IncobotulinumtoxinA for the treatment of lower-limb spasticity in children and adolescents with cerebral palsy: A phase 3 study

Florian Heinen, Petr Kaňovský, A. Sebastian Schroeder, Henry G. Chambers, Edward Dabrowski, Thorin L. Geister, Angelika Hanschmann, Francisco J. Martinez-Torres, Irena Pulte, Marta Banach and Deborah Gaebler-Spira

Abstract.

PURPOSE: Investigate the efficacy and safety of multipattern incobotulinumtoxinA injections in children/adolescents with lower-limb cerebral palsy (CP)-related spasticity.

METHODS: Phase 3 double-blind study in children/adolescents (Gross Motor Function Classification System – Expanded and Revised I–V) with unilateral or bilateral spastic CP and Ashworth Scale (AS) plantar flexor (PF) scores ≥ 2 randomized (1:1:2) to incobotulinumtoxinA (4, 12, 16 U/kg, maximum 100, 300, 400 U, respectively) for two 12- to 36-week injection cycles. Two clinical patterns were treated. Pes equinus (bilateral or unilateral) was mandatory; if unilateral, treatment included flexed knee or adducted thigh. Endpoints: Primary: AS-PF change from baseline to 4 weeks; Coprimary: investigator-rated Global Impression of Change Scale (GICS)-PF at 4 weeks; Secondary: investigator’s, patient’s, and parent’s/caregiver’s GICS, Gross Motor Function Measure-66 (GMFM-66).

RESULTS: Among 311 patients, AS-PF and AS scores in all treated clinical patterns improved from baseline to 4-weeks post-injection and cumulatively across injection cycles. GICS-PF and GICS scores confirmed global spasticity improvements. GMFM-66 scores indicated better motor function. No significant differences between doses were evident. Treatment was well-tolerated, with no unexpected treatment-related adverse events or neutralising antibody development.

CONCLUSION: Children/adolescents with lower-limb spasticity experienced multipattern benefits from incobotulinumtoxinA, which was safe and well-tolerated in doses up to 16 U/kg, maximum 400 U.

Keywords: Botulinum toxin, spasticity, cerebral palsy, all movement disorders, all paediatric, incobotulinum
of permanent disorders of movement and/or posture and of motor function, which are due to a nonprogressive interference, lesion, or abnormality of the developing/imature brain. The core motor dysfunction symptoms are often accompanied by impairments in sensation, perception, cognition, communication, and behaviour as well as epilepsy and secondary musculoskeletal problems [2].

Spasticity affects approximately 70–90% of children with CP [3–5]. The increased muscle tone due to spasticity results in a limited range of passive and active motion in joints and contributes to development of joint contractures, poor muscular control, and hyperactive reflexes [2]. In the lower limb (LL), it often manifests as pes equinus, a deformity associated with insufficient dorsiflexion of the ankle that prevents the heel from contacting the ground, which may mean that walking is done on the toes [6]. Pes equinus is most common in children with CP and LL spasticity [7], but a number of other spastic patterns including knee flexion and hip flexion/adduction may also present [8]. Pediatric spasticity has been associated with reduced health-related quality of life [9,10], which may be attributed in part to factors such as reduced mobility [11], inability to self-care [12], and pain [13–15].

A well-rounded treatment plan for a child with CP generally includes systematic rehabilitation, pharmacotherapy, physiotherapy, and perhaps surgical interventions [2,16] to reduce muscle spasms, facilitate mobility and dexterity, improve ease of care, improve posture, minimize contractures and deformity, reduce pain, and improve quality of life [17]. Factors to consider when determining a treatment plan to optimize function for a child with CP include age, stage of development, and distribution of muscle impairment, including the level (i.e., upper limb [UL] and/or LL), and pattern (i.e., unilateral vs. bilateral; pes equinus only or in combination with other muscle groups).

An individualized, multilevel, multipattern focal treatment approach to target specific muscle groups, especially those underlying particular functional deficits, can help address the diverse clinical presentation of spasticity in children with CP [18,19]. Botulinum neurotoxin type A (BoNT-A) is a recommended therapy for pediatric spasticity [20–24] that has been demonstrated to be effective and well-tolerated and can be well integrated in such a multimodal, multiprofessional, interdisciplinary treatment approach [25].

Three BoNT-A formulations, onabotulinumtoxinA, abobotulinumtoxinA, and incobotulinumtoxinA, are currently available in the European Union (EU) [26–28] and North America [29–31] for the treatment of spasticity. Specific indications for these BoNT-As vary by region and product in adults and in children/adolescents.

The phase 3 Treatment with IncobotulinumtoxinA in Movement (TIM) study investigated the efficacy and safety of three dose levels of incobotulinumtoxinA administered to a heterogeneous group of children and adolescents with LL spastic CP. TIM included patients who manifested all levels of CP disease severity and ambulatory ability. Each patient was treated in a multipattern approach with two clinical LL patterns. Depending on the investigator’s clinical judgement, treatment for pes equinus could be bilateral or unilateral and, if unilateral, include additional ipsilateral muscle groups. The study utilized several types of assessments to measure efficacy and considered the patients’, parents’/caregivers’, and clinicians’ perspectives. The unique study design allowed for individualization of treatment within standardized trial guidelines, reflective of real-world clinical patient needs.

