Amoxicillin Morbilliform Drug Eruption in Pediatric Male with Poor Feeding Treated with Ciproheptadine: A Case Report

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Abstract
Background: Ciproheptadine (CY) is an antihistaminic agent that is commonly used for symptom relief in skin conditions. The most common pattern of cutaneous drug eruption in children is the exanthematous type, with the penicillin family often cited. CY is also an antiserotoninergic agent with the side effect of appetite stimulation and has been used in children with poor feeding and poor weight gain. The Case: We report a case of a 31-month-old male patient seen in the outpatient setting with a diffuse morbilliform rash after use of amoxicillin for right otitis media. The patient was a post-operative congenital heart disease (CHD) patient, actively being treated with CY for feeding difficulties and low weight often seen in the CHD population. Amoxicillin was discontinued, while CY was continued. The patient did not encounter any pruritic symptoms during morbilliform rash, while weight gain of 3.1 kg occurred over a 6-month period, increasing patient from the 10th to 41st percentile. Conclusion: A review of studies on CY has shown antiallergic properties in histamine-mediated hypersensitivity reactions, most likely through H1 receptor antagonism. This mechanism may be used to address the pruritic symptoms during type IV T-cell mediated hypersensitivity cutaneous drug eruptions. CY also possesses 5-HT receptor antagonist properties with demonstrated ability to increase appetite in poor feeding pediatric patients. CY was successfully used for this purpose in our CHD patient.

Key Words: Amoxicillin; Exanthema; Ciproheptadine; Heart Defects; Congenital; Appetite (Source: MeSH-NLM).

Introduction
Pediatric drug-related cutaneous reaction patterns are numerous, but can generally be classified into five types: exanthemat, fixed drug eruptions, urticarial eruptions, serum sickness-like reactions, and photosensitve eruptions.3 The most common pattern of cutaneous drug eruption in children is the exanthematous type.4 The penicillin family is often implicated, with amoxicillin a frequent culprit.1

Drug reactions can occur in patients with co-morbidities, complicating the management. One potential comorbidity is failure to thrive, which may indicate the need for an appetite stimulant.5 Ciproheptadine (CY) is an antihistaminic and antiserotoninergic agent that has United States Food & Drug Administration (FDA) approval for indications of allergic conditions, with beneficial side effects of appetite stimulation.2

We present a patient being chronically treated with CY, who continued therapy to address a separate acute condition. The use of CY is reviewed for our patient’s comorbidities, an amoxicillin adverse drug reaction and feeding difficulties in repaired congenital heart disease (CHD) patient.

The Case
We evaluated a 31-month-old Asian male in the clinic with a chief complaint of rash. The rash onset began a day before with no particular pattern (i.e., cephalocaudal), and described as non-pruritic, non-tender, worsening rash all over the body. Parents were unaware of any previous penicillin exposure while reporting up-to-date immunization status. Parents deny history of fevers, chills, cough, nausea, vomiting, diarrhea, allergies, or any other previous rash. Family history not significant for allergies or similar rashes. Recent upper respiratory infection of acute otitis media (AOM) was diagnosed in the patient eight days earlier, but no recent history of pharyngitis noted. Past medical and surgical history involved CHD diagnosed at 4 months of age, specifically atrioventricular canal defect, hypoplastic aortic arch, and patent ductus arteriosus. The patient was hospitalized at the time of discovery for medical management of heart failure, pulmonary overload, and failure to thrive (admission weight 5.3 kg [2.3 percentile, Z-score = -2.0, CDC]) (Figure 1) with nutrition and diuresis optimized in anticipation of surgical repair. Complete repair performed at 6 months of age with no major issues in the operation room. Post-surgical complications developed for hypomobile left vocal cord, oral aversion, and poor weight gain.

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Medical interventions for low weight was initiated at 11 months of age (weight of 7.65 kg [0.4 percentile, z-score = -2.7, CDC]) with nutrition management, and consultations for dietitian, gastroenterology, speech therapy, physical therapy, and occupational therapy. At 19 months of age, weight gain progressed to 9.84 kg (3.4 percentile, z-score = -1.8, CDC) but increases faded 3 months later (22 months of age) at a weight of 10.25 kg (3.2 percentile, z-score = -1.8, CDC). At this point, the decision was made to start oral cyproheptadine. Medications at time of current clinic visit included amoxicillin (7 ml of a 400mg/5ml suspension, twice a day) on day 9 of a 10-day course for recent AOM and CY (2.1 ml of a 2mg/ml syrup, twice a day) for feeding difficulties. Besides current amoxicillin treatment regimen, no previous penicillin antibiotics noted in medical records. Vital signs at current visit showed temperature of 37.0 degrees Celsius, heart rate of 116 beats per minute, respiratory rate of 26 breaths per minute, and a recorded weight and height of 13.35 kg (41 percentile, z-score = -0.2, CDC), and 91.44 cm (44 percentile, z-score = -0.2, CDC), respectively, (Figure 1 and Figure 2).

