Staphylococcal food poisoning from cream-filled cake in a metropolitan area of South-Eastern Brazil

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PEREIRA, M. L. et al. Staphylococcal food poisoning from cream-filled cake in a metropolitan area of South-Eastern Brazil. Rev. Saúde Pública, 28: 406-9, 1994. Twelve people became ill with vomiting and diarrhoea approximately four hours after eating cake with a cream filling at a birthday party and on the day following. The cake had been prepared by a food handler who had long experience in preparing foods for such functions. Staphylococcus aureus that produced enterotoxin A was isolated from the nose, the fingernails, and a healed infection on the neck of the food handler, and from the cake. Enterotoxin A was detected in the remaining portion of the cake. The cake, while still warm, had been refrigerated for one hour after it was prepared before it was removed for the party; it was refrigerated after the party. The cake was large (6 kg) and hence it was not adequately cooled in the hour during which it was refrigerated before the party. The conclusion is that the cake was accidentally contaminated by the food handler and inadequately cooled before it was eaten.

Keywords: Staphylococcal food poisoning, epidemiology. Food handling. Enterotoxins, analysis.

Introduction

Staphylococcal food poisoning is much more common in Brazil than is recognized. Most cases are never reported or investigated. One paper has been published recently on several outbreaks occurring in S. Paulo (Cerqueira Campos et al., 1993). Several outbreaks have been investigated in the city of Curitiba, Paraná State (Bergdoll et al., 1992), but the major emphasis on investigating outbreaks has been in the city of Belo Horizonte, Minas Gerais State (Carmo and Bergdoll, 1990; Carmo et al., 1995). This has resulted in the establishment of a Staphylococci laboratory at the "Fundação Ezequiel Dias". This laboratory is primarily a research laboratory and with the help of graduate students has investigated problems related to the outbreaks in Minas Gerais State. Research was done on the involvement of cream-filled cakes (Anunciação et al., 1994) and a white cheese (Anunciação et al., 1994) characteristic of food poisoning in Minas Gerais State. These two foods have been the cause of almost all of the outbreaks reported in Minas Gerais State (Carmo and Bergdoll, 1990). One problem with the outbreaks is that they are seldom investigated as to the source of the causative organism. Most staphylococcal food poisoning outbreaks are a result of contamination by humans as they are common carriers of enterotoxigenic staphylococci.

The present work covers the investigation of a typical staphylococcal food poisoning outbreak in Minas Gerais State resulting from the consumption of a cream-filled cake at a birthday party and on the following day.

Description of outbreak

Three children and four adults became ill with vomiting and diarrhoea four and one-half hours after eating cream-filled cake at a birthday party. The following day four adults and one child became ill with vomiting and diarrhoea after eating some of the left-over cake which had been kept in the refrigerator. The child was hospitalized. All who became ill had eaten cake. Six adults who ate cake at the party did not become ill, whereas all of those who ate cake the following day became ill. Recovery occurred within 24h. The symptoms are typical of those observed in staphylococcal food poisoning.

The individual who made the cake (6 kg) placed the cream filling in the cake one hour before the party and put the cake in the refrigerator while it was still warm. The cream filling was prepared while the cake was being baked. The
cake was removed from the refrigerator for the party and was consumed two hours after the party began. It is doubtful that the cake was cooled sufficiently to stop the growth of the staphylococci, particularly in the inside of the cake, because of its an extra large size. At the time of the preparation of the cake, the individual preparing the cake had an infection on her neck, otherwise she was in good health. She had been preparing food for other occasions for several years without any problems.

**Material and Method**

**Bacterial cultures**

Swabs were taken from the neck, nares, throat, and fingernails of the food handler four days after the party.

