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Published in:
BMJ Open

DOI:
10.1136/bmjopen-2018-024340

Publication date:
2019

Document version
Final published version

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Citation for published version (APA):

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Botulinum toxin treatment of spasticity targeted to muscle endplates: an international, randomised, evaluator-blinded study comparing two different botulinum toxin injection strategies for the treatment of upper limb spasticity

Tiina Rekand,1 Bo Biering-Sørensen,2 Jun He,3 Ole Jakob Vilholm,4 Peter Brogger Christensen,5 Trandur Ulfarsson,6 Roger Belusa,7 Torbjörn Ström,7 Peter Myrenfors,7 Pascal Maisonneve,7,8 Torben Dalager9

ABSTRACT

Objectives The therapeutic effects of botulinum neurotoxin (BoNT) are well documented in upper limb spasticity. However, several factors may influence treatment efficacy, including targeting of neuromuscular junctions (NMJs). We examined whether NMJ-targeted BoNT injections were non-inferior, in terms of efficacy, to current injection practices.

Design Open-label prospective evaluator-blinded study.

Setting Conducted across 20 medical centres in Denmark, Finland, Norway and Sweden (24 September 2012 to 11 March 2015).

Participants Aged >18 years with upper limb spasticity (Modified Ashworth Scale [MAS] score of 2 or 3) following stroke or traumatic brain injury, had received ≥2 consecutive BoNT-A treatment cycles (the latest of which was abobotulinumtoxinA [aboBoNT-A]) and needed BoNT-A retreatment (same modality as previous cycle). Patients requiring aboBoNT-A doses >800 units were excluded. In total, 88 patients were randomised (intention-to-treat [ITT] population), most were male (n=58/88, 65.9%) and 54/88 (61.4%) completed the study (per-protocol [PP] population).

Interventions Randomisation (1:1) to receive a single dose of aboBoNT-A (≤800 U) according to either current clinical practice (300 U/mL) or as an NMJ-targeted injection (100 U/mL).

Primary outcome measure Proportion of patients with a ≥1 level reduction from baseline in MAS score at week 4 post-injection (responders).

Results In the ITT population, the proportion of responders at elbow flexors was 72.7% in the current practice group and 56.8% in the NMJ-targeted group (adjusted difference −0.1673 [95% CI: −0.3630 to 0.0284]; p=0.0886). Similar results were observed in the PP population (69.0% vs 68.0%, respectively, adjusted difference 0.0707 [−0.1948 to 0.3362]; p=0.6052).

Conclusions Owing to the limited number of participants, non-inferiority of NMJ-targeted injections could not be determined. However, there was no statistical difference between groups. Larger studies are needed confirm whether the two techniques offer comparable efficacy.

Trial registration number NCT01682148.

INTRODUCTION

Spasticity is a sensorimotor control disorder, resulting from an upper motor neuron lesion, presenting as intermittent or sustained involuntary activation of muscles.1 Patients with spasticity may suffer from decreased active movement, increased disability and impaired function.2 Intramuscular injections with botulinum neurotoxin (BoNT) is a documented efficacious and well-tolerated treatment
option for these patients, which has an established position in clinical practice. Studies with abobotulinumtoxinA (aboBoNT-A, Dysport) have demonstrated a beneficial effect on spasticity in terms of decreased resistance to passive movement, improvements in pain and goal attainment as well as in active function. The therapeutic effects of BoNT have been extensively studied in upper limb spasticity, which has been summarised in a number of recent reviews. Several factors can influence treatment efficacy, such as injection technique, targeting of neuromuscular junctions (NMJs), concentration of BoNT and volume of injected solution, as well as stimulation of BoNT uptake in target cells. Mapping of NMJ zones in the upper limb flexor muscles has enabled targeting of these zones to potentially increase the efficacy of BoNT injection in spasticity treatment. Indeed, results from studies in both animals and humans indicate that targeting NMJ zones can increase the effectiveness of BoNT injections. Gracies et al suggested that high volume or endplate targeted BoNT injections result in superior neuromuscular blockade, contraction and spasticity reduction than low volume, non-targeted injections. This study explores the hypothesis that the effect of a single high volume aboBoNT-A injection of low concentration per muscle, if targeted to the area of the NMJ zones, is equal to the effect of several injections using standard injection practices. An international randomised, multicenter, evaluator-blinded study was thus performed to evaluate non-inferiority between the two different injection strategies.

