Hand, Foot, and Mouth Disease Caused by Coxsackievirus A6

To the Editor: Coxsackievirus A6 (CVA6) is a human enterovirus associated with herpangina in infants. In the winter of 2012, we evaluated a cluster of 8 patients, 4 months–3 years of age, who were brought for treatment at Boston Children’s Hospital (Boston, MA, USA) with a variant of hand, foot, and mouth disease (HFMD) that has now been linked to CVA6 (Table, Appendix, wwwnc.cdc.gov/EID/article/18/10-0813-T1.htm). During this same period, the Boston Public Health Commission’s syndromic surveillance system detected a 3.3-fold increase in emergency department discharge diagnoses of HFMD. In the United States, HFMD typically occurs in the summer and early autumn and is characterized by a febrile enanthem of oral ulcers and macular or vesicular lesions on the palms and soles; the etiologic agents are most often CVA16 and enterovirus 71.

In contrast to the typical manifestation, the patients in the Boston cluster exhibited symptoms in late winter (Table, Appendix) and had perioral (Figure, panel A) and perirectal (Figure, panel B) papules and vesicles on the dorsal aspects of the hands and feet (Figure, panel C). Patients experienced a prodrome lasting 1–3 days, consisting of fever (8 patients), upper respiratory tract symptoms (4 patients), and irritability (7 patients). This prodrome was followed by the development of a perioral papular rash (8 patients), which was often impetiginized with secondary crusting; a prominent papulovesicular rash on the dorsum of the hands and feet (6 patients); and a perirectal eruption (7 patients). Half of the patients had intraoral lesions. Fever abated in most of the patients within a day after onset of the exanthem. The rash resolved over 7–14 days with no residual scarring. Samples from the oropharynx, rectum, and vesicles from these patients were sent to the Centers for Disease Control and Prevention (Atlanta, GA, USA) for analysis. Reverse transcription PCR and sequencing by using primers specific for a portion of the viral protein 1 coding region identified CVA6 (1) (Table, Appendix).

Outbreaks of HFMD caused by CVA6 have been described in Singapore, Finland, Taiwan, and most recently in Japan; most cases have occurred in the warmer months (2–6). Cases in the cluster described here are likely related to an emerging outbreak of CVA6-associated HFMD in the United States (7). The atypical seasonality of the outbreak, during the winter in Boston, could be related to the unusually mild temperatures in the winter of 2012.

Recent CVA6 outbreaks have been characterized by a febrile illness associated with an oral enanthem and lesions on the palms, soles, and buttocks. CVA6 infections in Taiwan during 2004–2009 were associated with HFMD in 13% of cases, with disease defined as oral ulcers on the tongue or buccal mucosa and vesicular rashes on the palms, soles, knees, or buttocks (2). In Singapore, where CVA6 accounted for 24% of HFMD cases, patients had oral lesions and <5 peripheral papules, placing them on a spectrum closer to the herpangina more typically observed in CVA6 infection (8).

The patients we report in this cluster most typically had perioral and perirectal papules in addition to vesicles on the dorsum of their hands. Two reports of CVA6-associated HFMD outbreaks describe cases that more closely resemble patients in the Boston outbreak. In a series from Finland in 2008, representative patients had both perioral lesions and vesicles on the dorsum of their hands (6). In a large series of patients with

HFMD in Taiwan in 2010, patients with CV A6 had perioral lesions in addition to an enanthem (3).

Outbreaks of CV A6-associated HFMD in Finland, Taiwan, and Japan were associated with onychomadesis, with the loss of nails occurring 1–2 months after initial symptoms (3,4,6). The association between more typical HFMD and onychomadesis has additionally been described in the United States and Europe but without a link to specific serotype or with a small percentage of CV A6-associated cases (9). Cases from the Boston epidemic may fit into an emerging clinical phenotype of CV A6, and it will be interesting to see whether nail loss develops in those patients.

Given the numerous CV A6 outbreaks in multiple countries in 2008 and a US population that may be relatively naïve to this serotype, CV A6 is likely to spread throughout North America. Clinicians should be aware that, although standard precautions are routinely recommended for managing enteroviral infections in health care settings, contact precautions are indicated for children in diapers to control institutional outbreaks (10). In addition, the presence of perioral lesions and peripheral vesicles on the dorsum rather than palmar/plantar surface of the hands and feet represents a unique phenotype of HFMD that could be confused with herpes simplex or varicella-zoster virus infections.

Because of the atypical presentation of CV A6-associated HFMD, clinical vigilance is needed to recognize emerging regional outbreaks. More detailed epidemiologic and genetic analyses will be required to characterize the role of CV A6 in US outbreaks of HFMD.