**Isolation of staphylococci**

Each swab was placed in a tube of tryptic soy broth containing 10% NaCl and incubated for 24h at 37°C. The cultures were streaked on Baird-Parker plates and incubated for 48 hours at 37°C. Five typical colonies (jet black to dark grey, smooth, convex, entire margins, off-white edge, and may show an opaque zone and/or a clear halo beyond the opaque zone) and three atypical colonies (gray and mucoid) were selected for further testing. Each colony was transferred to two test tubes containing 1 ml of brain heart infusion (BHI) broth and incubated for 24 hours at 37°C. Tests for coagulase and thermonuclease (TNase) production anaerobic fermentation of glucose and mannitol, and production of hemolysin using sheep blood were made. Any colonies that were positive for these characteristics were considered *S. aureus* and were tested for enterotoxin production.

**Determination of staphylococcal count**

Twenty-five grams of the cake filling was suspended in 225 ml of buffered peptone water; 0.1 ml was placed on Baird-Parker agar plates. Additional plates were prepared with 10-fold dilutions if necessary (Tatini et al., 1984). The plates were incubated for 48 hours at 37°C.

**Enterotoxin production**

For enterotoxin production, inocula were prepared by combining the five isolates from each site and incubating them in BHI broth overnight at 37°C. Membrane-over-agar plates were prepared with 25 ml of BHI-agar and covered with a membrane disk made from Spectra/Por membrane dialysis tubing, 6000-8000, 100 mm flat width (Thomas Scientific, Philadelphia, PA, USA) (Robbins et al., 1974). One-half milliliter of the inoculum was spread on the membrane and the plates were incubated at 37°C for 24h. The cultures were removed from the membranes by washing with 2.5 ml of 0.01 M Na2HP04 in three steps using 1 ml, 1 ml, and 0.5 ml of the phosphate buffer. The cultures were centrifuged and the culture supernatant fluids used for enterotoxin testing.

**Enterotoxin testing**

The optimum-sensitivity-plate (OSP) method was used (Robbins et al., 1974). In this method, 3 ml of agar (1.2%) is placed in 50 mm plastic petri plates with tight lids: wells are cut according to the original specifications. Specific antisera is placed in the center well, enterotoxin (4μg/ml) is placed in the two smaller wells, and culture supernatant fluids are placed in the four larger outer wells. Different plates are required for each enterotoxin (SEA, SEB, SEC, SED). The plates are placed in a humidified container and incubated overnight at 37°C. Positive reactions are determined from precipitin lines formed by the culture supernatant fluids that joined with the control lines.

**Enterotoxin detection in the cake**

Fifty grams of the cake containing the cream filling was homogenized in a Waring blender with 1 ml of 0.02 M NaHP04 in saline, pH 7.4, per gram of food (Freed et al., 1982). The pH of the supernatant fluid was readjusted to pH 7.4. The extract was tested for the presence of enterotoxin by use of the ELISA ball kit* (Fey and Pfister, 1983). This method is sensitive to 0.5 ng/ml and is the most reliable of the sensitive methods available for checking foods for enterotoxin (Wieneke 11, 1987, 1991). One antibody-coated ball for each enterotoxin (A-D) plus one ball coated with normal rabbit sera were placed in 20 ml of food extract and shaken gently overnight. Each ball was removed from the extract and washed with the wash solution and each placed in a color coded tube for treatment with

* Obtained from Dr. Bomelli AG, Stationstrasse 12, CH-3097 Libbeful-Bern, Switzerland
the conjugate. After 6h the conjugate was removed and the balls washed with the wash solution. One ml of the substrate was added and the color allowed to develop for 45 min. If any color developed, the extract was judged to contain the enterotoxins for which a color developed.

**Results**

Each of the five isolates from the food handler's throat, nasal passages, fingernails, and infection and from the cake was coagulase and TNase positive, fermented glucose and mannitol anaerobically, and were hemolytic. It was concluded that the isolates were *Staphylococcus aureus* because it is the only species carried by humans that is both coagulase- and TNase-positive and ferments mannitol anaerobically.

The staphylococcal count was $1.2 \times 10^8$ cfu/g of cake.

Examination by the OSP method showed isolates from all of the culture sites, except the isolates from the throat, and the ones from the cake produced enterotoxin A (SEA) (Table).