MATERIALS AND METHODS

Study design

The NMJ study was an international, multicenter, prospective, randomised, evaluator-blinded, parallel-group study, comparing two different injection techniques. Patients were injected with botulinum toxin (aboBoNT-A, Dysport) for the treatment of upper limb spasticity. Patients were randomised to 1 of 2 groups: (1) the current practice group receiving injections according to current clinical practice (without regard to NMJ) and (2) the NMJ-targeted group receiving injections according to a predefined NMJ targeting technique with lower concentration of the toxin (figure 1).

Patient and public involvement

In this study, the patient-centric research question was developed with the aim of limiting the injection points and thus limiting burden of injections. The patient’s experience of the different injection techniques was recorded by measuring injection pain using a Visual Analogue Scale (VAS). Evaluation of outcome measures was via the patient-centric Goal Attainment Scaling (GAS) score where the patient and treating physician agree on a goal that is meaningful for the patient, and evaluate the treatment by goal attainment.

Patients were asked whether they would like to participate in the study and decided if they wanted to participate or not after provision of informed consent. When completing the informed consent form, patients could indicate whether they wished to be informed of the study results or not. Patients wishing to know the results were informed by their investigator.

Patient recruitment

Medical centres in Finland, Denmark, Sweden and Norway (n=59) were included to recruit patients for participation in the study. Approval from the local ethical review board was granted in each separate country. The study conformed to the principles of the Helsinki Declaration and all patients signed an informed consent form ahead of participation.

Inclusion criteria included age >18 years, provision of written informed consent and suffering from upper limb spasticity position pattern type 1, 3 or 4 following stroke or traumatic brain injury. The position patterns were based on characteristic arm positions and postures recognised in upper limb spasticity. Spasticity was evaluated according to the Modified Ashworth Scale (MAS),

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Figure 1 Study design. *If no routine BoNT-A treatment was given at visit 3, max 24 weeks poststudy injection. NMJ, neuromuscular junction.
with a baseline score of 2 or 3 points at inclusion. Patients had received at least two consecutive previous treatment cycles with BoNT-A, of which at least the latest had to have been with aboBoNT-A with observed good efficacy according to investigator judgement. Furthermore, the investigator should have judged a need for a repeat of the same treatment modality as the previous cycle and the last BoNT-A treatment should have been dated >12 weeks before study entry. Subjects were randomised (1:1) to 1 of 2 treatment groups. The patients were stratified according to three different arm spasticity position patterns (1, 3 and 4) and country of residence, aiming at a balanced ratio regarding these parameters.

Exclusion criteria included a need for doses of >800 U aboBoNT-A in the upper limb, a likelihood of changing other spasticity treatment (such as physiotherapy) during the study period, other neurological disorders or use of agents known to interfere with neuromuscular transmission. For full inclusion/exclusion criteria, see ClinicalTrials.gov).

Recruitment dates were first subject in 24 September 2012, last subject in 16 December 2014 and last subject in 11 March 2015.

**Randomisation procedure**

A randomisation list was prepared by a statistician independent from the study, and was provided to the electronic version of the Case Report Form (eCRF) provider who incorporated it into the eCRF. The list was stratified for three arm spasticity position patterns (1, 3 and 4) and for participating countries, and generated with a balanced 1:1 treatment ratio. Once eligibility was confirmed, the investigator entered eligibility data into the eCRF. If the patient was eligible, based on the predefined randomisation list, the eCRF system would automatically assign the first not-used randomisation number corresponding to the patient’s spasticity pattern and country, to get the applicable treatment administration group allocation. The study drug was provided in 3 mL vials containing 500 U of aboBoNT-A and diluted by the investigator to the assigned concentration, 100 or 300 U/mL according to the assigned treatment group.

**Blinding**

This was an open study with respect to study drug. However, the treatment group assigned to a randomisation number was to be blinded prior to allocation. To optimise protocol blindness at each site, all study injections were to be performed by one of the investigators (unblinded) and all efficacy assessments were to be performed by another qualified study personnel, blinded to the injection technique. The eCRF provided two user profiles, one which was able to see the randomisation information (treatment arm) and another one for the evaluators, for which the treatment arm was hidden.