2. Methods

2.1. Participants

The TIM study recruited ambulant and non-ambulant children and adolescents, aged 2–17 years, with unilateral or bilateral LL spasticity due to CP. To be enrolled, patients were required to have a clinical need determined by clinicians for treatment of either unilateral pes equinus with treatment of ipsilateral flexed knee or adducted thigh, or bilateral pes equinus. In addition, patients had to have a clinical need for incobotulinumtoxinA 16 U/kg treatment of LL spasticity and an Ashworth Scale [32] of the plantar flexor (AS-PF) score of ≥ 2 (at least unilaterally; bilaterally for the treatment of bilateral pes equinus) at screening and baseline. Investigators evaluated the clinical needs of each patient and selected the appropriate treatment pattern. Patients could be BoNT treatment naïve (i.e., no BoNT treatment within 12 months prior to study treatment) or have previously received treatment with BoNT (i.e., > 16 weeks prior to study treatment).

Key exclusion criteria included fixed contracture or predominant forms of muscle hypertonia other than spasticity (e.g., dystonia) in the target limbs, surgery for pes equinus in the target limbs within 12 months prior to screening or planned within the study period, hip flexion requiring orthopedic management and/or BoNT-A injection or limitation of hip abduction to < 40°, or
Muscles treated for each pattern:
- Pes equinus: gastrocnemius (medial and/or lateral), soleus, tibialis posterior, and flexor digitorum longus/flexor hallucis longus. For both unilateral and bilateral injection of pes equinus, at least two of five muscles contributing to pes equinus were required to be injected in each treated LL.
- Flexed knee: semitendinosus, semimembranosus, biceps femoris, gracilis
- Adducted thigh: gracilis, adductor longus/brevis, adductor magnus

Fig. 1. Treatment according to (A) clinical patterns and (B) study design. *Study visits ± 3 days. Additional bi-weekly TC to check for eligibility for reinjection. c Additional control visits every 6 or 8 weeks from 14 weeks up to 36 weeks after each injection. Patients were randomized to one of three dose levels, and U is the maximum dose divided between the muscles at each site. The bilateral clinical pattern refers to treatment of pes equinus in both LLs, and the unilateral clinical pattern refers to treatment of pes equinus on one side and either ipsilateral adducted thigh or ipsilateral flexed knee. BW = body weight; IC = injection cycle; kg = kilogram; LL = lower limb; TC = telephone contact; U = Unit.

a prediagnosed migrational percentage > 30. Patients with a severe neurological diagnosis and comorbidity outside the spectrum of CP and those with pure dyskinetic CP or mixed CP with predominantly dyskinetic movements were also excluded.

2.2. Study design and treatment

The TIM study was a prospective, double-blind, randomized, multicenter, parallel-group, phase 3 study conducted in 45 sites across 14 countries worldwide. Eligible patients were randomized 1:1:2 to three parallel incobotulinumtoxinA dose groups, respectively:

- Low dose: 4 units/kilogram (U/kg) body weight (BW), maximum total dose 100 U; mid dose: 12 U/kg BW, maximum total dose 300 U; high dose: 16 U/kg BW, maximum total dose 400 U.

Two LL clinical patterns were selected for treatment for each patient, one of which was required to be pes equinus on one side of the body. The patterns chosen by the investigator reflected the patient’s clinical need for therapy, with consideration given to the severity of the involved spastic muscles of the clinical pattern, subject age/weight and muscle size, activity, and experience from previous BoNT treatments. In the bilateral group, patients were treated for pes equinus on both sides of
the body (Fig. 1A). In the unilateral group, patients were treated for pes equinus and ipsilateral flexed knee or adducted thigh. In this group, patients with an AS score \( \geq 2 \) in the flexed knee and/or adducted thigh had one pattern chosen for treatment based on the investigator’s judgement. Each clinical pattern was treated with half of the total incobotulinumtoxinA dose (2, 4, or 8 U/kg incobotulinumtoxinA with a maximum dose of 50, 150, and 200 U, respectively, per clinical pattern). The muscles treated for each clinical pattern are specified in Fig. 1A.

At the initial screening visit, each patient was evaluated medically for inclusion in the study, including Gross Motor Function Classification System (GMFCS) classification, AS score, and presence of pain; participants were also questioned about past and concomitant medications within the last 4 weeks, and prior BoNT-A medications. After a 14-day screening period which allowed investigators to check each subject’s eligibility for study participation, treatments were administered during two consecutive double-blind injection cycles, each followed by 12–36 weeks of observation (Fig. 1B), giving an overall study duration of 26–74 weeks. The injections were administered according to the study’s standardized treatment plans with predefined dose ranges and injection-site numbers for each muscle. Equal injection volumes were administered in all dose groups (total volume up to 8 mL; 4 mL/clinical pattern), with dose ranges and injection volumes adjusted for patients with < 25 kg BW. At least one form of technical guidance (ultrasound, electrical stimulation, or electromyography) was required for injections, and site-individualized local anesthesia and/or analgesia protocols could be employed as needed.

Eligibility for reinjection was assessed regularly from 12–36 weeks post-injection. The treatment plan defined for the first injection cycle was continued in the second injection. Patients were eligible for re-treatment if they had an investigator- and patient-agreed clinical need for reinjection in the LL(s) and clinical patterns chosen at the injection visit of injection cycle 1, and an AS score \( \geq 2 \) in the treated clinical pattern. For patients with an AS score of 1, the investigator decided whether to re-treat. The injection interval was flexible and based on clinical need. The time to reinjection for each of the three incobotulinumtoxinA dose groups was analyzed descriptively.

