Phenytoin-Induced Hypersensitivity Syndrome Presenting on the Palms and Soles of an 18-Year-Old Woman

Case Report

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Introduction

Anticonvulsant drug-induced hypersensitivity syndrome (DIHS) is a disorder that occurs in some patients taking anticonvulsant medications. Fever is the most common feature, seen in 90-100% of cases. It is also characterized by rash, hepatitis, and other multiorgan involvement. The mechanism is unknown. Implicated anticonvulsants include those in the phenytoin category, carbamazepine, and lamotrigine[1]. This syndrome is now most commonly known as Drug Reaction (or Rash) with Eosinophilia and Systemic Symptoms (DRESS), a term more descriptive of its presentation. It is also known as Drug-Induced Delayed Multorgan Hypersensitivity Syndrome (DIDMOHS) and Drug-Induced Pseudolymphoma. Anticonvulsant hypersensitivity syndrome is an older term that has lost favor due to the fact that, aside from antiepileptics, other medications such as sulfonamides and allopurinol have been implicated. We report an 18-year-old woman who presented with drug-induced hypersensitivity syndrome secondary to phenytoin.

Case Presentation

An 18-year-old woman with a newly diagnosed seizure disorder presented to the emergency room with nausea, vomiting, and an erythematous eruption on the face, neck, palms, and soles (Figures 1-4). Three weeks prior to her presentation, she was started on phenytoin 100 mg orally daily. Ten days later, she began to experience pruritus without rash. She visited an urgent care clinic and was prescribed a 5-day prednisone taper. This relieved the itching somewhat. However, she woke up two days later and noticed an erythematous eruption on her face, palms, and soles. She had associated nausea, profuse vomiting, and abdominal pain. The abdominal pain was diffuse, 8 out of 10 in severity, nonradiating, and slightly relieved with analgesics. She denied dysuria or hematuria. She denied sexual activity, recent travel, and recent new foods.

Cutaneous examination revealed erythematous papules coalescing into plaques on the perioral region involving the nasolabial folds (Figure 1). There was a similar distribution on the nape of her neck (Figure 2). She also presented with lesions on her palms and soles that were more focal (Figure 3-4). Her oral mucosa consisted of erosions without vesicles. She was found to have a temperature of 103.1°F, respiratory rate of 20, and mild leukocytosis of 11.3. Differential revealed 8.9 x 10³ granulocytes/microliter, 1.2 x 10³ lymphocytes/microliter, 0.9x x 10³ monocytes/microliter, and 0.3x x 10³ eosinophils/microliter. Her complete metabolic panel was negative. Chlamydia and gonorrhea DNA probes were negative. Monospot was negative. Throat culture revealed usual oral flora. Rapid plasma reagin (RPR) test was negative.

The patient was admitted with a working diagnosis of phenytoin-induced exanthem versus enterovirus infection with coxsackie A16 (“hand-foot-and-mouth syndrome”). Stevens-Johnson syndrome was also being considered. A punch biopsy of a right upper arm lesion was performed. The results revealed superficial perivascular dermatitis (Figures 5-6) with focal interface inflammatory component (Figure 7), favoring drug eruption over viral exanthem.

The patient was started on methylprednisolone 40 mg intravenously (IV) every eight hours and diphenhydramine 25 mg IV as needed for itching. She did not enter anaphylaxis and did not require epinephrine or albuterol. Her eruption regressed significantly over three days in the hospital. She was discharged with hydrocortisone 1% cream to be applied twice daily as well as oral diphenhydramine for pruritis. Her eruption completely cleared two weeks after discharge.

Discussion
Figure 1. An 18-year-old female presents with erythematous plaques in the perioral region involving the nasolabial folds shortly after starting phenytoin.

Figure 2. An 18-year-old woman developed erythematous papules coalescing into plaques shortly after starting phenytoin. The lesions on the back of her neck can be seen in this image.

Figure 3. An 18-year-old woman developed erythematous papules on her palms shortly after starting phenytoin.

