**Intrathecal baclofen trials: complications and positive yield in a pediatric cohort**

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**OBJECTIVE** Intrathecal baclofen (ITB) is an effective management option for childhood hypertonia. Given the potential complications of implanted ITB pumps, trials of ITB are usually performed as part of the workup for ITB pumps. Two methods are used for ITB trials, lumbar puncture (LP) and catheter insertion into the intrathecal space. Little has been written to date on the number of positive trials and complications in trials. This study aimed to report the outcomes and complications in ITB trials for childhood hypertonia (dystonia, spastic, or mixed).

**METHODS** A retrospective case notes review was conducted of all patients who underwent ITB trials at the Evelina London Children’s Hospital between 2005 and 2012 (inclusive). Positive trials were defined as a reduction in Modified Ashworth Scale by a minimum of 1 point in at least 2 muscle groups and improvement reported by the caregivers in the areas of goals agreed upon between professionals and the families.

**RESULTS** Our patient group comprised children with dystonia (n = 7), mixed spasticity/dystonia (n = 29), spasticity (n = 4), and pain (n = 1). A total of 47 trials were attempted in 41 children. Forty trials were successfully completed, with 39 being positive. Thirty-three were catheter trials, and 14 were LPs. The overall complication rate in the 47 attempted trials was 53%; 61% in catheter trials, and 36% in LP trials. This difference was not statistically significant.

**CONCLUSIONS** There is a high risk of minor self-limiting complications with ITB trials, which needs to be factored into the decision process of progression to trials. The rate of positive trials in this study was 98%, of which 21% did not progress to pump implantation. While the authors would still advocate for ITB trials prior to ITB pump insertion to aid parental decision-making, this figure suggests that with good patient selection, ITB pumps could be placed without a preceding trial.

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**KEY WORDS** dystonia; baclofen; intrathecal; children; complications; functional neurosurgery

**ABBREVIATIONS** CMDS = Complex Motor Disorders Service; GMFCS = Gross Motor Function Classification System; ITB = intrathecal baclofen; LP = lumbar puncture; MAS = Modified Ashworth Scale.


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There is growing literature regarding the efficacy of ITB pumps for high tone and the complications related to the surgery and ongoing care. To date, little published information is available regarding ITB trials, how they are performed, the incidence of positive/negative trials, and the potential complications or risks. The 2 methods for conducting ITB trials are lumbar puncture (LP) and an intrathecal catheter. LP is typically performed with the patient awake, which has the advantage of enabling the entire trial to be undertaken within a day without necessitating waiting for the patient to recover from general anesthesia. However, LP access can be traumatic for the child and difficult for the clinician, particularly in the presence of spinal deformity or extensor posturing. ITB trials may need to be repeated at higher doses to ensure a positive effect, such that a patient might require multiple LPs. A catheter trial, on the other hand, typically involves insertion of an intrathecal catheter with the patient under general anesthesia and administration of the test dose once the patient has recovered from the anesthesia. The catheter can be kept in place for repeat doses, if necessary, or it can be used for a continuous infusion via an external pump. Reports of outcomes and complications of ITB trials are currently limited.

The aim of this study was to report the outcome of ITB trials in a consecutive cohort of children and young people undergoing assessment at our center and to report the incidence and nature of complications arising in the cohort.

Methods
Study Participants
All patients undergoing an ITB trial between 2005 and 2012 (inclusive) were identified from the Complex Motor Disorders Service (CMDS) database at the Evelina London Children’s Hospital. Data were collected from case notes and entered on a standardized form. Data collected included sex, age at referral to CMDS, age at referral for the ITB trial, motor phenotype (dystonia, spasticity, or mixed),26 age at trial, underlying pathology, Gross Motor Function Classification System (GMFCS) level25 (in non-cerebral palsy cases a “GMFCS equivalent” score was applied as reported previously by our group24), the mode of the trial, dose of baclofen, method of scoring of the trial, trial result (positive or negative), all adverse events during admission for the trial and, if applicable, age at referral for pump, age at pump implant, and reason for not having a pump.

At the Evelina London Children’s Hospital, patients are offered ITB trials after a thorough assessment by a pediatric neurologist and experienced pediatric physiotherapist during an appointment in the CMDS clinic.

