1. Introduction

1.1 Background

From 1 January 2007 to 16 June 2016, Zika virus transmission has been documented in a total of 71 countries and territories. The spread of the virus has steadily broadened in the Americas and Caribbean since 2015, with mosquito-borne transmission reported from 39 countries and territories of this region, as well as from five countries and territories of the Pacific.

Microcephaly and other central nervous system malformations potentially associated with Zika virus infection or suggestive of congenital infection have been reported by 12 countries or territories (Brazil, Cabo Verde, Colombia, El Salvador, French Polynesia, Marshall Islands, Martinique, Panama, Puerto Rico, Slovenia, Spain and United States of America). Of the five cases linked to travel, four reported microcephaly in infants born to mothers with a recent travel history to Brazil (Slovenia, United States of America) and Colombia (Spain); and one involved travel to an undetermined country in Latin America. Since late 2015, increased rates of congenital microcephaly as high as 20-fold\(^1\) have been reported in north eastern Brazil.

On 1 February 2016, the World Health Organization (WHO), following a meeting of the International Health Regulations (IHR) Emergency Committee on Zika virus and the observed increase in neurological disorders and neonatal malformations, declared the clustering of microcephaly cases, Guillain-Barré syndrome (GBS) and other neurological conditions reported in some areas affected by Zika virus as a Public Health Emergency of International Concern.\(^{2-4}\)

1.2 Objectives

The aim of this document is to provide guidance on the screening, clinical assessment, neuroimaging and laboratory investigations of neonates and infants born to women residing in areas of Zika virus transmission. This document updates the WHO interim guidance *Assessment of infants with microcephaly in the context of Zika virus* published on 4 March 2016. Recommendations are provided regarding the management and follow-up of neonates and infants with, or at risk of, ‘congenital Zika virus syndrome’. The syndrome represents the spectrum of congenital abnormalities (see 2.1 and 2.2) that have been reported to date in association with fetuses exposed to Zika virus in utero. This update also includes narrative summaries of recent evidence underpinning the recommendations, as well as operational considerations for implementation.

This guidance is intended to inform the development of national and local clinical protocols and health policies that relate to neonatal and infant care in the context of Zika virus transmission. It is not intended to provide a comprehensive practical guide for the management of Zika virus infections or neonatal neurological conditions including microcephaly.

1.3 Scope

This guidance is relevant to all neonates born to women residing in areas of active Zika virus transmission, particularly those women with suspected or confirmed Zika virus infection during pregnancy. WHO guidance on pregnancy management in the context of Zika virus infection is provided in a separate document.\(^5\)

1.4 Target audience

The primary audience for this guidance is health professionals directly providing care to neonates and their families including paediatricians, general practitioners, midwives and nurses. This guidance is also intended to be used by those responsible for developing national and local health protocols and policies, as well as managers of maternal, newborn and child health programmes and policy-makers in regions affected by Zika virus.

2. Complications related to Zika virus infection in infants

2.1 Microcephaly

Microcephaly is a condition where a baby has a head that is smaller when compared with other babies of the same sex and age. An infant is considered to have microcephaly when the head circumference (also known as occipito-
Microcephaly can be caused by numerous genetic factors including chromosomal and metabolic disorders, and also non-genetic etiologies including congenital infections, intrauterine exposure to teratogens, perinatal injuries to the developing brain and severe malnutrition. Depending on the timing of insult, microcephaly may be present at birth (congenital) or may develop postnatally.

While microcephaly is a clinical sign and not a disease, congenital microcephaly (i.e. microcephaly present at birth) often indicates an underlying neurologic pathology and has been associated with a range of neurological complications including developmental delay, intellectual impairment, hearing and visual impairment and epilepsy. Microcephaly can be caused by numerous genetic factors including chromosomal and metabolic disorders, and also non-genetic etiologies including congenital infections, intrauterine exposure to teratogens, perinatal injuries to the developing brain and severe malnutrition. Depending on the timing of insult, microcephaly may be present at birth (congenital) or may develop postnatally.

