A case of vertical transmission of Chagas disease contracted via blood transfusion in Canada

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CASE REPORT

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Chagas disease is caused by the protozoan parasite Trypanosoma cruzi and is endemic in many countries in Latin America, where infected bugs of the Triatominae subfamily carry the parasite in the gut and transmit it to humans through fecal contamination of a bite. However, vertical transmission and transmission through blood transfusion and organ transplantation is well documented. Increasing immigration from endemic countries to North America has prompted blood operators, including Canadian Blood Services and Hema Quebec, to initiate blood donor testing for Chagas antibody. In the present report, an unusual case of vertical transmission from a mother, most likely infected through blood transfusion, and detected as part of a concurrent seroprevalence study in blood donors is described.

Key Words: Chagas disease; Transfusion transmission; Vertical transmission

Chagas disease is a zoonotic disease caused by the protozoan parasite Trypanosoma cruzi and is spread by Triatominae or Reduviidae vectors. It is endemic to Central and South America and Mexico, where an estimated 10 million people have been infected (1). Chagas disease can also be spread by blood transfusion, organ transplantation and through vertical transmission from mother to fetus. Cases of oral transmission via contaminated fruit, fruit and sugar cane juice, and water have also been described in endemic areas (2). In the acute stage of disease, approximately 5% of those infected will experience flu-like symptoms, including fatigue, loss of appetite and fever. The first characteristic sign may be a ‘chagoma’, an erythematous, indurated lesion that occurs at the site of entry of the parasite, which enters the bite site via the contaminated feces of the insect. Sometimes the conjunctiva is the site of entry, and the classic painless periorbital swelling known as Romana’s sign may develop (3). Acute illness more commonly occurs in children and elderly individuals in endemic areas. Treatment of acute infection with nifurtimox or benznidazole can alleviate symptoms, clear the parasite and moderate the risk of future sequelae (4).

Approximately 40% of individuals who become infected remain parasitic but asymptomatic, and 20% to 30% progress to develop chronic infection, which results when the parasite invades the smooth muscle (2). This may result in cardiomyopathy with arrhythmias and cardiac failure, megacolon or megaesophagus (3). The treatment of chronic Chagas disease remains controversial but is believed to be of benefit in patients younger than 50 years of age (5). With increasing immigration of individuals from endemic countries to North America, it is estimated that >300,000 individuals infected with T. cruzi live in the United States (US) (7).

Several blood donor seroprevalence studies conducted in the US in the early 1990s and in 1997, showed that in some areas, such as Los Angeles (California), New Mexico and Texas, >1.4% of donors were seropositive (8,9). With the US Food and Drug Administration licensure of an assay for blood donor screening, universal donor testing was implemented by most large US blood suppliers in early 2007. Many US blood suppliers have subsequently switched to a ‘test every donor once’ strategy. At present, >1600 donors have been confirmed positive for Chagas disease (10). In Canada, although the rates of immigration from endemic countries are less than that of the US, of the seven reported cases of transfusion-transmitted Chagas disease in North America before donor testing, two were reported in Canada in the province of Manitoba in 1989 and 2000 (11,12). A small seroprevalence study performed in asymptomatic immigrants from endemic areas in Toronto found one positive individual of 102 tested (13). In Canada, Canadian Blood Services (CBS) provides blood products to the entire country with the exception of Quebec, and collects approximately one million donations per year. CBS implemented selective Chagas disease donor testing based on risk (identified on the donor questionnaire) in 2010.

In conjunction with the selective donor testing program, CBS also set up a donor seroprevalence survey, in which a random selection of donors who expressed no risk (answered ‘no’ to the three risk questions on the donor questionnaire) for Chagas disease, lived mainly in urban areas and areas where Chagas disease had been previously identified in donors in the past were tested. To achieve significance with a 95% CI, assuming no positives were found, it was calculated that 60,000 donors would need to be tested. Subsequently, this number was increased to approximately 100,000 to be tested by study completion, which

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represents 10% of the donor population. At present, >27,000 'at risk' donors have been tested with 14 cases of confirmed Chagas disease being identified and >74,000 'no risk' donors have been tested in the study with one donor being identified with confirmed Chagas disease, who is described in the present report.

CASE PRESENTATION

In January 2011, a blood donor involved in the seroprevalence study tested repeat reactive in the Abbott PRISM Chagas assay for Chagas antibody (Abbott Laboratories, Germany). A specimen was sent to the National Reference Centre for Parasitology at McGill University in Montreal, Quebec, for confirmatory testing. This testing included additional inhouse ELISA testing, immunoblot and polymerase chain reaction (PCR). In addition, the specimen was sent to the Blood Systems Laboratories (Scottsdale, Arizona, USA) for testing using the Ortho Trypanosoma cruzi ELISA test (Ortho Clinical Diagnostics, USA) and to Quest Diagnostics (Virginia, USA) for the radioimmunoprecipitation assay, which is routinely used for screening and confirmatory testing by US blood operators. The results of these tests are shown in Table 1.

