Respiratory infections in X-linked hyper-IgM syndrome with CD40LG mutation: a case series of seven children in China

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Abstract

Background: X-linked hyper-immunoglobulin M (XHIGM), a primary immunodeficiency syndrome caused by mutations in the CD40 ligand gene (CD40LG), presents with recurrent respiratory infections in pediatric patients. We aimed to evaluate the spectrum of clinical features and respiratory pathogens in pediatric patients with XHIGM in China.

Methods: We retrospectively reviewed seven pediatric patients who were diagnosed with XHIGM and received follow-up treatment at the Guangzhou Women and Children's Medical Center between January 2010 and January 2021. We determined their clinical characteristics, causative pathogens, and prognosis by performing peripheral immunological and genetic tests.

Results: There were seven boys with age ranging from 4–20 months (median age, 13 months). Four of the seven respiratory infections were caused by Talaromyces marneffei (T. marneffei). Two patients had viral infections caused by cytomegalovirus (CMV) and human adenovirus respectively. One patient had a mixed infection caused by Pneumocystis carinii and CMV. Except for one child who died of respiratory failure, one patient received hematopoietic stem cell transplantation (HSCT) and recovered well, the other five patients survived with regular infusions of intravenous immunoglobulin (IVIg) during the follow-up period. Six patients had reduced antibody levels, especially IgG, IgA, and IgE levels. Increased serum IgM levels were detected in four cases, and three cases presented normal IgM levels at onset. All children were diagnosed with XHIGM with CD40LG variation. Three novel mutations were identified in the present study.

Conclusions: Our study suggests that respiratory infections usually begin within 2 years old, fungi and viruses are important pathogens causing respiratory infections in children with XHIGM. In endemic areas, T. marneffei is the common pathogen of respiratory tract infection in children with the disease.

Keywords: X-linked hyper-immunoglobulin M syndrome, Respiratory infection, Children, Pathogen, Mutation

Introduction

Hyper-immunoglobulin M syndrome (HIGM) is a rare primary immunodeficiency, in which defective B cell isotype switching leads to a phenotype characterized by low or absent serum levels of IgG, IgA and IgE, whereas the IgM concentration is either normal or increased [1–4]. This syndrome is known as immunoglobulin class switch recombination (Ig-CSR) deficiencies. Mutations in a
number of genes involved in CSR have been implicated in the development of HIGM profile, including CD40 ligand (CD40L), CD40, nuclear factor kappa-B essential modulator (NEMO/IKKγ), inhibitor of kappa light-chain gene enhancer in B cells, alpha (IkBa), nuclear factor kappa-B subunit 1 (NFKB1), activation-induced cytidine deaminase (AICDA/AID), and so on [2, 5, 6]. The X-linked Hyper-IgM syndrome (XHIGM), which is caused by CD40L mutations, is the most common form of HIGM and accounts for about 65–70% of all cases [7, 8]. The prevalence of XHIGM was approximately 1 in 1,000,000 live births. CD40L is expressed primarily on activated CD4+ T cells, and interacts with CD40 expressed on B cells, monocytes, macrophages, and dendritic cells [3]. Therefore, XHIGM is confirmed as the combined T and B immunological deficiency, and it is clearly illustrated by the susceptibility of patients with XHIGM to recurrent pyogenic and opportunistic infections [9].

Clinical manifestations of XHIGM are complex and severe. Patients with XHIGM due to the defective interaction between T and B. Therefore, they are more susceptible to infections that depend on IgA, IgG and IgE, with repetitive episodes of sinusitis, otitis, tonsillitis, cutaneous infections, giardiasis, helminthiasis, enterovirus meningitis, pneumonia, in addition to impairment of weight and height gain, and a tendency to autoimmune diseases and neoplasms [10]. The majority of patients with XHIGM develop recurrent pulmonary infections in the first 2 years of life [11]. Some patients present to medical attention with Pneumocystis carinii pneumonia during their first year of life [11, 12]. This retrospective study on seven children diagnosed with XHIGM preliminarily assessed the clinical presentation and unusual pathogens of respiratory infections in these patients and aimed to describe the respiratory infection characteristics of these patients.

Methods

Patients and ethics

The data of seven patients with XHIGM admitted to Guangzhou Women and Children's Medical Center between January 2010 and January 2021 were retrospectively analyzed. Clinical data including demographic characteristics (age and gender), clinical symptoms and signs, laboratory data, microbiological findings, imaging examinations, complications, treatments, and outcomes were collected and analyzed.