Investigation of infants with congenital microcephaly in settings of Zika virus transmission has detected transplacental transmission of Zika virus and, where the pregnancy has been terminated or resulted in a stillbirth, Zika virus has been recovered from fetal brain tissue. An autopsy study of a fetus with a history of Zika virus exposure in utero showed evidence of activated microglia and macrophages in the brain, suggesting that host immune responses may contribute to the pathogenesis of microcephaly. Zika virus is known to be highly neurotropic and may therefore adversely affect fetal development by directly infecting the fetal brain or indirectly, by infecting the placenta. In vitro studies have shown that Zika virus can infect neural progenitor cells and may affect their cell cycle regulation and survival.

2.2 Other complications associated with in utero exposure to Zika virus

In addition to congenital microcephaly and associated brain abnormalities, a range of other complications including craniofacial disproportion, spasticity, seizures, irritability, brainstem dysfunction, feeding difficulties and ocular abnormalities have been reported among neonates where there has been in utero exposure to Zika virus. In some cases these abnormalities have occurred without associated microcephaly and have become evident only following birth. In some cases these abnormalities have occurred without associated microcephaly and have become evident only following birth. Based on observational, cohort, and case control studies, there is a strong scientific consensus that Zika virus is a cause of microcephaly and other neurologic complications. Longer term clinical follow-up of infants born to women with a history of confirmed Zika virus infection at different times during pregnancy is needed. As a set of congenital abnormalities has been reported in a number of these infants, including abnormal neuroimaging findings, it suggests that a congenital syndrome, akin to congenital rubella or cytomegalovirus (CMV) infection, is attributable to in utero Zika virus infection.

3. Evidence and recommendations

3.1 Screening infants for congenital Zika virus syndrome

3.1.1 Initial history taking, clinical and anthropometric assessment

Microcephaly is defined as a head circumference of two standard deviations or more below the mean for age and sex. Severe microcephaly is present when the head circumference is more than three standard deviations below the mean for age and sex.

Increased rates of congenital microcephaly have been reported in settings of Zika virus transmission in Brazil beginning in late 2015 and French Polynesia from 2013-2015. However, not all children with congenital Zika virus syndrome present with congenital microcephaly. Some of these children with normal birth head circumference have appeared to have a disproportionately small head relative to the face or body (craniofacial disproportion), which may suggest relatively poor brain growth. According to an unpublished series in Recife, Brazil, up to 12% of infants with a history of maternal exposure to Zika virus in pregnancy and who developed neurological signs and symptoms in the months following birth had a head circumference at birth two standard deviations or more below the mean for age and sex and many had craniofacial disproportion (Vanessa van der Linden, personal communication, 2016).

Given the association between congenital microcephaly and other neurological morbidities such as cognitive delay, intellectual disability, cerebral palsy and epilepsy, a small
head circumference is an important clinical sign requiring further evaluation and follow-up. However, screening at birth for complications as a result of in utero Zika infection is presently hampered by diagnostic methods for determining Zika virus infection. Molecular methods can detect active infection in adults, but diagnostic technologies to establish prior infection such as during pregnancy are not available. Furthermore, it is estimated that up to 80% of Zika virus infections may be asymptomatic.\textsuperscript{21} Hence, routine measurement of head circumference of all infants born to mothers in areas of Zika virus transmission, in addition to evaluation for other possible signs or symptoms, is essential to screen for congenital Zika virus syndrome.

### 3.1.2 Head circumference cut-off values to determine microcephaly

Different head circumference cut-off values (i.e. the measurement used to determine if an infant has a small head or not) have been used for defining microcephaly. These have included: $<-2$ SD (i.e. more than 2 SD below the mean); $< 3^{rd}$ percentile (i.e. less than the 3$^{rd}$ percentile; and $<-3$ SD (i.e. more than 3 SD below the mean). Head circumference cut-offs of either $<-2$ SD or $< 3^{rd}$ percentile will therefore designate more infants as having microcephaly, whereas using a cut-off of $<-3$ SD will designate fewer infants having microcephaly, though these infants will have more severe microcephaly and will be more likely to have severe neurological or developmental abnormalities. Consistent agreed case definitions for congenital microcephaly is therefore important in order to standardise data.

