Distinctive Motor Stereotypy in a 4-month-old Infant with Congenital CMV Infection

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Abstract

Background: Abnormal stereotypy, although described in non-human primates with congenital cytomegalovirus infection, has not been described in human infants and neonates with central nervous system cytomegalovirus disease. Case Report: We present a case of a 4-month-old infant male with congenital cytomegalovirus infection who presented for complaints of repetitive and conscious stereotypy of the upper and lower extremities for the majority of the patient’s waking period. Conclusion: Asymptomatic neonates with central nervous system cytomegalovirus infection may go on to develop permanent neurologic sequelae of disease. The early recognition of congenital cytomegalovirus disease is critical to help initiate early treatment interventions. Clinicians should maintain a low threshold for suspicion for TORCH infections and neurologic disorders in infants with an abnormal stereotypy.

Keywords: Central Nervous System Infections, Cytomegalovirus, Infant, Muscle Hypotonia.

Introduction

Congenital cytomegalovirus (CMV) infection remains the principal cause of brain damage in children, as well as being the most common congenital infection in the United States [1,2]. An estimated 40,000 newborns are born with CMV annually in the United States, affecting approximately 1% of all live births [3]. An extensive number of studies have been undertaken to characterize the symptomatic presentation of congenital CMV disease, and various manifestations of central nervous system (CNS) disease have been described in CMV infection [4,5]. Although 50-90% of patients with symptomatic congenital CMV infection will develop long term deficits such as mental retardation or other cognitive defects, there is a substantial percentage of asymptomatic patients (approximately 7-25%) who go on to develop deficits later in life [6-8].

Some studies suggest asymptomatic CMV disease may be associated with a broad range of subtle neurodevelopmental sequelae [9]. This has brought on the need for close identification and follow-up of infants with subtle neurologic symptoms who have otherwise normal physical exams. Careful assessment of development and subtle physical exam findings are critical to identify those who require early interventions for corrective measures [10]. We present a rare case of a 4-month-old male with congenital CMV infection who demonstrated an abnormal repetitive and conscious motor stereotypy as the earliest clinical sign of disease. To the best of our knowledge, an abnormal stereotypy in an infant has not been described as a potential manifestation of intracranial CNS CMV disease.
Case Report

Our case involved a 4-month-old otherwise asymptomatic male infant with congenital CMV diagnosed by urine and blood PCR soon after birth. The patient was found to have intracranial calcifications on head ultrasound at an outside facility. The mother of the patient noted a 2-month history of a distinctive and persistent stereotypy involving high velocity and high amplitude synchronized flexion and extension of the lower extremities. The mother of the patient stated that she swaddled the patient for most of the day as this effectively suppressed the stereotypy. These movements on the mother’s estimation occurred “more than 90%” of the patient’s waking period if the patient was not swaddled to suppress the stereotypy.

Physical exam revealed a well-appearing infant with findings of mild diffuse hypotonia of the upper extremities, hypertonia of the lower extremities, and the above-described stereotypy during the entire duration of the approximately 35-minute clinic visit. Of note, this stereotypy did resolve with the momentary clapping of the feet by the provider during the physical exam and on placing the infant prone with a subsequent near-immediate resumption of the stereotypy on placing the infant back to supine position. The infant was in no apparent distress and was appropriately tracking objects with his eyes during the stereotypy. Findings were not consistent with seizure-like activity and EEG was deferred. A subsequent MRI study was ordered and demonstrated T2 hyperintense signal along the periventricular white matter consistent with CMV infection [Fig.1].

Discussion

The pediatric neurologic exam is an important cornerstone of the well-child visit. Components of the exam include evaluation of the spine for signs of underlying malformation such as tufts of hair and dimples. The pediatric neurologic exam also includes evaluating head size and shape to evaluate for findings such as microcephaly and macrocephaly. The evaluation of the cranial nerves and the reflexes of an infant also allows the provider to assess for appropriate neurologic development during a well-child visit. Finally, evaluation of motor activity primarily involves the assessment of normal spontaneous activity and assessing for deficits in movement.

A motor stereotypy is defined broadly as an involuntary and coordinated repetitive movement that is purposeless [11]. A complex motor stereotypy includes stereotypies that involve the arms and sometimes legs. Stereotypies are often rhythmic and last for seconds to minutes. They can occur in clusters and can be suppressed by distracting a patient or having the patient initiate another action [12]. With the exception of normal stereotypies such as thumb sucking, unique and abnormal stereotypies in infants are rare. Although stereotypies are poorly understood, there has been greater attention paid to stereotypies developing in early childhood due to their presentation in patients affected by autistic spectrum disorder [13].

Although stereotypies have been described in Rhesus macaque primates with CMV disease, multiple studies describing the range of symptomatic manifestations of CMV disease in humans have not listed abnormal stereotypies.
as a manifestation of congenital CMV disease in human infants and neonates [5,14-17]. Continuing studies have shown that approximately 13% of asymptomatic neonates may go on to develop permanent sequelae making the early recognition of congenital CMV disease in infants important to initiate early interventions [18].

Early interventions applied to infants affected with intracranial CMV infection are known to improve outcomes, without which neurological outcomes worsen [19,20]. A significant proportion of infants with CMV disease have subtle neurologic findings. Infections can also be clinically silent. Stereotypies are a poorly understood clinical entity, especially in the field of pediatric neurology.

Conclusion

We present our case as an opportunity to add to the list of neurologic manifestations of intracranial CMV disease. This case also illustrates the importance of assessing for normal motor activity during the neurologic assessment of the well-child visit.

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