Mimicking hypersensitivity pneumonitis as an uncommon initial presentation of chronic granulomatous disease in children

Hui Liu, Jinrong Liu, Huimin Li, Yun Peng and Shunying Zhao*

Abstract
Dry cough, dyspnea and diffuse centrilobular nodules in both lungs of radiologic findings similar to hypersensitivity pneumonitis (HP) are rare initial presentation in chronic granulomatous disease (CGD). CGD is remarkable for increased susceptibility to bacterial and fungal infections as well as high sensitivity to inciting antigens such as Aspergillus species due to dysregulated inflammation. We identified three children who had an initial presentation mimicking HP and were subsequently diagnosed as CGD. All patients developed invasive pulmonary A. fumigatus infection (IPAI) following systemic glucocorticoid therapy. Two of the three patients were found to have mutations in NCF1 gene and one patient in NCF2 gene. As HP is uncommon in children, we should consider the possibility of CGD in children with HP, even in mimicking HP patients with suggestive inhalation history and negative fungal cultures. A prompt diagnosis of CGD is essential to enable initiation of prophylactic antibacterial and antifungal therapies.

Keywords: Chronic granulomatous disease, Hypersensitivity pneumonitis, A. fumigatus, Glucocorticoid, Children

Letter to the editor

Chronic granulomatous disease (CGD) is characterized by recurrent and severe bacterial and fungal infections as well as excessive inflammation, which are most prominent in gastrointestinal and genitourinary tracts, such as granulomata mimicking Crohn’s disease [1]. An exuberant respiratory inflammation induced by the exposure to inciting antigens and clinically manifested as hypersensitivity pneumonitis (HP) or allergic bronchopulmonary aspergillosis (ABPA) has also been described in CGD [2–7]. However, HP as an initial presentation of CGD is uncommon and has never been reported. Here we review three children who had an initial presentation mimicking HP, developed invasive pulmonary A. fumigatus infection (IPAI) following systemic glucocorticoid therapy and were subsequently diagnosed as CGD.

Case presentation

Case 1
A 4-year-old boy was admitted to the hospital on September 8, 2011 after 3 weeks of dry cough, progressive dyspnea and fever. He lived in a fruit stall with many rotten fruits inside. He also had a history of pneumonia at 3 months old. He had a history of severe eczema and seasonal rhinitis at one year old.

On admission, his oxygen saturation at rest was 92%, and decreased to 86% after walking. Bilateral basilar rales were noted on auscultation. Chest high-resolution CT (HRCT) scan showed diffuse nodular opacities and slight ground-glass (Fig. 1a). Cultures revealed no evidences of mycobacteria, fungi and viruses. A specimen taken from video-assisted lung biopsy of the right lower lobe revealed bronchiolo centric lymphocytic, and non-necrotizing granulomas and no evidence of fungal or bacterial elements (Fig. 2). Bronchoalveolar lavage fluid (BALF) from his right middle lobe consisted of 46% macrophages, 2% eosinophils and 52% T cells, with a CD4+/CD8+ ratio of 0.62. Fungal and mycobacterial cultures of BALF were negative, as were T-cell interferon-γ release assays for tuberculosis and polymerase chain
reactions for pneumocystis jirovecii. After exclusion of infectious agents, diagnosis of mimicking HP probably due to inhalation of rotten fruit elements was made. Treatment with 1 mg/kg/day prednisone was initiated and clinical symptoms improved after 3 days.

Three weeks after the beginning of tapered prednisone, he developed fever and cough with purulent sputum. HRCT found consolidation in left upper lobe and cavity in right upper lobe (Fig. 2a). Sputum culture was positive for *A. fumigatus* three times. Parenteral voriconazole therapy for 2 months followed by oral voriconazole was administered for 6 months until lung lesions disappeared completely. In consideration of the patient’s progressive course, he was referred to immunological test. Dihydrorhodamine-1,2,3 (DHR) test showed the absence of neutrophil oxidative burst consistent with CGD. Gene mutation analyses revealed compound heterozygous mutations (c.278A > T and c.475delA) in *NCF2* gene, indicating autosomal recessive CGD [8]. Continuous prophylactic treatment with trimethoprim-sulfamethoxazole and itraconazole were administered, and no infection recurred in a follow-up period of 4 years.

