CASE REPORT

Aggressive treatment in paediatric or young patients with drug-induced hypersensitivity syndrome (DiHS)/drug reaction with eosinophilia and systemic symptoms (DRESS) is associated with future development of type III polyglandular autoimmune syndrome

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SUMMARY

We experienced a 6-year-old case of drug-induced hypersensitivity syndrome (DiHS)/drug reaction with eosinophilia and systemic symptoms (DRESS) with subsequent development autoimmune thyroiditis (Hashimoto’s thyroiditis), type 1 diabetes with antithyroglobulin, thyroid peroxidase, insulinoma-associated antigen and anti-insulin antibodies at 4 months, alopecia at 7 months, vitiligo, uveitis due to Vogt-Koyanagi-Harada disease at 8 months after clinical resolution of the DiHS/DRESS. He was diagnosed as type III polyglandular autoimmune syndrome (PASIII) after DiHS/DRESS. Prompted by this case, we sought to determine which triggering factors were responsible for later development of PASIII in previously published cases with autoimmune sequelae. In the literature review, five patients with DiHS/DRESS were found to develop autoimmune sequelae consistent with PASIII. All cases with PASIII were much younger than those without them. Four out of the five patients were treated with intravenous immunoglobulin or pulsed prednisolone in the acute stage, although effective in short-term outcomes.

BACKGROUND

Type III polyglandular autoimmune syndrome (PASIII) is a disorder characterised by the coexistence of more than two glandular autoimmune diseases, such as type 1 diabetes.1 Little information is available on critical triggers in the pathogenesis of PASIII in genetically predisposed individuals. Of note, very frequent component autoimmune diseases of PAS III reportedly occur as sequelae of a distinct type of severe drug eruption, drug-induced hypersensitivity syndrome (DiHS)/drug reaction with eosinophilia and systemic symptoms (DRESS); this syndrome is characterised by sequential activations of herpes viruses due to functional defects in regulatory T cells (Tregs).2–5 Paediatric DiHS/DRESS is often misdiagnosed as Kawasaki disease due to similar clinical presentations and treated with intravenous immunoglobulin (IVIG).

We describe a child case of DiHS/DRESS with subsequent development of PASIII. In this case, IVIG used would have served to accelerate progression to PASIII, although beneficial in short-term outcomes. Surprisingly, all previously published DiHS/DRESS cases with autoimmune sequelae had clinical features consistent with PASIII and most had been treated with pulsed prednisolone and/or IVIG in the acute stage.

CASE PRESENTATION

Case report

A 6-year-old boy presented with erythematous lesions on his trunk and extremities. Two months prior to this visit, he suffered a fracture of the right upper arm and received a surgical operation. After the operation, he developed osteomyelitis and was treated with sulfamethoxazole–trimethoprim. Forty-five days later, erythematous plaques began on his extremities and trunk. Sulfamethoxazole–trimethoprim was withdrawn, but his conditions became worse and the patient was referred for further diagnostic workup and treatment to our hospital.

On examination, the patient had numerous 2–5 cm purplish and erythematous plaques on the back and extremities (figure 1A). The patient was hospitalised with fever of 38°C. Physical examination revealed leucocytosis (14.5X109/L) with atypical lymphocytosis (2.9X109/L), but no eosinophilia (0.5X109/L), liver dysfunction (alanine aminotransferase (2.9X103/L), but no eosinophilia (0.5X103/L), liver dysfunction (alanine aminotransferase (ALT) 357 U/L) and cervical lymphadenopathy (figure 2). A skin biopsy revealed a spongiosis change of the epidermis with scattered necrotic keratinocytes and a mild perivascular infiltrate. The diagnosis of DiHS/DRESS was made and oral prednisolone, 30 mg/day, was begun. The patient, however, once again experienced the similar erythematous plaques with a high-grade fever and a marked increase in liver enzyme levels (ALT 1857 U/L). On the 12th day of hospitalisation, he received IVIG, at a dose of 1g/kg/day for 2 days given his low immunoglobulin (Ig)G levels (386 mg/dL). After starting IVIG, he showed clearing of
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lesions and rapid improvement in liver enzyme levels. Because of frequent flaring of liver dysfunction and rashes, we decided to taper prednisolone more gradually over a prolonged period of time (5 mg/day per 2–3 week).

