Case Report

Variant of X-Linked Chronic Granulomatous Disease Revealed by a Severe Burkholderia cepacia Invasive Infection in an Infant

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Chronic granulomatous disease (CGD) is a primary immunodeficiency characterized by increased susceptibility to bacteria and fungi since early in life, caused by mutations in any of the five genes coding for protein subunits in NADPH oxidase. X-linked variant CGD can be missed during routine evaluation or present later in life due to hypomorphic mutations and a residual superoxide production. The case of a 10-month-old boy who died of pneumonia is reported. The isolation of Burkholderia cepacia from his lung, together with a marginally low nitroblue tetrazolium reduction assay (NBT), made us suspect and pursue the molecular diagnosis of CGD. A postmortem genetic analysis finally demonstrated CGD caused by a hypomorphic missense mutation with normal gp91phox expression. In a patient being investigated for unusually severe or recurrent infection, a high index of suspicion of immunodeficiency must be maintained.

1. Introduction

Chronic granulomatous disease (CGD) is a rare primary immunodeficiency that affects microbial killing by phagocytes, resulting in bacterial, fungal, and/or mycobacterial infections since early life [1, 2]. The superoxide production by NADPH oxidase is markedly reduced or absent due to mutations in any of the five genes coding for protein subunits of the enzymatic complex [3]. Mutations in CYBB, coding for gp91phox, result in the most common X-linked CGD (65%–70% of all cases) [4]. Hypomorphic mutations (Xgp91+ and Xgp91−) may result in X-linked variant CGD [5, 6]. Patients with variant CGD express the gp91phox protein and produce decreased but detectable superoxide, which allow the defect to manifest later in life with a milder history of infections. By far, the most common micro-organisms causing infections in CGD are Staphylococcus aureus and Aspergillus species; other agents include Pseudomonas, Serratia, Salmonella, and Candida species. Burkholderia cepacia infection is frequently associated to CGD diagnosis (6–8). Here, we present the case of a patient who died of Burkholderia cepacia lung infection, in whom the diagnosis of X-CGD could only be attained postmortem due to residual superoxide production and normal protein expression.
2. Case Report

A 10-month-old boy, the first child of nonconsanguineous parents living in the Tahiti archipelago (French Polynesia), was referred for severe pneumonia. The father is from Europe and the mother is from Oceania; there was no relevant family history. During the first months of life, the patient had experienced some infections, mostly of the upper airways, as well as bronchitis and diarrhea. He received all the immunizations according to his age (including BCG) with no adverse events. He developed a failure to thrive at the age of 3 months. One month before admission he had a severe lung infection with fever, cough, dyspnea, and diarrhea, unresponsive to empiric oral macrolide (josamycin). Upon admission to his local hospital, he had fever (39.5°C), mild respiratory distress, and crackles on auscultation. Oxygen saturation was 95% in room air. Complete blood count (CBC) reported marked leukocytosis (36,600/mL) with neutrophilia (29,000 g/dL) and anemia, elevated serum ferritin, and Gram stain reported 1,100 cells (97% PMN) and abundant Gram negative bacteria that grew from the trachea and bronchi. Bronchoalveolar lavage (BAL) and Gram stain reported 1,100 cells (97% PMN) and abundant Gram negative bacteria that grew Burkholderia cepacia (10⁶ CFU/mL). Antibiotherapy was then switched to IV rifamycin and trimethoprim/sulfamethoxazole. Despite intensive supportive care, including broad-spectrum antibiotics and daily granulocyte transfusions, his lung infection worsened, and he finally died of acute respiratory distress and multiorgan dysfunction in the intensive care unit. Permission to perform an autopsy was refused by his parents.

The clinical presentation and the impaired NBT reduction assays of this boy were consistent with a primary phagocyte defect. We assessed superoxide (O₂⁻) production in PMNs from the patient as measured by the cytochrome-c reduction assay, compared to another patient with known X-linked CGD (−) and a healthy control (+), following stimulation with phorbol myristate acetate (PMA). Residual NADPH oxidase activity was detected in the PMNs of the patient (Figure I(a)). In addition, 123-dihydrorhodamine (DHR) oxidation assay by flow cytometry revealed a partial deficiency of ROS production in the patient's PMN, while his mother had two granulocyte populations: one strongly rhodamine-positive (reactive) and the other rhodamine-low fluorescence intensity (Figure I(b)). These results again suggested that our patient had a partial defect in the respiratory burst. We next investigated the H₂O₂ production upon milder activation, involving priming with TNF-α, IL-1β, or cytochalasin B, followed by FMLF (formyl-methionyl-leucyl-phenylalanine) stimulation. PMNs from the patient produced detectable but low H₂O₂ (Figure I(c)). Genomic sequencing of CYBB revealed a hemizygous A > G substitution in exon 9, generating the replacement of a histidine by an arginine residue (H338R) in the FAD binding domain (FADBR), a probably damaging substitution according to the PolyPhen-2 prediction website (http://genetics.bwh.harvard.edu/pph2/). The patient’s mother was heterozygous, and his brother (born after the patient’s death) was hemizygous for the mutation. The mutation was confirmed also in cDNA from the patient (c.1013A > G). We investigated the molecular basis of the germline H338R mutation through detection of flavocytochrome b₅₅₈ expression by flow cytometry, using the monoclonal antibody 7D5 (MLB, Nagasaki, Japan), which recognizes residues¹⁶⁶IKN¹⁶⁸ and ²²⁶RIVRG²³⁰ on gp91phox in the presence of p22phox. Protein expression in Epstein-Barr virus transformed B cells (EBV-B cells) from the patient was similar to the healthy control (Figure 2).

3. Discussion

The isolation of Burkholderia cepacia from lung secretion or blood of a previously healthy patient is strongly suggestive of CGD. Aside from it, lung infections caused by Burkholderia species can be seen in patients with existing bronchiectasis (lung epithelial damage is a prerequisite for Burkholderia invasiveness), including notably patients with cystic fibrosis [7] and in some immunocompromised and hospitalised patients [8, 9]. In a child being investigated for recurrent infections, isolation of Burkholderia should always raise the suspicion of CGD [10–12]. For some patients with normal gp91phox expression and residual superoxide production as measured by conventional assays, a milder activation assay with FMLF might be needed to demonstrate low ROS production.
Figure 1: Continued.
Missense mutations beyond amino acid 309 of gp91phox usually allow normal protein expression but result in null superoxide production. The patient’s residual ROS generation is thus different from the thorough survival analysis by Kuhns et al. [3]. Also, given this infant’s residual superoxide production, a severe course with early demise is surprising.

In conclusion, we identified postmortem a point mutation in a CGD causing gene from a 10-month-old boy who presented with a *Burkholderia* spp. overwhelming lung infection. X-CGD diagnosis was delayed because of initial normal results. A high index of suspicion for CGD must be maintained in patients with *Burkholderia* isolates and close to normal values of usual CGD diagnostic tests such as NBT.
An early and accurate diagnosis can lead to genetic counselling, to family screening, and to a timely intervention.

**Abbreviations**

CGD: Chronic granulomatous disease  
PID: Primary immune deficiency  
NADPH: Nicotinamide adenine dinucleotide phosphate hydrogen  
BCG: Bacillus Calmette-Guérin.

**Conflict of Interests**

The authors declare no conflicts of interest.

**Authors’ Contributions**

Saul Oswaldo Lugo Reyes and Nizar Mahlaoui equally contributed to this work.

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**References**