**Case 2**

An 8-year-old girl was admitted to the hospital on February 15, 2015 because of high spiking fever and chills, dry cough, progressive dyspnea and chest stuffy for 20 days. Twenty-four days ago she had burned decayed cornhusks with her brother (case 3) for 4 h. She had a history of severe eczema and seasonal rhinitis at 3 years old.

On admission, her oxygen saturation at rest was 93%, and decreased to 84% after walking. Bilateral basilar rales were noted on auscultation. HRCT scan showed the presence of bilaterally diffuse centrilobular nodules and slight ground-glass (Fig. 1b). BALF from her right middle lobe consisted of 41% macrophages, 3% eosinophils and 56% T cells, with a CD4+/CD8+ ratio of 0.65. Fungal and mycobacterial cultures of BALF were negative. The diagnosis of mimicking HP was made. A treatment of 1 mg/kg/day prednisone was initiated, and clinical symptoms improved after 2 days.
As prednisone being tapered after 3 weeks, she developed fever, coughing with purulent sputum and chest stuffy. Repeated HRCT showed multi-nodules bilaterally distributed along bronchi and part of multi-nodule fused into pieces more in upper lung (Fig. 2b). Specific IgG antibody to *A. fumigatus* was positive. Both culture of sputum and BALF from right middle lobe found *A. fumigatus*. Amphotericin liposome B was taken for 2 months followed by oral voriconazole for one year. The nodes disappeared and the pieces shrank. In consideration of the fact that his brother had a history of perianal abscess and was diagnosed as CGD by means of an abnormal DHR test, she was referred to DHR test, which was consistent with CGD. Sequencing of genomic DNA revealed homozygous mutation (c.541delG) in *NCF1* gene. As prednisone being tapered after 3 weeks, she developed fever, coughing with purulent sputum and chest stuffy. Repeated HRCT showed multi-nodules bilaterally distributed along bronchi and part of multi-nodule fused into pieces more in upper lung (Fig. 2b). Specific IgG antibody to *A. fumigatus* was positive. Both culture of sputum and BALF from right middle lobe found *A. fumigatus*. Amphotericin liposome B was taken for 2 months followed by oral voriconazole for one year. The nodes disappeared and the pieces shrank. In consideration of the fact that his brother had a history of perianal abscess and was diagnosed as CGD by means of an abnormal DHR test, she was referred to DHR test, which was consistent with CGD. Sequencing of genomic DNA revealed homozygous mutation (c.541delG) in *NCF1* gene, indicating autosomal recessive CGD [9]. Continuous prophylactic antibiotic treatment was administered, and no symptoms recurred anymore.

**Case 3**

A 5-year-old boy with high spiking fever and chills, dry cough and progressive dyspnea was admitted to the hospital at the same time with his sister (case 2). He had the same exposure to decayed cornhusks. He had a history of perianal abscess and severe eczema at 3 months old, and no history of previous pneumonia, asthma, or exercise intolerance.

On admission, his oxygen saturation at rest was 95%, and decreased to 87% after walking. Bilateral basilar rales were noted on auscultation. HRCT revealed bilaterally diffuse ill-defined centrilobular nodules and slight ground-glass (Fig. 1c). Lymphocytosis of 50% with a CD4+/CD8+ ratio of 0.73, 48% macrophages and 2% neutrophils were shown in BALF. Fungal and mycobacterial cultures for BALF were negative. The diagnosis of mimicking HP was made. Treatment with 1 mg/kg/day oral prednisone was initiated and clinical symptoms improved after 3 days. A slow taper of prednisone was continued after 3 weeks.

Despite the lack of aggravated symptoms, specific IgG antibody to *A. fumigatus* was positive and his chest HRCT revealed a nodular consolidation with halo sign in left upper lobe (Fig. 2c). Oral voriconazole was taken for 4 months. The halo disappeared and the consolidation shrank. In consideration of his history of perianal abscess, he was referred to DHR test, and the result was consistent with CGD. He was subsequently confirmed to have the same *NCF1* gene mutation as his sister. Prophylactic antibiotic treatment was continuously administered. No extra symptoms occurred anymore.

The clinical data relating to HP and IPAI of the 3 cases are summarized in Table 1. Antecedent history and genetic tests for CGD are summarized in Table 2.

