The treatment of childhood hydrocephalus represents a relevant share of pediatric neurosurgery. Various techniques to balance the disturbed CSF dynamics are described, such as endoscopic fenestration for establishing intracranial CSF communication, transient CSF drainages by reservoirs or external drainages, CSF diversion by shunting, or the combination of CSF diversion and endoscopic fenestration. The debate about the optimal therapy for infants with hydrocephalus continues, because they represent a challenging cohort of patients with high complication rates. Primary treatment and possible complications have a relevant impact on the long-term outcome in these patients, whose brains are still developing.

The most common complications in shunt therapy of pediatric hydrocephalus are infections, shunt occlusion, under- and overdrainage, or malpositioning of implants. In particular, overdrainage is relevant in children as a long-term complication that might lead to microcephaly, slit ventricle syndrome, hyperostosis, and chronic headaches. Differing valve designs, including either adjustment units or antisiphon devices, have addressed this problem and have been investigated over the years. The adjustable differential pressure (DP) valves attempt to adjust the resistance of CSF drainage according to the individual circumstances of anatomy, activity, and CSF imbalance. So-called antisiphon devices address posture-dependent overdrainage, which

OBJECTIVE The use of adjustable differential pressure valves with gravity-assisted units in shunt therapy of children with hydrocephalus was reported to be feasible and promising as a way to avoid chronic overdrainage. In this single-center study, the authors’ experiences in infants, who have higher rates of shunt complications, are presented.

METHODS All data were collected from a cohort of infants (93 patients [37 girls and 56 boys], less than 1 year of age [mean age 4.1 ± 3.1 months]) who received their first adjustable pressure hydrocephalus shunt as either a primary or secondary implant between May 2007 and April 2012. Rates of valve and shunt failure were recorded for a total of 85 months until the end of the observation period in May 2014.

RESULTS During a follow-up of 54.2 ± 15.9 months (range 26–85 months), the Kaplan-Meier rate of shunt survival was 69.2% at 1 year and 34.1% at 85 months; the Kaplan-Meier rate of valve survival was 77.8% at 1 year and 56% at 85 months. Survival rates of the shunt were significantly inferior if the patients had previous shunt surgery. During follow-up, 44 valves were exchanged in cases of infection (n = 19), occlusion (n = 14), dysfunction of the adjustment unit (n = 10), or to change the gravitational unit (n = 1).

CONCLUSIONS Although a higher shunt complication rate is observed in infant populations compared with older children, reasonable survival rates demonstrate the feasibility of using this sophisticated valve technology. The gravitational unit of this valve is well tolerated and its adjustability offers the flexible application of opening pressure in an unpredictable cohort of patients. This may adequately address overdrainage-related complications from early in treatment.

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KEY WORDS adjustable gravitational valve; infant; hydrocephalus; shunt survival; valve survival

ABBREVIATIONS DP = differential pressure.


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* Ms. Gebert and Dr. Schulz contributed equally to this work.
results from the hydrostatic forces in the standing position when the CSF drains from the elevated level of the intracranial compartment toward the lower level of the peritoneum.\textsuperscript{3,13,19,21,46} We previously reported on the use of an adjustable DP valve with gravitational assistance, the proGA\textsuperscript{\textregistered} (MIETHKE, B. Braun, Aesculap), and were able to show the feasibility of this device for the treatment of childhood hydrocephalus, as well as its acceptance by patients and caretakers.\textsuperscript{18,48}

The treatment of hydrocephalus in infants is especially challenging due to their less developed immune status, delicate skin condition, rapidly changing cerebral dimensions, closure of the fontanelle and cranial sutures, and the wide range of intracranial and intraabdominal pressure conditions being influenced (e.g., by crying and abdominal pressure). Only a few studies have addressed treatment for hydrocephalus exclusively in an infant population.\textsuperscript{23,31,32,42,55} Thus, it seems reasonable to present our long-term experience in infants with hydrocephalus treated with CSF-diverting shunts using the proGA\textsuperscript{\textregistered} adjustable, gravity-assisted valve, which has been routinely used at our institution since 2007.

