Case Report: Pancytopenia as an indicator for lysosomal storage disease (Gaucher’s Disease) [version 2; peer review: 1 approved]

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Abstract

Introduction: Lysosomal storage disorders are a rare group of genetic diseases, with Gaucher Disease (GD) among the most common. GD type 1 is the most common form of this condition, and patients with this variant may present with hematologic unexplained cytopenias, in addition to hepatosplenomegaly, multi-faceted bone involvement, and, late in adult life, with other cases neurological disorders such as Parkinsonism and peripheral neuropathy. A case of a Because cases of Gaucher disease are uncommon in Ecuador where information about GD is very limited, we present an illustrative patient with thrombocytopenia, leukopenia, and hepatosplenomegaly. A patient is presented, whose results showed thrombocytopenia and leukopenia in addition to hepatosplenomegaly whose diagnosis was not suspected until performance of a bone marrow biopsy.

In Ecuador, there are very few reported cases of this clinical entity, and information on this disease is very limited. Our aim is to emphasize the importance of early recognition of pathology suggestive of GD type 1 so that this disease is routinely included in the differential diagnosis in patients with unexplained cytopenias and hepatosplenomegaly. Once suspected, the diagnosis may be established non-invasively by testing for lysosomal glucocerebrosidase (acid β-glucosidase) deficiency either in dried blood spots (DBS) or in peripheral blood leukocytes and further elucidated by sequencing the GBA gene for pathogenic variations (mutations). Timely, early diagnosis should be followed by appropriate treatment (either intravenous enzyme replacement [ERT] or oral substrate synthesis inhibition ["SRT"] directed at preventing irreversible complications and morbidity and improving health-related quality of life.

Case: We present a case of a patient diagnosed with GD, who presented with thrombocytopenia and leukopenia in addition to hepatosplenomegaly, with the aim of emphasizing the importance of early recognition of this pathology, especially in patients with unexplained cytopenia’s or hepatosplenomegaly’s. In suspicion of GD, enzymatic quantification of β-glucocerebrosidase was performed, showing its deficit in addition to alteration in the GBA gene.

Conclusion: We believe that a high index of suspicion together with enzymatic and genetic testing are essential for an early specific diagnosis, which will allow the administration of timely treatment and avoid irreversible complications in order to reduce morbidity and improve the clinical aspects of the patient.

Keywords

Gaucher disease, lysosomal storage disorders, cytopenia, β-glucocerebrosidase.

Revised Amendments from Version 2 (NJW)
Introduction

Lysosomal storage disorders (LSDs) are rare, generally recessive, hereditary monogenic diseases. Because of total or partial functional loss of specific lysosomal enzymes or their cofactors, undegraded and sometimes water-insoluble substrates accumulate in cellular lysosomes of various tissues causing multi-organellar and cellular dysfunctions that extend to immediate cellular and extracellular microenvironments. These processes lead to immune and inflammatory pathophysiology that is expressed as signs and symptoms of clinical disease.¹²

Gaucher disease (GD) is one of the most common LSDs with an estimated incidence of 1:40,000 to 1:86,000 new births in the general world population. There is no known racial or national proclivity for GD with the exception of Jews who trace their last 1,000 year ethnicity to regions of eastern, northern and central Europe referred to in Jewish tradition as Ashkenaz. Most Ashkenazi Jews currently reside in Israel, Western Europe, Russia, North America and a number of Latin American countries. The incidence of GD among Jews with four Ashkenazi grandparents is estimated to be 1:450 births although the recognized prevalence may be only 1:850 due to low penetrance genotypes.³ In 1950, there were approximately 5,000 Jews in Ecuador of mixed ethnicity, with no relevant family or personal history, except a history of cancer in the family? How long is the family living in Loja?

GD is characterized by defective function of lysosomal glucocerebrosidase that leads to accumulation of glucocerebroside in the lysosomes of different cells, but primarily in tissue macrophages resident in bone marrow, spleen and liver.⁴ The sphingolipid storage process, that includes minor but pathogenic substrates such as glucosylsphingosine, ultimately leads to pancytopenia, hepatosplenomegaly, bone involvement (including loss of bone mineral, medullary infarction, osteonecrosis, osteolysis and pathologic fractures) and, sometimes neurological alteration early in life.⁵ There are three major clinical subtypes, which are classified by the absence or presence and progression of central neurological involvement: type 1 or non-neuropathic form; type 2, the acute neuropathic form of infantile onset; and type 3, the neuropathic form of juvenile onset.⁶ However, there is phenotypic heterogeneity within each subtype, partly but not totally determined by genotype. In symptomatic patients, GD is usually a progressive disease that, if left untreated, can cause irreversible organ damage, severe morbidity, reduced quality of life and even premature death.⁷