The study was conducted in accordance with the Declaration of Helsinki, and the protocol was approved by the Ethics Committee of Guangzhou Women and Children's Medical Center of Guangzhou Medical University (Grant No. 2021084A01). Written informed consent was obtained from the children's parents or legal guardians for the use of their clinical and laboratory data from their medical reports.

Diagnostic methods

Patients were diagnosed with XHIGM based on the clinical characteristics and genetic test results of the patients. The updated classification of primary immunodeficiency disorders (PIDs) was performed to the standards as diagnostic criteria by the Primary Immunodeficiency Expert Committee of the International Union of Immunological Societies [13]. Selective PID panel testing (MyGenostics, Inc., China), a molecular diagnosis system based on targeted region sequencing of candidate genes, was performed on our patients.

Nasopharyngeal swab and sputum, bronchoalveolar lavage fluid (BALF), pleural fluid, and/or serum samples were collected for tests including immunofluorescence staining, antigen and antibody tests, nucleic acid detection, and bacterial culture. Pathogen high-throughput genome sequencing (KingMed Diagnostics, Inc., China) of BALF was performed in four patients to screen for bacteria, viruses, fungi, parasites, Mycobacterium tuberculosis complex, Mycoplasma, and Chlamydia.

Results

Clinical characteristics and pathogens

The clinical features of the children are summarized in Table 1. There were seven boys with age ranging from 4–20 months (median age, 13 months). All children had respiratory infections, including fever and cough. All patients had at least one episode of respiratory tract infection before diagnosis at first hospitalization in our center, four (P1, P5, P6, and P7) had Talaromyces marneffei (T. marneffei) infection and two (P2 and P3) were identified to have cytomegalovirus (CMV) and human adenovirus (HAdV) infections, respectively; Only P2 was confirmed to have a mixed infection caused by P. carinii and CMV. From a pulmonary imaging perspective, four patients developed lymphadenopathy and diffuse infiltration with T. marneffei infection on high-resolution computed tomography (HRCT) (Fig. 1A). The other three patients (P2, P3, and P4) showed ground-glass opacities on HRCT (Fig. 1B). Specifically, seven patients were followed up for 3–10 years. During the follow-up period, only P3 died of respiratory failure. P4 underwent hematopoietic stem cell transplantation (HSCT), and quite a good recovery of hemopoiesis was observed 6 months after HSCT. In addition, the others survived with regular infusions of intravenous immunoglobulin (IVIg).

Hematological and immunological parameters

Immunological evaluations of the seven patients at the time of diagnosis are listed in Table 2 and Table
S1. Almost all patients had reduced serum IgG levels, except for one child (P6) treated with IVIg before the examination. Serum IgM levels of P3, P4, and P5 were slightly elevated, while serum IgM levels of the other patients (P1, P2, P6, and P7) were normal. High or normal T and B lymphocyte counts were observed in all patients. The counts of $CD19^+$ B cells were elevated in P1, P2, P3, P4, and P7, whereas the other two patients (P5 and P6) had normal $CD19^+$ B cell counts. Nitro blue tetrazolium test results were normal in all patients. Four children (P1, P4, P5, and P7) were found temporary neutropenia, the ratio of neutrophils decreased in most patients (P1, P2, P3, P4 and P7).

**Table 1** Clinical features of respiratory infection in the children with XHIGM

| Patients | Age (months) | Gender | Respiratory symptoms and signs | Pathogens | Pulmonary imaging | Prognosis |
|----------|--------------|--------|--------------------------------|-----------|------------------|-----------|
| 1        | 13           | M      | Cough, fever, polypnea         | $T. marneffei$ (BALF, Airway mucosal biopsy) | Multiple patchy, Hilar lymphadenopathy | Survival |
| 2        | 6            | M      | Cough, fever, polypnea         | $P. carinii$, CMV (BALF) | Ground-glass opacity | Survival |
| 3        | 4            | M      | Cough, fever, polypnea         | HAdV (BALF, Nasopharyngeal swabs) | Ground-glass opacity | Died     |
| 4        | 5            | M      | Cough, polypnea                | CMV (Sputum) | Ground-glass opacity | Survival |
| 5        | 19           | M      | Cough, fever                   | $T. marneffei$ (Blood) | Multiple nodules, Hilar and mediastinal lymphadenopathy | Survival |
| 6        | 15           | M      | Cough, fever, wheeze, polypnea | $T. marneffei$ (Blood) | Hydrothorax, Hilar and mediastinal lymphadenopathy | Survival |
| 7        | 20           | M      | Cough, fever                   | $T. marneffei$ (Airway mucosal biopsy) | Multiple nodules and patchy, Hilar lymphadenopathy | Survival |

