

Monocytosis in a patient with a novel GATA2 mutation

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ABSTRACT

GATA2-associated disorders include: (*i*) monocytopenia with mycobacterial infections (MonoMAC); (*ii*) dendritic cell, monocyte, B and NK lymphoid deficiency; (*iii*) familial myelodysplastic syndrome (MDS) and acute myeloid leukemia; and (*iv*) congenital deafness with lower limb lymphedema deficiency (Emberger syndrome). Markedly reduced or absent monocytes have been considered as the hallmark of the disease. Here we report on a patient that presented in infancy with hearing loss and lymphedema. By 4 years of age the patient developed acne, disseminated warts, lymphadenopathy, and MDS, yet with increased monocyte as well as normal NK- and B-cell numbers. The patient was found to have a novel mutation in *GATA2* that was predicted to disrupt the C-terminal zinc finger. Importantly, and in contrast to common concepts, *GATA2*-associated syndromes might present with monocytosis.

Statement of novelty: We describe a novel mutation in GATA2 associated with monocytosis.

Introduction

The *GATA2* gene, located at chromosome 3q21.3, encodes a dual zinc-finger transcription factor, known as GATA binding protein 2 that binds to the promoter and regulatory regions of target genes in the nucleus. *GATA2* regulates transcription of genes involved in the development and proliferation of diverse cell lineages, including hematopoietic, endothelial, placental, neuronal, etc. (Bresnick et al. 2010). The impaired function of hematopoietic stem cells from *GATA2*^{+/-} mice (Rodrigues et al. 2005) and the association of hematological malignancies with translocation of *GATA2* enhancer elements (Gröschel et al. 2014) suggest a dose-dependent effect for *GATA2* on hematopoiesis. Indeed, heterozygous *GATA2* mutations were recently identified among patients with diverse hematological and nonhematological manifestations. These conditions have been characterized as: (i) monocytopenia and mycobacterial infections (MonoMAC) syndrome (Hsu et al. 2011); (ii) dendritic cell, monocyte, B and NK lymphoid (DCML) deficiency (Dickinson et al. 2011); (iii) familial myelodysplastic syndrome (MDS) and acute myeloid leukemia (AML) (Hahn et al. 2011); and (iv) primary lymphedema, congenital deafness, and MDS (Emberger syndrome) (Ostergaard et al. 2011). Additionally, GATA2 mutations predispose for pulmonary alveolar proteinosis, disseminated warts and human papillomavirus-associated squamous cell cancer (Hsu et al. 2011), childhood neutropenia (Pasquet et al. 2013), as well as NK-cell deficiency (Mace et al. 2013). Recent analysis of 57 patients with GATA2 mutations found a high correlation (p < 0.001) between monocytopenia, B, NK, and CD4 lymphocytopenia and the

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presence of disease (Spinner et al. 2014), emphasizing the importance of measuring monocyte numbers for the diagnosis of this condition. Indeed, 78.9% of this cohort had reduced monocyte numbers, 15.7% had normal monocyte numbers, and only 5.3% of the patients had increased monocyte numbers (Spinner et al. 2014). However, this cohort as well as other reports of *GATA2*-associated disorders included mostly teenagers and adult patients (Pasquet et al. 2013; Dickinson et al. 2014).

Here we report on a patient that presented in infancy with severe hearing loss and lymphedema. By 4 years of age the patient developed acne, disseminated warts, lymphadenopathy, and myelodysplastic syndrome. This patient had long-standing monocytosis with a novel *GATA2* mutation.

Case report

The Research Ethics Board at the Hospital for Sick Children, Toronto, Ontario, approved the study, and a written informed consent for the research was obtained from the parents. The patient was the second daughter born to nonconsanguineous parents after a full-term pregnancy that was complicated by maternal heart arrhythmia. Immediately after birth, apnea episodes were noted, and the patient was treated for 7 days for presumed pneumonia and sepsis with a gentamicincontaining regimen. At 2 months of age the patient was diagnosed with profound sensory neural hearing deficit, for which she received cochlear implants. Analysis of several genes associated with hearing loss, including GJB2, GJB6, SLC26A4, MTRNR1, and MTTS1 was normal. The patient also had pulmonary artery narrowing that resolved. At 3 months of age lymphoedema was noted, which initially involved the left foot and right hand and progressed to all extremities by 6 months of age. A lymphogram demonstrated an absent migration of lymph in the upper left extremity as well as slow migration in the right upper and left lower extremities. A micro-array demonstrated a 458 kb insertional duplication of chromosome 9Q34.3 of unknown significance. At 4 years of age, because of unusual facial acne (Figure 1A), multiple planter warts (Figure 1B), recurrent respiratory and ear infections, as well as persistent lymphadenopathy the patient was referred for immune evaluation. Laboratory findings included normal immunoglobulin levels, production of antibodies, T-, B-, and NK-cell numbers as well as normal T-cell response to stimulation (Table 1). At that time, the patient was also noted to have intermittent neutropenia, markedly