Figure 4. The 18-year-old woman also presents with erythematous papules on her soles.

Figure 5. This low-powered hematoxylin and eosin (H&E) stain demonstrates superficial perivascular dermatitis.

Figure 6. This high-powered H&E stain further exhibits the superficial perivascular dermatitis.
DIHS has been implicated in many drugs. The most common culprits are antiepileptics (phenytoin, phenobarbital, carbamazepine, oxcarbazepine, and lamotrigine), 64% of which were estimated in one study [2]. Other causes included allopurinol, dapsone, vancomycin, leflunomide, and nitrofurantoin. Multiple other drugs have been described in the literature. One 14-month-old child developed DIHS due to chloral hydrate [3].

The incidence of DIHS is unknown due to its unpredictable clinical presentation. The onset and timing of DIHS is extremely variable, which has made its report in the literature difficult. Some individuals develop signs and symptoms as early as two weeks, while others may not experience them until 3 months after starting a new antiepileptic medication [1]. Nevertheless, this syndrome has a much longer latency than morbilliform drug eruptions (mean of 10 days) and type-1 hypersensitivity syndromes (within a day). The liver is the most common organ involved in DIHS, seen in 80% of patients. This is followed by the kidneys, lungs, heart, muscles, and pancreas [4]. Liver involvement in the form of hepatitis is usually mild but can be severe. The mortality rate is between 18 and 40% if hepatitis is present [1]. Liver function tests (LFTs) may be extremely elevated and continue to rise even after the drug has been discontinued. It may take up to one year for them to return to baseline. Half of all patients with DIHS have hematological abnormalities. The most common is atypical lymphocytes (63%), followed by cosinophilia (52%), lymphocytopenia (25%), thrombocytopenia (25%), and lymphocytosis (25%) [4].

The cutaneous findings of DIHS usually begin similarly to a morbilliform drug eruption: erythematous, diffuse macules and papules. These start on the face and trunk. Erythematous facial edema occurs in half of patients. Mucosal involvement of the mouth and pharynx may occur; erosions are rarely seen. 50% of the body surface area is usually involved [5].

One study of the pharmacogenetics of antiepileptic-induced DIHS investigated associations between HLA alleles and several antiepileptic drugs in diverse populations. The study showed that HLA-B*15:02 was associated with carbamazepine, lamotrigine and phenytoin-induced Stevens-Johnson syndrome in Asian populations. The authors suggest to pretreat such individuals prior to the start of antiepileptic medications [6]. In addition, the pathogenesis of DIHS has been closely linked with herpesvirus reactivation. In a study of 100 patients with DIHS, an increase in the antibody titer against HHV-6 was detected in 60% of patients two to four weeks after symptom onset [7]. Another study found that EBV, HHV-6, and HHV-7 reactivation occurred in 29 of 40 patients with DIHS; CD8+ lymphocytes specific to these viruses were found in the blood, skin, and liver [8]. This suggests that DIHS may be due to viral reactivation and cross-reaction of T-cells with the offending drug.

Under light microscopy, DIHS usually reveals nonspecific spongiosis, a perivascular lymphocytic infiltrate with or without cosinophils (Figures 5 – 6), interface inflammation (Figure 7), and dermal edema [9]. Due to the presence of atypical lymphocytes in the infiltrate, the differential may include cutaneous lymphoma. Biopsies of involved lymph nodes may also demonstrate atypical lymphocytes, but the features are usually benign. In the case of liver involvement, biopsy reveals lobular inflammation, portal inflammation, scattered foci of necrotic hepatocytes, and granulomatous infiltrates containing cosinophils. Portal inflammation and cholestasis also may be seen. Confluent hepatocyte necrosis and lobular disarray due to inflammation and regenerative changes are seen in severe cases. Kidney biopsy reveals tubulointerstitial nephritis (TIN) patterns.