Selection criteria for ITB trials were generalized or segmental hypertonia, predominantly in the lower limbs, interfering with comfort, care, or function; insufficient response to addressing exacerbating factors and other treatment options, such as physiotherapy, occupational therapy, medications, and botulinum toxin; and realistic expectations/goals for intervention.

Caution is applied in cases of unrealistic expectations, significant central hypotonia, respiratory concerns related to upper airway or frequent chest infections, identified risk factors for poor compliance, and in very small patients (i.e., weight below 14–15 kg). Primary dystonia is preferentially considered for deep brain stimulation.1

Trial Method
Decisions about the trial mode were typically made in the clinic based on clinical assessment and family preference. LP was offered when it was believed to be clinically feasible to perform the procedure with the child awake. Insertion of the catheter under general anesthesia was planned when there was significant spinal deformity or it was anticipated that the child would be uncooperative and difficult to position, when an anesthetic for brain MRI was planned prior to the trial, or when the family was averse to an awake LP. As observed in our practice, sedative medications or general anesthesia can reduce hypertonia for more than 2 hours, particularly in cases of dystonia; therefore, to avoid unclear trial results, we do not administer test ITB doses with the patient under sedation or general anesthesia (also a reported practice1,17). Patients were referred to the neurosurgical team for catheter insertion with radiographic guidance when severe spinal deformity was anticipated to prohibit successful access or in the presence of comorbidities such as known hydrocephalus.

Both LP and catheter insertions were performed as sterile procedures following careful skin preparation and using standard LP needles and epidural catheters available at our pediatric hospital. Trials were conducted according to the local protocol by a pediatric neurologist and physiotherapist on the pediatric neurology ward, with an experienced nurse allocated to the patient. LP was performed following application of topical anesthesia using a 22-gauge Quincke spinal needle with patients lying on their side. Once CSF flow was observed from the needle, a dose of ITB was administered from a syringe as a slow push over 1–2 minutes.

For the sterile catheter procedure, an epidural catheter (most commonly 18-gauge, or in smaller patients 24-gauge) was inserted into the intrathecal space while the patient was under general anesthesia and secured with sterile dressing on the skin. The catheters were not tunneled. Most of the catheters were inserted by a pediatric neurologist without available radiographic guidance. In some cases, the catheter was inserted by a neurosurgeon with radiographic guidance. Once the child recovered from the anesthesia, usually the following morning, the catheter site was inspected, and the catheter access port was then aseptically accessed for aspiration of 1 ml of CSF to ensure appropriate catheter position and integrity. The baclofen dose was then slowly administered (over 1–2 minutes) followed by a slow flush of 1–2 ml of 0.9% NaCl, and then the port was secured. Following drug administration, patients were placed supine with the head elevated 20° to 30°.

The baclofen dose was weight dependent. If the child weighed less than 20 kg, a starting dose of 25 µg was used; for children over 20 kg, the starting dose was 50 µg. If no response was documented, the dose was escalated by 25 µg and repeated daily, aiming for a maximum of 3 administrations. A dose of 12.5 µg was used in 1 patient.
who weighed 14.5 kg. We have not used continuous infu-
sion via an external pump.

A pediatric neurologist and a physiotherapist assessed
the patients prior to administration of the dose and 2, 4,
and 6 hours afterward.

Response to the ITB test dose was evaluated using the
Modified Ashworth Scale (MAS) score, passive range of
movement, presence and quality of deep tendon reflexes,
and subjective comments from children and caregivers
e.g., related to comfort, ease of care, and positioning). The
MAS score was used pragmatically for the assessment of
our whole cohort. With the majority of patients having
dystonia as a contributing feature, evaluation of their
hypertonia with MAS presented assessment challenges. Two
experienced professionals performed assessments inde-
dependently, and a positive response was documented when
detected by both raters. A positive trial was defined as the
reduction in the MAS score by a minimum of 1 point in
at least 2 muscle groups and improvement reported by the
patients/caregivers in the areas of goals agreed upon be-
tween professionals and the families.