Enterotoxin A (SEA) was detected in the cake implicated in the food poisoning outbreak (Table).

**Discussion**

This outbreak was a typical staphylococcal food poisoning outbreak resulting from the eating of a food served at a function involving a number of people. Although the twelve who fell ill was not a large number, all had eaten the same food, a cream-filled cake. Six people who ate the cake at the party did not become ill, indicating that those eating the cake on the day it had been made did not receive as much toxin as those who ate the cake on the following day. Although the cake had been refrigerated after the party, the staphylococci would continue to grow in the cake and produce enterotoxin until adequate cooling, took place. The cream filling was placed in the cake while the cake was still warm and the cake was refrigerated for one hour before the party. It is unlikely that the cake cooled adequately, especially because of its size, to prevent the staphylococci from continuing to grow until it was consumed. However, the amount of enterotoxin in the cake at the time of the party was probably small as it required over 4h after the cake was eaten before the symptoms developed, considering that symptoms may develop between one and 6 hours after consumption of food containing enterotoxin. Although refrigeration is the recommended method for preventing staphylococcal food poisoning, if food is exposed to room or warmer temperatures for 2 to 3 hours without refrigeration, enough growth and enterotoxin production could take place to result in food poisoning. Another concern is the size of the food item refrigerated. In this case the cake was extra large which made it doubt ful that the interior of the cake could have cooled sufficiently to prevent the staphylococci to continue growing. If possible food should be refrigerated in small quantities.

The majority of staphylococcal food poisoning outbreaks result from the contamination of the food during its preparation by the food handler. Humans are common carriers of enterotoxigenic staphylococci in the nose and throat, or on the skin. In addition many infections can be caused by staphylococci, which is a more dangerous situation because they are present in large numbers and can easily be transferred to the food being handled. Many staphylococcal food poisoning outbreaks have resulted from foods prepared by food handlers with infections. It is recommended that foods being prepared for groups should not be prepared by a food handler with any type of infection. The wearing of surgeon’s gloves during the preparation of the food would add insurance against contamination of the food, providing the food handler avoided touching any other part of the body with the hands during the preparation of the food.

**Table. Biological characteristics and enterotoxin production of isolates from staphylococcal cake poisoning.**

<table>
<thead>
<tr>
<th>Isolates</th>
<th>Coag.</th>
<th>TNase</th>
<th>Mann.(an)</th>
<th>Glu.(an)</th>
<th>Hemol.</th>
<th>Enterotoxin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nose*</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td></td>
<td>SEA</td>
</tr>
<tr>
<td>Throat*</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td></td>
<td>Neg</td>
</tr>
<tr>
<td>Neck*</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>SEA</td>
</tr>
<tr>
<td>Fingernail*</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>SEA</td>
</tr>
<tr>
<td>Cake†</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>SEA</td>
</tr>
</tbody>
</table>

* - From food handler who prepared the cake
† - $1.2 \times 10^8$ cfu/g of cake

PEREIRA, M.L. et al. Intoxicação estafilocócica causada por bolo recheado em área metropolitana do sudeste do Brasil. Rev. Saúde Pública, 28:406-9, 1994. Doze pessoas foram acometidas de vômito e diarréia aproximadamente 4 horas após haverem ingerido bolo recheado, servido em uma festa de aniversário, e no dia seguinte à festa. *Staphylococcus aureus* produtor de enterotoxina A foi isolado no bolo, fossa nasal, leito subungueal e, essencialmente, em uma ferida em fase de cicatrização, localizada na nuca da manipuladora, que dispunha de longa experiência na área de produção de alimentos. O bolo, de cerca de 6 kg, quando ainda quente, foi levado ao refrigerador, por uma hora antes de ser servido não tendo...
portanto, sido convenientemente resfriado. Esses dados permitiram concluir que o referido alimento foi, acidentalmente, contaminado pela manipuladora e inadequadamente resfriado antes de ser ingerido.


References


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