The non-hallucinogen 2-bromo-lysergic acid diethylamide as preventative treatment for cluster headache: An open, non-randomized case series

Matthias Karst¹, John H Halpern², Michael Bernateck¹ and Torsten Passie¹

Date received: 10 September 2009; accepted: 24 January 2010

Introduction

Cluster headache (CH) is a stereotyped primary headache characterized by strictly unilateral severe orbital or periorbital pain and categorized as either episodic or chronic (1,2). Its prevalence is 0.1% (3). Oxygen and sumatriptan are the treatments of choice for individual attacks, whereas verapamil, lithium, corticosteroids and other neuromodulators can suppress attacks during cluster periods (1). All standard medication treatments may be ineffective. Surgical treatment may be an option for medication non-responders, including deep brain (4) or occipital nerve stimulation (5). However, serious complications from brain surgery, including death, can occur (6).

An Internet survey of 53 CH patients reported on claims that psilocybin is better at aborting acute attacks than either oxygen or sumatriptan and that LSD and psilocybin are both better at triggering and extending remission than standard drugs (7). However, due to hallucinogenicity and the absence of established medical indication, these drugs are criminalized and placed within the most restrictive Schedule I of the Controlled Substances Act, which sanctions only limited research use. Although the hallucinogenic properties of LSD and psilocybin are undesirable from both regulatory and patient safety perspectives, it was unclear to us at the outset whether a non-hallucinogenic analog could also provide meaningful relief to CH patients. To address the question of whether the CH relief associated with these two structurally diverse compounds is related to the mechanisms triggering intoxication, we decided to investigate the efficacy of a non-hallucinogenic analog of LSD. LSD’s hallucinogenic effects are completely lost when the double bond in the D ring is saturated and with substitution at R² (e.g. by bromination in 2-bromo-LSD) (BOL-148) (8). BOL-148 has been studied in volunteers (up to 20 mg per os) (9) and in patients suffering from vascular headaches but not, apparently, in patients with CH (9,10). These past studies concluded that BOL-148 is non-toxic and non-hallucinogenic. Only very mild side effects, if any, have been observed, when given in the dose range used in our project (30 μg/kg/body weight) (9). No long-term behavioral or psychological effects from BOL-148 have been reported from past studies with more than 300 healthy, normal subjects (11), and 30 mg BOL-148 administered daily over four to five weeks failed to alter active psychosis in chronically ill schizophrenic women (12).

Case series

Patients referred to Hannover Medical School’s Pain Clinic were identified with CH if they met the respective diagnostic criteria of the International Classification of Headache Disorders (2). All patients, who were seriously affected by the disease, were non-responders to verapamil (or could not tolerate its side effects at higher doses) and to some extent to other prophylactic medications as well, although not all medication alternatives (e.g. topiramate or prednisone), or more invasive procedures (e.g. intravenous dihydroergotamine or occipital nerve stimulator implantation), had been attempted.

¹Hannover Medical School, Germany.
²McLean Hospital and Harvard Medical School, USA.

Corresponding author:
Matthias Karst, Department of Anesthesiology, Pain Clinic,
Carl-Neuberg-Str. 1, 30625 Hannover, Germany.
Email: karst.matthias@mh-hannover.de
All patients signed an informed consent that declared their agreement to participate in this project on the compassionate use of BOL-148 for CH. It was approved by the local ethics committee in accordance with German law. Patients kept a standardized daily diary of CH symptoms (see www.clusterbusters.com for a copy) starting at least two weeks prior to BOL-148 administration. BOL-148 was manufactured by THC pharm GmbH (Frankfurt am Main, Germany). A purity of >99.2% was identified by high-performance liquid chromatography (HPLC) and other analytical tests. BOL-148 30 µg/kg/body weight was dissolved in distilled water and then given once every five days for a total of three doses per os. BOL-148 was administered in the presence of two of the authors (MK, TP). Alterations in consciousness, thought disturbances, and vital signs (blood pressure, heart rate) were measured during a three-to-four-hour observational period, as BOL-148 is typically active for two to three hours. Patients were asked to continue completing daily headache diaries for at least one month or until they experienced three days of attacks, starting a new cluster series.

Results are summarized in Table 1 and Figure 1. One patient (S2) with episodic CH, who was in an active attack period, and four patients with the chronic form participated. All but one patient (S1) had experienced symptoms for more than 10 years. Patient S2’s cluster period terminated after BOL-148 with a long-lasting remission period of six months (at last follow-up) and continuing. Patients S3 and S5 reported pronounced reduction of attack frequency, including full remission for more than one month, indicating transition from a chronic to an episodic form. Cluster attacks resumed after a two-month remission for patient S5. In nine months since BOL-148 treatment, patient S3 describes ongoing remission of cluster period, reporting only a few solitary sporadic attacks. Patient S4 reported a profound reduction in attack frequency, although without one full month of remission and attack frequency increasing approximately six months after BOL-148 treatment. In addition, patients S3 and S4 found the pain intensity of remaining occasional attacks so improved that they no longer administered an acute intervention, as they had prior to BOL-148. Although patient S1 did not experience pronounced attack reduction similar to the other four patients, he indicated a decrease of attack intensity of about 30% within the first four months. It is likely relevant that patient S1 continued to drink alcohol (contrary to advice), a known and common trigger for attacks.

