) compared with EVD alone. Nonetheless, the 30- and 90-day mortality rates and the GCS scores were not significantly different between the EVD and endoscopic surgery groups (24).

Several small randomized and observational studies of endoscopic surgery, EVD alone, or EVD with intraventricular fibrinolysis studies in IVH were assessed in a meta-analysis conducted by Li et al. of 680 patients from 11 studies. This meta-analysis demonstrated that compared with EVD with intraventricular fibrinolysis, endoscopic surgery with EVD significantly reduced mortality and shunt rates and increased hematoma evacuation rate. Therefore, the authors suggested that endoscopic surgery with EVD may be better than EVD with intraventricular fibrinolysis in the management of IVH (68).

In brief, the available evidence suggests that mechanically-based techniques can safely and effectively evacuate IVH. Mechanically-based techniques have the potential to improve functional outcomes in some patients with IVH, but more evidence is needed. Zhu et al. are performing a prospective, randomized, controlled, multicenter clinical trial to compare the prognosis of patients undergoing endoscopic surgery vs. those undergoing EVD for moderate-to-severe IVH. This clinical trial will enroll 956 patients with moderate-to-severe IVH who will be randomly assigned 1:1 to the endoscopic surgery group or the EVD group and recruitment began in 2020 (69). The results from this trial will provide high-quality evidence for mechanically-based MIS treatment for IVH.

From the abovementioned clinical studies concerning the pharmacologically- or mechanically-based MIS for ICH and IVH, it is clear that MIS is safe and effective in removal of hematoma compared with conventional surgery and medical treatment. However, MIS is not suitable for patients with unstable hematoma. Additionally, phases II and III clinical trials do not supply definitive evidence that MIS can improve the prognoses of patients. There are several explanations for this. First, extended clot dissolution time offsets the main goal to minimize secondary injury induced by ICH, which begins only a few hours after bleeding and progresses over time. Second, removing the bleeding foci by dissolving the blood clot without properly managing the suspected blood vessel may lead to further expansion of the hematoma. Third, each MIS method lacks standardization and, thus, it is difficult to teach technology and generalize the results. Last but not least, surgical performance may be different during MIS. Catheter manipulation problems, surgical protocol deviations, and greater surgeon/site experiences influence hematoma evacuation, while the volume of residual hematoma is associated with functional outcomes. Therefore, generalization of optimal performance with hematoma evacuation will require focused surgeon education, emphasizing technical nuances, better demonstration of experience, and a rigorous definition of benchmarks for successful tasks. Future research should focus on the best way to achieve this goal and case selection may need to be customized separately for various technologies. Besides, since ultra-early surgery results in a higher rate of rebleeding, the timing of surgery may be an important part of the effectiveness of MIS and this factor should be included in the trial design.

NEW MIS TECHNOLOGIES

Three-Dimensional Printing Technology

In the recent years, 3D printing technology has become a hot field of biomedical research; it is a rapid prototyping technology based on the digital model file, which is used to print solid objects with complex geometry through layered processing. For ICH, 3D printing technology is based on the original Digital Imaging and Communications in Medicine (DICOM) data of CT. This technology can help to analyze the shape and location of the hematoma of the patient and design surgical approaches that avoid important areas such as the venous sinuses, functional areas, and frontal sinuses. In addition, during hematoma puncture, the puncture angle and puncture position can be fixed by 3D-printed guide holes to reduce tissue damage caused by inaccurate positioning and repeated punctures. Zhang et al. retrospectively analyzed 12
patients with basal ganglia hemorrhage who underwent 3D-printed model-guided endoscopic evacuation for hematoma. They found that the average evacuation rate of hematoma was 97.2% and all the patients had good outcomes evaluated by functional independence measure scores at 6 months, which suggested that 3D-printed model-guided endoscopic evacuation was effective and safe (70). Additionally, Wang et al. applied a 3D-printed navigation mold in puncture drainage for brainstem hemorrhage and found that 3D-printed technology made the puncture drainage more individualized and precise (71).

Diffusion Tensor Imaging and MIS
Diffusion tensor imaging has been used to assess the integrity of the white matter tract and predict motor functional outcomes in patients with ICH (72, 73). Wu et al. found that the changes of the cortical spinal tract could be observed intuitively through DTI in patients with ICH and MIS can reduce the damage to the cortical spinal tract led by hematoma and even could restore the cortical spinal tract, which was oppressed and displaced by hematoma (74). Moreover, preoperative DTI can help doctors to clarify the positional relationship between hematoma and white matter tracts and, thereafter, to determine the MIS approach and surgical trajectory. Therefore, DTI may be combined with different MIS technologies to optimize MIS approaches and explore the efficacy of MIS.

Robot for Neurosurgery
The robot for neurosurgery has been used in clinical studies. It establishes 3D coordinates based on CT or MRI. Then, the mapping relationship between the computer and actual images is established through positioning marks and stereotactic surgery is planned and operated virtually. Finally, the auxiliary positioning and navigation by multisensor intelligent mechanical arms are realized. In addition, the robot can display 3D visualization of cerebral vessels based on multimodal image infusion, which could help to accurately locate the hematoma and plan the hematoma puncture trajectory, avoiding vessels. Wang et al. retrospectively analyzed 17 patients with hypertensive ICH who received robot-assisted frameless stereotactic MIS. They found that the average positioning error was 1.28 ± 0.49 mm and the functional outcome of patients was improved at 3 months (75).

Xiong et al. conducted a systematic review to assess the safety and efficacy of robotic surgery for ICH and suggested that compared with conventional surgery or conservative treatment, robotic surgery is safer and more effective (76).

CONCLUSION
Spontaneous ICH, especially those related to IVH, is still one of the most fatal and disabling diseases, causing a high-cost burden on society. Extensive practice mode, vague practice standard, lack of proven therapy, and persistent adverse results cast a shadow on its management. So far, it has not been clearly shown that neurosurgical interventions can improve the prognosis of patients with ICH. However, new emerging results of MIS for ICH and IVH have been very encouraging: MIS can reduce the secondary injury after ICH and IVH, thus potentially reducing mortality and dependence. During the process of MIS, surgical performance is important, since it is associated with the outcomes of patients. Additionally, new technologies could improve the accuracy and safety of MIS. Currently, trials of several new minimally invasive approaches for ICH and IVH evacuation are being evaluated and more clear evidence is expected in the near future. Moreover, ongoing clinical trials are expected to change clinical practice by optimizing case selection and surgical tasks and integrating these new tools into treatment.

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ZZ and QW contributed to the literature review and wrote the manuscript. JL contributed to writing and modifying the manuscript. SS read and checked the manuscript. All authors contributed to the article and approved the submitted version.

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