All children remained in the hospital for a minimum of
24 hours after the trial. Fluid intake was carefully moni-
tored, with intravenous fluids given if there was reduced
fluid intake or vomiting. Patients were encouraged to lie
flat in bed for a minimum of 12 hours after LP and fol-
lowing catheter removal, as well as with the catheter in
situ. Any adverse event documented as arising during and
following the trial was considered a complication of the
trial. Furthermore, any adverse experiences reported by
the families within a few days following discharge that
were believed to be related to the trial were considered
complications.

Results

A total of 41 children (16 female, 25 male) were identi-
fied as having had an ITB trial between 2005 and 2012
(inclusive). The mean age in years at trial was 9, with a
range of 3.3 to 17.6 years. Table 1 summarizes the clinical
features of the patients; Table 2 summarizes the GMFCS
scores.

A total of 47 trials were attempted in 41 patients. Of
the 47 trials, 40 (85%) were successfully completed, with
34 patients (83%) having completed the trial on the first
attempt. Of the 7 trials that failed on first attempt, 1 (2%)
was a failed LP that was subsequently successful with
catheter placement, and 3 (6%) were failed catheter place-
ments by a pediatric neurologist that were later completed
after catheter placement under radiographic guidance by
a neurosurgeon. In 2 patients, the catheter became dis-
lodged before the trial was started. In both patients, the
catheter method had been chosen, because the patients
had required general anesthesia for brain and spinal MRIs
prior to the trial. These were both successfully converted
to LP trials. One catheter was removed due to continued
excessive CSF leakage, and the trial was not reattempted
at parental request.

The intrathecal catheter was used more commonly than
LP bolus. A total of 7 patients required catheter insertion
by a neurosurgeon, including the 3 patients with previ-
ously failed attempts and 4 patients referred directly to the
urosurgeon (3 due to significant lumbar lordosis and 1
due to hydrocephalus with a ventriculoperitoneal shunt).
The complication rate and types of complications for both
methods are summarized in Tables 3 and 4.

Although 61% of patients in the catheter trials and 36%
of patients in the LP trials had complications, the differ-
eance in complication rate was not statistically significant
(Fisher’s exact test, p = 0.4971).

Of the 40 patients who successfully completed the trial,
1 had a negative response and 39 had positive responses
(98%), with 31 (79%) of these progressing to ITB pump
placement. Eight patients did not progress to the ITB
pump despite a positive trial. Reasons included respira-
tory concerns identified during a sleep study after the trial
although no respiratory complications were noted at trial
(n = 2), medications trial/alternative therapies improved
condition (n = 2), parental concern following ITB trial
complication of meningitis (n = 1), body habitus too small
to accommodate a pump (n = 1), awaiting pump placement
at the time of data collection (n = 1), and unknown (n = 1).

The child for whom the ITB trial was considered nega-
tive presented with evidence of leukodystrophy on MRI of
the brain. Clinically, he exhibited minimal hypotonicity
of the lower limbs with some upper motor neuron signs
present, and pain as the predominant symptom for which
ITB was tested. Tone improved following ITB bolus but
no improvement in pain was noted according to a numeri-
cal rating scale for self-report of pain, and a significant
underlying weakness was unmasked. This was therefore
classified as a negative response.

The majority of trials were concluded with 1 dose of
baclofen, with 11 patients receiving a second higher dose.
A third dose was not required in our cohort. A dose of 100
µg was used once and resulted in deep sedation.

Discussion

In a consecutive cohort of 40 children completing an
ITB trial at our institution, we found positive responses
to the trial in all but 1 patient and a complication rate of
53% (61% for catheter trials and 36% for LP trials). In all

<table>
<thead>
<tr>
<th>TABLE 1. Clinical phenotype of patients</th>
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<tbody>
<tr>
<td>Phenotype</td>
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<tr>
<td>Mixed spasticity/dystonia</td>
</tr>
<tr>
<td>Dystonia</td>
</tr>
<tr>
<td>Spasticity</td>
</tr>
<tr>
<td>Pain</td>
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<tr>
<th>TABLE 2. Gross Motor Function Classification Score of patients</th>
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<tr>
<td>GMFCS</td>
</tr>
<tr>
<td>-------</td>
</tr>
<tr>
<td>3</td>
</tr>
<tr>
<td>4</td>
</tr>
<tr>
<td>5</td>
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Albright and Ferson\textsuperscript{4} described the use of ITB infusion via external pump for trials in patients with dystonia; however, they reported that for younger and smaller children a single bolus might be sufficient to determine a response. We have not used continuous infusions but were ready to consider this method in dystonic patients if a response to bolus was negative.