### 3.1.3 Choice of growth standards for head circumference measurements

The WHO Child Growth Standards (WCGS),\textsuperscript{22} derived from the WHO Multicentre Growth Reference Study (MGRS) describe optimal growth trajectories of infants and children from birth for whom there are no apparent barriers to growth.\textsuperscript{23} The WCGS provide mean and median values for weight, length/height and head circumference by sex and age, and describe their distributions according to either percentiles or standard deviations. However, measurements less than the 1$^{st}$ percentile cannot be further classified to indicate the severity of microcephaly. For example, head circumferences of 31.0 cm and 30.4 cm in a term boy are both less than the 1$^{st}$ percentile; but 31.0 cm is between -2 SD and -3 SD, and 30.4 cm is below -3 SD. Standard deviation measurements can also be aggregated to provide a mean or median Z score for a specific population, whereas head circumference values based on percentiles cannot be summarised in the same way. However, the WCGS only provide values for term infants (i.e. from 37-42 weeks gestation) and were not disaggregated within this range. A single head circumference standard is therefore applied for all neonates considered term from 37 weeks to 41 weeks and 6 days ($37^{+0}$ to $41^{+6}$).

The INTERGROWTH-21\textsuperscript{st} project (IG-21) adopted a similar methodology to the WHO MGRS to describe normal fetal growth and birth anthropometric measurements for weight, length and head circumference.\textsuperscript{24} However, the IG-21 Size at Birth Standards are disaggregated by sex and gestational age (including between 37-42 weeks), and also provide standards for very preterm infants.

The choice of standard used – WCGS or IG-21 – should also reflect the availability and reliability of gestational age assessments. Accurate gestational age is difficult to ascertain unless an ultrasound assessment has been performed early in the first trimester. Dates of last menstrual period are commonly unreliable and estimated dates of delivery may vary widely when these are used to determine ‘term’ for any pregnancy.

The WCGS provides an appropriate reference standard for term neonates where gestational age is not reliably known. However, when the gestational age is accurately known it is preferable to use a standard appropriate for that gestational age. Otherwise, it is possible that microcephaly will be over-diagnosed. For example, an infant boy born at 37 weeks gestation with a head circumference of 31.0 cm (between -1 SD and -2 SD based on IG-21 standards) will be considered to have microcephaly based on -2 SD WCGS for boys (i.e. 31.9 cm). Similarly, an infant girl at 38 weeks gestation with a head circumference of 31.0 cm (between -1 SD and -2 SD based on IG-21 standards) will also be considered to have microcephaly based on -2 SD WCGS for girls (i.e. 31.5 cm).

In some regions, large numbers of women experience unfavourable conditions before and during pregnancy and their offspring therefore are at greater risk of fetal growth restriction. In these populations, using either WCGS or IG-21 standards, more neonates may be identified as having microcephaly. For example, in parts of Kenya up to 10% of neonates may have head circumference less than -2 SD of median for age (Charles Newton, personal communication, 2016).
Table 1. Comparison of head circumference standards – WCGS and IG-21 by sex and gestational age

<table>
<thead>
<tr>
<th>Gestational age (weeks)</th>
<th>Standard deviation</th>
<th>IG-21 size at birth (cm)</th>
<th>WHO Child Growth Standards</th>
</tr>
</thead>
<tbody>
<tr>
<td>BOYS</td>
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<tr>
<td>37</td>
<td>0</td>
<td>33.02</td>
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<td>30.54</td>
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<td>38</td>
<td>0</td>
<td>33.47</td>
<td>WCGS provides a single set of head circumference values from 37 weeks to 41 weeks and 6 days gestational age</td>
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<tr>
<td></td>
<td>-2 SD</td>
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<td></td>
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<td>0</td>
<td>33.90</td>
<td>0 SD = 34.5 cm</td>
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<td>-2 SD</td>
<td>31.54</td>
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<td>-3 SD</td>
<td>30.19</td>
<td>-3 SD = 30.7 cm</td>
</tr>
<tr>
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<td>0</td>
<td>34.31</td>
<td>3rd percentile = 32.1 cm</td>
</tr>
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<td>-2 SD</td>
<td>32.00</td>
<td>1st percentile = 31.5 cm</td>
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<tr>
<td></td>
<td>-3 SD</td>
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<tr>
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<td>-3 SD</td>
<td>31.14</td>
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<td>GIRLS</td>
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<td>37</td>
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<td>32.61</td>
<td>WCGS provides a single set of head circumference values from 37 weeks to 41 weeks and 6 days gestational age</td>
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<tr>
<td></td>
<td>-2 SD</td>
<td>30.24</td>
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<tr>
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<td>28.85</td>
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<td>0 SD = 33.9 cm</td>
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<td>3rd percentile = 31.7 cm</td>
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</table>