With confirmation of the presence of Chagas antibody, the donor was contacted and informed of the results, and subsequently completed a detailed risk questionnaire. The results of this testing were communicated to her physician, who referred the donor to an infectious diseases specialist for further testing and follow-up. The donor questionnaire contained detailed questions regarding the risk for Chagas disease, including birth or residence in an endemic country, birth or residence of a mother or maternal grandmother in an endemic country (to identify vertical transmission risk), long-term travel in an endemic country (>6 months) and history of previous blood transfusion. Extensive questioning of this donor revealed no identifiable risk factor for Chagas disease. Several days after the donor was interviewed, she informed CBS that her mother (a very astute nurse) wondered whether she may be the source of her daughter's infection because she had been transfused in Manitoba in 1978 and 1983 – the latter around the time of the donor's birth. The mother was subsequently tested for Chagas disease at the National Reference Centre for Parasitology using the inhouse ELISA, immunoblot, PCR and hemoculture. She was found to be positive for Chagas disease by ELISA and immunoblot; however, the PCR and hemoculture results were both negative. An extensive history revealed no risk factors for Chagas disease other than the blood transfusions received in 1978 and 1983. Unfortunately, because records of donors or recipients of blood transfusions from the 1970s were not retained by the Canadian Red Cross or the hospital or clinic where the transfusion was performed, donors and recipients of co-components could not be traced. For the transfusions in 1983, three of four donors were located and were found to be negative for T cruzi antibody. Incidentally, the donor's sibling, who was born before the transfusions, tested negative for Chagas disease. The donor's son was also screened for Chagas disease and also tested negative.

The donor and her mother were referred to a regional, specialty tropical medicine clinic where they were evaluated for signs and symptoms of chronic Chagas disease. Both patients denied any history or symptoms of intestinal obstruction or constipation. The donor reported a long history of gastroesophageal reflux, which was controlled with a proton pump inhibitor. Neither patient reported symptoms of heart failure or arrhythmia, including peripheral or pulmonary edema, palpitations, syncope or chest pain. Physical examinations and routine biochemistry, hematology, and renal and hepatic function tests were normal. Investigations for infection with T cruzi were described above. Both patients underwent electrocardiography, echocardiography and 24 h Holter monitoring that revealed no cardiac or conductive abnormalities. Examination of barium swallows revealed normal esophageal function. It was recommended that the donor undergo treatment to reduce the risk of progression to cardiac disease. The donor's mother was not offered treatment due to her age.

The donor underwent 60 days of nifurtimox therapy (8 mg/kg/day), as recommended by the WHO. During therapy, she experienced daily nausea and vomiting and developed mild leucopenia. Her leukocyte values returned to normal after completion of treatment and the nausea also resolved.

DISCUSSION

Most North American physicians do not have experience in the management of chronic Chagas disease or the potential adverse effects of drug therapy. While all patients with acute and subacute Chagas disease should be treated, the role of treatment in chronically infected patients remains somewhat controversial. Patients with advanced age or advanced cardiac disease are unlikely to benefit from treatment (5,14,15). Although most experts would agree that patients younger than 50 years of age without evidence of cardiac disease should be treated, access to the drugs required for treatment is limited, and the drugs have numerous, some being serious, potential side effects (5). In Canada, until recently, only nifurtimox was available, and only through the Health Canada Special Access Programme. Benznidazole, widely shown to be better tolerated and to have fewer side effects (5,15) was out of production until early 2012. The management of Chagas disease also requires lifelong evaluation for its most common complications, which include mainly cardiac conduction anomalies, megaesophagus and megacolon. At the very least, electrocardiography should be performed yearly, regardless of the therapeutic regimen (14).

The present article describes the first reported case in Canada of vertical transmission of Chagas disease acquired through blood transfusion. Cases of vertical transmission in endemic countries are estimated to occur at rates of 0.13% to 19%, depending on the country (16,17). In the US, based on the prevalence rates in the country of origin of the mother and a 5% risk of vertical transmission, it has been estimated that there are 166 to 638 infected newborns (18).

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REFERENCES


### Table 1

<table>
<thead>
<tr>
<th>Test</th>
<th>Result</th>
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<tbody>
<tr>
<td>ELISA (inhouse, TESA antigens)</td>
<td>Positive</td>
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<tr>
<td>Immunoblot (inhouse, TESA antigens)</td>
<td>Positive</td>
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<tr>
<td>Inhouse polymerase chain reaction assay</td>
<td>Negative</td>
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<tr>
<td>Trypanosoma cruzi primers*</td>
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</tr>
<tr>
<td>Ortho Trypanosoma cruzi ELISA†</td>
<td>Positive</td>
</tr>
<tr>
<td>Radioimmunoprecipitation assay‡</td>
<td>Positive</td>
</tr>
</tbody>
</table>

† Trypanosoma cruzi infection of squirrel monkeys: comparison of blood smear examination, commercial ELISA and polymerase chain reaction analysis as screening tests for evaluation of monkey-related injuries (18). 1 Blood Systems Laboratories (Scottsdale, Arizona, USA), 2 Quest Diagnostics (Virginia, USA). TESA Trypomastigote excreted/secreted antigens.