Methods

This single-center, retrospective study included 93 patients less than 12 months old, who received a CSF-diverting shunt including an adjustable DP valve with a gravitational unit (proGA\textsuperscript{\textregistered}) for the treatment of hydrocephalus. All proGA\textsuperscript{\textregistered} systems (either primary implants or secondary implants used at shunt revision between May 2007 and April 2012) were identified and followed for an additional 2 years until the end of the observation period in May 2014, for a maximum of 85 months.

In the present study, the overall proGA\textsuperscript{\textregistered} survival and shunt survival rates were compared after 12 months and at the end of the observation period (a maximum of 85 months). An adverse event (i.e., failure) was considered any event leading to a surgical revision of the shunt system, including infection. The infection rate was calculated as the number of documented infections divided by the number of surgeries of the whole patient cohort during the observation period. Data were retrospectively collected by a review of the operative database and the patient’s medical records.

ProGA\textsuperscript{\textregistered} System

The programmable gravity-assisted valve combines an adjustable DP valve with a gravitational unit, connected by silicon tubing. The adjustment unit operates by a ball-in-cone mechanism, which can be magnetically changed by adjusting the position of a rotor, which in turn alters the tension of a spring toward the ball. To readjust the DP valve, a magnetic pen is used to apply pressure in the center of the unit to loosen the locking mechanism and magnetically transmit the selected setting. An opening pressure between 0 and 20 cm H\textsubscript{2}O can be chosen, which represents the permanent resistance level independent of the patient’s position. The gravitational unit gradually adds further resistance, which is defined by the weight of a tantalum ball, when the valve is erected from 0° to a 90° upright position. The gravitational unit cannot be adjusted after implantation; a unit with a resistance of 20 cm H\textsubscript{2}O was always used for infants at our institution. The setting of the adjustable DP unit depends on the individual surgeon’s decision in every case but was set initially between 5 and 8 cm H\textsubscript{2}O, according to our hospital’s guidelines. The ventricular catheter was placed in a frontal precoronal position in all patients to allow the valve to be placed in a strictly vertical position subcutaneously behind the ear.\textsuperscript{43} The ventricular catheter was connected to a bur hole reservoir. Antibiotic-impregnated silicon tubes (Bactiseal; Johnson & Johnson) were used for all cases from 2009 on, and the tubes were fixated with nonresorbable ligatures. The peritoneal catheter was inserted, with a minimum length of 35 cm.

Readjustments of the valve setting were performed in cases of clinical symptoms or radiological signs of over- or underdrainage. An elevated setting of the valve (by 2 cm H\textsubscript{2}O above the normal routine resistance setting) was chosen for patients with intensive crying or activity, according to our hospital’s standard protocol. Patients were seen for clinical evaluation 4–8 weeks after discharge and then every 3 months. Follow-up with MRI was arranged every 12 months and, for complex cases, 3 months after surgery.

Statistical Analysis

The software used for graph design and statistical analysis was GraphPad Prism 6 (GraphPad Software, Inc.). All values are given as the mean ± SD. Comparison of Kaplan-Meier survival curves was performed using the log-rank (Mantel-Cox) test. A p value less than 0.05 was considered statistically significant.

Results

The gestationally corrected mean age of the 93 patients (37 girls and 56 boys) at the time of shunt insertion was 4.1 ± 3.1 months (Table 1). One girl who was born extremely prematurely (at 24 weeks of gestation) died from multiple organ failure 1 month after implantation of the valve and was therefore excluded from follow-up. The mean follow-up was 54.2 ± 15.9 months. Among the 93 patients, 46 were preterm infants (less than 37 weeks of gestation). The etiology of hydrocephalus was posthemorrhagic (n = 43), myelomeningocele (n = 18), idiopathic (n = 9), tumor

<table>
<thead>
<tr>
<th>Table 1. Demographic data from 93 infants who received a primary or secondary proGA\textsuperscript{\textregistered} implant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variable</td>
</tr>
<tr>
<td>Time period of implantation</td>
</tr>
<tr>
<td>No. of proGA\textsuperscript{\textregistered} implants</td>
</tr>
<tr>
<td>No. of pts</td>
</tr>
<tr>
<td>Mean age, mos (gestationally corrected for premature infants)</td>
</tr>
<tr>
<td>Mean follow-up, mos (range)</td>
</tr>
<tr>
<td>No. of primary implants</td>
</tr>
<tr>
<td>No. of secondary implants (revisions)</td>
</tr>
</tbody>
</table>
The most common etiology of hydrocephalus was posthemor-

rhagic (n = 32).