Participants were allowed to maintain prior usual and concomitant therapies. These included nonpharmacological therapies such as physical therapy, orthotic management other than casting and rehabilitation, and pharmacological treatments, such as muscle relaxants and antidepressants. Patients who completed the TIM study had the option of enrolling in the open-label Treatment with IncobotulinumtoxinA in Movement Open Label (TIMO) study with 4 further injection cycles.

2.3. Standard protocol approvals, registrations, and patient consent

The study was conducted in accordance with the ethical principles of the Declaration of Helsinki and registered on clinicaltrials.gov (NCT01893411). The study protocol, informed consent forms, and other appropriate study-related documents were reviewed and approved by the local independent ethics committees and institutional review boards. Parents/guardians of all patients provided written informed consent, and patients provided assent if applicable.

2.4. Efficacy endpoints

The primary efficacy endpoint was the change from baseline in the AS-PF on the primary body side chosen for treatment at week 4 of injection cycle 1. The 5-point AS-PF scale was used, ranging from 0 (no increase in muscle tone) to 4 (limb rigid in flexion or extension). The coprimary efficacy endpoint was the investigator’s Global Impression of Change of Plantar Flexor Spasticity Scale (GICS-PF) score on the primary body side chosen for treatment at week 4 compared with the condition before the last injection. The GICS-PF is a 7-point Likert scale from –3 (very much worse function) to +3 (very much improved function) for assessing functional change due to treatment of PF spasticity. Investigators were trained to perform and rate the AS evaluation. Based on their clinical experience-based review and judgment, investigators rated changes only of PF function compared with the condition before the last injection. Patients receiving bilateral treatment of pes equinus were assessed with the GICS-PF separately on each side. The same investigator was required to make the baseline and week 4 assessments of the first injection cycle of both coprimary endpoints and every attempt was made to have the same investigator perform the AS at all other study visits.

Secondary endpoints included AS scores for the knee flexors and thigh adductor muscles and the investigator’s Global Impression of Change Scales (GICS) to assess global changes after treatment. For the latter, overall LL spasticity change was assessed on a 7-point Likert scale from –3 (very much worse) to
+3 (very much improved) and rated by the investigator, the child/adolescent (if applicable), and the parent/caregiver to gain different perspectives. The GICS has proven to be a useful assessment tool for spasticity [33].

Change in a patient’s gross motor function over time was assessed using the Gross Motor Function Measure (GMFM)-66 (scores range from 0 to 100, with higher scores reflecting better function). The GMFM-66, a shorter version of the GMFM-88, is a standardized observational instrument designed and validated to measure change in gross motor function over time in people with CP [34–36]. It has been shown to be a reliable and valid tool in children with different disabilities and is useful to assess children aged < 3 years [37].

2.5. Safety

Safety endpoints assessed throughout the study included the occurrence of treatment-emergent adverse events (TEAEs), TEAEs of special interest (TEAE-SIs) potentially indicating distant toxin spread, and treatment-emergent serious adverse events (TESAEs). BoNT antibody testing was conducted in patients ≥ 21 kg BW at the screening and end-of-study visits. Blood samples were screened for BoNT antibodies using the fluorescence immunoassay for detecting antibodies (FIA-AB), and positive samples were validated using the mouse ex vivo hemidiaphragm assay (HDA) to test for neutralising antibodies (NABs).

2.6. Statistical analysis

Efficacy data were analyzed descriptively in the full analysis set (a subset of patients in the safety evaluation set [SES] who had at least a baseline AS-PF score or the investigator’s GICS-PF at week 4 of injection cycle 1 available). Safety variables were analyzed descriptively in the SES, which included all patients who received at least one study treatment.

Comparison of dose groups was the primary statistical analysis. The primary and coprimary efficacy endpoints, the change from study baseline in AS-PF and investigator’s GICS-PF for the primary side at week 4, were analyzed using a mixed-model for repeated measures (MMRM; two-sided, significance level $\alpha = 0.05$) or an analysis of covariance, respectively. Least squares mean values were compared using a two-step hierarchical test procedure to detect differences between incobotulinumtoxinA high and low dose in step 1, and between mid and low dose in step 2. If one of two hierarchical tests did not yield a statistically significant result, the consecutive test(s) were still performed but were considered descriptive only.

Secondary and other analyses included the change from baseline in AS-PF and investigator’s GICS-PF for the primary side at week 4 of injection cycle 2 and at additional times during injection cycle 1 and injection cycle 2 and changes from baseline in secondary endpoints at various times during both injection cycles. Statistical analyses were performed using the statistical analysis system (SAS®) software package, version 9.2 or higher (SAS Institute, Cary, NC, USA).

Safety data were analyzed descriptively.

2.7. Sample size calculations and study power

It was estimated that the sample size of $N = 300$ participants would provide 94% power (product of the single power calculations for both the primary and the coprimary efficacy variables for the high- vs. low-dose treatment comparison) to identify a statistically significant difference between the high- and low-dose groups. A power of 83.3% was provided for the treatment comparison of the mid versus the low dose in both primary efficacy variables. Using a randomization ratio of 2:1:1, at least 150 subjects were planned to be randomized to the high-dose treatment group and at least 75 subjects each were planned to be randomized to the mid- and low-dose treatment groups.