After 4 months, the patient was again admitted to our hospital for nausea, fatigue and excessive thirst. Laboratory tests revealed a marked increase in fasting blood sugar, 580 mg/dL, haemoglobin A1c, 10.2%, anti-insulinoma-associated antigen 2 (IA-2) antibody titre 1.0 U/mL (normal <0.4 U/mL) and anti-insulin antibody titre 50.0 U/mL (<0.4 U/mL). He was diagnosed as having type 1 diabetes mellitus and treatment with insulin was initiated. Because laboratory tests also revealed an increase in antithyroglobulin antibody 36 IU/mL (<28 IU/mL) and antithyroid peroxidase antibody titre 113 IU/mL (<16 IU/mL), he was also diagnosed with Hashimoto’s disease. The doses of prednisolone were gradually reduced and eventually withdrawn on day 180.

On days 215–243, he developed alopecia and vitiligo on his back and upper extremities (figure 1B). On day 243, he complained of impaired vision in both eyes. Fundus examination showed multiple areas of neurosensory detachments and focal areas of multiple confluent yellow-coloured placoid choroidal lesions. A diagnosis of concurrent development of Vogt-Koyanagi-Harada disease and prednisolone was again increased to a dose of 5 mg/day. He developed alopecia totalis with generalised vitiligo 21 months after onset of disease (figure 1C).

INVESTIGATIONS
Prompted by this case, we investigated whether autoimmune diseases with various autoantibodies occurring in reported DiHS/DRESS cases could have clinical features consistent with PASIII. As shown in table 1, five published cases sequentially developing PASIII including this case, were found, although none have reported these autoimmune diseases in the name of PASIII. The mean age of these five cases (21±13.4 years) was much younger than that in our 55 institutional cases with DiHS/DRESS (54.5±2.7 years) (manuscript submitted): four of the five cases were categorised as paediatric (<18 years) DiHS/DRESS. Among the three subcategories of PASIII, the present case met the criteria for A and C type of PASIII.

DIFFERENTIAL DIAGNOSIS
There are a number of differential diagnostic considerations in children; they include Kawasaki disease and viral eruptions, which show unexplained illness that includes rash, fever and lymphadenopathy. In particular, there is an emerging recognition that some children with DiHS/DRESS have been misdiagnosed...
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as Kawasaki disease and erroneously treated by high doses of IVIG within the first 10 days of fever onset.

**TREATMENT**

Although clinical experience suggested that high doses of IVIG or pulsed corticosteroids were effective in abrogating the acute inflammation of DiHS/DRESS, our case and previously reported cases with PASIII suggested that these aggressive therapies in paediatric patients paradoxically increased the risk of future development of autoimmune sequelae, particularly PASIII. The potential reward should outweigh the potential risk, especially in a long-term outcome.

**OUTCOME AND FOLLOW-UP**

Most importantly, pulsed corticosteroids or IVIG was used for treatment of DiHS/DRESS during the acute stage in four of the five reported DiHS/DRESS cases who eventually developed PASIII, except for Case 1, in whom detailed information is not available. Although current literature recommends, DiHS/DRESS cases treated with these aggressive therapies in the acute stage, especially paediatric cases, developed PASIII several months after successful resolution of initial clinical symptoms.9-12

**DISCUSSION**

Which triggering factors are responsible for later development of PASIII in genetically predisposed individuals after clinical resolution of DiHS/DRESS? Of note, the development of PASIII occurred after administration of a short course of pulsed prednisolone in three of the five reported cases (table 1). Some clues may come from studies of immune reconstitution inflammatory syndrome (IRIS), where a rapid immune recovery on withdrawal or reduction of immunosuppressive agents, paradoxically causes extensive pathology.13 Because a rapid and great reduction of prednisolone doses, for example, from 1 g/day to 60 mg/day over a period of 2 days, is necessarily needed in pulsed prednisolone therapy, thereby increasing the risk of IRIS, pulsed prednisolone therapy for DiHS/DRESS may serve to accelerate aggressive cytotoxic immune responses causing extensive pathology.