**Treatments**

Optimal dose of aboBoNT-A (Dysport) for each subject was determined by the investigator at the first visit to optimise treatment efficacy prior to inclusion (see the Patient recruitment section). Maximum dose injected in the spastic arm was 800 U. Patients in the current practice group received injections according to current clinical practice (toxin concentration of 300 U/mL). The NMJ-targeted group were injected using an NMJ targeted technique (low toxin concentration of 100 U/mL). All injections were targeted to the same muscles of the arm and performed with support of electromyography (EMG) guiding and/or ultrasound (US). The study treatment was performed using the same individual dose as determined at the pre-study treatment visit.

The two different injection patterns are outlined in table 1. Each patient visited the clinic on at least three occasions. The first visit was devoted to screening, randomisation and treatment with aboBoNT-A. The second visit was scheduled 4 weeks after the first injection. Lastly, the third visit was a follow-up at 12 weeks after the first injection. Additional treatment with aboBoNT-A was given if necessary.

<table>
<thead>
<tr>
<th>Injection site</th>
<th>Current practice</th>
<th>NMJ-targeted</th>
</tr>
</thead>
<tbody>
<tr>
<td>m. biceps brachii*</td>
<td>2–4</td>
<td>1</td>
</tr>
<tr>
<td>m. brachialis*</td>
<td>2–4</td>
<td>1</td>
</tr>
<tr>
<td>m. brachioradialis*</td>
<td>1–2</td>
<td>1</td>
</tr>
<tr>
<td>m. flexor carpi radialis/m. flexor carpi ulnaris*</td>
<td>2–3</td>
<td>1</td>
</tr>
<tr>
<td>Volume of injection</td>
<td>0.1–0.7 mL per muscle</td>
<td>0.4–2.0 mL per muscle</td>
</tr>
<tr>
<td>Concentration of aboBoNT-A</td>
<td>300 U/mL</td>
<td>100 U/mL</td>
</tr>
</tbody>
</table>

*Number of injection points. aboBoNT-A, abobotulinumtoxinA.

**Assessments and outcome measures**

The aim of the study was to assess non-inferiority of a treatment strategy using fewer injections of low concentration BoNT, as compared with current clinical practice. The primary endpoint was percentage of patients with at least a 1 level decrease in MAS (scored between 0 and 4). Baseline MAS score in the elbow flexors was...
compared with MAS score measured 4 weeks post injection. A ≥1 level decrease in MAS from baseline was considered clinically meaningful. Secondary endpoints included change in MAS score in the elbow flexors from baseline to week 12, intensity of spasticity related pain as measured by a VAS (scored 0–100) at baseline, week 4 and week 12, pain from injection measured by a VAS at first visit, score on the GAS (scored from −2 to 2) at week 4 or week 12, subject global evaluation of treatment effect at the study end and investigator preference of injection technique.

Sample size and statistics

Statistical methods: both the primary endpoint (MAS responders at week 4) and the non-inferiority design were taken into account for the sample size calculation. A two-sided 95% CI for the difference in clinical success rates was calculated using a generalised linear model with treatment, spasticity position patterns, country and baseline MAS score as factors. Assuming a responder rate (π) of 63% in the reference group, a sample size of 122 patients per group would provide 80% power to detect a clinically relevant delta (Δ) of 17% at an alpha (α) level of 2.5% (one-sided). Assuming a dropout rate of 10%, 136 patients per group were required to be randomised (ie, 272 patients in total).

The statistical analyses were performed in accordance with ICH E9 guideline and were based on the pooled data from the individual study sites, unless otherwise stated. All statistical analyses were performed by PCG using SAS V9.4.

RESULTS

Patient population

The study was terminated early owing to slow recruitment, meaning the final population size in the two treatment arms did not reach the preplanned population size of 272 subjects. The main reasons for this were recruitment difficulties and protocol violations. In all, 20 centres included a total of 88 patients who were subsequently randomised in the study (intention-to-treat [ITT] population). However, only 54 of these completed the study without protocol violation (per protocol [PP] population) (online supplementary figure 1). Violation of protocol was recognised for 34 patients. Of these, 23 were not injected as PP and 11 had other violations, for example, 5 patients had missing MAS evaluations. Baseline characteristics for the ITT and PP populations for both aboBoNT-A treatment regimens (current practice vs NMJ-targeted) are shown in table 2.

