Protocol for bilateral application of botulinum toxin type A to avoid asymmetry during treatment of hemifacial spasms

Protocolo de aplicação bilateral de toxina botulínica tipo A para evitar assimetria no tratamento de espasmo hemifacial

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ABSTRACT

Introduction: Hemifacial spasm (HFS) is characterized by the involuntary tonic-clonic movement of the muscles of the hemiface. It is usually treated with botulinum toxin (BTX). The classically described unilateral application of BTX results in an asymmetry similar to facial paralysis. The aim of this study was to standardize the treatment of HFS by applying BTX bilaterally to prevent the occurrence of iatrogenic facial asymmetry. Methods: The outcomes of 66 applications in 15 patients were analyzed according to the protocol of the facial paralysis service, to which pretarsal sites were added on the HFS side. On reassessment 15 days later, a complementary dose was administered to patients who exhibited some residual degree of spasm or asymmetry with the aim of determining the dose required to achieve satisfactory spasm control without causing facial asymmetry. Results: The total mean dose was 20.2 U at the contralateral side and 28.4 U at the spasm side (a total dose of 48.6 U per application). There was a significant difference between the doses applied to the zygomaticus, orbicularis oris, and orbicular oculi muscles on each hemiface. Conclusions: The proposed bilateral BTX application technique was effective in controlling HFS and prevented iatrogenic asymmetry. In general, application should be performed at a ratio of 1:1.5 U in the orbicularis oculi (lateral portion) and 1:2 U in the orbicularis oris. In the remaining muscles, the same dose should be administered on both sides and an additional dose can be applied 15 days later if some degree of spasm is present. The pretarsal region of the orbicularis oculi muscle is the only area for which BTX application on the healthy side is unnecessary.

Keywords: Botulinum toxin type A; Facial paralysis; Facial asymmetry/therapy; Hemifacial spasm

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Protocol for bilateral application of botulinum toxin type A

Introdução: O espasmo hemifacial (EHF) caracteriza-se por movimento tônico-clônico involuntário da musculatura de uma hemiface. O tratamento tem sido realizado com aplicação de toxina botulínica (TxB). A aplicação unilateral classicamente descrita resulta em assimetria semelhante à paralisia facial. O objetivo desse trabalho foi normatizar o tratamento do EHF bilateralmente com TxB, a fim de prevenir a ocorrência de assimetria facial iatrogênica.

Método: Foram analisadas 66 aplicações em 15 pacientes, seguindo o protocolo do serviço para paralisia facial, acrescentado de pontos pré-tarsais no lado com EHF. Foi feita dose complementar na reavaliação após 15 dias nos pacientes que apresentavam algum grau residual de espasmo ou assimetria, buscando-se a dose necessária para alcançar controle satisfatório do espasmo sem causar assimetria facial. Resultados: A dose média total foi 20,2 U do lado não acometido e 28,4 U do lado acometido, totalizando 48,6 U por aplicação. Houve diferença significante entre as hemifaces na dose para os músculos zigomático, orbicular da boca e orbicular dos olhos. Conclusões: A técnica proposta de aplicação bilateral de TxB controlou adequadamente o EHF e evitou assimetria iatrogênica. Como regra geral, a aplicação deve ser feita na proporção de 1:1,5 U no orbicular dos olhos (porção lateral) e 1:2 U no orbicular da boca. Nos demais músculos, a dose nos dois lados deve ser a mesma, realizando-se dose de reforço em 15 dias caso permaneça algum grau de espasmo. O único local com pontos exclusivos do lado acometido é a região pré-tarsal do músculo orbicular do olho.

RESUMO

Descritores: Toxina botulínica tipo A; Paralisia facial; Assimetria facial/terapia; Espasmo hemifacial.

INTRODUCTION

Hemifacial spasm (HFS) is characterized by an involuntary tonic-clonic movement of the muscles in the hemiface. It affects the facial expression muscles innervated by the ipsilateral facial nerve. HFS can be divided into two types: primary and secondary. The etiology of the former has not been fully determined. The latter can be caused by trauma, infection, or post-paralytic facial syndrome.

The treatment of HFS with oral drugs (for example, carbamazepine, gabapentin) has not resulted in satisfactory long-term outcomes, nor has extracranial surgery (rhizotomy, myotomy, neural anastomosis) or intracranial surgery such as microvascular decompression of the facial nerve. Currently, the first therapeutic choice is the direct application of botulinum toxin type A to the facial muscles.

Since Elston treated HFS with botulinum toxin for the first time in 1985, the effectiveness of this treatment has been demonstrated consistently. Several studies reported the successful treatment of 76-100% of patients.