Acknowledgments

We thank Richard Rossi and Renee Roy for clinical sample preparation and processing and Kenneth McIntosh for critical review of this manuscript.

A.A.A. is supported by National Institutes of Health grant no. 5 K08 AI093676-02.

Kelly Flett,1 Ilan Youngster,1 Jennifer Huang, Alexander McAdam, Thomas J. Sandora, Marcus Rennick, Sandra Smole, Shannon L. Rogers, W. Allan Nix, M. Steven Oberste, Stephen Gellis, and Asim A. Ahmed

Author affiliations: Children’s Hospital Boston, Boston, Massachusetts, USA (K. Flett, I. Youngster, J. Huang, A. McAdam, T.J. Sandora, S. Gellis, A.A. Ahmed); Boston Public Health Commission, Boston (M. Rennick); Massachusetts Department of Public Health, Jamaica Plain, Massachusetts, USA (S. Smole); and Centers for Disease Control and Prevention, Atlanta, Georgia, USA (S.L. Rogers, W.A. Nix, M.S. Oberste)

References


1These authors contributed equally to this article.
Duffy Phenotype and Plasmodium vivax infections in Humans and Apes, Africa

To the Editor: Benign tertian malaria, caused by Plasmodium vivax, has long been considered absent, or at least extremely rare, in western and central Africa. In these regions, 95%–99% of humans are of the Duffy negative phenotype, a condition that is thought to confer complete protection against the parasite during the blood stages of its life cycle (1,2). Sporadic reports throughout the latter half of the 20th century, however, have hinted at the presence of the parasite in these regions, the most convincing of which were the steady and consistent numbers of non-African travelers who returned to their countries of origin infected with malarial parasites that were subsequently identified as P. vivax (2).

More recently, evidence has emerged regarding the transmission of P. vivax in regions of Africa where the local human population is predominantly Duffy negative (3–6). In 4 (3.5%) of 155 patients from western Kenya (6), 7 (0.8%) of 898 persons from Angola (4), and 8 (8.2%) of 97 persons from Equatorial Guinea (4), P. vivax parasites were detected in the blood of apparently Duffy-negative persons, suggesting that the parasite might not be as absolutely dependent on the Duffy receptor for erythrocyte invasion as previously thought. These findings are supported by a report from Madagascar (where the human population is composed of a mixture of Duffy-positive and Duffy-negative persons), in which 42 (8.8%) of 476 Duffy-negative persons who had symptoms of malaria were reported to be positive for P. vivax by both microscopy and PCR (7). The prevalence of P. vivax in Duffy-negative persons was significantly lower than its prevalence in Duffy-positive persons residing in the same area, suggesting that Duffy negativity is a barrier to the parasite to some degree. Given the extremely high rates of malaria transmission in western and central Africa, a P. vivax parasite that could efficiently invade Duffy-negative erythrocytes would, presumably, become highly prevalent very rapidly.

With the exception of the cases reported from Angola and Kenya, which lie outside the area where the proportion of the population with Duffy negativity is highest, the reports of the transmission of P. vivax within predominantly Duffy-negative populations all come from regions inhabited by chimpanzees and gorillas (i.e., Democratic Republic of the Congo [3], Uganda [4], and Equatorial Guinea [5]). During our seroepidemiologic study from the Democratic Republic of the Congo, in which P. vivax sporozoite–specific antibodies were detected in ≈10% of the population, we found that women were significantly more likely than men to have been exposed to P. vivax sporozoites (3). Women in this region typically spend more time than men near the forest fringe, where they work in crop fields. This forest is within the known habitat range of the chimpanzee Pan troglodytes and the gorilla, Gorilla gorilla gorilla, both of which have been reported to be natural hosts of the malaria parasite P. schweizti, which is a P. vivax–like or P. ovale–like parasite that might also be unable to invade the erythrocytes of persons who are Duffy negative (8). These animals have recently been shown to be infected occasionally with parasites that have mitochondrial genomes closely resembling those of P. vivax (9,10).

We have argued that, given the high malaria transmission rates in sub-Saharan Africa, it is plausible that the 1%–5% of the human population who are Duffy positive might maintain the transmission of the parasite (2). The discovery of P. vivax parasites (or P. vivax–like parasites) in the blood of African great apes leads to a question: could nonhuman primates in Africa be acting as Duffy-positive reservoirs of P. vivax in regions where the human population is almost entirely susceptible? This possibility warrants further investigation. Given the increasing rarity of the great apes, however, their capacity to act as zoonotic reservoirs could be limited. It would be informative, in any case, to determine how the regions that P. vivax–positive travelers visit during their stay in Africa correspond with the ranges of chimpanzees and gorillas.

If African great apes do, indeed, constitute a zoonotic reservoir of P. vivax parasites, what are the