No changes to heart rate and blood pressure were observed during BOL-148 treatment. Most of the patients recorded some kind of “flabby” or “light drunk” feelings. Patient S2 noted a “funny” feeling, tense muscles, and sweaty palms. These mild subjective effects lasted from one to two hours. No visual hallucinations or distortions occurred, nor was there any evidence of delusional thinking or overt psychosis.

Discussion

The results show that three single doses of BOL-148 within 10 days can either break a CH cycle or considerably improve the frequency and intensity of attacks, even resulting in changing from a chronic to an episodic form, with remission extending for many months or longer. While for patients S3, S4, and S5 the remission is very likely due to BOL-148 treatment, for S1, who charted in his diary continued attacks with reduced pain, and S2, who suffered from episodic CH, the observed effects may also be due to the natural course of the disease, despite S1 and S2’s impression that their cluster attack cycle improved in ways they had not experienced before BOL-148. Except for very mild alterations of subjective state and mild to no sympathetic reactions for about two hours, no other side effects were observed.

Sicuteri et al. used LSD and some of its derivatives (with BOL-148 among them) in the treatment of migraine and other vascular headaches (10). Because those studies were entwined with the task of identifying the pathophysiological mechanism of vascular headaches (13), they lack exact documentation and follow-up results of the exposed subjects. Especially considering the results we report, no evidence has been found that BOL-148 was administered specifically for active CH in these earlier trials. A sufferers-driven interest in the clinical effects of LSD and psilocybin for CH did not develop until recently, from anecdotal observations to Internet-based discussions to the published Internet survey (7) and subsequent science-media interest. Interestingly, those reports describe a single dose or a few doses resulting in long-lasting effects, which we now also demonstrate from BOL-148. Taken together and in regard to failure of other more direct explanations, especially for the long-range remission extension, these results indicate that BOL-148, psilocybin, and LSD may influence the expression of genes (epigenetics), which are responsible for the biological clock of the organism (14). However, prolonged administration of BOL-148 does not result in cross-tolerance to LSD (15). This, in turn, suggests that BOL-148’s mechanism of action for CH is unrelated to those receptor systems thought to be involved with hallucinogenicity: 5-HT-1A and 5-HT-2A (16). Similarly, psilocybin and LSD’s treatment effects for CH also, then, may have little to do with their capacity to induce hallucinogenic effects. The ergotamines (including BOL-148, LSD, dihydroergotamine, and methysergide) likely have positive treatment effects for CH through serotonin-receptor-mediated vasoconstriction. BOL-148 was
**Table 1.** Demographic data and clinical aspects

<table>
<thead>
<tr>
<th>Subject</th>
<th>S1</th>
<th>S2</th>
<th>S3</th>
<th>S4</th>
<th>S5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex (M/F)</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
<td>M</td>
</tr>
<tr>
<td>Age (years)</td>
<td>46</td>
<td>28</td>
<td>47</td>
<td>41</td>
<td>41</td>
</tr>
<tr>
<td>Body weight (kg)</td>
<td>83</td>
<td>68</td>
<td>106</td>
<td>105</td>
<td>74</td>
</tr>
<tr>
<td>Body height (cm)</td>
<td>180</td>
<td>168</td>
<td>188</td>
<td>195</td>
<td>174</td>
</tr>
<tr>
<td>Years of illness</td>
<td>3</td>
<td>10</td>
<td>10</td>
<td>33</td>
<td>32</td>
</tr>
<tr>
<td>Cluster headache form</td>
<td>Chronic</td>
<td>Episodic</td>
<td>Chronic since 2005</td>
<td>Chronic since 2001</td>
<td>Chronic since 2007</td>
</tr>
<tr>
<td>Attacks per week in the pre-assessment week</td>
<td>6</td>
<td>7</td>
<td>10</td>
<td>15</td>
<td>19</td>
</tr>
<tr>
<td>Mean intensity of attacks (VAS) in the pre-assessment week</td>
<td>8.4</td>
<td>8.3</td>
<td>5.5</td>
<td>6.4</td>
<td>7.0</td>
</tr>
<tr>
<td>Treatments (acute)</td>
<td>Sumatriptan 20 mg IN</td>
<td>100% oxygen 15 l/min</td>
<td>100% oxygen 15 l/min</td>
<td>100% oxygen 15 l/min</td>
<td>100% oxygen 15 l/min</td>
</tr>
<tr>
<td>Treatments (prophylactic)</td>
<td>Verapamil 240 mg/day*</td>
<td>Verapamil 240 mg/day*</td>
<td>Frovatriptan PO (up to 2.5 mg TID)</td>
<td>Methysergide unknown dose (for 1 year)</td>
<td>Verapamil 960 mg/day (for several months)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Prednisolone 80 mg (for 5 days)</td>
<td>4 cycles with prednisolone starting with a daily dose of 100 mg</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Verapamil 320 mg/day (for 3 months)</td>
<td>Lithium 450 mg/d (for 14 days)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Lithium unknown dose (for 3 months)</td>
<td>Doxepine 10 mg/day (for several months)</td>
</tr>
<tr>
<td>BOL-148 (30 μg/kg) three times within 10 days (days 1, 5, and 10)</td>
<td>2.5 mg</td>
<td>2.0 mg</td>
<td>3.1 mg</td>
<td>3.1 mg</td>
<td>2.2 mg</td>
</tr>
<tr>
<td>Side effects</td>
<td>“Flabby feeling” for about 2 h</td>
<td>“Funny feeling” for about 2 h</td>
<td>“Slightly tipsy” for about 2 h</td>
<td>“Slightly tipsy” for about 2 h</td>
<td>“Slightly tipsy” for about 2 h</td>
</tr>
<tr>
<td>Vital signs</td>
<td>Unchanged</td>
<td>Unchanged</td>
<td>Unchanged</td>
<td>Unchanged</td>
<td>Unchanged</td>
</tr>
</tbody>
</table>