Beyond this point, the literature is controversial. Some studies were predominantly based on cases of blepharospasm and asymmetry of the upper third of the face, in which the unilateral application of toxin was the standard method. However, when unilateral application was also used in the lower third, it exacerbated muscle weakness in the treated hemiface and caused asymmetry of the facial expression. In 1991, Borodic et al. initiated studies in which doses of toxin were applied to both the upper third and two lower thirds of the face to soften facial asymmetry. The author noted a common complication - iatrogenic unilateral facial asymmetry - when the toxin was applied only on one hemiface. This complication was mostly observed when treating the middle and lower thirds of the face.

This asymmetry gives patients an appearance similar to that of facial paralysis, with the smile slanting to the contralateral side (not treated with botulinum toxin), which causes great social discomfort. At this point in the evolution of HFS treatment using botulinum toxin type A, the discussion has moved to the...
standardization of sites for the bilateral application of botulinum toxin since the first application\textsuperscript{9,18,20}. The spasm would thus be controlled by the application of the toxin to the spasm side and iatrogenic asymmetry would be avoided by applying the toxin to the contralateral side.

**OBJECTIVE**

The objective of this study was to standardize the treatment of HFS with the bilateral application of botulinum toxin type A (from the first application) to standardized muscle sites of the face to prevent the occurrence of iatrogenic facial asymmetry.

**METHODS**

The present study was prospective from the start and was conducted between May 2005 and June 2012 (the end of data collection). The study was approved by the Ethics Committee of our institution (nº 393/09). Patients were treated by the Plastic and Aesthetics Surgery Group of the Faculty of Medicine, University of São Paulo. The PubMed database of the US National Library of Medicine of the National Institutes of Health was searched for relevant literature using the following keywords: botulinum toxin type A, facial paralysis, facial asymmetry/therapy, and facial spasm.

Sixty-six applications of botulinum toxin in 15 patients were analyzed. The treatments were administered with 5- to 7-month intervals between applications. All patients gave their written informed consent. All patients were interviewed and their HFS etiologies were investigated. The mean patient age was 63.2 years. The cohort was 78.5% female and 21.5% male. The left side was affected in 60% of cases.

All study patients were administered botulinum toxin type A (Botox\textsuperscript{®}; Allergan, Inc., Irvine, California, USA). The sites of application were determined according to the protocol of the facial paralysis service, standardized by Salles\textsuperscript{21} and Salles et al.\textsuperscript{22}, 0.5 or 1 U were applied to the pretarsal sites on the HFS side to control spasms of the orbicular oculi (Figure 1).

The toxin was applied to all muscles in which spasms were observed at a dose of 1-2 U. Symmetrical sites were marked on the contralateral side, in the muscles that cause asymmetry in the upper, middle, and lower thirds of the face.

The muscles analyzed for toxin application were as follows: frontal (2-5 sites), corrugator/procerus (1-3 sites), orbicularis oculi (lateral portion) (2-3 sites), pretarsal (2-4 sites), nasalis (1 site), levator labii superioris (1 site), zygomaticus (1-3 sites), risorius (1-2 sites), depressor labii inferioris (1-3 sites), mentalis (1-2 sites), and orbicularis oris (1-2 sites). The platysma muscle was excluded from this analysis.

All patients were evaluated objectively using standardized photos of the front view, side view, and three-fourth view with a resting face and smiling face, while contracting the corrugator and nasalis muscles, and while protruding the lip and depressing the lower lip. Patients were also filmed both statically and while moving their facial expression muscles. They were also evaluated subjectively at the start of the treatment and after each application, and the degrees of satisfaction and symmetry were always assessed by the same professional. Patients were followed up 15 days after toxin application, and at each visit, patients were reassessed by the first author and photographed to compare the treatment effects over time.

A complementary dose was administered 15 days after the application of botulinum toxin type A only in patients who exhibited a residual degree of HFS or static or dynamic asymmetry. These patients were reassessed after another 15 days to determine the results and side effects. The total dose was defined as the dose needed to achieve satisfactory spasm control without causing facial asymmetry.
RESULTS

During the 6-year study period, the mean number of applications was 4.7 per patient (varying between 4 and 16 applications). The interval between applications was 5 to 7 months.

The cause of spasm was idiopathic in 13 cases (86.6%), trauma in one case, and stroke in one case.

The mean doses (UI) of botulinum toxin type A used per application and per muscle group in each side of the face are listed in Table 1.