2.8. Data availability statement

Key elements of the study protocol, design, and statistical analysis plan were deposited in the U.S. National Library of Medicine database (www.clinicaltrials.gov, NCT01893411) and EU Clinical Trials Register (https://eudract.ema.europa.eu/, 2012-005054-30). All relevant information is contained within this manuscript and the supplementary material.

3. Results

A total of 338 patients were screened, of whom 311 were randomized and treated with incobotulinumtoxinA. Of these, 78 patients were treated with low-dose incobotulinumtoxinA (4 U/kg, maximum 100 U), 77 were treated with mid-dose incobotulinumtoxinA (12 U/kg, maximum 300 U), and 156 were treated with high-dose incobotulinumtoxinA (16 U/kg, maximum 400 U). A total of 278 (89.4%) patients completed both injection
cycles. The discontinuation rate was low and similarly distributed across all incobotulinumtoxinA dose groups (Fig. 2).

Demographics were similar across dose groups (Table 1). Patients were generally young (mean age 6.6 years). Three-quarters of patients had low- to mid-level ambulatory gross motor impairment (GMFCS - Expanded and Revised [GMFCS-E&R] Level I–III) and one-quarter was more severely impaired and non-ambulant (GMFCS-E&R Level IV–V). Most patients presented with bilateral LL spasticity (78.8%), with 11.9% presenting with unilateral right-sided symptoms and 9.3% with unilateral left-sided symptoms. Almost 70% of participants had at least one concomitant disease. The most frequently reported were epilepsy (42.8%), strabismus (18.6%), intellectual disability (16.1%), hypokinesia (14.8%), and foot deformity (10.9%). More than one-half (61.4%) of patients received a BoNT treatment before study enrolment, with a mean (standard deviation [SD]) of 3 (2.9) pre-treatments.

During the study, the majority of patients were treated for bilateral pes equinus (72.7% injection cycle 1, 73.5% injection cycle 2). Of those receiving unilateral treatment, more patients were treated for unilateral pes equinus and flexed knee in injection cycle 1 and injection cycle 2 (19.3% and 18.5%, respectively) than unilateral pes equinus and adducted thigh (8.0% in both
Table 1
Patient demographics and baseline characteristics, SES/FAS

| Characteristic                      | Low dose 4 U/kg, maximum 100 U | Mid dose 12 U/kg, maximum 300 U | High dose 16 U/kg, maximum 400 U | Total |
|------------------------------------|---------------------------------|---------------------------------|---------------------------------|-------|
|                                    | n = 78                          | n = 77                          | n = 156                         | n = 311 |
| Sex, n (%)                         |                                 |                                 |                                 |       |
| Male                               | 42 (53.8)                       | 44 (57.1)                       | 83 (53.2)                       | 169 (54.3) |
| Female                             | 36 (46.2)                       | 33 (42.9)                       | 73 (46.8)                       | 142 (45.7) |
| Age, years; mean (SD)              | 7.1 (4.6)                       | 6.6 (3.8)                       | 6.4 (3.9)                       | 6.6 (4.1) |
| Weight, kg; mean (SD)              | 24.6 (16.0)                     | 22.7 (11.9)                     | 22.3 (11.8)                     | 22.9 (13.0) |
| GMFCS-E&R*, n (%)                  |                                 |                                 |                                 |       |
| Level I                            | 14 (17.9)                       | 17 (22.1)                       | 34 (21.8)                       | 65 (20.9) |
| Level II                           | 24 (30.8)                       | 25 (32.5)                       | 50 (32.1)                       | 99 (31.8) |
| Level III                          | 19 (24.4)                       | 16 (20.8)                       | 33 (21.2)                       | 68 (21.9) |
| Level IV                           | 17 (21.8)                       | 12 (15.6)                       | 24 (15.4)                       | 53 (17.0) |
| Level V                            | 4 (5.1)                         | 7 (9.1)                         | 15 (9.6)                        | 26 (8.4) |
| AFFECTED BODY SIDE, n (%)          |                                 |                                 |                                 |       |
| Unilateral right                   | 11 (14.1)                       | 11 (14.3)                       | 15 (9.6)                        | 37 (11.9) |
| Unilateral left                    | 9 (11.5)                        | 5 (6.5)                         | 15 (9.6)                        | 29 (9.3) |
| Bilateral                          | 58 (74.4)                       | 61 (79.2)                       | 126 (80.8)                      | 245 (78.8) |
| Baseline AS-PF score               |                                 |                                 |                                 |       |
| Mean (SD)                          | 2.7 (0.6)                       | 2.7 (0.5)                       | 2.8 (0.5)                       | 2.7 (0.5) |
| Median (interquartile range)       | 3.0 (2.0, 3.0)                  | 3.0 (2.0, 3.0)                  | 3.0 (2.0, 3.0)                  | 3.0 (2.0, 3.0) |
| BoNT pretreatment, n (%)           |                                 |                                 |                                 |       |
| Yes                                | 40 (51.3)                       | 54 (70.1)                       | 97 (62.2)                       | 191 (61.4) |
| No                                 | 38 (48.7)                       | 23 (29.9)                       | 59 (37.8)                       | 120 (38.6) |
| Concomitant diseases, n (%)        |                                 |                                 |                                 |       |
| Patients with at least one         |                                 |                                 |                                 |       |
| Most common*                       |                                 |                                 |                                 |       |
| Intellectual disability            | 19 (24.4)                       | 9 (11.7)                        | 18 (11.5)                       | 46 (14.8) |
| Epilepsy                           | 13 (16.7)                       | 14 (18.2)                       | 23 (14.7)                       | 50 (16.1) |
| Strabismus                         | 10 (12.8)                       | 13 (16.9)                       | 35 (22.4)                       | 58 (18.6) |
| Foot deformity                     | 8 (10.3)                        | 10 (13.0)                       | 16 (10.3)                       | 34 (10.9) |

