Outbreak of wound botulism in injecting drug users

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SUMMARY

Between October and December 2005, 16 cases of wound botulism were notified to the health authorities of North Rhine-Westphalia, Germany. All patients were injecting drug users (IDU) and the epidemiological investigations suggested contaminated injection drugs as the most probable source of infection. Clostridium botulinum was cultivated from clinical samples of six patients and molecular typing revealed that the different isolates were clonally identical. Two samples of heroin, one of them provided by a patient, were examined but C. botulinum could not be isolated. This outbreak demonstrates that IDU are at risk for acquiring wound botulism by injecting contaminated drugs. A greater awareness of this disease is needed by physicians and a close cooperation between public health authorities, street workers, operators of sheltered injecting facilities, and medical centres focusing on IDU is essential to prevent and manage outbreaks in IDU in a timely manner.

Key words: Clostridium botulinum, drug abuse, heroin, outbreak investigation, parenteral molecular typing, wound botulism.

INTRODUCTION

Botulism is a rare disease, caused by neurotoxin release from the anaerobic spore-forming bacteria Clostridium botulinum. Three main, naturally occurring epidemiological types of botulism can be distinguished: foodborne, intestinal colonization (infant form) and wound botulism. In Germany, botulism is a notifiable disease. Between 2001 and 2004, 33 cases of botulism were notified to the German health authorities (6–11 per year) of which 29 were considered to be foodborne in adults, three were infant botulism, and one wound botulism [1].

In the days leading up to 3 November 2005, nine cases of wound botulism in injecting drug users (IDU) were notified in North Rhine-Westphalia (NRW). An investigation was initiated to identify the extent and the source of the outbreak in order to prevent the occurrence of more infections. In this paper we describe the epidemiological and laboratory investigations conducted.

METHODS

Epidemiological investigations

Case-patients were defined as current IDU, resident in Germany, presenting with acute onset of flaccid
paralysis or cranial nerve palsy, from 1 October 2005 onwards. Once their clinical condition allowed, patients who gave informed consent were interviewed by health authority personnel or trained social workers with a detailed standardized questionnaire containing questions on clinical symptoms, consumption of food known to bear a risk for transmission of botulism, drug-consumption habits during the 4 weeks prior to hospital admission, and whether the patients knew about other cases. Patients were also asked whether they bought their heroin in Germany or in the adjacent Netherlands and whether they knew from which country or region the heroin originated.

On 14 November 2005 a message was circulated to all public health departments and emergency physicians in NRW via email networks maintained by the Institute for Public Health, NRW and the local public health departments of NRW. Via the German national Epidemiological Bulletin physicians and public health authorities in other German federal states were informed about the occurrence of wound botulism in IDU in NRW [2, 3]. Through organizations offering outreach, sheltered injecting facilities, and via street workers attempts were made to inform the drug-using communities in Germany about the clinical symptoms of botulism as well as possible routes for acquiring the disease. IDU were asked to seek medical advice if symptoms (flaccid paralysis, difficulties in swallowing, blurred vision) were present. Physicians and laboratories were reminded to immediately notify any cases of botulism.

Due to possible cross-border spread of the outbreak, information was also distributed through the European Early Warning and Response System as well as Eurosurveillance [4].

Laboratory investigation

Wound swabs obtained from six patients were sent to the National Reference Laboratory for Anaerobes (NRL-A) for further identification.

As illicit injection drugs were the most strongly suspected source of infection, patients were asked for samples of heroin they were using in order to investigate it for contamination with Clostridium spores.

Culture and identification of clinical samples

At NRL-A wound swabs were analysed employing Columbia agar (Oxoid Ltd, UK) supplemented with 5% sheep blood (Oxoid GmbH, Germany), vitamin K1 (Sigma Chemical Co., USA), and haemin (Serva Feinbiochemica, Germany), as well as standard appropriate supplemented broth media under anaerobic conditions to cultivate anaerobes. Standard appropriate solid and broth media were used for aerobic cultures to control possible aerobe growth. The anaerobic cultures were incubated in an anaerobic chamber (Heraeus, Germany) containing 80% N₂, 15% CO₂ and 5% H₂ at 37 °C. Obligate anaerobic strains were identified by the Rapid ID 32A system (bioMérieux, France) and by using pre-reduced anaerobic systems (PRAS; Anaerobe Systems, USA).