<table>
<thead>
<tr>
<th>MUSCLE GROUP</th>
<th>Mean dose</th>
<th>Mean dose</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Affected side</td>
<td>Contralateral</td>
</tr>
<tr>
<td>Pretarsal</td>
<td>3.6</td>
<td>0.0</td>
</tr>
<tr>
<td>Orbicularis oculi (lateral)</td>
<td>6.1</td>
<td>4.3</td>
</tr>
<tr>
<td>Frontal</td>
<td>2.6</td>
<td>2.6</td>
</tr>
<tr>
<td>Corrugators</td>
<td>1.2</td>
<td>1.0</td>
</tr>
<tr>
<td>Depressor labii inferioris</td>
<td>2.1</td>
<td>1.9</td>
</tr>
<tr>
<td>Levator labii superioris</td>
<td>1.8</td>
<td>1.8</td>
</tr>
<tr>
<td>Zygomaticus</td>
<td>4.0</td>
<td>3.2</td>
</tr>
<tr>
<td>Risorius</td>
<td>3.3</td>
<td>2.8</td>
</tr>
<tr>
<td>Orbicularis oris</td>
<td>2.8</td>
<td>1.3</td>
</tr>
</tbody>
</table>

The total mean dose of botulinum toxin per application was 20.2 U at the contralateral side and 28.4 U at the spasm side, for a total mean dose of 48.6 U per application considering the whole face. Figure 2 shows an example of the outcome of toxin application (static and dynamic); pretarsal sites are seen on the side with HFS.

The following complications were observed: slight speech disturbance (50%), slight difficulty in swallowing liquids (35.7%), dry eyes (14.2%), and mild ectropion (6%). These lasted for a maximum of 18 days. All patients were satisfied with the treatment, both with regard to spasm control and facial symmetry.

Table 2 shows a statistical comparison of the two sides of the face. The muscle groups with a p-value > 0.05 were not statistically different, and therefore the dose was the same on both sides of the face.

Table 3 shows the proposed protocol, based on a statistical analysis of the doses needed to treat the spasm without causing iatrogenic asymmetry in this sample.

DISCUSSION

There is consensus in the literature regarding the efficacy of botulinum toxin for the treatment of various disorders. The toxin is endocytosed by the presynaptic neuron and inhibits the release of acetylcholine at the neuromuscular junction, which leads to temporary complete or incomplete paralysis of the muscles in the injected area.

Since Elston treated HFS with botulinum toxin type A for the first time in 1985, studies have reported facial spasm control results of 76-100%.

However, currently available studies on the method of application vary considerably with regard to sample size, ranging from 6 cases to 332 cases. Moreover, most studies only consider the upper third of the face.

In addition, unilateral toxin application is used by most authors, with the exception of Borodic et al. and Colakoglu et al., both of whom studied a small number of patients. There is no standardized protocol for the bilateral application of botulinum toxin in patients with HFS in the literature. This study is therefore unique in that it analyzed 66 bilateral applications, and included muscles of the upper, middle, and lower thirds of the face.
Table 2. Statistical comparison of doses used on the two hemifaces. The affected side was treated for hemifacial spasm and the contralateral side was treated to avoid asymmetry.

<table>
<thead>
<tr>
<th>MUSCLE GROUP</th>
<th>Total dose Spasm</th>
<th>Total dose Contralateral</th>
<th>Chi-square test for proportions</th>
<th>Statistical test results</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>chi²</td>
<td>p</td>
</tr>
<tr>
<td>Frontal</td>
<td>173</td>
<td>169</td>
<td>50.58%</td>
<td>49.42%</td>
</tr>
<tr>
<td>Corrugators</td>
<td>78</td>
<td>62</td>
<td>55.71%</td>
<td>44.29%</td>
</tr>
<tr>
<td>Depressor labii inferioris</td>
<td>140</td>
<td>124</td>
<td>53.03%</td>
<td>46.97%</td>
</tr>
<tr>
<td>Levator labii superioris</td>
<td>116</td>
<td>118</td>
<td>49.57%</td>
<td>50.43%</td>
</tr>
<tr>
<td>Risorius</td>
<td>218</td>
<td>183</td>
<td>54.36%</td>
<td>45.64%</td>
</tr>
<tr>
<td>Zygomaticus</td>
<td>266</td>
<td>214</td>
<td>55.42%</td>
<td>44.58%</td>
</tr>
<tr>
<td>Orbicularis oris</td>
<td>184</td>
<td>83</td>
<td>68.91%</td>
<td>31.09%</td>
</tr>
<tr>
<td>Orbicularis oculi</td>
<td>400</td>
<td>281</td>
<td>58.74%</td>
<td>41.26%</td>
</tr>
<tr>
<td>Pretarsal</td>
<td>239</td>
<td>0</td>
<td>100.00%</td>
<td>0.00%</td>
</tr>
</tbody>
</table>

Table 3. Suggested dose for avoiding asymmetry from the first application. The column “ratio” indicates the difference in dose between the two sides.