DIHS is a clinical diagnosis. Many criteria have been proposed for diagnosis. RegiSCAR criteria include at least 3 of the following 7 characteristics: 1) skin eruption, 2) fever (>38°C), 3) lymphadenopathy of at least 2 sites, 4) involvement of at least 1 internal organ, 5) lymphocytosis (>4×10³/µL) or lymphocytopenia (<1.5×10³/µL), 6) blood cosinophilia (>10% or 700/µL), and 7) thrombocytopenia (<120×10³/µL). The 18-year-old woman presented with fever, lymphopenia, and the exanthem, thus satisfying DIHS by RegiSCAR criteria.

Bocquet's criteria require meeting the following 3 features: 1) skin eruption, 2) blood cosinophilia (>1.5×10³/µL) or the presence of atypical lymphocytes, and 3) internal organ involvement, including lymphadenopathy (>2 cm in diameter), hepatitis (liver transaminases values > twice the upper normal limit), interstitial nephritis, and interstitial pneumonia or carditis.

The Japanese consensus group diagnostic criteria for DIHS includes the following features: 1) eruption of macules and papules developing at least 3 weeks after starting the medication, 2) prolonged clinical symptoms 2 weeks after discontinuing the drug, 3) fever (>38°C), 4) elevation of liver enzyme (alanine aminotransferase [ALT] >100 U/L) or involvement of other organs, 5) leukocytosis (>11×10³/µL), atypical lymphocytosis (>5%) or cosinophilia (>1.5×10³/µL), 6) lymphadenopathy, and 7) human
herpesvirus (HHV)-6 reactivation. Diagnosis of typical DIHS requires the presence of all 7 criteria. Atypical DIHS is diagnosed in patients with the first five listed criteria.

One study compared patients according to the three criteria. A total of 60.4% of patients satisfied RegiSCAR definitive criteria and 77.1% satisfied Bocquet's criteria. Only 18.8% satisfied atypical DIHS criteria from the Japanese group. They concluded that Bocquet's criteria is most appropriate because of its efficiency in clinical practice. Lymphocyte and eosinophil counts, as well as creatinine and ferritin levels, could be useful early prognostic factors.


The primary management of DIHS is to remove the offending drug. If the manifestation is not severe enough to result in end organ involvement, then symptomatic treatment is preferred: antihistamines for pruritis and topical corticosteroids for skin manifestations. The most effective treatment for DIHS with organ involvement is systemic corticosteroids. In one study of 17 patients, 13 patients treated with oral prednisone recovered completely, two developed post-inflammatory hyperpigmentation (PIH), one developed renal failure, and one patient developed liver failure.

It is important to note that systemic corticosteroids have had no proven benefit for liver manifestations of DIHS. Nor has a treatment duration been definitively determined.

Post-treatment monitoring has not been clearly outlined in the literature, but it is suggested to perform weekly blood work to assess for end organ involvement and resolution: complete blood count with differential and complete metabolic panel. The prognosis of this disease is usually favorable, with only a five to 10% mortality that is often related to severe end organ involvement.

**Conclusion**

Anticonvulsant drug-induced hypersensitivity syndrome (DIHS) is a rare but sometimes fatal disorder. It is characterized by fever, rash, hepatitis, and other multiorgan involvement. Implicated anticonvulsants include phenytoin, carbamazepine, phenobarbital, and lamotrigine. The incidence of DIHS is unknown due to its unpredictable clinical presentation. Onset occurs between 2 weeks and 3 months of starting the offending agent. Studies of genetics and heritability propose an association with HLA-B*15:02. Three diagnostic criteria include RegiSCAR, Bocquet's, and the Japanese consensus group diagnostic criteria for DIHS. Removal of the offending agent and symptomatic treatment are the mainstay of management. An 18-year-old woman presented with nausea, vomiting, and an erythematous eruption on the face, neck, palms, and soles two weeks after starting phenytoin. Her symptoms resolved after a short hospitalization, systemic corticosteroids, and subsequent symptomatic topical treatment.

**References**