During the inclusion period, 79 infants received the first

valve and shunt (79 primary implants). Fourteen infants,

who were referred from other hospitals and already had

shunts, underwent revisions during the inclusion period

and received a proGAV shunt system (14 secondary im-

plants). During the inclusion period, 19 children underwent

revision(s) of the valve while they were still in their infant-

cy; among those infants, 24 proGAV valves were implant-

ed (additional 24 secondary implants). Therefore, a total

of 117 proGAV valves (primary and secondary implants)

were implanted in patients during infancy and the survival

of the valves was followed. The type of CSF diversion was

ventriculoperitoneal for all primary implants; among the

secondary implants were 2 ventriculoatrial shunts.

A total of 83 revisions had to be performed in 53 pa-

tients, among them 30 revisions in the preterm subgroup.

In summary, a total of 176 surgeries (including primary

surgeries and revision surgeries) were performed dur-

ing the entire follow-up period. A revision addressed the

nonfunctioning parts of the shunt system (e.g., ventricular

catheter, valve, or distal catheter; either alone or in com-

bination). In all cases of infection, the complete shunt sys-

tem was revised. A total of 19 shunts were revised due to

a documented shunt infection (infection rate 10.8%). The

ventricular catheter had to be revised 65 times in 41 chil-

dren (among them were 21 preterm infants and 19 infec-

tions), and the distal catheter had to be revised 28 times in

25 children (among them were 14 preterm infants and 19

infections). Forty-four valve exchanges were necessary in

37 patients; 20 of them were in the preterm subgroup.

A shunt infection was the reason for valve revision in

19 of 44 patients (among them 13 preterm infants) a mean

of 3.47 ± 6.7 months after implantation. Ten infections oc-

curred within the first month and a total of 16 infections

occurred during the first 3 months after implantation.

Three of the preterm infants needed 2 valve exchanges
each due to recurring infections. In these patients, explana-
tion, externalization, and administration of intravenous

antibiotics was performed. However, retrospective analysis
showed that the implantation of the system was sched-

uled too early, i.e., before complete CSF clearance of the

infection could be reached. Among the 6 infants who were

born at full term, there was only 1 recurrent infection.

An occlusion of the proGAV was seen in 14 revisions

after a mean of 28 ± 21.6 months; among them were 9

children with either a postinfectious or posthemorrhagic

hydrocephalus. No patient had more than 1 occlusion of

the valve, and the presence of protein clots was seen in

all valves. A dysfunction of the valve (i.e., the inability to

alter the setting of the valve) was found in 10 cases after

a mean of 33 ± 18.6 months. In 2 cases, the dysfunction

was due to improper use of the DP unit prior to implant-

tation. In those cases, the outer titanium membrane was
deviated by immoderate pressure and the blocking mecha-

nism of the rotor was incapacitated. In all other cases, the

manufacturer’s investigation revealed that protein debris

formed by immoderate pressure and the blocking mecha-
nism of the rotor was incapacitated. In all other cases, the

manufacturer’s investigation revealed that protein debris

was due to improper use of the DP unit prior to implant-

tation. In those cases, the outer titanium membrane was
deviated by immoderate pressure and the blocking mecha-
nism of the rotor was incapacitated. In all other cases, the

manufacturer’s investigation revealed that protein debris

around the rotor was blocking the adjustment mechanism.

One valve was exchanged to alter the setting of the gravi-
tational unit (Table 2).