*GMFCS-E&R level: I, walks without limitations; II, walks with limitations; III, walks using a handheld mobility device; IV, self-mobility with limitations, may use powered mobile; V, transported in a manual wheelchair. \(^{b}\)n = 306 and 310 observations for patients treated in left and right PFs, respectively. \(^{c}\)'Most common' defined as >10% in any group. \(^{d}\)The SES and the FAS include the same patients and are thus interchangeable. AS-PF = Ashworth Scale of the Plantar Flexors; FAS = full analysis set; GMFCS-E&R = Gross Motor Function Classification System – Expanded and Revised; SD = standard deviation; SES = safety evaluation set.

Injections). IncobotulinumtoxinA doses administered overall and by pattern and muscle group during each injection cycle are summarized in the Supplementary Table.

The median time to reinjection or discontinuation was 14.3 (range 4–37) weeks overall in injection cycle 1 and 14.6 (range 11–38) weeks overall in injection cycle 2. The median (range) time to reinjection was similar in all three dose groups (incobotulinumtoxinA 4, 8, and 16 U/kg [maximum 100, 300, and 400 U]) in injection cycle 1, being 14.1 (4–37), 14.4 (8–36), and 14.3 (8–37) weeks, respectively, with a mean (SD) time to reinjection of 15.7 (5.9), 15.9 (5.7), and 15.3 (4.6) weeks, respectively. Injection intervals were 12 to <16 weeks in most patients (59.0%, 49.4%, and 66.0%, respectively). Injection intervals of ≥16 weeks were noted in 30.8%, 33.8%, and 26.3%, respectively, with intervals of 16 to <20 weeks observed in 19.2%, 19.5%, and 14.1% and of 20 to <30 weeks in 6.4%, 9.1%, and 9.6%, respectively. In total, 12 patients (3.9%) had injection intervals of ≥30 weeks (5.1%, 5.2%, and 2.6%, respectively, per dose group). Compliance, as measured by the total volume of incobotulinumtoxinA injected, was near 100% for both injection cycles.

3.1. Efficacy

3.1.1. Primary and coprimary endpoint results

Patients in all three incobotulinumtoxinA dose groups (total body doses of 4, 8, and 16 U/kg [maximum 100, 300, and 400 U]) experienced significant improvements in AS-PF scores on the primary body side chosen for treatment 4 weeks post-injection in injection cycle 1 versus the study baseline (p < 0.0001 for all, MMRM) (Fig. 3). In patients who received incobotulinumtoxinA 2, 4, and 8 U/kg (maximum 50,
Fig. 3. The effect of incobotulinumtoxinA on mean change from baseline at week 4 on the AS-PF on the primary body side, FAS, OC. AS score: 5-point scale from 0 (no increase in muscle tone) to 4 (limb rigid in flexion or extension). The change in the AS-PF from baseline to week 4 was the primary efficacy variable. ***p < 0.0001 versus study baseline. AS = Ashworth Scale; AS-PF = Ashworth Scale of the plantar flexors; BW = body weight; FAS = full analysis set; IC = injection cycle; kg = kilogram; OC = observed cases; SE = standard error; U = unit.

150, and 200 U) injected into the primary pes equinus site, AS-PF score improvements of 1–3 points from the study baseline to 4 weeks post-injection were generally observed in more than one-half of patients in injection cycle 1: 55.1%, 55.8%, and 59.6%, respectively. No worsening of AS-PF scores was observed. Likewise, investigator’s GICS-PF scores indicated a clinically meaningful response to incobotulinumtoxinA treatment in all dose groups 4 weeks post-injection in injection cycle 1 (Fig. 4). Spasticity improvement (responses of “minimally improved,” “much improved,” and “very much improved”) due to treatment was determined in more than 93% of patients in all dose groups. As similar improvements in spasticity were observed in all dose groups, no statistically significant differences between dose groups were demonstrated in the primary or coprimary efficacy analyses.

Greater improvement was associated with higher baseline AS scores. Of note, treatments produced comparable improvements in AS-PF scores across all GMFCS-E&R levels.

3.1.2. Further results

3.1.2.1. AS

Consistent improvements for all patterns with each injection cycle were seen at weeks 4 and 8. AS-PF scores of the primary body side further improved from baseline to week 4 of the second injection cycle in all three dose groups (Fig. 3). The treatment effects were greater 4 and 8 weeks following each injection, after which changes decreased by week 12. Following unilateral LL treatment, all doses of incobotulinumtoxinA resulted in improvements from baseline in AS scores of the knee flexors (Fig. 5A) and the thigh adductors (Fig. 5B) 4 weeks following each injection. Wherever estimable, all doses of incobotulinumtoxinA produced comparable improvements.