**Discussion**

In CGD patients, hypersensitivity to *Aspergillus* clinically manifested as HP and ABPA occurs after exposure to a variety of antigens, and is characterized by the constitutional symptoms and non-necrotizing granulomas in walls of alveoli and airways. However, HP as the first manifestation of CGD is rare in children.

Our cases illustrate that they had the clinical presentations similar to HP, including the history of exposure to potential inciting antigens, lymphocytosis in BALF, compatible image features, poorly formed granulomas on lung biopsy in one patient (case 1), and a favorable response to systemic glucocorticoids. However, these patients did not fully meet the diagnostic criteria for HP published by Venkatesh and Wild [10].

Lung biopsy in one patient (case 1) revealed non-necrotizing granulomatous pneumonitis with variable airspace organization. It is necessary to rule out the pulmonary granulomatous process related to the underlying CGD. Granulomas in patients with CGD are typically small, containing central neutrophil micro-abcesses surrounded by epithelioid histiocytes and giant cells and spreading in airways or lung parenchyma [11, 12]. Unlike CGD, the granulomas of our patient were loosely formed and relatively larger, spread through airway to lymph tract, and did not contain central micro-abcesses, similar to the adult CGD patient reported by Katsuya and colleagues [5].

Recently, Esenboga et al. reported that a 16-year-old patient atypically presented with chronic HP caused by close contact with pigeons and exposure to their allergens before the diagnosis of CGD caused by homozygous deletion mutation in *NCF1* [13]. Like the present cases, the patient recovered by allergen avoidance combining with the use of anti-inflammatory drug. Meanwhile in another study that followed up 33 patients with X-linked CGD for 10 years, four patients developed interstitial lung disease (ILD). Two of the four patients, one 20 years old and the other 23 years old, were diagnosed with HP through CT images and histopathological examination, probably caused by breathing in dust in a plant and an unknown antigen, respectively [14]. However, the clinical symptoms in the two patients were mitigated only by allergen avoidance, suggesting that CGD patients are more susceptible to hyperinflammation resulting from inhalation of antigens.

The combination of HP/ABPA features has been described in 2 CGD patients with a gradually progressive course over many years [6, 7]. Interestingly, case 2 had the ABPA features of elevated total serum IgE (3000 IU/mL). However, in ABPA patients, acute course or exacerbation may present nodular pulmonary infiltrates, centrilobular nodules and bronchiectasis on HRCT [15]. Lymphocytosis in BALF is uncommon in ABPA.
Table 1 Clinical, physiological, radiographical, and pathological data relating to mimicking HP and IPAI in the 3 CGD patients

| CGD patient | Case 1 | Case 2 | Case 3 |
|-------------|--------|--------|--------|
| Age (yr)    | 4      | 8      | 5      |
| Sex         | male   | female | male   |
| Data of mimicking HP |          |        |        |
| Exposure to an offending environment | Antigens probably relating to rotten fruits | Antigens probably in musty cornhusks | Antigens probably in musty cornhusks |
| Symptom/sign | Dry cough, dyspnea, fever, bilateral basilar rales | Spiking fever with chills, dry cough, dyspnea, chest stuffy, bilateral basilar rales | Spiking fever with chills, dry cough, dyspnea, bilateral basilar rales |
| Pulmonary function | FEV1, 0.48 L (51.2% predicted); FVC, 0.65 L (68.3% predicted) | FEV1, 0.8 L (58.7% predicted); FVC, 0.92 L (58.9% predicted); DLCO, 6.46 ml/min/mmHg (46.6% predicted) | FEV1, 0.59 L (57.3% predicted); FVC, 0.72 L (70.5% predicted) |
| Chest HRCT scan | Diffuse nodular opacities and slight ground-glass in bilateral inferior field | Bilaterally diffuse ill-defined centrilobular nodules and slight ground-glass | Bilaterally diffuse ill-defined centrilobular nodules and slight ground-glass |
| BALF cells | AM: 46%; Lym: 52%; Eos: 2%; CD4+/CD8+: 0.62 | AM: 41%; Lym: 56%; Eos: 3%; CD4+/CD8+: 0.65 | AM: 48%; Lym: 50%; Neu: 2%; CD4+/CD8+: 0.73 |
| Lung biopsy | Bronchiolo centric lymphocytic, non-necrotizing granulomas and no evidence of fungal or bacterial elements | Not available | Not available |
| Bacterial/viral/fungal cultures | Negative | Negative | Negative |
| Treatment | 1 mg/kg/d prednisone | 1 mg/kg/d prednisone | 1 mg/kg/d prednisone |
| Data of IPAI |          |        |        |
| HRCT scan finding | Consolidation in left upper lobe and cavity in right upper lobe | Multi-nodules bilaterally distributed along bronchi and part of multi-nodules fused into pieces more in upper lung | A nodular consolidation with halo sign in left upper lobe |
| Bacterial/viral/fungal cultures | A. fumigatus | A. fumigatus | A. fumigatus |
| Treatment | Infusion of voriconazole for 2 months followed by oral voriconazole for 6 months | Infusion of amphotericin liposome B for 2 months followed by oral voriconazole for one year | Oral voriconazole for 4 months |