Another reason for an operative revision of a shunt sys-

tem was impaired wound healing, which occurred in 5 in-

fants (2 of them preterm). This resulted in shunt infection

in 1 patient, 2 others had further surgeries including ven-

tricular catheter revision and valve occlusion, and 2 cases

received only wound revision as additional surgery after

shunt implantation.

**Shunt and Valve Survival Rates**

The Kaplan-Meier rate of shunt survival of the whole

patient cohort was 69% at 12 months and 34% at 85

months; for the valve only, the survival rate was 78% at 12

months and 56% at 85 months (Fig. 1A, Table 3). In con-

trast, the corresponding shunt rate for premature infants

was 61% at 12 months and 27% at 85 months, compared

with a 79% shunt survival rate at 12 months and 46% at

81 months in full-term infants (p = 0.054; Fig. 1B). The

survival rate of the valves in premature infants was 69%

at 12 months and 58% at 85 months compared with 89%

at 12 months and 53% at 81 months in full-term infants (p

= 0.26). No differences in survival rates were seen when

shunt implantation was performed in younger infants (age

≤ 100 days after the estimated day of delivery) compared

with older ones (> 100 days).

However, there was a significant difference if a previ-

ous surgery was performed before implantation of a new

proGAV system. The survival rates for primary implanted

shunts (without previous surgery) at the end of the observa-

tion period revealed a shunt survival of 78% at 12 months

and 39% at 85 months, whereas valve survival was 85% at

12 months and 59% at 85 months. This was observed to be

significantly better compared with infants with secondary

implanted shunts, in whom the implantation of the system

was performed as a revision (shunt: 50% at 12 months

and 28% at 79 months, p = 0.006; valve: 63% at 12 months

and 53% at 79 months, p = 0.07; Fig. 1C and D).

The Kaplan-Meier shunt and valve survival rates of the

2 largest etiological groups of hydrocephalus, posthemor-

rhagic and in association with a myelomeningocele, were

**TABLE 2. Number and sites of surgical revisions**

<table>
<thead>
<tr>
<th>Total No. of Revisions (n = 83)</th>
<th>Site of Revision</th>
<th>No. of Revisions</th>
<th>No. of Children (all/preterm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Combined revisions of:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ventricular catheter</td>
<td>65 (infection, 19; other, 46)</td>
<td>41:21</td>
<td></td>
</tr>
<tr>
<td>Distal catheter</td>
<td>28 (infection, 19; other, 9)</td>
<td>25:14</td>
<td></td>
</tr>
<tr>
<td>Valve (proGAV)</td>
<td>44 (infection, 19; occlusion, 14; dysfunction, 10; other, 1)</td>
<td>37:20</td>
<td></td>
</tr>
<tr>
<td>Wound</td>
<td>5</td>
<td>5:2</td>
<td></td>
</tr>
</tbody>
</table>
compared with the rest of the patient cohort. The shunt survival rates among patients with posthemorrhagic hydrocephalus compared with other causes for hydrocephalus were 30% versus 40% (p = 0.26) and the valve survival rates were 50% versus 62% (p = 0.11) at the end of the observation period. The valve survival rates among patients with hydrocephalus in association with a myelomeningocele compared with other causes for hydrocephalus were 68% versus 53% (p = 0.051; Fig. 1E) at the end of the observation period.

Discussion

Several studies have shown significant complications in the treatment of infants with hydrocephalus. Possible reasons for this are fragile skin conditions, immature immune response, soft and thin calvarium, and the open fontanelle leading to possible wound-healing problems, CSF collections, shunt infection, involution of the fontanelle, or subdural hygroma. Special pediatric implants were designed to be smaller and less voluminous, lack the option to adapt the amount of CSF diversion according to the patient’s activity and changing anatomy, and might need later surgical exchange to avoid overdrainage. The new technology of adjustable valves demonstrates the advantage of optionally adapting the drained CSF volume to the individual conditions of the patients, which might be of particular importance in very young infants who will outgrow the pressure setting of a fixed-pressure, low-resistance valve. Antisiphoning devices were introduced to overcome posture-dependent hydrostatic overdrainage.