M, male; F, female; VAS, visual analog scale; BOL-148, 2-bromo-LSD; IN, intranasal; PO, per os; SC, subcutaneous; h, hours; min, minutes; *Higher doses not tolerated.
specifically created as a completely non-hallucinogenic form of LSD, but methysergide was developed to have even more potency at serotonin receptors (and less hallucinogenic effects than LSD) (17). While methysergide, an often effective preventative compound if taken on a daily basis for up to six months (18), does not generally induce remissions, the repetitive intravenous and subcutaneous application of 1 mg dihydroergotamine for up to three weeks has been shown in an open retrospective trial to sometimes break a cluster period (19). However, dihydroergotamine is not approved for intravenous or subcutaneous injection in Germany. In addition, BOL-148 seems to exert its effects in a totally different way, as outlined above. Although, after extended and chronic use, both methysergide and dihydroergotamine may be associated with an increased risk for fibrotic complications (such as retroperitoneal fibrosis), this risk is unknown for BOL-148 and seems to be more unlikely from the limited, non-chronic dosing regimen of BOL-148 we employed. Pointedly, there are no pre-clinical studies linking LSD to fibrosis, and, despite an extensive history of illicit use, only one case report is identified in the PubMed database describing prior use of LSD in two individuals with “idiopathic” retroperitoneal fibrosis (20). None of the approved ergot-based medications for CH realize the type of profound and lasting treatment response we report from just three oral doses of BOL-148 or in the prior case series of LSD and psilocybin use (7). BOL-148 apparently also differs from methysergide in that prior research indicates methysergide is a less effective preventative for chronic CH than for episodic forms (21).

The results of this case series must be regarded as preliminary, in that they are unblinded and uncontrolled. In acute attack treatment trials, the frequencies of placebo responders were up to 42% while in chronic CH a placebo response as low as 14% was reported in one trial (which employed a very strict endpoint of cessation of attacks), but no placebo response (for efficacy) was noted in five of seven controlled trials (22). Especially since chronic CH patients appear “to have a relatively modest placebo response” (22), the extended durability of response to three doses of BOL-148 administered over ten days is unlikely to be an artifact. An additional limitation to this report is that not all known prophylactic alternatives had been tried with our patients to confirm their extent of treatment resistance, but all five subjects did respond to BOL-148. In contrast to the compassionate use setting in this case series, follow-up research with more specific inclusion criteria (e.g. prior verapamil trial of at least 500 mg/day, separation of evaluation of BOL-148 for either episodic or chronic forms) will allow more specific conclusions to be drawn about BOL-148 as a potential treatment for CH. Given that the current standard of care involves interventions that break single headache attacks and reduce pain duration, frequency and intensity of attack cycles, and that identified treatments that extend remission are lacking, the potential breakthrough treatment of BOL-148 warrants wide dissemination of these early findings to encourage aggressive development to randomized controlled trials.

**Acknowledgements**

We are grateful for the kind support of Clusterbusters, Lombard, IL, USA. Clusterbusters were neither involved in the conduct of the compassionate use treatment in CH nor collection, management, analysis, or interpretation of the data.
or preparation, review, or approval of the manuscript. Dr Halpern co-holds a patent on BOL-148 for CH with Dr Passie, which has not been licensed and has not generated any royalties or other payments. Drs Bernateck and Karst report no conflicts of interest.

References