Although our patient’s young age is unusual in patients with DiHS/DRESS and PASIII, another unusual aspect was its low levels of IgG followed by a marked increase in the levels 1 month later, at which time its levels usually return to normal; this suggests that marked rebound of IgG synthesis may have occurred around the time of IVIG administration in this case: thus, the timing of IVIG administration was likely to coincide with the timing of the rebound and with the peak of liver dysfunction. In this regard, we previously demonstrated that both severe liver damage and defective T reg function in combination could be needed for the subsequent generation of autoantibodies in patients with DiHS/DRESS.14 In our patient, given the clinical course of the disease, IVIG would have been administered at or immediately after the peak of liver damage and rebound in IgG, A and M synthesis. This timing of IVIG administration may allow autoreactive B cells to function in an uncontrolled fashion: IVIG administration at the peak of severe liver damage could further accelerate the rapid recovery of B cells, thereby causing the subsequent expansions of autoreactive B cells with the specificity to a variety of epitopes. Supporting this hypothesis, De Grandmont et al observed that the addition of IVIG in culture of CD40L-stimulated B cells accelerated the differentiation of part of peripheral B cells into IgG-producing plasma cells in the presence of interleukin (IL)-4 and IL-10: importantly, secreted IgGs were shown to be reactive with antigens such as nucleoprotein, dsDNA, tetanus toxin and human IgG F(ab')2 fragments. The action of IVIG on DiHS/DRESS may be different depending on the state of activation of the B cells exposed to IVIG.15

In conclusion, paediatric patients with DiHS/DRESS exhibit an increased risk for subsequently developing PASIII. We
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**Table 1** Characteristics of patients who developed PAS after drug-induced hypersensitivity syndrome/drug reaction with eosinophilia and systemic symptoms

| Patient no. | Age (y) | Sex | Suspected medications | Systemic corticosteroid therapy (mg/day) | Additional therapy | PAS type | Symptoms of PAS | Autoantibodies | Occurrence of autoimmune sequelae and detection of autoantibodies (weeks or months after onset) |
|------------|---------|-----|-----------------------|-----------------------------------------|-------------------|---------|----------------|----------------|----------------------------------------------------------------------------------------------------------------------------------|
| 1	extsuperscript{a} | 15/F    | Minocycline | PSL 40 mg; escalation dose from 10 mg to 40 mg for 4 days | None | IIIA | Autoimmune thyroiditis, autoimmune-associated type 1 diabetes | Anti-TPO, TG, TSI, TSHR, GAD, IA2, ANA, 5m and SS-A antibodies | 7 weeks: autoimmune thyroiditis in the thyrotoxic phase with anti-TG and TPO antibodies. 5 months: Graves disease with anti-TSI and TSHR antibodies. 7 months: type 1 diabetes with anti-GAD and IA2 antibodies. ANA, anti-5m and SS-A antibody titres were elevated. |
| 2	extsuperscript{a} | 17/F    | Zonisamide | Pulsed PSL; 1 g of mPSL for 3 days followed by 45 mg of PSL | None | IIIA +C | Autoimmune thyroiditis, immune-mediated type 1 diabetes, alopecia, vitiligo. Photophobia, vision impairment, uveitis due to Vogt-Koyanagi-Harada disease | Anti-TPO, TG and TSHR antibodies | 2 months: type 1 diabetes, autoimmune thyroiditis (Hashimoto’s thyroiditis) with anti-TPO, TG and TSHR antibodies. 5 months: alopecia. 6 months: vision impairment. 9 months: vitiligo. 10 months: Vogt-Koyanagi-Harada disease. |
| 3	extsuperscript{a} | 21/F    | Lamotrigine | mPSL 160 mg (40 mg every 6 hours) followed by 80 mg of PSL | IVIG | IIIA | Autoimmune thyroiditis, immune-mediated type 1 diabetes | Anti-TPO, anti-insulin and IA2 antibodies | 5 months: type 1 diabetes with anti-insulin and IA2 antibodies. 8 months: autoimmune thyroiditis. |
| 4	extsuperscript{a} | 46/F    | Zonisamide | Pulsed PSL, 1 g of mPSL for 3 days | IVIG | III | Alopecia, autoimmune thyroiditis, Raynaud’s phenomenon, sclerodermatous lesions | ANA, RF, anti-ss-DNA, centromere, cardiolipin and TSHR antibodies | 6 months: alopecia. 3 years: thyroid dysfunction, Raynaud’s phenomenon. 4 years: sclerodermatous lesions with increased titres of ANA. |
| Five present case | 6/M     | Sulfamethoxazole/trimethoprim | PSL 36 mg (1.5 mg/kg/day) | IVIG | IIIA +C | Autoimmune thyroiditis, immune-mediated type 1 diabetes, alopecia, vitiligo, uveitis | Anti-TG, TPO, IA-2 and anti-insulin antibodies | 4 months: autoimmune thyroiditis (Hashimoto’s thyroiditis), type 1 diabetes with anti-TG, TPO, IA-2 and anti-insulin antibodies. 7 months: alopecia. 8 months: vitiligo, uveitis due to Vogt-Koyanagi-Harada disease. |