GD is a rare disorder with which many physicians, including hematologists, are unfamiliar. As a

Description of the case

A 29-year-old housewife from Loja-Ecuador, of mixed ethnicity, with no relevant family or personal history, was evaluated at the outpatient service of the Isidro Ayora General Hospital, Loja, Ecuador. She presented with a pulsating holocranial headache of slight intensity. Concomitantly, she related a history of heartburn which responded positively to self-medication with omeprazole; additionally, she mentioned she was experiencing progressive loss of weight, fatigue and pain in the lower limbs and hands. Physical examination showed ecchymosis and petechiae scattered...
throughout the body. The liver and spleen were palpably enlarged. Routine laboratory results are summed up in Table 1.

Table 1. Laboratory results of patient’s parameters and their normal range.

<table>
<thead>
<tr>
<th>Patient’s parameters</th>
<th>Normal range</th>
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<tbody>
<tr>
<td>Leukocytes (White blood cell)</td>
<td>3400 mm³</td>
</tr>
<tr>
<td>Neutrophils</td>
<td>1.9 K/μL</td>
</tr>
<tr>
<td>Lymphocytes</td>
<td>1.5 K/μL</td>
</tr>
<tr>
<td>Red blood cell</td>
<td>4.3 K/μL</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>12.5 g/dL</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>38%</td>
</tr>
<tr>
<td>Platelet</td>
<td>59 K/μL</td>
</tr>
<tr>
<td>PCR (C-Reactive Protein (CRP))</td>
<td>16.1 mg/L</td>
</tr>
<tr>
<td>VSG (erythrocyte sedimentation rate (ESR))</td>
<td>33 mm/h</td>
</tr>
<tr>
<td>Total bilirubin</td>
<td>3 mg/dL</td>
</tr>
<tr>
<td>BD (Direct bilirubin)</td>
<td>0.48 mg/dL</td>
</tr>
<tr>
<td>BI (Indirect bilirubin)</td>
<td>2.52 mg/dL</td>
</tr>
</tbody>
</table>

Viral serology tests for hepatitis B, C and human immunodeficiency virus were negative. An ultrasound of the upper abdomen was performed, where there was evidence of splenomegaly and an enlarged liver
without other abnormalities [Figure 1]. Hepatosplenomegaly was observed in the CT-Scan in addition to degenerative osteoarthritis of the dorsal spine.

A bone marrow biopsy demonstrated abundant cells between the bone trabeculae, consisting of hematopoietic tissue, with a marked decrease in megakaryocytes of hypolobular nuclei, and myeloid-erythroid relationship conserved. In the inter and paratrabecular spaces, many clusters of large macrophages with small, regular nuclei displaced to the periphery and “crumpled paper” cytoplasm were observed as found in classical Gaucher cells [Figure 2]. Now, suspecting GD, measurement of WBC glucocerebrosidase was performed that showed deficient activity (0.27 µmol/L/h (normal range: 2.3 – 12 nmol/h/mL). Sequential analysis of the GBA1 gene showed the presence of an apparently homozygous pathogenic alteration in the GBA1 gene (Nucleotide change: c.1226A>G; Amino acid change: G p. Asn409Ser).

Commented [TW2]: As Dr Sidransky had previously asked and suggested, what diagnoses were considered at this point that prompted the performance of the bone marrow biopsy? ITP? How did you explain the elevated total and indirect bilirubin? For example, was hereditary spherocytosis considered? Gilbert’s? Were there any gall stones on either the US or CT scans? The decrease in megakaryocytes is somewhat unusual for GD.
Figure 2. Bone marrow biopsy: Cuts show bony trabeculae, including extensive medullary replacement by histiocytic reticulum-like cells, with eosinophilic, pale cytoplasm and eccentric nuclei, some cells have a vacuolated cytoplasm with folded appearance on crumpled paper.

Unfortunately, enzymatic replacement or substrate reduction therapy are not available at our hospital. At the time of writing the patient is regularly monitored until ERT can be obtained. The patient remains in good general condition, without worsening signs of symptoms. It should be mentioned that although she continues to have thrombocytopenia, leukopenia and hepatosplenomegaly, her condition is expected to remit once the enzymatic treatment has been initiated.

Discussion

This clinical case is representative of many clinical, biochemical and genetic characteristics of GD type 1, a disorder that is characterized by variability in its signs, symptoms, severity and progression. The most common signs and symptoms are: splenomegaly, hepatomegaly, radiological bone disease, thrombocytopenia, anemia, and bone pain.