Primary endpoint

The proportion of MAS responders (defined as ≥1 level decrease in MAS from baseline) in the ITT population for elbow flexors at week 4 post-injection was 72.7% in the group treated as current practice and 56.8% in the group treated with NMJ-targeted injections (figure 2). The adjusted difference in clinical success rates between the two treatment regimens (group treated with NMJ-targeted injections minus group treated as current practice) was −0.1673 (95% CI [−0.3630 to 0.0284], p=0.0986) in favour of the group treated as current practice. Mean (SD) changes in MAS score for elbow flexors at week 4 compared with baseline are presented in figure 3 (ITT population).

Table 2 Baseline characteristics

<table>
<thead>
<tr>
<th>Variable</th>
<th>ITT population</th>
<th>PP population</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Current practice</td>
<td>NMJ-targeted</td>
</tr>
<tr>
<td>Number of subjects</td>
<td>44</td>
<td>44</td>
</tr>
<tr>
<td>Age, years (mean±SD)</td>
<td>60.3±14.4</td>
<td>57.1±11.4</td>
</tr>
<tr>
<td>Male, n (%)</td>
<td>28 (−63.60)</td>
<td>30 (−68.20)</td>
</tr>
<tr>
<td>Spasticity patterns % (I, III, IV)</td>
<td>9.1</td>
<td>11.4</td>
</tr>
<tr>
<td></td>
<td>54.5</td>
<td>59.1</td>
</tr>
<tr>
<td></td>
<td>36.4</td>
<td>29.5</td>
</tr>
<tr>
<td>Years since stroke/injury (median, range)</td>
<td>4 (1–21)</td>
<td>4 (1–24)</td>
</tr>
<tr>
<td>Total aboBoNT-A units (mean±SD)</td>
<td>566.4±169.8</td>
<td>588±200.3</td>
</tr>
<tr>
<td>Injection guidance n (%)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Electric stimulation</td>
<td>10 (22.7%)</td>
<td>8 (19.0%)</td>
</tr>
<tr>
<td>Electromyography</td>
<td>34 (77.3%)</td>
<td>33 78.6%</td>
</tr>
<tr>
<td>Ultrasound</td>
<td>0</td>
<td>1 (2.4%)</td>
</tr>
<tr>
<td>Missing</td>
<td>0</td>
<td>2</td>
</tr>
</tbody>
</table>

aboBoNT-A, abobotulinumtoxinA; ITT, intention-to-treat; NMJ, neuromuscular junction; PP, per protocol.
Proportion of responders (defined as ≥1 level decrease in MAS from baseline) in the PP population for elbow flexors at week 4 was 69.0% in the group treated as current practice and 68.0% in the group treated with NMJ-targeted injections (95% CI of the adjusted difference [−0.1948 to 0.3362], p=0.6052), as outlined in figure 2. There were no significant differences observed between groups. Non-inferiority was not shown in either population. Mean (SD) changes in MAS score for elbow flexors at week 4 compared with baseline are presented in figure 3.

Secondary endpoints
Secondary endpoints evaluated were MAS responders at 12 weeks, improvement in GAS score as well as spasticity pain between baseline and 4 weeks and injection related pain at first visit. All secondary endpoints were measured on the ITT population. No secondary endpoint showed any significant difference between the two groups. Results from the secondary endpoints are summarised in table 3.

Safety
The adverse events reported were consistent with the known safety profile of aboBoNT-A and no new adverse effects were observed. Treatment emergent adverse events (TEAEs) were experienced in 11 subjects in the group treated as current practice (25.0%; 16 events were reported) and 15 in the group treated with NMJ-targeted injections (34.1%; 24 events were reported). For all subjects (n=88), the TEAEs reported by more than one subject were urinary tract infection (reported by four subjects, 4.5%), headache and epilepsy (both reported by three subjects, 3.4%) and pain in extremity, epididymitis and fatigue (all three reported by two subjects, 2.3%). Regarding investigator assessed TEAEs related to treatment, three events were reported in two subjects in the group treated as current practice (4.5%) and one event was reported in the group treated with NMJ-targeted injections (2.3%). In the current practice group, these concerns one patient with injection site hypersensitivity, which was assessed as mild and study treatment related, and one patient with dysphagia and fatigue, both of which were assessed as mild and study treatment related. In the NMJ-targeted group, one patient experienced fatigue, assessed as moderate and related to the study treatment. Overall, the majority of events reported were mild to moderate in intensity. Additionally, in the NMJ-targeted group, three subjects experienced severe TEAEs which were assessed as non-related to aboBoNT-A treatment by the investigator. Two subjects in each group experienced serious adverse events judged by the investigator as non-related to the treatment. There were no reported TEAEs resulting in withdrawal or death.