Figure 1. Patient Weight-For-Age Clinical Growth Chart (CDC) and Initiation of Cyproheptadine. *Initiation of Cyproheptadine at 22 Months of Age.

History and physical exam, including timeline of amoxicillin administration, lead to a clinical diagnosis of adverse reaction to penicillin. Management involved consideration for discontinuation of medications. Amoxicillin was stopped due to high suspicion as the causative agent of the rash. Further antibiotic treatment deemed unnecessary as resolution of AOM was noted per physical exam. With a low index of suspicion for CY as the underlying etiology of the rash, the decision to continue CY was made. CY outpatient therapy was maintained for management of the dermatologic symptoms, including itch while sustaining enhanced appetite stimulation and weight gain effect. Patient’s parents were in agreement to therapeutic plan as they reported good compliance with CY, positive response in terms of improved weight gain and absence of pruritic symptoms. Weight at current encounter was a 1.95 kg increase from the time of CY initiation. Per discussion with patients at a subsequent unrelated encounter, rash continued for 2-3 days after discontinuing amoxicillin and cleared approximately a week later.

Figure 3. Morbilliform Rash on Patient’s Ventral Trunk.

No bullous lesions or desquamation was noted. Differential diagnosis included measles, rubella, scarlet fever, infectious mononucleosis associated amoxicillin rash, and unlikely, Kawasaki disease, anaphylaxis, and Stevens-Johnson syndrome - toxic epidermal necrolysis.
Still, antihistamines are commonly used in the management of type IV adverse drug reactions for dermatological symptoms. A likely benefit is the antihistamines sedative effects leading to a reduction in itch, especially at night. The natural course of an amoxicillin rash is a self-limiting variable, with resolution of rash reported 7-15 days after therapy cessation. Our patient’s rash continued for approximately 10 days after amoxicillin withdrawal, consistent with normal rash resolution. Additional consideration for an unclear diagnosis of a rash that may be a type I IgE-mediated hypersensitivity, could benefit from CY and it’s FDA indication as adjunctive therapy, in addition to standard measures.

In regard to off-label appetite stimulant evidence found by De Bruyne et al for CY in pediatric patients, there were two retrospective studies. A retrospective chart review found 82 children with low appetite and poor growth who took CY regularly, where 96% of parents reported a positive change in mealtime and feeding behaviors. Additionally, significant improvement in mean weight-to-age Z scores was observed after starting CY. A retrospective open-label study for the efficacy of CY in dyspeptic children showed positive overall response in 55% of patients. A search of the medical literature did not reveal any reported cases of CY use in a post-operative CHD patient. CHD patients can have increased energy and nutrient needs, complicated by feeding difficulties. As our patient had a multi-disciplinary team addressing nutrition, development/behavioral, psychosocial issues over a course of a year with failure to reach catch-up growth, adjunctive therapy in the form of CY was initiated before considering escalating management to a pediatric feeding program. Cyproheptadine’s antiserotonergic properties is thought to play a major role in appetite stimulation, though the mechanism is not well understood. A hypothesis for the mechanism of action include anticholinergic and antiserotonergic receptors at the ventromedial hypothalamus and CY influence on growth hormone and insulin-like growth factor axis. Either or both of these mechanisms could play a role in increasing appetite and causing weight gain in post-operative CHD patients.

Cyproheptadine has been reported as a safe medication in the pediatric population. Aside from expected increased appetite, reported adverse effects included self-limited somnolence, irritability, and abdominal pain. Our patient’s feeding difficulties resolved after the first few days of therapy. Our case report of CY toxicity in a 5-year-old female described signs and symptoms consistent with anticholinergic toxicity, urinary incontinence, and tachycardia. Our patient had no reported adverse effects from the medication.

In conclusion, this case report reviews studies of CY as an antiallergic and orxigenic agent and CY’s application to a post-operative CHD patient with new-onset amoxicillin morbilliform drug eruption in the background of poor weight gain. Cyproheptadine’s H1 and H2 receptor antagonist properties are the mechanism’s most strongly supporting CY’s use in histamine mediated hypersensitivity reactions and appetite stimulation. We suspect CY may provide some benefit specifically for delayed type IV T-cell mediated hypersensitivity drug reactions and increased feeding in CHD patients. Clinicians can utilize the dual mechanisms of action in their pediatric patients for various allergic indications and/or difficulty feeding and poor weight gain.
References

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