The proGAV has been used at our institution since 2007 as the standard valve for all children. It provides both the opportunity to respond to changing pathophysiological conditions by adjusting the DP unit and a complementary system to counteract the siphoning effect of the gravitational unit. The valve was introduced in selected pediatric patients before 2007. After promising early results, the standard regimen was changed to include infant patients. In the present study, we report the shunt and valve survival rates and the reasons for valve exchanges exclusively for the infant cohort. The retrospective investigation of revisions and complication rates in this young patient cohort shows limited shunt and valve survival rates. However, we conclude that implantation of the proGAV is feasible in the infant age group and that implantation of the gravitational unit is tolerated. Its use adds the option of being able to adjust the valve settings early in the therapeutic regimen.

No significant differences were observed in the overall shunt and valve survival rates between patients younger or older than 100 days at the time of implantation. However, the inferior rate of shunt survival in patients who received the proGAV as part of a revision of their shunt system stresses the importance of the necessary optimal primary implantation. Especially within the first year after implantation, a significant loss of implants is observed, which might be influenced by a higher infection rate in this cohort compared with older children. The long-term decrease in the valve survival curve is rather related to obstruction or dysfunction of the valve. This is mostly associated with elevated protein CSF levels that are usually recognized when the valve needs a readjustment, which is not feasible at this stage.
Perhaps the most relevant data from the study include the survival rate of primary implants in infants without any previous interventions (n = 79), which was 85% at 12 months and 59% at 85 months for the valve and 78% at 12 months and 39% at 85 months for the shunt; these data may have the strongest value for comparison with data from the literature. In our previous report, which examined 237 children with hydrocephalus who had a proGAV implant, 62 infants with a mean follow-up of 21 months (6–38 months) were included. The Kaplan-Meier valve survival rate after 12 months for all children in this study was similar to the present data (valve, 81% and shunt, 72%). This previous study clearly demonstrated the higher risk of shunt failure in infants compared with older children (≤1 year, 77% vs > 1 year, 90.1%). The importance of a longer follow-up is apparent, because shunt failure becomes increasingly likely with longer follow-up.

Among other studies, Robinson et al. reported the complication rates in 158 infants with shunts, who received either no valve, a low-pressure valve, or a medium-to-high-pressure fixed DP valve (Table 4). With a total of 246 revisions during a follow-up period of 9 years, they reached the study end point with overall shunt survival rates of 52%, 48%, and 37% after 1, 2, and 5 years, respectively. Dividing the subgroups did show a significantly lower survival rate for the no- or low-pressure valve group, especially during the first 6 months after shunt insertion (shunt failure rates: 1 year 58% and 5 years 72% with no valve or a low-pressure valve, 1 year 31% and 5 years 47% with a medium- or high-pressure valve). Twenty-five revisions were necessary due to overdrainage. The relevant conclusion from this observation is that no- or low-pressure rates might cause major long-term complications due to significant overdrainage, which is mostly compensated for by infants and young children but to a lesser extent by adolescents. The authors concluded that a low valve opening pressure is an important contributing factor for shunt complications.

With a smaller cohort, the results from Weinzierl et al. underline the necessity to focus on overdrainage-related problems early in childhood. Their cohort included 15 infants < 6 months old, who were treated with an adjustable Hakim DP valve (Codman) without a siphon-compensating mechanism. The development of slit ventricles as a result of overdrainage, which were observed by ultrasound within the first 12 months, could not be sufficiently prevented by increasing the DP valve from an initial pressure of 10 cm H$_2$O to 15–20 cm H$_2$O. Two other available devices were used by Martínez-Lage et al. in an infant cohort. They implanted the adjustable valve Sophy SM3 as well as the subsequent valve types, SM8 and Polaris (Sophysa), in a cohort of 100 babies (40 preterm and 60 full term; < 2 months old). In this study, 31 revisions due to proximal obstruction, infection, technical problems, or “miscellaneous” reasons were reported. The death of 1 child was related to shunt malfunction. After a

