What does congenital Zika syndrome look like?

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Transmission electron microscope image of Zika virus (red), isolated from a microcephaly case in Brazil. The virus is associated with cellular membranes in the center. Credit: University of California - San Diego

Even as the Zika virus becomes more prevalent—the Centers for
Disease Control reports that the number of U.S. infants born with microcephaly and other birth defects is 20 times over the normal rate—researchers are still trying to fully pin down the identifying consequences of the viral infection.

In a new paper published this week in the *American Journal of Medical Genetics*, first author Miguel del Campo, MD, PhD, associate professor in the Department of Pediatrics at University of California San Diego School of Medicine, and colleagues in Brazil and Spain, describe the phenotypic spectrum or set of observable characteristics of congenital Zika (ZIKV) syndrome, based upon clinical evaluations and neuroimaging of 83 Brazilian children with presumed or confirmed ZIKV congenital infections.

"These findings provide new insight into the mechanisms and timing of the brain disruption caused by Zika infection, and the sequence of developmental anomalies that may occur," said del Campo, who also serves as a medical geneticist at Rady Children's Hospital-San Diego.

The physical characteristic most associated with Zika infection is microcephaly, a birth defect in which the baby's brain does not develop properly resulting in a smaller than normal head. In this case, prenatal brain development is initially normal, but then disrupted by the viral infection of neural progenitors. The research team reported that evidence of microcephaly and related skull abnormalities was present in 70 percent of the infants studied, though often it was subtle.

"Some cases had milder microcephaly or even a normal head circumference," said del Campo.

Besides the redundant scalp and abnormal cranial shapes resulting from the arrest in brain growth, other physical features consistent with ZIKV infection reflect immobility of the joints, resulting from altered brain
function *in utero*. These included deep and multiple dimples (30.1 percent), distal hand or finger contractures (20.5 percent), feet malpositions (15.7 percent) and generalized athrogryposis involving multiple joints (9.6 percent).

Neurologically, the primary features seen most often and most evidently in infants were alterations in motor activity, reflected in body tone, posture and motility or movement; severe hypertonia (abnormal muscle tension and contraction); abnormal neurobehaviors, such as poor or delayed response to visual stimuli, and excitability. Babies cried excessively but monotonously, and were often inconsolable.

Brain imaging revealed combinations of characteristic abnormalities, such as calcifications, poor gyral patterns and underdevelopment of the brainstem and cerebellum. There was a marked decrease in both gray and white matter volumes.

Del Campo said the findings suggest these children will have severe disabilities, a reality with strong implications for future clinical care. They highlight the variable severity of ZIKV brain damage and other characteristics, depending upon onset of maternal infection. Earlier detection of maternal and prenatal infection, he said, is critical to developing remedies to prevent or ameliorate ZIKV effects.

**More information:** *American Journal of Medical Genetics*, DOI: 10.1002/ajmg.a.38170