DISCUSSION
In the current study, both injection techniques using aboBoNT-A resulted in a high proportion of responders, while no statistically significant difference between the injection strategies was observed. However, the number of subjects included in the study was a limiting factor and may explain the failure to demonstrate non-inferiority statistically. Previously, the injection strategy used for BoNT treatment has been identified as important for optimising treatment outcomes. Indeed, NMJ-targeted strategies have the advantage of requiring fewer injection points, potentially minimising pain associated with administration and are expected to be preferable for both patients and physicians. This study contributes to the discussion about the significance of targeted injections.
Open access

Assessment of the ability for passive movements, as measured by MAS, did not show any significant differences between the groups, at 4-week or at 12-week follow-up. Generally, injection techniques based on a lower number of injection points, as in the NMJ strategy, are expected to be preferable for both patients and physicians.

Gracies et al.20 suggested that high volume or endplate targeted BoNT injections results in superior neuromuscular blockade, contraction and spasticity reduction than low volume, non-targeted injections.20 In the Gracies study, the effect was demonstrated using the Tardieu scale in contrast to the MAS used in the current study. Hence, the results are not directly comparable. The lack of statistically significant differences between the strategies in our study might, however, be explained by the different outcome measures used. It might not be possible to capture a statistical difference with the MAS scale in this limited study group, due to lack of power. The GAS, which is a more sensitive scale than MAS to investigate functionally important changes after injection,29 generated similar results between the strategies used in our study.

The experienced pain related to spasticity or injection was mild or moderate. A reduction in pain intensity after 4 weeks was observed in both treatment strategies. Pain associated with injections of BoNT could hypothetically influence optimal treatment results and goal attainment by affecting compliance. An inverse relationship between injection site pain and compliance to BoNT treatment has, for instance, been reported for hyperhidrosis.30

The study did not disclose any previously unknown adverse events. Furthermore, there was no difference in the observed frequency of adverse events between the two injection strategies, as assessed in the ITT population.

The major limitation of this study was the low number of included subjects, as compared with the number needed according to the calculation of statistical power conducted before the study. Most likely, this prevented statistical demonstration of non-inferiority. The main reasons underlying the slow recruitment were difficulties in finding patients who were receiving continuous treatment who also had a MAS score of 2 or higher in the elbow flexors. Also, patients treated successfully with BoNT-A did not fluctuate to the extent that they increased a full level on the MAS between injections. A further limitation was the fact that although patient priorities and preferences were considered when developing the study, patients were not consulted specifically on its design. However, given the focus of the study, this was unlikely to have had a major impact on the overall findings.

The study compared current clinical practice with a NMJ targeting/higher volume technique. As a result, both the targeted injection and the increased volume (resulting in

Table 3 Evaluation of secondary endpoints in the ITT population.

<table>
<thead>
<tr>
<th></th>
<th>Current practice</th>
<th>NMJ-targeted</th>
<th>n=44</th>
<th>n=44</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>MAS response at 12 weeks</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n (%)</td>
<td>19 (48.7%)*</td>
<td>13 (33.3%)†</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adj difference</td>
<td>−0.1324; 95% CI (−0.3531 to 0.0884)</td>
<td>0.24</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>GAS score improvement</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>n (%)</td>
<td>25 (61.0%)‡</td>
<td>21 (63.6%)§</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P value</td>
<td>0.5747</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Spasticity related pain, VAS (mean±SD)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Baseline</td>
<td>14.32±22.64</td>
<td>21.66±28.97</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Week 4</td>
<td>10.30±19.99¶</td>
<td>14.05±27.50**</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean change</td>
<td>−4.35±12.29</td>
<td>−5.8±23.07</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Decrease of mean (%)</td>
<td>30.4</td>
<td>26.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Injection-related pain VAS (at first visit)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean±SD</td>
<td>30.68±27.33**</td>
<td>25.67±25.37††</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Difference</td>
<td>−5.01</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P value</td>
<td>0.406</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*5 patients missing 12-week MAS evaluation. †5 patients missing 12-week MAS evaluation. ‡3 patients missing GAS evaluation. §11 patients missing GAS evaluation. ¶1 patient missing VAS evaluation. ‡‡7 patients missing VAS evaluation. ††8 patients missing VAS evaluation.