<table>
<thead>
<tr>
<th>MUSCLE GROUP</th>
<th>Ratio</th>
<th>Status da chance</th>
<th>Healthy : Spasm application in practice</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frontal</td>
<td>1.02</td>
<td>Apply the same dose on both sides</td>
<td>1 UI : 1 UI</td>
</tr>
<tr>
<td>Corrugators</td>
<td>1.26</td>
<td>Apply the same dose on both sides</td>
<td>1 UI : 1 UI</td>
</tr>
<tr>
<td>Depressor labii inferioris</td>
<td>1.13</td>
<td>Apply the same dose on both sides</td>
<td>1 UI : 1 UI</td>
</tr>
<tr>
<td>Levator labii superioris</td>
<td>0.98</td>
<td>Apply the same dose on both sides</td>
<td>1 UI : 1 UI</td>
</tr>
<tr>
<td>Risorius</td>
<td>1.19</td>
<td>Apply the same dose on both sides and, if necessary, add a complementary dose after 15 days</td>
<td>1 UI : 1 UI</td>
</tr>
<tr>
<td>Zygomaticus</td>
<td>1.24</td>
<td>Apply the same dose on both sides and, if necessary, add a complementary dose after 15 days</td>
<td>1 UI : 1 to 1 : 1.5 UI</td>
</tr>
<tr>
<td>Orbicularis oris</td>
<td>2.22</td>
<td>Apply higher dose on the side affected by HFS</td>
<td>1 UI : 2 UI</td>
</tr>
<tr>
<td>Orbicularis oculi</td>
<td>1.42</td>
<td>Apply higher dose on the side affected by HFS</td>
<td>1 UI : 1.5 UI</td>
</tr>
<tr>
<td>Pretarsal</td>
<td>infinite</td>
<td>Apply dose only on the side affected by HFS</td>
<td>0 UI : 1 UI</td>
</tr>
</tbody>
</table>

Complications described in the literature include paralysis of the orbicularis oris with difficulty in oral continence (38.8% of cases), dry eyes (19.8%), eyelid ptosis (between 10.9% and 52.7%), eyelid ptosis (between 2% and 27.7%), and diplopia (between 2% and 27.7%). Complications in the present study were similar to those described above and occurred in similar proportions: slight speech disturbance (50%), slight difficulty in swallowing liquids (35.7%), dry eyes (14.2%), and mild ectropion (6%). The maximum duration was 18 days.

The mean dose required to completely control HFS decreased over time. The intervals between applications showed an increasing trend. These subjective findings require further confirmation by larger studies.

The present study showed that the pretarsal region should only be treated on the side affected by spasm. There was a statistical difference between the doses required to control spasm without causing iatrogenic asymmetry in 3 muscle groups: the dose administered to the zygomaticus on the affected side was 1.24 times higher than that administered to the contralateral side, the dose administered to the orbicularis oris muscles was 2.22 times higher, and the dose applied to the orbicularis oculi muscles was 1.42 times higher.
There was also a statistical difference with regard to the remaining analyzed muscle groups (frontal, corrugator, depressor labii inferioris, levator labii superioris, and risorius). These exhibited dose differences as well, with the spasm side requiring higher doses of toxin; however, the differences were not statistically significant ($p > 0.05$).

The physician should start the injection by determining the dose needed in each muscle with dystonia on the spasm side, using doses of 0.5 to 2 units. On the contralateral side, the sites are marked according to the dose adjustment proposed in Table 3. A complementary dose can be administered 15 days later if residual spasm or asymmetry is present.

**CONCLUSION**

The technique proposed for the bilateral application of botulinum toxin was effective in controlling HFS and prevented iatrogenic asymmetry of the facial muscles, a complication observed in cases in which unilateral application is used to solely treat dystonia.

There was a statistically significant difference between the doses applied to each side of the face with regard to the zygomaticus, orbicularis oris, and orbicularis oculi muscles. The analysis should be performed on a case-by-case basis, but in general, the toxin should be applied at a 1:1.5 U ratio at the orbicularis oculi (lateral portion) and a 1:2 U ratio at the orbicularis oris. In the remaining muscles (frontal, corrugator, depressor labii inferioris, levator labii superioris, and risorius), the same dose should be applied on both sides. The zygomaticus should receive a similar dose, and a complementary dose can be administered 15 days later if there is some degree of spasm. There is the option of using an asymmetrical dose of 1:1.5 U if the spasm is very intense in this area. The pretarsal region of the orbicularis oculi is the only area for which botulinum toxin application on the healthy side is unnecessary.

There was a low incidence of complications and these were mild and self-limited. The cost of a mean dose of 48.6 U per session is feasible, considering that the mean annual cost of the medication to treat one dose of 48.6 U per session is feasible, considering that these were mild and self-limited. The cost of a mean dose of botulinum toxin on the healthy side is unnecessary.

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