3.1.2.2. Investigator’s, child/adolescent’s, and parent/caregiver’s GICS scores

Whether from the perspective of the investigator, the parent/caregiver, or the patient, GICS scores confirmed a consistent and global improvement in LL spasticity at 4 weeks post-injection for all three incobotulinumtoxinA dose groups and across both injection cycles (Fig. 6A–C). Across both injection cycles, investigators indicated improvement rates of > 90% when taking the sum of “minimally improved,” “much improved,” and “very much improved” frequencies. Parents/caregivers reported improvements in more than 80% of patients at these times. GICS scores were available from 150 of 311 and 135 of 287 children/adolescents (48% and
47.0%) at injection cycle 1 and injection cycle 2, respectively; of those who responded, > 80% reported improvement after each injection cycle.

3.1.2.3. Motor functioning

GMFM-66 scores indicated that motor function improved in all three treatment groups from the end of injection cycle 1 to the end-of-study visit 8 weeks later. Mean (SD) GMFM-66 scores increased from baseline by 1.8 (2.8), 1.2 (3.5), and 1.4 (3.1) at the end of injection cycle 1 and by 3.1 (3.4), 3.3 (4.5), and 2.8 (4.1) at the end-of-study visit in patients who received incobotulinumtoxinA 4, 8, and 16 U/kg (maximum 100, 300, and 400 U), respectively.

3.2. Safety

IncobotulinumtoxinA treatment was generally well-tolerated over both injection cycles, with TEAEs reported in 42.8% of patients overall (Table 2). The incidence of TEAEs was slightly higher in patients who received the highest dose of incobotulinumtoxinA, followed by the lowest dose, and then the mid dose. Most TEAEs were mild or moderate in intensity. TEAEs assessed by investigators as treatment-related also occurred at a low frequency (4.8% of patients overall). These events were localized muscular weakness (n = 5 patients), injection-site pain (n = 4 patients), injection-site erythema, pain in an extremity, pyrexia (n = 2 patients each), fall, hematoma, influenza-like illness, injection-site inflammation, injection-site warmth, and rash (n = 1 patient each).

TEAESIs were reported in 2.3% of patients overall. The most frequent TEAESI, localized muscular weakness, was reported in five patients and, in all cases, affected the treated LL and was considered by the investigator to be related to treatment. Given the low incidence of TEAESIs, no meaningful conclusions could be drawn concerning any relationship between CP severity and TEAESIs.

TEAESIs occurred at low frequencies in all dose groups for each injection cycle (high dose: 2.6% and 2.1%; mid dose: 0.0% and 1.4%; low dose: 3.8% and 4.1% in the first and second cycles, respectively). No treatment-related TESAE or fatal TEAE was reported. One patient in the high-dose group discontinued the study because of a non-treatment-related nonserious TEAE (asthma).

At the screening visit, 19 of 127 (15.0%) subjects eligible for testing were tested positive on the FIA-AB. Three patients subsequently tested positive for NABs according to HDA test results; all three patients had been pretreated with other BoNT-As. Of these, one had a negative HDA test at the end-of-study visit, and the other two patients did not undergo further HDA testing. Based on a protocol deviation, ten patients at screening and two patients at the end-of-study visit were FIA-AB positive but were not HDA tested (HDA missing). All patients with positive HDA or HDA missing based on protocol deviation responded to treatment, based on the investigator’s GICS-PF assessment.

4. Discussion

The TIM study is one of the most extensive BoNT-A studies of its kind investigating incobotulinumtoxinA
### Table 2
Summary of TEAEs by treatment over two injection cycles, SES

| Adverse events                  | Low dose 4 U/kg, maximum 100 U | Mid dose 12 U/kg, maximum 300 U | High dose 16 U/kg, maximum 400 U | Total* |
|--------------------------------|-------------------------------|-------------------------------|---------------------------------|--------|
|                                 | n = 78                        | n = 77                        | n = 156                         | n = 311|
| Any TEAE, n (%)                | 30 (38.5)                     | 26 (33.8)                     | 77 (49.4)                       | 133 (42.8) |
| Mild                            | 19 (24.4)                     | 14 (18.2)                     | 41 (26.3)                       | 74 (23.8) |
| Moderate                        | 10 (12.8)                     | 11 (14.3)                     | 33 (21.2)                       | 54 (17.4) |
| Severe                          | 1 (1.3)                       | 1 (1.3)                       | 3 (1.9)                         | 5 (1.6) |
| Treatment-related               | 2 (2.6)                       | 2 (2.6)                       | 11 (7.1)                        | 15 (4.8) |
| Any TEAE, n (%)                | 1 (1.3)                       | 1 (1.3)                       | 5 (3.2)                         | 7 (2.3) |
| Treatment-related               | 0 (0.0)                       | 0 (0.0)                       | 4 (2.6)                         | 5 (1.6) |
| Any TESAE, n (%)               | 6 (7.7)                       | 1 (1.3)                       | 7 (4.5)                         | 14 (4.5) |
| Treatment-related               | 0 (0.0)                       | 0 (0.0)                       | 0 (0.0)                         | 0 (0.0) |
| Any TEAE leading to discontinuation*, n (%) | 0 (0.0) | 0 (0.0) | 1 (0.6) | 1 (0.3) |
| Treatment-related               | 0 (0.0)                       | 0 (0.0)                       | 0 (0.0)                         | 0 (0.0) |
| Any fatal TEAE, n (%)          | 0 (0.0)                       | 0 (0.0)                       | 0 (0.0)                         | 0 (0.0) |

*Total relates to number of patients with TEAEs in any injection cycle. **TEAEs leading to discontinuation in injection cycle 1, moderate asthma (unrelated to treatment). SES = safety evaluation set; TEAE = treatment-emergent adverse event; TESAE = TEAE of special interest (potentially indicating distant toxin spread); TESAE = treatment-emergent serious adverse event.