<table>
<thead>
<tr>
<th>Variable</th>
<th>No. of Pts</th>
<th>Type</th>
<th>12 Mos</th>
<th>24 Mos</th>
<th>36 Mos</th>
<th>48 Mos</th>
<th>60 Mos</th>
<th>After Max Observation Period (mos)</th>
<th>% Survival</th>
<th>p Value*</th>
</tr>
</thead>
<tbody>
<tr>
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<td>Valve</td>
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<td>77</td>
<td>71</td>
<td>66</td>
<td>59</td>
<td>56 (85)</td>
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<tr>
<td></td>
<td></td>
<td>Shunt</td>
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<td>61</td>
<td>49</td>
<td>39</td>
<td>32</td>
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<td>77</td>
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<td>56 (85)</td>
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<tr>
<td></td>
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<td>Shunt</td>
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<td>61</td>
<td>49</td>
<td>39</td>
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<tr>
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<td>78</td>
<td>77</td>
<td>71</td>
<td>67</td>
<td>61</td>
<td>56 (85)</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>Shunt</td>
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<td>60</td>
<td>49</td>
<td>47</td>
<td>44</td>
<td>38 (85)</td>
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<td>69</td>
<td>64</td>
<td>58</td>
<td>58</td>
<td>58 (85)</td>
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<tr>
<td></td>
<td></td>
<td>Shunt</td>
<td>61</td>
<td>53</td>
<td>42</td>
<td>35</td>
<td>32</td>
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<td>79</td>
<td>77</td>
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<tr>
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<td></td>
<td>Shunt</td>
<td>79</td>
<td>70</td>
<td>58</td>
<td>53</td>
<td>46</td>
<td>46 (81)</td>
<td>0.046</td>
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<td>Valve</td>
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<td>63</td>
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<td>50</td>
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<td>Valve</td>
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<td>84</td>
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<td></td>
<td>Shunt</td>
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<td>70</td>
<td>57</td>
<td>51</td>
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<td>Valve</td>
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<td>66</td>
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<td>36</td>
<td>30 (85)</td>
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<tr>
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<td>Valve</td>
<td>83</td>
<td>81</td>
<td>81</td>
<td>75</td>
<td>62</td>
<td>62 (81)</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>Shunt</td>
<td>73</td>
<td>64</td>
<td>57</td>
<td>48</td>
<td>40</td>
<td>40 (81)</td>
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</tr>
<tr>
<td>MMC</td>
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<td>Valve</td>
<td>90</td>
<td>90</td>
<td>90</td>
<td>90</td>
<td>68</td>
<td>68 (81)</td>
<td>0.051</td>
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<tr>
<td></td>
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<td>Shunt</td>
<td>85</td>
<td>80</td>
<td>62</td>
<td>46</td>
<td>46</td>
<td>46 (81)</td>
<td>0.24</td>
<td></td>
</tr>
<tr>
<td>Others</td>
<td>97</td>
<td>Valve</td>
<td>75</td>
<td>74</td>
<td>67</td>
<td>62</td>
<td>56</td>
<td>53 (85)</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>Shunt</td>
<td>66</td>
<td>57</td>
<td>46</td>
<td>42</td>
<td>37</td>
<td>32 (85)</td>
<td>0.24</td>
<td></td>
</tr>
</tbody>
</table>

Max = maximum; MMC = myelomeningocele; NA = not applicable; PH = posthemorrhagic.
* Versus corresponding group.
Shunt survival in infants with hydrocephalus

Mean follow-up of 55 months, they performed 91 readjustments. Thirty-three of those became necessary after unintentional MRI-related readjustments (Polaris excluded), and 62% of all patients who underwent MRI needed 1 or more valve pressure resettings. Because we avoid routine CT scanning in children to minimize radiation and use MRI as the routine elective for imaging, we appreciate the advantage of the proGAV or the Polaris valve, which are inert to magnetic field exposure up to 3 T.\(^3,4,28–30,34,35\)