$T. marneffei$ Talaromycetes marneffei, $P. carinii$ Pneumocystis carinii, BALF Bronchoalveolar lavage fluid, HAdV Human adenovirus, CMV Cytomegalovirus

**Mutation studies**

Using selective PID panel testing, all patients were found to have mutations in the $CD40LG$ gene (Table 3) and were diagnosed with XHIGM. In this cohort, hemizygous mutations were found in five patients (P1, P2, P3, P4, and P7), and microdeletions were found in two others (P5 and P6). The mutations were mostly inherited via the maternal lineage (P1, P2, P5, P6, and P7). However, parents of P3 and P4 did not carry the same mutation as their children. Two novel hemizygous mutations (c.424_436del and c.488delT) in exon 5 were detected in P1 and P3, respectively, whereas a novel hemizygous mutation (c.92_104del) in exon 1 was found in P2. These mutations
Table 2 Immunoassays findings of the children with XHIGM\textsuperscript{a}

| Patients | IgG (g/L) (Normal range) | IgA (g/L) (Normal range) | IgM (g/L) (Normal range) | IgE (IU/ML) (Normal range) | CD19 + Abs (cells/ul) (Normal range) | CD3 + 4 + (cells/ul) (Normal range) | CD3 + 8 + (cells/ul) (Normal range) | CD56 + (cells/ul) (Normal range) | NBT test |
|----------|--------------------------|--------------------------|--------------------------|-----------------------------|-------------------------------------|-------------------------------------|-------------------------------------|-------------------------------------|----------|
| 1        | 0.52 (3.52–10.69)        | 0.1 (0.13–1.38)          | 0.53 (0.38–2.03)         | < 5 (0–60)                  | 2279.28 (90–660)                  | 1746.12 (410–1590)                  | 431.18 (190–1140)                  | 161 (90–590)                  | Normal   |
| 2        | 0.33 (3.6–92)            | 0.07 (0.08–0.56)         | 0.48 (0.38–1.26)         | < 5 (0–60)                  | 3440.47 (240–1317)                | 3470.7 (345–2350)                  | 1628.6 (314–2080)                 | 561.13 (210–1514)                | Normal   |
| 3        | 0.88–1.26 (3.06–7.74)    | 0.07 (0.07–0.41)         | 2.63–5.45 (0.24–0.88)    | < 5 (0–15)                  | 2295.16 (240–1317)                | 6745.74 (345–2350)                 | 2372.63 (314–2080)                | 217.89 (210–1514)                | Normal   |
| 4        | 0.71 (3.06–7.74)         | 0.07 (0.07–0.41)         | 1.34 (0.24–0.88)         | < 5 (0–15)                  | 3441.95 (240–1317)                | 2365.98 (345–2350)                 | 1824.7 (314–2080)                 | 234.14 (210–1514)                | Normal   |
| 5        | <0.33 (3.82–10.58)       | 0.17 (0.14–1.14)         | 2.65 (0.4–1.28)          | < 5 (0–60)                  | 364.43 (90–660)                   | 1458.66 (410–1590)                 | 347.75 (190–1140)                 | 262.96 (90–590)                  | Normal   |
| 6        | 9.79 (3.82–10.58)        | 0.07 (0.14–1.14)         | 0.58 (0.4–1.28)          | < 5 (0–60)                  | 395.23 (90–660)                   | 1973.05 (410–1590)                 | 1706.35 (190–1140)                | 247.34 (90–590)                  | Normal   |
| 7        | 1.49 (3.82–10.58)        | 0.07 (0.14–1.14)         | 0.48 (0.4–1.28)          | < 5 (0–60)                  | 1945.34 (90–660)                  | 3323.59 (410–1590)                 | 1350.24 (190–1140)                | 479.35 (90–590)                  | Normal   |

\textsuperscript{a} The normal ranges were presented because they changed with different kit and different age

\textsuperscript{b} The tests were conducted after IVIg treatment

NBT Nitro blue tetrazolium test
have never been reported in gnomAD and were predicted to be disease-causing mutations using PolyPhen2, Mutation Taster, PROVEAN, and SIFT software.