Figure 1: Dermatological abnormalities at 4 years of age. Facial acne (A), and multiple planter warts (B).

increased mean corpuscular volume and mean corpuscular hemoglobin, and elevated hemoglobin F, whereas monocytes were persistently increased or normal (Table 2). Bone marrow examinations revealed mildly hypocellular particles with adequate trilineage hematopoiesis and no increase in blasts. Cytogenetic analysis by G banding and interphase fluorescent in situ hybridization showed monosomy 7 in the majority of the cells. A diagnosis of MDS was made. Because of the patient's constellation of symptoms, the *GATA2* gene was interrogated in the patients' peripheral blood mononuclear

Table 1: Patient's immune evaluation at 4 years of age.

	Patient	Normal range (age matched)	
IgM (g/L)	0.8	0.5–1.9	
IgA (g/L)	0.6	0.3–1.5	
IgG (g/L)	5.0	3.2–11.5	
Anti-tetanus toxoid IgG (IU)	0.3	>0.1	
Anti-pneumococcal IgG (mg/mL)	>270	>100	
Anti-measles IgG	Positive	Positive	
Anti-mumps IgG	Positive	Positive	
Anti-rubella IgG	Positive	Positive	
Anti-varicella zoster IgG	Positive	Positive	
CD4+ cells/µL	1700	500-2400	
CD4+ cells (%)	49	23–48	
CD8+ cells/µL	1200	300–1600	
CD8+ cells (%)	34	14–33	
CD20+ cells/µL	210	200–2100	
CD20+ cells (%)	6	5–20	
CD3-CD16+CD56+ cells/µL	217	100–1000	
CD3-CD16+CD56+ cells (%)	6	4–23	
Stimulation index to phytohemoagglutinin	274	>200	

Index	36 months	48 months	50 months	52 months	54 months	56 months	Normal range
Neutrophils × 10 ⁹ /L	1.6	0.7↓	0.7↓	2.1↓	0.2↓	0.1↓	1.5–8.5
Lymphocytes × 10 ⁹ /L	2.0	2.5	1.6 ↓	2.1	2.7	2.5	2.0-8.0
Monocytes × 10 ⁹ /L	0.5	1.4 ↑	2.3 ↑	1.5 ↑	0.2	0.3	0.05-0.8
Mean corpuscular volume	90↑	104 ↑	103	102 ↑	102 ↑	105 ↑	70–86
Mean corpuscular hemoglobin	31	36 ↑	35 ↑	36 ↑	35 ↑	34 ↑	24–31
Hemoglobin fetal (%)		8.3 ↑					0.1–1.8

Table 2: Patient's hematological indices.

Note: ↓, reduced; ↑, increased

cells. Sequencing demonstrated a heterozygous novel c.1041delC mutation (Figure 2) immediately upstream to the C-terminal zinc-finger domain. The mutation is predicted to cause a reading frame shift and to later stop codon 39 amino acids (Figure 3). The mutation was not present in the patient's parents or sister. Although the mutation was not demonstrated to be a germ line, the rarity of such somatic mutations in MDS and the constitutional features of the patient suggest that the *GATA2* mutation was the cause of the disease.

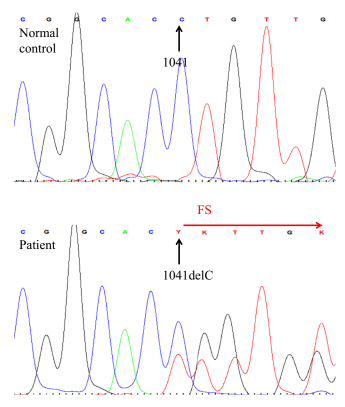


Figure 2: Novel 1041delC in the patient's *GATA2* gene. Electropherogram of the patient's *GATA2* gene (bottom) demonstrating the heterozygous C nucleotide deletion at position 1041 (arrow) resulting in frame shift (FS), in comparison with sequencing of a normal control (top).