Although the complication rate did not statistically differ between the 2 trial methods, catheter trials caused patients to commonly have more than 1 complication. The profile of complications differed between the 2 trial types. In the LP bolus method, only 1 child had more than 1 complication, and the only complications described were vomiting, infection (not meningitis or LP-site infection), and sedation associated with a small reduction in blood pressure (short lived with no respiratory compromise). For the catheter group, a greater range of complications was noted, including infection (minor in 2 patients, severe in 1 patient [meningitis]), symptoms related to low intracranial pressure from CSF leakage, and side effects from baclofen itself (e.g., dizziness/drowsiness).

The most severe complication, meningitis, occurred following a catheter trial. The patient developed symptoms of meningitis (fever, headache, vomiting, photophobia, and neck stiffness) within 24 hours of catheter removal. The tip of the catheter is routinely sent to the microbiology department for testing and in this case was positive for \textit{Staphylococcus aureus}. The patient was treated with intravenous antibiotics for 14 days according to sensitivity and made a full recovery. Although the trial had been positive, this complication discouraged the patient's parents from pursuing the ITB pump. Deterioration of dystonia prior to and shortly after the trial was recorded in 2 patients. We believe this deterioration was related to discomfort caused by the presence of a spinal catheter. In both cases, reduction in tone and dystonic spasms was observed following administration of the baclofen test dose.

The patient group reported here is complex, with dystonia contributing to the high tone in the vast majority of the patients. Only 4 patients with pure spasticity were included. In the presence of dynamic posturing, mechanical complications such as catheter displacement may be more likely. The small numbers of patients in different groups reported here preclude a more detailed analysis.

Statistical comparison of the complication rate between each group is limited by the small sample size and the disproportion between group sizes. To be powered to detect a difference in proportions of complications in the 2 groups as demonstrated in our sample at the level of statistical significance, a sample size of 75 in each group would have been required. Assuming that the complication rate in the catheter group represents the “true” complication rate, given the sample size in our study, the analysis was powered only to reach statistical significance if the complication rate in the LP group was below 16% or greater than 95%. We cannot, therefore, conclude that there is no difference in complication rates, although our results raise the possibility of a trend in this direction.

Complications in ITB trials have been previously reported in articles focusing on the efficacy of ITB, including unilateral radicular pain,\textsuperscript{29} symptoms of low intracranial pressure,\textsuperscript{4,17,23} infection at the skin level at the catheter site, and more severe complications such as positive blood cultures,\textsuperscript{3,28} dizziness,\textsuperscript{17,28} and drowsiness.\textsuperscript{3,17,19,28} Deep vein thrombosis and pulmonary embolism have also been published as a case report.\textsuperscript{12}

In a retrospective report by Vloeberghs et al.\textsuperscript{29} that focused on the outcomes of ITB pump implantation in spasticity due to cerebral palsy, 10% of 85 patients undergoing a trial had spinal tap syndrome with no other complications reported. All of these were catheter trials. Hoving et al.\textsuperscript{17} in a prospective study reported 14 of 17 (82%) patients in their cohort with catheter trials had symptoms

<table>
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<tr>
<th>Complications by trial method Variable</th>
<th>Catheter</th>
<th>LP Bolus</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of patients</td>
<td>33</td>
<td>14</td>
</tr>
<tr>
<td>No. of patients w/ complications (%)</td>
<td>20 (61)</td>
<td>5 (36)</td>
</tr>
<tr>
<td>Back pain</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Vomiting</td>
<td>7</td>
<td>2</td>
</tr>
<tr>
<td>Dehydration</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>CSF leak</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Infection</td>
<td>3 (1 meningitis)</td>
<td>1</td>
</tr>
<tr>
<td>Dizziness</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Drowsiness</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Headache</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>Deep sedation</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td>Worsened dystonia prior to &amp; after trial</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>Hypotension</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Catheter removal prior to use</td>
<td>1</td>
<td>NA</td>
</tr>
<tr>
<td>Catheter displacement</td>
<td>2</td>
<td>NA</td>
</tr>
<tr>
<td>Catheter placement failed</td>
<td>3</td>
<td>NA</td>
</tr>
<tr>
<td>LP failed</td>
<td>NA</td>
<td>1</td>
</tr>
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\textit{NA} = not applicable.
of lowered CSF pressure, with other complications being reported. Three of 17 patients in whom the catheter was tunneled did not have symptoms of low CSF pressure.