Cultivation of heroin samples

Two heroin samples were evaluated for microbiological culture. The samples were divided into several aliquots and thereafter added to brain heart infusion broth (Oxoid Ltd) supplemented with vitamin K1 and haemin. Samples were either untreated or heat-shocked and incubated at 36 °C for 3 weeks under aerobic and anaerobic conditions. Subcultures were performed on days 7, 14, and 21 or after visible growth on appropriate agar plates as described above and incubated under aerobic and anaerobic conditions.

Characterization and comparison of isolates

16S rDNA amplification was performed with the primers 609V and 699R as described by Ackermann et al. [5] resulting in a PCR product of ~300 bp. Amplification sequences were compared using the NCBI BLAST database (http://www.ncbi.nlm.nih.gov/blast/Blast.cgi).

For PCR, DNA was prepared using the Qiagen tissue kit (Qiagen GmbH, Germany) according to the manufacture’s instructions. PCR assays were separately performed for C. botulinum type A, B, E, and F neurotoxin genes with the specific primers AS-11 and AS-22 (toxin type A; sequences described by Takeshi et al. [6]) to produce a PCR product of ~283 bp and the specific primers CBMLB1, CBMLB2, CBMLE1, CBMLE2, CBMLF1, and CBMLF2 (toxin types B, E, and F; sequences described by Lindström et al. [7]) to produce a PCR product of ~205 bp (toxin type B), 389 bp (toxin type E), and 543 bp (toxin type F). After 2 min of denaturation at 95 °C, 35 cycles of the PCR mix with a final volume of 50 μl were run as follows: 30 s at 94 °C, 30 s at 55 °C for toxin types B, E, and F, and 30 s at 53 °C for toxin type A, respectively, 1 min
at 72 °C and a final extension cycle of 3 min at 72 °C. 

**C. botulinum** reference strains (REB 1750, toxin type A; REB 89, toxin type B; REB 1072, toxin type E; and REB 1718, toxin type F) were used as control for each toxin type.

For the antibiogram the susceptibility of the isolates to penicillin, clindamycin, piperacillin/tazobactam, imipenem, and metronidazole was determined by Etest (AB Biodisk, Sweden), performed according to the manufacturer’s instructions [8].

Serum samples from eight patients were tested in bioassays on mice. Briefly, 0.5 ml serum of a patient with suspected wound botulism was administered intraperitoneally to two mice. In the control group of two other mice, each was initially administered intraperitoneally 0.5 ml antitoxin (Botulism Antitoxin Behring; 1 ml contains: 750 IU **C. botulinum** type A antitoxin, 500 IU **C. botulinum** type B antitoxin, 50 IU **C. botulinum** type E antitoxin; Chiron Behring GmbH & Co., Germany) followed by 0.5 ml serum of the patient 30 min later. The mice were observed until day 7 post-injection for survival and clinical signs of botulism.

**PFGE** was performed by contour-clamped homogeneous electric field electrophoresis (CHEF-DR III apparatus; Bio-Rad, Germany) using the GenePath Group 1 Reagent kit (Bio-Rad Laboratories, USA) according to the protocol described previously for the GenePath Group 3 Reagent kit and SpeI [9]. Restriction digestion was performed with *Sma* (Bio-Rad Laboratories). For PFGE interpretation the criteria described by Tenover et al. were applied [10].

Different control strains were used: a **C. botulinum** isolate carrying the neurotoxin A gene and additionally a **C. botulinum** strain carrying the neurotoxin B gene (reference strain REB 89). A **C. sporogenes** strain served as control for closely related other *Clostridium* species. The **S. aureus** strain (NCTC 8325) was used as a culture control with the GenePath Group 1 kit.

**RESULTS**

**Epidemiological investigation**

Between 13 October and 5 December 16 cases occurred. After information on the outbreak had been distributed at local, national, and European level, six cases which initially had not been notified came to the attention of the German health authorities. One patient self-reported following hospitalization with classical symptoms retrospectively classified as botulism, but misdiagnosed as a drug overdose. Cases from outside Germany were not reported.

Of the 16 case-patients 12 were male and four female. Median age was 33 years (range 20–50 years). Cases resided in eight different districts of NRW which are in close proximity to each other (Fig. 1).