Discussion
XHIGM is a primary combined immunodeficiency syndrome that results from mutations in the CD40LG, which encodes for the CD40L protein [2]. This retrospective analysis of seven pediatric patients aimed to collect data on the clinical presentation, treatment, and causative respiratory pathogens of patients with XHIGM and use this information to gain further understanding of clinical features and the pathogens of respiratory infections in pediatric patients with XHIGM.

The most prominent clinical feature observed in XHIGM patients is recurrent respiratory infection, the symptoms usually develop in infancy and second year of life [3]. In our study, all seven boys suffered respiratory infections by the age of 2, and they had at least one episode of respiratory tract infection before a diagnosis is made. Furthermore, XHIGM results in combined immune deficiency with a defect in Ig isotype switching and enhanced susceptibility to opportunistic infections [14]. According to the European and US cohorts, the most common pathogen causing opportunistic infections is P. carinii, while others include Bartonella, Cryptosporidium parvum, Histoplasma, Mycobacterium avium, Salmonella dublin, Herpes family viruses, etc. [3, 14, 15]. In this study, only P2 had P. carinii infection, as previously reported, while four patients (P1, P5, P6, and P7) were found to have T. marneffei infection in the lungs. T. marneffei is an opportunistic pathogen that infects immunodeficient patients in Southeast Asia as a dimorphic fungus, and T. marneffei infection is a severe disease that can lead to high mortality rates of greater than 50% in children in previous reports [16]. In adults, T. marneffei infection has been considered to be exclusively associated with acquired immunodeficiency virus (HIV) infection [17]. In contrast to adults, pediatric patients with PIDs are more susceptible to T. marneffei according to previous reports [16, 18]. Therefore, pediatricians should be alert to T. marneffei respiratory infection in XHIGM patients in high-prevalence areas.

It should be noted that viral infections have also been linked to the development of respiratory infections in these children [3, 12]. In our study, three patients presented with interstitial pneumonia with extensive ground-glass opacity on HRCT, due to P. carinii and other viral infections. Of these, a unique case of HAdV infection (P3) progressively worsened and the patient eventually died of respiratory failure. What needs to pay attention is that all three patients began with the respiratory symptoms within 6 months of birth as first infection. Of note, interstitial lung disease in infancy should be paid more attention to XHIGM by pediatric clinicians even for the first time.

XHIGM with CD40LG mutations has been classified as combined T and B immunodeficiency, and the overall prognosis of these patients is poor, with an overall survival rate of 20–28.2% [9, 14, 19, 20]. Earlier management with IVIg replacement therapy and antibiotic prophylaxis could improve the quality of life of patients by reducing the incidence of life-threatening infections [3]. In this study, most confirmed cases were regularly administered IVIg replacement therapy, and no serious infections occurred in them during the follow-up years. Hematopoietic stem cell transplantation (HSCT) represents the only curative choice available for patients with XHIGM [21]. However, the presence of severe pre-transplant infections, organ damage (usually affecting the liver and/or lung), lack of a matched donor, or socioeconomic constraints, while age at transplantation seems to be the major challenge. Recent research found a more favorable outcome and greater event-free survival among patients who underwent transplantation before age 6 years in comparison with patients 6 years or older.

### Table 3 Genetic detection of the children with XHIGM

| Patient | Gene        | Genetic variation | Source | Nucleotide variation | Amino acid mutation | Amino acid variation |
|---------|-------------|-------------------|--------|----------------------|---------------------|---------------------|
| 1       | CD40LG      | Hemizygote        | Mother | c.424_436del         | Frameshift mutation | p.E142Tfs*3         |
| 2       | CD40LG      | Hemizygote        | Mother | c.92_104del          | Frameshift mutation | p.F31Lfs           |
| 3       | CD40LG      | Hemizygote        | Spontaneous mutation | c.488del | Frameshift mutation | p.R165Dfs*26       |
| 4       | CD40LG      | Hemizygote        | Spontaneous mutation | c.761C>T | Missense mutation | p.T254M           |
| 5       | CD40LG      | Microdeletion     | Mother | > 132 kb             | -                   | -                   |
| 6       | CD40LG      | Microdeletion     | Mother | c.1978+1G>A          | -                   | -                   |
| 7       | CD40LG      | Hemizygote        | Mother | c.598A>T             | Nonsense mutation   | p.R200*           |
In our cohort, only one patient has been performed HSCT and recovered well, while the other patients had not been treated with HSCT for the matched donor’s lack or economic reason. IVIg replacement therapy is still the first choice for XHIGM patients to prevent infection before HSCT.