3.1.4 When to measure the head circumference

In order to obtain the most accurate comparison with either WCGS or IG-21 standards, head circumference measurements should be taken within the first 24 hours to be compatible with the time intervals used in the respective studies.22, 24

No matter which standard is used and when it is measured, it is essential to meticulously follow recommended methods to avoid measurement errors.25
3.1.5 Recommendations

1. Neonates should have their head circumference measured in the first 24 hours of life:
   a. For term neonates (37-42 weeks), WHO Child Growth Standards for size at birth should be used to interpret measurements. If accurate gestational age is known, Intergrowth-21 Size at Birth Standards are preferred.
   b. For preterm neonates, Intergrowth-21 Size at Birth Standards for gestational age and sex should be used to interpret measurements.

2. All mothers should be asked about clinical signs and symptoms suggestive of Zika virus infection and/or laboratory confirmation of Zika virus infection during pregnancy, including when the possible infection occurred (first, mid or final trimester).

3. Neonates should be examined to assess whether the head appears disproportionately small relative to the face or body (craniofacial disproportion).

Operational considerations

- If head circumference cannot be measured during the first 24 hours, it should be measured within the first 72 hours.
- Health practitioners should be trained in the use of these growth standards in areas where they are not in routine use.

3.2 Clinical assessment of neonates for congenital Zika virus syndrome

3.2.1 Etiology of congenital microcephaly

Microcephaly is associated with numerous genetic etiologies, including chromosomal and metabolic disorders and also non-genetic causes. Non-genetic causes include congenital infections including the TORCH infections (toxoplasmosis, rubella, cytomegalovirus and herpes), syphilis, varicella–zoster, parvovirus B19 and human immunodeficiency virus (HIV). Other non-genetic causes include intrauterine exposure to teratogens such as alcohol and ionizing radiation, pre- and perinatal injuries to the developing brain (hypoxia-ischaemia, trauma, stroke), and severe malnutrition. Depending on the timing of insult, the onset of microcephaly may be prenatal or postnatal.

3.2.2 Congenital microcephaly and neurodevelopmental outcomes

When all forms of microcephaly are considered, there appears to be general correlation between the degree of microcephaly and the likelihood of neurological impairment. A study based on the National Institute of Neurological Disorders and Stroke Collaborative Perinatal Project found that among children with birth head circumference between -2 SD and -3 SD, about 11% had an intellectual quotient (IQ) less than 70; and among children with birth head circumference -3 SD or below, 51% had IQ < 70 at seven years of age. However, some children in the general population with head circumference between -2 SD and -3 SD may still have normal development.

Studies of children with congenital infections report frequent microcephaly in children with symptomatic congenital CMV and congenital rubella syndrome. However, children with congenital CMV and without microcephaly may still have cerebral cortical malformations that lead to neurologic impairments such as intellectual disability and epilepsy. In the context of a congenital infection, microcephaly is often predictive of worse neurodevelopmental outcomes.

Congenital infections may also be associated with other neurological consequences ranging from isolated sensorineural hearing loss to severe destructive brain lesions. Congenital infections, particularly CMV, are one of the most common causes of hearing impairment. Postnatal onset of hearing impairment and a progressive course are also common. Children with congenital microcephaly of unspecified etiologies also demonstrate an increased incidence of sensorineural hearing loss. Microcephaly may similarly be associated with eye and vision abnormalities. One large study found that 30% of children with microcephaly of heterogeneous etiologies had disorders of the eyes. Chorioretinitis and other visual abnormalities are frequently reported in children with congenital CMV.

3.2.3 Zika virus exposure in utero and neurological consequences

Reports from areas with Zika virus transmission note that children with congenital Zika virus syndrome commonly, but not always, have congenital microcephaly. Evidence from animal models, fetal autopsy, placental studies and cerebrospinal fluid (CSF) examination of affected infants have found the presence of Zika virus in such neonates, supporting the conclusion that Zika virus can have a major deleterious effect on developing brain structure. Early reports suggest that children with congenital Zika virus syndrome may also have sensorineural hearing loss; however, due to the limited duration of follow-up among index cases to date, the prevalence and clinical course of this abnormality are not yet fully known. Ocular findings such as bilateral macular and perimacular lesions as well as optic nerve abnormalities have also been reported in children with congenital Zika virus syndrome.