Our overall complication rate was 53%. In our retrospective study, we included all adverse events recorded in case notes, regardless of whether alternative explanations could be suggested as to the cause of the reported adverse feature. For example, for 2 of the potential infections, short-lasting pyrexia alone was reported, which might relate to other aspects of the hospital stay that exposed our patients to potential infections.

The differences in complication rates can be related to the different trial methods, study design, methodology of data collection, and definition of complications (e.g., individually recorded vomiting, headache, and CSF leak in our study versus spinal tap syndrome\(^{23}\)).

An important finding of this study is the high positive response rate. Of the 40 successfully completed trials, all but 1 child demonstrated a positive response. One patient had a negative trial, for whom reduction of pain was the primary goal rather than tone, the latter responding to baclofen. This suggests a highly selective process for progression to trial. However, 21% of those did not progress to the ITB pump implant for different reasons, and 1 patient’s family refused progression to the pump due to the complications experienced during the trial. There is a similar report by Murphy et al. of parents declining pumps due to complications experienced during the trial of ITB therapy in a pediatric cohort with cerebral palsy.\(^{22}\)

A high incidence of positive ITB trials has been reported, and 100% positive trial results have been found in smaller studies. Some centers in the US and Europe have abandoned ITB trials due to expected positive response and high complication rates. Performing ITB trials was related to complications but was found not to have an effect on risk of dissatisfaction with ITB pump implantation, although more studies are required to confirm this finding. The expense of trials is significant due to required hospital admission and involved procedures.

Our findings add evidence to support progressing directly to ITB pump implantation in carefully selected patients with hypertonia in specialized centers. However, ITB trials might be useful in ambulant patients and those with complex hyperkinetic movement disorders, where benefits of ITB therapy are less well documented. The ITB trial may remain a useful tool to help patients/families make an informed decision, particularly given the high rate of complications from baclofen pumps and the long-term commitment to meticulous follow-up for pump refills. Additionally, the level of response to the test dose guides the initial ITB daily infusion rate, usually calculated as twice the effective trial dose. Documenting the individual response at the trial might be a useful reference tool if loss of effect from ITB therapy is suspected.

It is important to acknowledge the limitations of this study. This is a relatively small sample that is heterogeneous in terms of etiology and the dominant movement disorder phenotype. It is also a relatively severely affected cohort as seen by the GMFCS levels. Data were retrospectively collected and consequently adverse events attributed to the ITB trial may have had alternative explanations.

Conclusions

We have reported the outcomes of ITB trials in a consecutive cohort of patients undergoing assessment at our center. A relatively high complication rate was observed. Overall, the complication rate did not statistically differ between the 2 methods used in the ITB trial, although children receiving the catheter demonstrated the potential for multiple complications. The most severe complication, meningitis, occurred in a catheter trial. All but 1 child demonstrated a positive response. These results support proceeding directly to ITB pump implants in patients carefully selected by an experienced multidisciplinary team. Trials might be useful to demonstrate to families the potential gains from ITB therapy. Trials continue to play an important role in the selection process for ambulant patients and hyperkinetic as opposed to hypertonic movement disorders.

The final decision of whether or not to perform the trial should be made together by the involved clinical team and the patient/family carefully considering the indications for the trial while being mindful of the common complications encountered.

References

Complications of ITB trials in childhood


Disclosure

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Author Contributions

Conception and design: Sayer, Lumsden. Acquisition of data: Sayer. Analysis and interpretation of data: Sayer, Lumsden. Drafting the article: Sayer. Critically revising the article: Kaminska, Lumsden, Perides, Tustin, Bassi. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Kaminska. Statistical analysis: Sayer, Lumsden, Lin. Study supervision: Kaminska, Lumsden.

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