HP hypersensitivity pneumonitis, IPAI invasive pulmonary A. fumigatus infection, HRCT high-resolution computer tomography, BALF bronchoalveolar lavage fluid, AM Alveolar macrophages, Lym lymphocytes, Neu neutrophils, Eos eosinophils, FEV1 forced expiratory volume in one second, FVC forced vital capacity, DLCO decreased lung diffusion of carbon monoxide

Table 2 Clinical, radiographical, laboratory and genetic data for CGD of the 3 patients

| CGD patient | Case 1 | Case 2 | Case 3 |
|-------------|--------|--------|--------|
| Antecedent history | Pneumonia at 3 months old, eczema and seasonal rhinitis at one year old | Eczema and seasonal rhinitis at 3 years old | Perianal abscesses and eczema at 3 months old |
| Igs | IgG 12.2 g/L, IgM 1.72 g/L, IgA 2.59 g/L, IgE 598.9 IU/mL | IgG 26.8 g/L, IgM 1.12 g/L, IgA 4.55 g/L, IgE 3000 IU/mL | IgG 26.6 g/L, IgM 1.29 g/L, IgA 4.23 g/L, IgE 365.9 IU/mL |
| Lymphocyte subsets in peripheral blood | NK cells (9.3%), B cell (11.9%); CD4* cells (43.2%), CD8* cells (32.85%) | NK cells (4%), B cell (14%); CD4* cells (39%), CD8* cells (37%) | NK cells (11%), B cell (17%); CD4* cells (33%), CD8* cells (36%) |
| DHR test | No uptake in neutrophil oxidative burst after phorborolymristate acetate stimulation | No uptake in neutrophil oxidative burst after phorborolymristate acetate stimulation | No uptake in neutrophil oxidative burst after phorborolymristate acetate stimulation |
| Gene mutation | NCF2 gene (compound heterozygosity mutation c.278 A > T and c.475del(A) | NCF1 gene (homozygous mutation c.541del(G) | NCF1 gene (homozygous mutation c.541del(G) |

CGD chronic granulomatous disease, DHR dihydrorhodamine-1,2,3, Igs immunoglobulins, NK natural killer
The mechanisms underlying this hyperinflammation are still under investigation. A plausible explanation is that the reduced ROS cannot adequately inhibit the production of inflammatory cytokines [16], and that this ROS deficit in CGD allows for the continuous production of inflammatory cytokines, resulting in immune dysregulation or hyperinflammation. Thus, one of the effective therapeutic approaches for such hyperinflammation is the use of corticosteroids or immunosuppressive drugs [17]. Indeed, infliximab, a chimeric antibody against tumor necrosis factor-α (TNFα), has shown therapeutic efficacy for refractory CGD colitis, as TNFα is thought to play a critical role in granuloma formation in CGD.

Although the symptoms of HP were improved rapidly after systemic corticosteroid treatment, the present patients suffered worsening disease after 3 weeks after systemic corticosteroid treatment, the present patient with newly diagnosed HP or mimicking HP. This is thought to play a critical role in granuloma formation in CGD.