Because of the MDS with progressive neutropenia and ominous cytogenetics, the patient underwent bone marrow transplantation using a 10/10 human leukocyte antigen (HLA) matched unrelated donor at 56 months of age. The myeloablative conditioning regimen comprised fludarabine (160 mg/m²) and busulfan (3.2 mg·kg⁻¹·dose⁻¹, adjusted to achieve area under the curve of 3600–6000 micromol/litre/minute × 4 doses).

Discussion

Many patients with *GATA2* mutations have reduced lymphocytes and severe monocytopenia. Therefore, low monocyte numbers have been traditionally considered the sine qua non for the diagnosis of *GATA2* deficiency. Appropriate *GATA2* expression is critical for the maturation of hematopoietic cells (Bresnick et al. 2010); hence, the common hypothesis is that developmental abnormality results in reduced cell numbers. However, the patient described here had elevated monocyte counts as well as the normal NK- and B-cell numbers. It is possible that the monocytosis is related to the development of the myelodysplasia, as already suggested in a recent cross-sectional analysis of 24 patients' with

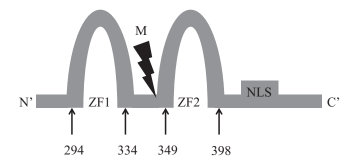


Figure 3: Putative effects of 1041delC mutation on the C-terminal finger and nuclear localization signal of *GATA2*. Diagram demonstrating GATA2 protein N-terminal and C-terminal zinc fingers (ZF1 and ZF2, respectively), the nuclear localization signal (NLS), and the location of the patient's 1041delC mutation (M).

GATA2 deficiency (Dickinson et al. 2014), although monocytopenia were common and profound among patients with GATA2 defects even in the setting of MDS (Spinner et al. 2014). Also, the normal monocyte numbers observed in our patient at 36 months of age, prior to establishing the diagnosis of MDS, might have been a harbinger of the bone marrow abnormalities, as reflected by the increased mean corpuscular volume at that age. Regardless of the etiology of monocytosis, our patient illustrates the importance of investigating GATA2 even in the absence of monocytopenia, and considering myelodysplasia among patients with GATA2 defects who have normal or increased monocyte numbers.

The original publication by Emberger et al. (1979), as well as subsequent studies of patients with Emberger syndrome and GATA2 deficiency, described lymphedema that often appeared in childhood and adolescence (Mansour et al. 2010). In contrast, lymphedema was already detected in our patient at 3 months of age and she suffered from disseminated warts and significant hematopoietic abnormalities by 4 years of age. Although a clear genotype-phenotype correlation has not been established in GATA2 deficiency, a recent report suggested that mutations causing frame shift and disruption of the C-terminal zinc finger, such as the novel one identified in our patient, are associated with a younger age of presentation and higher clinical score (Dickinson et al. 2014). Future studies involving a larger number of patients will help clarify whether patients with frame-shift mutations and haplo-insufficiency of the C-terminal zinc finger should be managed differently than patients with amino acid substitutions.

Optimal treatment for GATA2 deficiency has not been established. Our patient and many other GATA2deficient patients suffered from repeated infections and disseminated warts. Yet, T-cell numbers and functions as well as production of antibodies are often preserved. Hence, antibiotic and (or) immunoglobulin prophylaxis are unlikely to be of significant clinical benefit. It is possible that monocyte function might be improved by use of a granulocyte-macrophage colony stimulating factor, similar to the benefit observed among patients with other macrophage defects (Wang et al. 2005; Greenhill and Kotton 2009). However, this strategy has not been reported in GATA2-deficient patients and might be associated with significant risk because of the potential to accelerate propensity for MDS or AML in genetically altered stem cells, as suggested for other growth factors and patients with marrow abnormalities (Beekman and Touw 2010). Hematopoietic stem cell transplantations (HSCT) with myeloablative or nonmyeloablative conditioning have been utilized in GATA2-deficient patients, primarily because of hematological malignancies and progressive immunodeficiency (Cuellar-Rodriguez et al. 2011; Pasquet et al. 2013; Grossman et al. 2014). Our patient received HSCT at a very young age, before the development of malignancy and severe infections, albeit after the appearance of diffuse lymphedema, a severe hearing defect, and MDS with monosomy 7. Accordingly, longterm follow up of our patient will provide a unique opportunity to assess whether HSCT can prevent the complications associated with GATA2 deficiency including pulmonary alveolar proteinosis and gynecological malignancies.

In conclusion, GATA2 defects should be investigated in patients suffering from early hearing loss, lymphedema, recurrent infections, and (or) hematological abnormalities, particularly MDS or familial AML, even if monocytes and lymphocyte subpopulations are not affected. The best treatment for GATA2-deficient patients needs to be determined.

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