GAS, Goal Attainment Scaling; MAS, Modified Ashworth Scale; NMJ, neuromuscular junction; VAS, Visual Analogue Scale.
wider spread) could contribute to the effect on the spastic muscle in the NMJ group. Both treatment arms experienced a high response rate to treatment, suggesting that a much larger study population is needed to be able to detect any difference, if present at all. Different injection guidance techniques (electric stimulation and EMG) were allowed, to reflect the situation in real-world clinical practice. In an attempt to increase recruitment, a protocol amendment was also made allowing use of US guidance techniques; however, only one patient was injected using this technique. Given that the two main injection guidance techniques well balanced between groups, the specific technique used during the study was likely to have had only a limited impact on the overall results.

In summary, the current study provides further insights in the discussions of the effectiveness of BoNT injections targeted to NMJs. Since this strategy enables patients to be treated using fewer injections, the approach may minimise the potentially painful injection experience while having a similar effect on spasticity. Furthermore, the high number of responders with both injection techniques indeed confirms aboBoNT-A to be an effective treatment option for spasticity. Thus, any measures that could have an effect on implementation and/or compliance are important.

This study was conducted on flexors in the spastic arm, and the effect on other spastic muscles using NMJ-targeted injections has yet to be substantiated. Several previous publications suggest injections near NMJs, but an implementation barrier is the limited number of muscles with known localisation of muscle endplates. However, in light of the expanding support for NMJ-targeted injections, further studies adding information to the field are warranted.

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**Acknowledgements** The authors thank the investigators and patients participating in this study. The authors thank PGI for assisting in data management. Medical writing support was provided by Nordic Health Economics and funded by Ipsen AB. The Article Processing Charge was paid by Ipsen AB.

**Contributors** TR and BB-S performed the experiments, analysed the data and wrote the paper. TU and TD conceived and designed the experiments, performed the experiments and analysed the data. RB conceived and designed the experiments, analysed the data and wrote the paper. JH, OJV and PBC performed the experiments. TS conceived and designed the experiments, analysed the data. PM analysed the data. PM conceived and designed the experiments, analysed the data and contributed analysis tools.

**Funding** The study was sponsored by Ipsen.

**Competing interests** The authors have, during the last 3 years, received educational support from the sponsor, Ipsen AB and/or served as consultants for the sponsor. The sponsor contributed to the design of the study; in the collection, analyses/interpretation of data; in the writing of the manuscript and in the decision to publish the results.

**Patient consent for publication** Not required.

**Ethics approval** The publication received ethical approval from the committee for Scientific Ethics (De videnskapselte Komitee for Region Hovedstaden Kongens Vange 2, 3400 Høje Taastrup, Denmark (21 February 2012, 20 June 2012 and 04 July 2014); Central Finland Hospital District, Ethical Commission (Keski-Suomen sairaanhoitopiiri, Ettekin toimikunta), Sairaanhoitopiirin toimisto Rak. 6/2, Keskuussairaantie 19, 40620 Jyväskylä, Finland (27 January 2012 and 27 August 2012); the regional Ethics Committee in Stockholm (Regionala Etikprövningsnämnden i Stockholm), FE289, 171 77 Stockholm, Sweden (01 February 2012, 20 June 2012 and 04 July 2014) and the Regional Committee for Medical and Health Ethics south east Norway (Regional komité for medisinsk og, helsefaglig forskningsetikk REX, sør-øst- Norge), PO-box 1130, 0316 Oslo, Norway (07 August 2012, 01 November 2012 and 04 July 2014). The study was registered on clinicaltrials.gov on 12 September 2012 (NCT01682149).

**Provenance and peer review** Not commissioned; externally peer reviewed.

**Data sharing statement** Trial data analysed during the reported study are available to qualified researchers. Reasonable requests should be directed to the corresponding author.

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