For the current patient no. 1, the patient was treated with minocycline, and the systemic corticosteroid therapy included prednisolone (PSL) at a dose of 40 mg, escalation dose from 10 mg to 40 mg for 4 days. Additional therapy included none. The PAS type was IIIA, and the symptoms of PAS included autoimmune thyroiditis, autoimmune-associated type 1 diabetes. The autoantibodies detected were anti-TPO, TG, TSI, TSHR, GAD, IA2, ANA, 5m and SS-A antibodies. The occurrence of autoimmune sequelae and detection of autoantibodies were 7 weeks, 5 months, and 7 months, respectively.

For the current patient no. 2, the patient was treated with zonisamide, and the systemic corticosteroid therapy included pulsed PSL, 1 g of mPSL, followed by 45 mg of PSL. Additional therapy included none. The PAS type was IIIA +C, and the symptoms of PAS included autoimmune thyroiditis, immune-mediated type 1 diabetes, alopecia, vitiligo. Photophobia, vision impairment, and uveitis due to Vogt-Koyanagi-Harada disease. The autoantibodies detected were anti-TPO, TG and TSHR antibodies. The occurrence of autoimmune sequelae and detection of autoantibodies were 2 months, 5 months, 6 months, and 9 months, respectively.

For the current patient no. 3, the patient was treated with lamotrigine, and the systemic corticosteroid therapy included mPSL, 160 mg (40 mg every 6 hours) followed by 80 mg of PSL. Additional therapy included IVIG. The PAS type was IIIA, and the symptoms of PAS included autoimmune thyroiditis, immune-mediated type 1 diabetes. The autoantibodies detected were anti-TPO, anti-insulin and IA2 antibodies. The occurrence of autoimmune sequelae and detection of autoantibodies were 5 months and 8 months, respectively.

For the current patient no. 4, the patient was treated with zonisamide, and the systemic corticosteroid therapy included pulsed PSL, 1 g of mPSL, followed by 45 mg of PSL. Additional therapy included IVIG. The PAS type was III, and the symptoms of PAS included alopecia, autoimmune thyroiditis, Raynaud’s phenomenon, sclerodermatous lesions. The autoantibodies detected were ANA, RF, anti-ss-DNA, centromere, cardiolipin and TSHR antibodies. The occurrence of autoimmune sequelae and detection of autoantibodies were 6 months, 3 years, and 4 years, respectively.

For the current patient no. 5, the patient was treated with sulfamethoxazole/trimethoprim, and the systemic corticosteroid therapy included PSL, 36 mg (1.5 mg/kg/day). Additional therapy included IVIG. The PAS type was IIIA +C, and the symptoms of PAS included autoimmune thyroiditis, immune-mediated type 1 diabetes, alopecia, vitiligo, uveitis. The autoantibodies detected were anti-TG, TPO, IA-2 and anti-insulin antibodies. The occurrence of autoimmune sequelae and detection of autoantibodies were 4 months, 7 months, and 8 months, respectively.

*ANA, antinuclear antibodies; F, female; GAD, glutamic acid decarboxylase; IA2, insulinoma-associated antigen-2; IVIG, intravenous immunoglobulin; M, male; PAS, polyglandular autoimmune syndrome; PSL, prednisolone; TG, thyroglobulin; TPO, thyroid peroxidase; TSHR, thyroid stimulating hormone receptor; TSI, thyroid stimulating immunoglobulin.*
recommend that patients with DiHS/DRESS, especially paediatric patients, be screened for the generation of various autoantibodies and that at-risk patients be advised to avoid therapies that can accelerate rapid recovery of B cells and T cells, such as IVIG and pulsed prednisolone in the acute stage.

Learning points

► In literature review, five patients with drug-induced hypersensitivity syndrome (DiHS)/drug reaction with eosinophilia and systemic symptoms (DRESS) were found to develop autoimmune sequelae consistent with type III polyglandular autoimmune syndrome (PASIII) 2–7 (average 4.8) months after onset. Four out of the 5 patients were treated with intravenous immunoglobulin or pulsed prednisolone in the acute stage.

► Paediatric or younger patients with DiHS/DRESS have the greatest risk for subsequently developing PASIII, when treated with intravenous immunoglobulin or pulsed prednisolone that can accelerate rapid recovery of B and T cells in the acute stage, although beneficial in the short-term outcome.

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