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**Fig. 6.** The effect of incobotulinumtoxinA on (A) investigator, (B) parent/caregiver, and (C) child/adolescent GICS scores† at week 4, FAS, OC. †GICS scores were available from 150 of 311 and 135 of 287 children/adolescents (48% and 47.0%) at IC1 and IC2. The proportion of children/adolescents responding was attributed to the respondents’ young age or their cognitive abilities. ***P < 0.0001. BW = body weight; FAS = full analysis set; GICS = Global Impression of Change Scale; IC = injection cycle; kg = kilogram; OC = observed cases; SE = standard error; U = unit.
in pediatric and adolescent patients with CP-related LL spasticity across all GMFCS levels (ambulatory and non-ambulatory), reflecting real-world CP distribution. Consequently, this study fills an important gap in the literature and existing clinical evidence. IncobotulinumtoxinA treatment with doses of 4, 12, and 16 U/kg (maximum 100, 300, and 400 U, respectively) per patient divided over two clinical patterns showed consistent improvement in all LL clinical patterns treated, as measured by the AS. These results were supported by better global and functional outcomes from the perspective of patients, parents/caregivers, and physicians, all of whom reported notable improvements in GICS scores in 80% or more of patients. Furthermore, sustained and cumulative improvement in spasticity and functional outcomes was observed with repeated incobotulinumtoxinA treatment to the same clinical patterns.

No significant differences between doses were observed for either the primary or the secondary outcomes, and so the primary efficacy analyses of the study were not met. In planning the TIM study, it was assumed that the higher incobotulinumtoxinA doses could be differentiated from the lower, but ultimately the results demonstrated a measurable response for patients receiving all doses tested. For example, in our study, the AS-PF LS-mean change was -0.70 for the high-dose group and -0.66 for the low-dose incobotulinumtoxinA group \( p = 0.650 \). Other BoNT-A LL studies that compared active treatment and placebo using the modified AS (MAS) also reported no significant differences between lower and higher doses (abobotulinumtoxinA 10 vs. 15 U/kg/leg [38]; onabotulinumtoxinA 4 vs. 8 U/kg [39]).

The original AS and the MAS [40] are routinely used to assess the treatment effects of BoNT on spasticity in adults [41–43] and children with CP [25,44]. The MAS, a 6-point scale, would be expected to yield a greater effect size than the original 5-point AS scale. However, the changes in muscle tone from baseline in our study using the original AS were comparable to the LL spasticity effect sizes in studies that used the MAS and included placebo controls [38,39].

As already reported, incobotulinumtoxinA (total doses of 2–16 U) was also equally efficacious at producing partial paralysis of the extensor digitorum brevis muscle of the foot in a double-blind study in healthy volunteers [45]. Furthermore, demonstration of significant differences between dose groups can be more difficult in LL spasticity than in UL spasticity. Indeed, the recently published IncobotulinumtoxinA in aRM treatment in cerebral pAlsy (XARA) trial (NCT02002884) demonstrated that incobotulinumtoxinA 8 U/kg BW produced AS scores that were significantly superior to those of 2 U/kg BW for UL CP-related spasticity [46].

The results also indicate that incobotulinumtoxinA had a favorable safety and tolerability profile at doses of up to 16 U/kg (maximum 400 U) for patients in all GMFCS severity groups. No new or unexpected safety concerns were identified over two injection cycles (up to 72 weeks).

Overall, the incobotulinumtoxinA treatments were well-tolerated, as evidenced by the high patient retention rate (89.4% completed the study). Our results also align with a previous retrospective report that found incobotulinumtoxinA to be safe and effective for CP-related spasticity in children [47].

Despite using an incobotulinumtoxinA dose of up to 400 U and repeated treatments, no secondary nonresponse to treatment due to NABs occurred. This is of importance for the anticipated long-term treatment of spastic CP as a chronic condition occurring in young patients [18]. In a study of BoNT-A use in patients with a variety of neurological impairments, 13.9% of 596 patients developed measurable NABs [48]. The rate of NABs varied between BoNT-A formulations: 6% for abobotulinumtoxinA and 7% for onabotulinumtoxinA compared with 0% for incobotulinumtoxinA. It should be noted that not all patients who did not respond to BoNT-A therapy had high NAB levels, and some patients with high NAB levels still responded to BoNT-As [48]. The current results add to the growing body of evidence that incobotulinumtoxinA may carry less risk of inducing an immunogenic response relative to other BoNT-As – and hence less risk of nonresponse to therapy – because it is a highly purified formulation of BoNT-A [48] that does not contain BoNT-A accessory proteins [18]. For this reason, it may be the preferred choice for a chronic condition in a pediatric population [50].