In CGD, the presentation of IPAI is critical to recognize. Firstly, patients suffering from profound immunodeficiency are prone to develop characteristic invasive fungal infections due to Aspergillus specie. Secondly, prolonged steroids therapy makes a contribution to occurrence of IPAI by increasing the patient's susceptibility to infection. Thirdly, the risk of Aspergillus infections in the present patients is higher than that of other pathogens infections because of the exposure to rotten fruits or decayed cornhusks which are considered as Aspergillus spores-rich environment. The occurrence of IPAI makes it easier to diagnose CGD, but which would be detected much earlier in the inflammatory period.

There are two previous reports of 10 patients who developed fulminant mulch pneumonitis (FMP) as emergency presentation of CGD [19, 20]. Unlike our series, the 10 patients presented acute, rapid, often fatal, invasive aspergillosis symptoms after an identifiable exposure to organic material; culture results were positive for Aspergillus before systemic treatment of glucocorticoids, and 5 of the 10 patients died despite the treatment of antifungals and steroid. Based on the disease duration of the present patients, similar inflammatory responses characterized by lung granuloma formation, lymphocytosis in BALF and Aspergillus specific IgG antibodies could be viewed as the subacute end of the acute FMP spectrum caused by inhalation of variable doses of Aspergillus spores.

The incidence of HP is probably much higher in CGD patients, especially in children. CGD should be first excluded by diagnostic DHR testing before steroids therapy in patient with newly diagnosed HP or mimicking HP. Furthermore, it should be noted that the combined application of antifungal drugs and steroids or other anti-inflammatory drugs is more appropriate from the beginning of the treatment in HP patients with suspected CGD, especially in patients with a history of fungal exposure.

Conclusions

CGD should be considered in children with HP, even in mimicking HP patients with suggestive inhalation history and negative fungal cultures before steroids therapy. A prompt diagnosis of CGD is essential to enable initiation of prophylactic antibacterial and antifungal therapies.

Abbreviations

ABPA: Allergic bronchopulmonary aspergillosis; BALF: Bronchoalveolar lavage fluid; CGD: Chronic granulomatous disease; DHR: Dihydrorhodamine-1,2,3; HP: Hypersensitivity pneumonitis; HRCT: High-resolution computer tomography; IPAI: Invasive pulmonary A.fumigatus infection; TNFα: Tumor necrosis factor-α

Acknowledgements

We would like to thank all of the patients and their families for their kind cooperation.

Funding

The study received no funding.

Availability of data and materials

Please contact author for data requests.

Authors’ contributions

HLi supervised the patient care, conceptualized and designed the report, and reviewed and revised the manuscript; JZL supervised the patient care, conceptualized and designed the report, and reviewed and revised the manuscript; YP supervised the patient care, collected the data and revised the article; HLi supervised the patient care and collected the data; YP supervised the radiological images; SZ supervised the patient care, conceptualized the report, and reviewed and revised the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate

The research was approved by the Ethics Committee of Beijing Children’s Hospital. All patient guardians gave informed consent to the study.

Consent for publication

All patient guardians gave informed consent to the publication of this study.

Competing interests

The authors declare that they have no competing interests.

Publisher’s Note

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details

1Department of Respiratory Medicine, Beijing Children’s Hospital, Capital Medical University, National Center for Children’s Health, Nanlishi Road 56, Xicheng District, Beijing, China. 2Imaging Center, Beijing Children’s Hospital, Capital Medical University, National Center for Children’s Health, Nanlishi Road 56, Xicheng District, Beijing, China.

Received: 19 June 2017 Accepted: 6 October 2017

References

1. Marks DJ, Miyagi K, Rahman FZ, Novelli M, Bloom SL, Segal AW. Inflammatory bowel disease in CGD reproduces the clinicopathological features of Crohn’s disease. Am J Gastroenterol. 2009;104(1):117–24.
2. Morgenstern DE, Gifford MA, Li LL, Doenschuk CM, Dinauer MC. Absence of respiratory burst in X-linked chronic granulomatous disease mice leads to...
abnormalities in both host defense and inflammatory response to Aspergillus fumigatus. J Exp Med. 1997;185(2):207–18.

3. Eppinger TM, Greenberger PA, White DA, Brown AE, Cunningham-Rundles C. Sensitization to Aspergillus species in the congenital neutrophil disorders chronic granulomatous disease and hyper-IgE syndrome. J Allergy Clin Immunol. 1999;104(6):1265–72.