XHIGM is a PID caused by mutations in the \textit{CD40LG} gene. \textit{CD40LG} is located on the X chromosome, which encodes CD40L (CD154), a member of the tumor necrosis factor family made-up of three functional domains: intracellular, transmembrane and extracellular domains \cite{23}. Since CD40L is necessary for T lymphocytes to induce B lymphocytes to undergo class switching from IgM to IgG, IgA, and IgE, patients with XHIGM syndrome have reduced levels of IgG, IgA, and IgE, and normal or elevated levels of IgM \cite{8, 9}. In this study, our patients had typical immune parameters of XHIGM at onset, which indicated the possibility of XHIGM and the necessity for genetic testing. As expected, mutations in \textit{CD40LG} were found in all seven pediatrics patients. Two patients were diagnosed with XHIGM due to microdeletions of the \textit{CD40LG} gene (P5 and P6). Interestingly, three cases of novel hemizygote mutations have not been reported in gnomAD so far (P1, P2, and P3), where other variants at the same position are found. Based on the clinical manifestations and pedigree analysis of the patients according to the American College of Medical Genetics variation classification guidelines \cite{24}, the three mutations can be taken as evidence for pathogenicity. Future research could focus on functional verification by animal experimentation.

**Strengths**

This study emphasized fungi and viruses are important pathogens of respiratory infections in pediatric patients with XHIGM. In our cohort, patients were more susceptible to \textit{T. marneffei}, which suggested \textit{T. marneffei} infections could be a clinical clue to realize XHIGM for pediatricians. For infant, interstitial lung disease could be the first symptom of respiratory infection in XHIGM patients. In addition, three novel mutations in \textit{CD40LG} were first reported in our cohort.

**Limitations**

This study has some limitations that should be noted. This was a single-center study with a limited number of cases. CD40 ligand expression levels in in vitro-activated lymphocytes should be assayed for the patients, but it has not yet been carried out in our hospital, it is great regret that our patients failed to get tested. Nevertheless, this study provides a valuable reference for respiratory infections, especially fungal and viral infections, in children with XHIGM.

**Conclusions**

XHIGM patients usually presents respiratory infections within 2 years old, and fungi and viruses are important respiratory pathogens. \textit{T. marneffei} respiratory infections were more common in children with XHIGM in endemic areas. Infants who were diagnosed with interstitial lung disease should be alert to XHIGM. In this study, we presented three novel mutations in \textit{CD40LG} that have not been reported previously.

**Abbreviations**

XHIGM: X-linked hyper-immunoglobulin M; \textit{T. marneffei}: Talaromyces marneffei; CMV: Cytomegalovirus; IVIg: Intravenous immunoglobulin; PID: Primary immunodeficiency; HIGM: Hyper-immunoglobulin M; Ig-CSR: Immunoglobulin class-switch recombination; CD40L: CD40 ligand; BALF: Bronchoalveolar lavage fluid; HAdV: Human adenovirus; \textit{P. carinii}: Pneumocystis carinii; HRCT: High-resolution computed tomography; AIDS: Acquired immunodeficiency syndrome; HSCT: Hematopoietic stem cell transplantation.

**Acknowledgements**

We would like to thank the children who participated in this study and their families.

**Authors’ contributions**

Kunling Shen, Gen Lu and Huifeng Fan helped contributed to the conception and design of the study, and write the manuscript; Li Huang, Diyuan Yang, Changhao Zhang, Qiang Zeng contributed to collection and analysis of data; Genquan Yin participated in the critical review of the manuscript; All authors read and approved the final manuscript.

**Funding**

Not applicable.

**Availability of data and materials**

The datasets generated and/or analyzed during the current study are not publicly available due individual privacy of patients could be compromised, but are available from the corresponding author on reasonable request.

**Declarations**

**Ethics approval and consent to participate**

This study protocol was conducted in accordance with the Declaration of Helsinki and approved by the Ethics Committee of Guangzhou Women and Children’s Medical Center of Guangzhou Medical University (Grant No. 2021084A01). Written informed consents were signed during hospitalization. The data used in this study was anonymised before its use.

**Consent for publication**

Not applicable.

**Competing interests**

The authors have no conflicts of interest to report.

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Received: 24 June 2022  Accepted: 31 October 2022
Published online: 22 November 2022

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