Although the majority of participants in TIM required bilateral injections for pes equinus, almost 20% were treated for unilateral pes equinus and ipsilateral flexed knee and 8% for unilateral pes equinus and ipsilateral adducted thigh. IncobotulinumtoxinA therapy demonstrated efficacy in all these patterns, adding to evidence supporting the use of incobotulinumtoxinA in multipattern spasticity.

The TIM study is part of a large, international phase 3 pediatric study program investigating the efficacy and safety of incobotulinumtoxinA for the treatment of CP-related spasticity. All of these studies, including the TIM study, were designed to incorporate important as-
pects of individualized treatment within the perspective of a class I phase 3 study based on distribution of spasticity, i.e., multiple affected clinical patterns. The physician was able to tailor treatment using standardized dose ranges for the complete set of muscles of a pattern, defining injection sites based on anatomy, and adjust injection intervals. Overall, the reports from this and two other trials (TIMO: NCT01905683; and XARA: NCT02002884) confirm that incobotulinumtoxinA is safe and effective for multipattern spasticity in children and adolescents with CP [46,51,52], including those with severe symptoms [53].

The mean time to reinjection was 15.6 (5.2) weeks, although 29.3% of patients did not need a second dose until 16 weeks or later. Currently, minimal intervals of 12–16 weeks are recommended for the treatment of spasticity in children [29–31], although other reports have suggested longer injection intervals may be appropriate at times [54,55]. These results from the TIM study suggest that some flexibility in time to reinjection based on a patient’s clinical needs may be warranted in clinical practice.

The changes in motor function, as assessed by GMFM-66 scores over time, were consistent with those of previous studies of BoNT-A for LL spasticity [56–58]. Although GMFM-66 improvements should be considered in the context of the patient’s global rehabilitation plan and cannot be solely attributed to incobotulinumtoxinA treatment, improvements after a single injection in the current study were within changes being reported as minimum clinically important differences by Oeffinger et al. [59], who defined medium and large effects as changes of 0.7–1.7 and 1.2–2.7, respectively, in ambulatory children with CP.

Spasticity is just one factor among many others that could interfere with gross motor development [60], even in children with severe forms of spastic CP. For instance, strength (impaired by the underlying paresis) [61] and selective motor control [62] have also been found to be contributory. A meaningful improvement, as shown here, is thus a promising result, especially when considering that the GMFM was assessed only at the end-of-cycle visits (when the effect was waning) and not at week 4 visits (the time point of expected maximum effect).

4.1. Strengths and limitations

The strengths of the TIM study include its patient-centric approach to treatment. Individualized treatment options within standardized, multipattern treatment protocols for LL spasticity were available to address the real-world needs of a large, heterogeneous population of children and adolescents with CP. Participants represented a broad age range (2–17 years) and all levels of disease severity (GMFCS-E&R levels I–V). The efficacy and safety of incobotulinumtoxinA were assessed across a wide dose range. The study design allowed for the treatment of several muscle groups at once, giving physicians the freedom to adjust dose, injection site, number of injections per site, and time to reinjection as required, based on individual patient need. Additional strengths were the generation of a comprehensive dataset, which included efficacy and safety data for other LL muscle groups in addition to the PFs and the uni/bilateral combination treatment. The outcomes provided data on the effect of incobotulinumtoxinA on muscle resistance, global improvement, and gross motor function, from the point of view of the patient, parent/caregiver, and physician.

The lack of a placebo control in the TIM study may be viewed as a limitation, but use of placebo in this vulnerable young patient population raises concerns in terms of ethical aspects and feasibility. Furthermore, the effect sizes, as discussed earlier, were within established ranges, and the current patient-centric approach allowed for a thorough investigation of long-term efficacy and safety across a broad dose range of incobotulinumtoxinA in children and adolescents with spastic CP.

5. Conclusion

This large phase 3 study demonstrated that incobotulinumtoxinA used according to an individualized treatment plan within standardized guidelines is beneficial in improving muscle tone and motor function for children with spasticity-related CP. IncobotulinumtoxinA total doses up to 16 U/kg BW (maximum 400 U) were effective and well-tolerated for the multipattern treatment of LL spasticity due to CP for ambulant and non-ambulant children and adolescents presenting with all levels of disease severity. IncobotulinumtoxinA had a good safety and tolerability profile and did not induce NAB formation.

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**Ethical considerations**

The study protocol, informed consent forms and other appropriate study-related documents were reviewed and approved by the local independent ethics committees and institutional review boards. Parents/guardians of all patients provided written informed consent, and patients provided assent if applicable.

**Conflict of interest**

Florian Heinen has received speaker’s honoraria from Allergan plc. Desitin, Ipsen Biopharmaceuticals, Merz Pharmaceuticals, and Novartis and unrestricted educational grants from Allergan and Merz Pharmaceuticals. Petr Kaňovský has received speaker’s honoraria from Desitin, Ipsen Biopharmaceuticals, Merz Pharmaceuticals, and Medtronic. A. Sebastian Schroeder has received speaker’s honoraria from and participated in advisory boards for Allergan plc, Ipsen Biopharmaceuticals, and Merz Pharmaceuticals. Henry G. Chambers serves as a consultant for Orthopediatrics Corp and Allogeneic. Thorin L. Geister is an employee of Merz Pharmaceuticals GmbH. Marta Banach has served as a consultant for Teva and Kashiva.

**Supplementary data**

The supplementary files are available to download from http://dx.doi.org/10.3233/PRM-210040.

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