Another aspect is to compare our data with those of nonadjustable, gravity-assisted valves collected in a multicenter prospective study of 169 patients, including 86 infants (< 1 year) who received the paediGAV;\(^20\) this study reported shunt survival rates of 65.6% and 57.4% and valve survival rates of 72.4% and 71.1% after 12 and 24 months, respectively. However, the comparison of a single-center retrospective study with a multicenter prospective study is limited. In another multicenter study by Hanlo et al., a population of 557 patients from all age groups underwent implantation of the nonadjustable Orbis-Sigma II valve.\(^21\) A significant difference was observed in the overall shunt survival rate between patients younger than 6 months (55%) and those who were older than 6 months (63%). The authors also reported that survival differences in infants occurred within the first few months after surgery, which is consistent with our results and those of other studies in which a failure rate of 48%–58% within the first year is described.\(^8,14,23,36,42,45\) This is also emphasized by Reinprecht et al.,\(^41\) who reported results from a cohort of 42 preterm infants with posthemorrhagic hydrocephalus and shunt implantation. They noted occlusion of the distal catheter as the primary cause for revision, which occurred mainly during the first few months.

Although our demonstrated survival rates of the implanted proGAV shunt system are comparable to the rates reported in the literature, limitations of our study include its retrospective, single-center setup without a control group. Furthermore, the observation period is still too short to adequately assess signs and symptoms of chronic overdrainage and to answer the question of whether a gravitational unit with a resistance of 20 cm H\(_2\)O is enough for long-term follow-up. Nevertheless, the feasibility of using this shunt system in an infant population is demonstrated, which has similarly been shown for a general pediatric population.\(^20,43,48\) The use of the gravitational unit of the system to try to counteract severe overdrainage is also well tolerated by infants. Clinical observation has shown that during long-term follow-up, signs and symptoms of overdrainage (e.g., decreased ventricular size on imaging, diminished development of head circumference, and headaches) may develop with the advancing growth and activity of the child. Therefore, the adjustability of the DP unit of proGAV is important because it adds the option to gradually increase the resistance of the shunt system.\(^18\)

To generally improve shunt survival in infants, therapeutic approaches are needed. These include the strict protocol to avoid shunt infections and, possibly, surgical approaches to decrease CSF protein concentration before shunt implantation. In this regard, we have recently introduced neuroendoscopic lavage of the ventricular system, which our institution routinely uses for intraventricular

<table>
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<th>Table 4: Literature review of valve types and outcomes</th>
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<td><strong>Authors &amp; Year</strong></td>
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<tr>
<td>Present study</td>
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<tr>
<td>Robinson et al., 2002</td>
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<td>Haber et al., 2003</td>
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<td>Notarianni et al., 2009</td>
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<td>Golmard et al., 2012</td>
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<td>GAV = gravity-assisted valve, GU = gravitational unit.</td>
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hemorrhage and postinfectious conditions. For this study, however, it has a limited impact because only 5 infants with posthemorrhagic and 2 infants with postinfectious hydrocephalus who were receiving neuroendoscopic lavage were included. The outcome data of this approach are currently being evaluated to quantify survival rates with this enhanced protocol. A further challenge will be to investigate the long-term neurofunctional outcome in these infant patients to prove that counteracting chronic overdrainage is beneficial.

Conclusions

Hydrocephalus in infants represents a special cohort of patients in whom avoiding shunt complications remains challenging. The survival rates of the proGAV system, which is used routinely in our institution for all children, demonstrate good applicability of this system in this age group, but do not show much difference compared with other studies and other valve systems as published in the literature. The gravitational unit with a pressure setting of 20 cm H₂O is well tolerated by all infants without showing symptoms of underdrainage and may avoid problems of overdrainage from early in treatment. Whether this will apply for much longer durations of follow-up cannot be answered by this study. The adjustable DP unit aims to apply flexibility in a cohort of patients for whom the optimal opening pressure is hard to predict. The rationale is that the initial use of a sophisticated technology for infants with hydrocephalus might achieve long-term quality of care. Further studies are necessary to prove the benefit of this approach and to collect data on neurofunctional outcome.

References