Eleven of the cases lived in cities (Cologne, Bonn, Aachen, Krefeld), the remainder in adjacent rural districts. The most common symptoms reported were difficulties in breathing and swallowing [14 cases (88%) and 9 cases (56%), respectively], and blurred vision [11 cases (69%)]. Cranial nerve palsy, ptosis, and palsy of peripheral nerves were observed in nine (56%), six (38%), and four (25%) cases, respectively. All patients were hospitalized (median duration 26 days) and recovered fully. Assisted ventilation was required by five (31%), and administration of antitoxin also by five patients. All patients were IDU, injecting heroin on a regular basis. Twelve reported injecting drugs subcutaneously and/or intramuscularly. The other four denied this route of injection, but nevertheless clinical examination of their skin revealed changes (circular scars, swellings, and redness) which could be due to subcutaneous injection (‘skin popping’).

Nine cases (six males) were interviewed with the study questionnaire while the other seven refused to participate. One of the nine questionnaires was excluded from analysis because it largely contained missing or indeterminate answers. Two respondents answered that they knew about further cases; however, they were unable to provide names or addresses for these persons. No meals or gatherings were attended by two or more patients. Food items known to bear a risk for transmission of botulism, including home-bottled or vacuum-packed foods, had not been consumed recently. Six patients had been using drugs for a median length of 12 years (range 2–35 years). Two patients did not specify how long they had used drugs previously. However, one of them had used drugs for ‘several years’ and both indicated that they had injected at least daily the month before the infection.

The heroin consumed by the patients before onset of symptoms came from different sources and was reported to be of different colours (ranging from pale to dark brown) and consistencies – two patients described a black-brown resinous consistency. Five respondents had used dark-coloured heroin within the 4 weeks before symptom onset, whereas the other three said, that they used ‘normal’ heroin without
further specification. A traceback was not possible. Two patients reported buying heroin in The Netherlands themselves, and two others had bought their heroin from dealers who had bought it in The Netherlands. The other four had recently changed dealers and did not know the origin of the heroin sold to them. None of the patients had recently shared needles or syringes. None had changed their typical sources of water, acidifying agents (ascorbic acid and lemon juice) and other ingredients for the preparation of the drugs.

Laboratory investigation

In six patients *C. botulinum* were isolated from wound swabs. In one case bacteria were isolated from different wounds and appeared macroscopically as two morphologically distinct isolates. However, by PFGE analysis the strains were indistinguishable. In three wound swabs, aside from *C. botulinum*, cultures of normal skin flora were obtained, i.e. viridans streptococci and coagulase-negative *Staphylococcus* spp. From one specimen *S. aureus* was cultivated. No other *Clostridium* spp. were detected. The MIC ranges ($\mu$g/ml) of the *C. botulinum* strains were: 0.06–0.25 penicillin, 4.0–8.0 clindamycin, 0.5–4.0 piperacillin/tazobactam, 0.06–0.125 imipenem, and 0.25–4.0 metronidazole.

The neurotoxin B gene was detected by PCR in all of these strains whereas PCR was negative for neurotoxin A, E, and F genes. All seven strains investigated by PFGE exhibited an identical clonal profile (Fig. 2). In contrast, the two *C. botulinum* control strains showed distinct patterns.

Serum samples were obtained for mouse bioassays from three patients with a positive culture of *C. botulinum* and from additional five patients with either a negative culture result or without available material for culture. None of the bioassays were unequivocally positive.

Only one patient was willing to submit a heroin sample from his personal stock for testing. A second sample was obtained from the police. This sample had been confiscated in Cologne and although it is unclear whether this sample bears any relation to the outbreak, both samples were examined but were repeatedly culture negative for *C. botulinum*. Only *Propionibacterium* spp., *Enterococcus* spp., and
DISCUSSION

We describe the first outbreak of wound botulism in Germany. Although the definitive source could not be determined, the finding of clonality in PFGE analysis of seven *C. botulinum* isolates derived from six patients strongly suggests a common source of the outbreak. Contaminated heroin seems to be most probable vehicle since other links between the cases (such as shared needles or dilution materials) could not be identified.

The negative culture result of the single heroin sample obtained from a case does not exclude heroin as a probable vehicle because street heroin has been described as having an antibacterial effect and the number of positive cultures depends on the pre-analytical preparation methods in the laboratory and the culture conditions of the heroin samples [11, 12]. In addition, the sample may not be from the same batch as the heroin at the source of the outbreak.

The investigation of this outbreak was hampered by legal restrictions concerning the usage of confiscated heroin samples for purposes other than criminal prosecution. A critical review of this issue by the district government responsible came to the conclusion that even heroin confiscated in the context of a criminal prosecution cannot be readily provided for other purposes such as microbiological investigations.

