Combination of local Botulinum toxin A injections and standard oral drug treatment allows improvement of long-term pain management in severe post-herpetic neuralgia: A case report

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Abstract

Systematic reviews showed that Botulinum toxin A (BoTN A) may be helpful in controlling neuropathic pain like in Post-herpetic Neuralgia (PHN). We introduce a case report on the effects of a long-term management with intracutaneous BoTN A injections combined with classic drug treatment in a patient with severe PHN. The combination of regular BoTN A therapy every three month and oral drug treatment allows to reduce pain rating with VAS by 50% and suggested that this combination may be helpful for long term management of severe PHN.

Introduction

PHN is a very painful neuropathic status as the consequence of nerve injuries (e.g. demyelinisation, loss of axon, small-fiber-degeneration, reorganization in dorsal horn, neuroplastic central changes) due to herpes-zoster-infection. Definition includes duration of pain for more than 3 months [1,2]. First line treatment of PHN is anticonvulsant in combination with antidepressant, second or third line treatment today is using opioids, and local therapy with lidocaine or capsaicin. However, patients over 50 years of age often show severe side effects from first line oral treatments and opioids and are not satisfied by the duration of effects of current local treatment options. Usually aim of these chronic symptomatic therapies for PHN is to reduce about 30- 50% of pain [3-5].

After the first report on the positive effect of BoTN A for pain in dystonia published by Brin et al. 1987, many studies on positive effects on chronic pain including modulation and management of spasticity associated pain and neurogenic pain, have been published [6-10]. The mechanism of pain reduction from BoTN A injections is still matter of discussion and included the effect on reducing release of various inflammation-mediated substances (substance P, glutamate, calcitonin gene – related peptide) by blocking exocytosis from BoNT A [6-10]. The recent meta-analysis of international association for the study of Pain (IASP) showed positive evidence of BoTN therapy for treatment of neuropathic pain and suggested intracutaneous BoTN A injections as an effective therapy for focal neurogenic pain [11]. Some studies in PHN suggested positive short term effects of BoTN A therapy as alternative treatment in non-manageable PHN by classic medications [12-14]. Although the effects of multifocal intracutaneously injected BoTN A for the treatment of PHN have been known, long term efficacy and safety data of this local therapy approach in PHN is lacking and however this treatment in most countries still is used as “off-label” option yet and no reimbursement is established, i.g. in Germany and other European countries.

Case

The 56-years old male showed up first time in August 2013 and complained severe pain due to PHN in the area of left shoulder and ventral thorax. He stated that severe pain has begun following skin changes due to herpes zoster in June 2011. He experienced at that time pimples within several hours in the painful area between the spinal processes of Th3 - Th 11 and the medial line of left scapulae as well as on the body of sternum and some left skin areas of it (Figure 1). The constant severe pain niveau were judged by the patient with 7 - 8 of 10 Visual Analog Scale (VAS), interrupted by severe shooting and electric-like pain episodes over seconds with levels up to 8 - 9 VAS even under regular oral treatments. Any accompanied diseases were unknown.

At the second day of symptoms with pimples herpes virus infection was diagnosed and only treated with Zn – ointment, no other treatment was introduced by the dermatologist. The scabbing and crusting of the skin injuries needed several weeks, but the severe pain, scars, pigments remained as post-herpetic neuralgia persisted. The severe...
PHN was treated at first by dermatologist and later by a pain specialist, an anesthiologist. The patient had taken for PHN oral pregabalin up to 450mg per day. Morphine up to 80 mg per day, additionally local treatment was introduced with capsaicin ointment and lidocaine plasters were applied by the therapist. He also took mirtazapin (NaSSA: Noradrenergic and Specific Serotonergic Antidepressant), and citalopram (SSRI), but all drugs applied had no positive effect for him. Serial lidocaine plaster also were applied locally on the affected skin, but had only a short time-relief of pain and therefore helped only a little. Additionally 6 acupuncture therapies for 3 weeks were introduced but had no reduction of pain intensity, but following acupuncture the very severe pain up to 8-9 VAS disappeared, but the constant pain level had been remained on levels of 7 - 8 of 10 VAS even under constant drug treatment with morphine 80 mg per day.

In August 2013 we introduced local intracutanous multi-point injections of BoNT A (Incobotulinum Toxin, product name:Xeomin©) with dosage per treatment session of 150 IU (IU=international units, dilution in normal saline to 100 IU / 2 ml) to a total of up to 70-75 intracutanous injection points (one injection site within every 2cm) and a dosage per injection point of 2 - 2.5 IU every 3 months (Figure 2). We used very small needles with 30 gauge and small tuberculin (1 ml volume) syringeous.

Following the first and the following 20 injection cycles usually three weeks following intracutanous BoTN A injections patient reported pain reduction from VAS 7 - 8 to pain levels VAS 4 - 5. Pain level reduction allows him to reduce Morphine to 40 mg per day (half of the initial dose) for the following years. During the first and following injection cycles the patient reported pain reduction for up to 11 weeks and then increasing pain levels up to the pre-injection levels before follow-up injections.

Summing up the intracutanous injection of BoTN A were effective in reducing the intensities of PHN pains and enabled to reduce oral administrations of morphine, so that the patient asked for continuation of BoTN A injections. Up to now we have performed 20 injection sessions. No side effects of BoTN injections were found over the reported treatment regime of 3 years. When comparing the costs of drug including BoTN A injections with the costs before this additional local therapy regime with BoNT A we found an slight elevation in costs of 30%, which represents in Germany an elevation of 96 euros per month.

Discussions
To our opinion and in concordance with recent reports and systematic reviews, severe neuropathic pain like PHN often been found as one of the not completely resolved tasks in pain treatment for clinicians. In recent RCTs sub- and intracutanous injections of BoNT A showed effectiveness in the treatment of neuropathic pain [11-14]. According to the German guideline for diagnosis and therapies in Neurology, the aim of therapies for PHN is to reduce 30-50% of pain and the effectiveness of therapies should be estimated after 2-4 weeks under sufficient dosage [3]. As in our patient, Apalla et al. observed effectiveness of BoTN A treatment for PHN 2-3 weeks after injection. All reported studies for PHN treatment estimated the effectiveness of BoTN A injections with effect size of greater than 50% pain reduction [12-14].

We have performed now over 20 therapies with BoTN A focal injections in the patient reported as additional local therapy in the painful areas and found comparable neurogenic pain reduction of up to 50%, from 7-8 / 10 VAS to 4 - 5 / 10 VAS. Additionally, we were able to reduce dosage of morphine therapy to 50%. We observed very good effectiveness starting mean 3 weeks after injections and last for up to 11 weeks, on average pain was reduced by 50%. Although the positive effect of regular serial intracutanous BoTN A injections was observed in this patient with severe PHN for about 5 years, the pain never was completely eliminated by this treatment, but markedly reduced both subjectively and objectively and the patient never showed any side effect.

Conclusion
Our observations suggest that intracutanous BoTN A injections could be used effectively as one important adjunctive local therapy for non treatable PHN. The costs for symptomatic management might be increase slightly by introducing BoNT A, but the additional effect results in major improvement and justify its use.

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