GD is caused by mutations in the GBA1 gene, located on chromosome 1 (1q21), leading to markedly decreased activity of the lysosomal enzyme, β-glucocerebrosidase, which hydrolyzes the glycolipid.
glucocerebrosidase activity to develop classical manifestations of GD. Our patient had sufficiently decreased glucocerebrosidase activity to develop classical manifestations of GD. However, the identified genotype (N409S/N409S) was unexpected as this DNA variant is most often found among Ashkenazi Jews and is uncommon in the non-Jewish Latin American populations. For example, although the N409S allele was the most prevalent mutation found among 25 GD1 patients from Colombia (which borders Ecuador and which has a current Jewish population of approximately 13,000 of whom many are Ashkenazim who immigrated after World War 2), only two (8%) were homozygous for this variant, one Mestizo and one of unknown ethnicity. Although comparable data for Ecuador are not published, it is therefore improbable that these very rare N409S/N409S GD1 patients in Columbia and Ecuador (ours being the index case) are associated with the 20th century Ashkenazi immigrants.

Rather, it is more likely that the descent of these patients can be traced to the time of the original Spanish conquests in Latin America. Haplotype analysis suggests that the original N409S allele arose in non-Jewish Europeans, was subsequently introduced into the Iberian Jewish (Sephardi) population sometime in the middle ages pre-dating the appearance of N409S in the Ashkenazi Jews. Due to internal and external prohibition of intermarriage and sporadic bottleneck events, genetic diversity decreased in the Iberian Jewish population and N409S frequency increased. The prevalence of N409S homozygous GD1 patients in the modern Spanish Gaucher Registry population (N=178) is 14.6% and likely reflects the eventual integration of large numbers of Iberian Jews who converted to Catholicism (conversos) into the general Spanish population. However, the inexorable persecution the Holy Office of the Inquisition caused great peril to the New Christians. Many illegally emigrated to the Spanish Portuguese new world colonies where, for a time, the power of the Inquisition was less pronounced. It is likely that the N409S allele was introduced to Latin America by these Spanish-Portuguese emigrants of the 16th century, some of whom eventually settled in Loja and its vicinity in what is now modern Ecuador. It is conceivable that our patient is a descendant of converso settlers of Loja. This hypothesis could be further examined by testing for the presence of a GBA1 intronic polymorphism, 5470G→A that is present only in N409S alleles of Iberian origin.

Glucocerebrosidase deficiency causes accumulation of glucosylceramide in macrophages, transforming them to Gaucher cells that under optical microscopy are conspicuous as enlarged cells with eccentric nuclei and condensed chromatin, and cytoplasm with an appearance as “crumpled paper or napkin.” These cells were abundant in the bone marrow biopsy specimen obtained from our patient. The presence of apparent Gaucher cells (pseudo-Gaucher cells) has also been described in the absence of GD in patients with various hematologic malignancies, hemolytic anemias such as thalassemia, and mycobacterial infections. Thus, the Gaucher cell is not pathognomonic, and only demonstration of absent or reduced glucocerebrosidase activity is the gold standard for the diagnosis of all GD variants.

The usual signs and symptoms associated with GD are shared many other more common illnesses including chronic liver disease, hematologic malignancies, and, in relevant parts of the world with infectious diseases such as malaria, leishmaniasis, and mycobacterioses. Failure to include GD in the differential diagnosis of a patient with classic manifestations can lead to unnecessary invasive diagnostic procedures, misdiagnosis resulting in inappropriate treatment such as splenectomy, and to delayed initiation of effective therapy causing largely preventable irreversible complications and permanent disabilities especially from skeletal disease. Therefore, it is essential to continue to disseminate knowledge about GD including its worldwide prevalence, methodologies for diagnosis, and current and evolving guidelines for management and therapy.
Conclusion

Despite its relative rarity, GD should always be considered in the differential diagnosis of patients with unexplained pancytopenia and/or hepatosplenomegaly. The ease of obtaining and transporting stable dried blood spot samples for diagnosis makes it feasible to detect GD and many other rare LSDs even in remote and medically deprived regions of the world. A high index of suspicion together with enzymatic and genetic testing are essential for an early specific diagnosis followed by the administration of timely treatment, thus minimizing the risk for irreversible complications and chronic morbidity that detract from health-related quality of life and even shorten life expectancy.

Consent

Written informed consent for publication of their clinical details and clinical images was obtained from the patient.

Data availability

Underlying data

All data underlying the results are available as part of the article and no additional source data are required.

References