The negative mouse bioassay results of the serum samples of eight patients might be explained by late collection of serum samples, although the samples were collected prior to treatment with antitoxin. This is known to be an important factor, because the toxin cannot be detected in serum once it permanently becomes bound to its cell receptors [13]. Furthermore, toxin type A is more aggressive than toxin type B [14]. Thus, the free and unbound amount of toxin present in the serum was probably below the level of detection. Toxin testing was positive only for 37% of sera collected from 309 persons with clinically diagnosed foodborne botulism reported to CDC from 1975 to 1988 [15]. Critchley et al. reported on 27 patients in an outbreak of foodborne botulism in North West England and Wales in 1989. All serological tests of botulism in the affected patients were negative [16]. On the other hand, a study in the UK showed that 33/88 cases of wound botulism in heroin injectors (37.5%) reported in the UK and Ireland between 2000 and 2004 were laboratory confirmed by the detection of botulinum neurotoxin in serum [17]. Furthermore, Akbulut et al. reported the detection of botulinum neurotoxin by bioassay in the sera of 9/16 (56.3%) patients with suspected botulism who were IDU [18].

In the late 1990s wound botulism had become an increasing problem in California and subsequently became apparent elsewhere in the USA [19, 20]. In Europe, an increase of soft-tissue infections with spore-forming bacteria (*Clostridium* and *Bacillus* spp.) was reported from the UK starting in 2000 [13, 21]. In those reports the disease was linked to IDU injecting drugs by muscle or skin ‘popping’ [21]. In the USA, the intramuscular or subcutaneous usage of ‘black tar heroin’ (BTH), a dark, gummy and frequently impure heroin product primarily from Mexico and countries south of the USA, was described to be particularly associated with wound botulism [19, 20, 22, 23]. Due to its tarry consistency, this form of heroin is injected mainly subcutaneously or intramuscularly which provides excellent environmental conditions for the spores to germinate to bacteria. BTH does not seem to be the reason for the increase of disease in the UK since the heroin distributed in Europe primarily originates from Asia [24, 25]. However, different forms of heroin may be injected subcutaneously or intramuscularly, especially

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**Fig. 2.** PFGE pattern of seven *C. botulinum* strains isolated from six patients during the outbreak and control strains. Lane 12, λ-ladder (50–1000 kb); lane 11, *S. aureus* control strain of the PFGE kit; lanes 1, 10, *C. botulinum* control strains (lane 1, neurotoxin A gene-carrying strain; lane 10, neurotoxin B gene-carrying strain); lane 3, *C. sporogenes*; lanes 2, 4–9, *C. botulinum* strains from the six patients.
by IDU experiencing difficulties in injecting intravenously. Acidifying agents required to dissolve the heroin can cause tissue damage and thus create favourable conditions for the growth of anaerobic bacteria.

All of the IDU in this outbreak either reported subcutaneous or intramuscular injection of heroin or showed clinical signs of this practice. Five survey respondents mentioned using heroin which appeared to them to be of an unusually dark colour. However, this information could not be verified and therefore does not allow us to conclude whether this heroin was actually BTH or not.

Parenteral drug use is well known for carrying the risk of transmission of several infectious diseases [26–29]. Wound botulism is of special importance for two reasons: first, the clinical course is often severe and life-threatening, and second, the rarity of the disease renders diagnosis difficult. Furthermore, in IDU the disease is often misdiagnosed as drug-related complications, a problem which also occurred in this outbreak. The correct and timely diagnosis of botulism in IDU is often complicated by multiple health problems due to malnutrition, poor hygiene and underlying co-infections. In addition to the harmful effect on the health of those concerned, the need for intensive care and treatment costs also have repercussions on the health-care system. A greater awareness of this disease is needed by physicians, as early diagnosis and prompt therapy are vital for the reduction of both the risk for death and duration of hospitalization [30]. If IDU choose to inject drugs into muscle or skin they should be made aware of the dangers of wound botulism in order to help detect the disease early on.

Illicit drug use remains an important issue for public health authorities in the future [31, 32]. Close cooperation between public health authorities, street workers, operators of sheltered injecting facilities, and medical centres focusing on IDU is essential to establish a network able to spread information about newly emerging health threats swiftly and to establish hazard reduction measures quickly in this high-risk group for infectious diseases.

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DECLARATION OF INTEREST

None.

REFERENCES