4. Konig G, Baur X, Fruhmann G. Sarcoïdosis or extrinsic allergic alveolitis? Respiration. 1981;42(3):150–4.

5. Katsuya Y, Hojo M, Kawai S, Kawai T, Onodera M, Sugiyama H. Chronic granulomatous disease with pulmonary mass-like opacities secondary to hypersensitivity pneumonitis: a case report. J Med Case Rep. 2014;8:242.

6. Segerer F, Morbach H, Hassold N, Kleinert S, Tony HP, Roessler J, et al. A 58-year-old man with respiratory insufficiency after a 50-year history of hypersensitivity pneumonitis and pulmonary Aspergillus infections. J Allergy Clin Immunol Pract. 2013;1(6):677–80.

7. Patiroglu T, Gungor HE, Lazaroski S, Unal E. Chronic granulomatous disease with markedly elevated IgE levels mimicking hyperimmunoglobulin E syndrome. Acta Microbiol Immunol Hung. 2013;60(2):155–62.

8. Patino PJ, Rae J, Noack D, Erickson R, Ding J, de Olarte DG, et al. Molecular characterization of autosomal recessive chronic granulomatous disease caused by a defect of the nicotinamide adenine dinucleotide phosphate (reduced form) oxidase component p67-phox. Blood. 1999;94(7):2505–14.

9. Roos D, de Boer M, Köker MY, Dekker J, Singh-Gupta V, Ahlin A, et al. Chronic granulomatous disease caused by mutations other than the common GT deletion in NCF1, the gene encoding the p47phox component of the phagocyte NADPH oxidase. Hum Mutat. 2006;27(12):1218–29.

10. Venkatesh P, Wild L. Hypersensitivity pneumonitis in children. Clinical features, diagnosis, and treatment. Pediatr Drugs. 2005;7(4):235–44.

11. Moskaluk CA, Pogrebniak HW, Pass HI, Gallin JJ, Travis WD. Surgical pathology of the lung in chronic granulomatous disease. Am J Clin Pathol. 1994;102(5):684–91.

12. Corrin B, Nicholson AG. Chronic granulomatous disease. In: Corrin B, Nicholson AG, editors. Pathology of the lungs. 3rd ed. London: Churchill Livingstone; 2011. p. 484–5.

13. Essenboga S, Emiraloglu N, Cagdas D, Erman B, De Boer M, Oguz B, et al. Diagnosis of interstitial lung disease caused by possible hypersensitivity pneumonitis in a child: think CGD. J Clin Immunol. 2017;37(3):269–72.

14. Kawai T, Watanabe N, Yokoyama M, Nakazawa Y, Goto F, Uchiyama T, et al. Interstitial lung disease with multiple microgranulomas in chronic granulomatous disease. J Clin Immunol. 2014;34(8):933–40.

15. Stevens DA, Moss RB, Kurup VP, Knutsen AP, Greenberger P, Judson MA, et al. Allergic bronchopulmonary aspergillosis in cystic fibrosis–state of the art: Cystic Fibrosis Foundation consensus conference. Clin Infect Dis. 2003;37(Suppl 3):S225–64.

16. van de Veerdonk FL, Smeekens SP, Joosten LA, Kullberg BJ, Dinarello CA, van der Meer JW, et al. Reactive oxygen species independent activation of the IL-1beta inflammasome in cells from patients with chronic granulomatous disease. Proc Natl Acad Sci U S A. 2010;107(7):3030–3.

17. Seger RA. Modern management of chronic granulomatous disease. Br J Haematol. 2008;140(3):255–66.

18. Kelly J, Pinto A, Whitelaw W, Borstad O, Bowen T, Matheson D. Fatal Aspergillus pneumonia in chronic granulomatous disease. Am J Clin Pathol. 1986;86(2):235–40.

19. Siddiqui S, Anderson VL, Hilligoss DM, Abinun M, Kuijpers TW, Masur H, et al. Fulminant mulch pneumonitis: an emergency presentation of chronic granulomatous disease. Clin Infect Dis. 2007;45(6):673–81.

20. Ameratunga R, Woon ST, Vyas J, Roberts S. Fulminant mulch pneumonitis in undiagnosed chronic granulomatous disease: a medical emergency. Clin Pediatr (Phila). 2010;49(12):1143–6.