Other clinical signs and symptoms commonly noted in neonates with congenital microcephaly where maternal Zika virus infection in pregnancy was either suspected or confirmed include arthrogryposis, early-onset spasticity, hyperirritability, swallowing difficulties and seizures.
3.2.4 Assessing a neonate with congenital microcephaly

Identifying the underlying cause of microcephaly has implications for the child’s prognosis, and is also important to monitor and manage potential complications and to counsel future pregnancies. Some causes of microcephaly may be suspected or diagnosed by history (e.g. fetal alcohol syndrome or maternal malnutrition), physical and neurological examination (e.g. syndromes with dysmorphisms) or a combination of both (e.g. congenital infections). Ancillary tests including neuroradiological and laboratory investigation often aid etiological diagnosis (see 3.3 and 3.4).

3.2.5 Recommendation

4. In neonates with congenital microcephaly or in whom the head appears disproportionately small relative to the face or body, a full history and physical and neurological examination, including assessment of hearing and vision, should be performed in order to detect additional malformations potentially associated with Zika virus infection.

Operational considerations

- The clinical history and full physical and neurological examinations may help to differentiate congenital infections from other causes of congenital microcephaly such as genetic or environmental.
- It is essential that all neonates, especially those born in areas with active Zika virus transmission, are screened for hearing loss at the earliest possible opportunity, preferably before they are discharged from hospital after birth.
- Hearing screening should be performed according to the WHO guiding principles for newborn and infant hearing screening. Screening can be done using automated auditory brainstem responses (ABR) or otoacoustic emissions (OAE) screening procedures. In places where it is not possible to undertake physiological tests to identify hearing loss, assessment can be undertaken through behavioural measures;
- Accurate assessment of vision by clinical examination during the newborn period may be difficult. Where possible an ophthalmologist should perform an ocular examination.

3.3 Neuroimaging of neonates for congenital Zika virus syndrome

3.3.1 Neuroimaging and microcephaly

Neuroimaging abnormalities are common in children with congenital microcephaly, especially where there are associated neurological signs or symptoms. These findings may help to determine the underlying cause of microcephaly. In settings without Zika virus, neuroimaging abnormalities were noted in 80% of children with head circumference <-3 SD by computed tomography (CT) or magnetic resonance imaging (MRI) and 88% of these children by MRI. Most of the children had neurological signs or symptoms, though not necessarily present at birth.

Neuroimaging data of infants with congenital microcephaly in settings of Zika virus transmission is limited; cerebral calcification has commonly been detected in such children and is often subcortical in location. Other reported findings include brain atrophy and ventriculomegaly, cerebellar and brainstem anomalies, cortical gyral abnormalities and callosal abnormalities. Gyral abnormalities are described as polymicrogyria, pachygyria or lissencephaly and are often frontal predominant. The presence and pattern of these gyral abnormalities suggest that Zika virus directly interferes with early brain development, as opposed to destroying the brain later in development.

These neuroimaging abnormalities can also be found in infants with other congenital infections such as CMV syndrome. For example, intracranial calcifications have been identified in 14.2-58.5% of children with symptomatic congenital CMV though these calcifications tend to be subependymal rather than subcortical. Congenital CMV infection can also cause brain malformations such as polymicrogyria, pachygyria, atrophy and other anomalies similar to those described in infants with congenital Zika virus syndrome. However, emerging evidence suggests that the neuroimaging findings in congenital Zika virus syndrome may be more striking than those with other congenital infections.

3.3.2 Magnetic resonance imaging, computerized axial tomography or ultrasound examination

Cerebral calcification may be more readily identified by CT compared to MRI. However MRI has a higher resolution and better ability to delineate abnormalities such as those of the cerebral cortex and posterior fossa. The available literature and limited clinical experience suggest that either CT or MRI is sufficient to identify typical radiological features of congenital Zika virus syndrome.

The utility of postnatal cranial ultrasound in congenital Zika virus syndrome is unknown. In congenital CMV infection, cranial ultrasound is often useful to detect pathological findings including calcification, ventriculomegaly and cystic changes. However, the distribution of brain calcification in congenital Zika virus syndrome appears to be more superficial, making it potentially more difficult to detect by postnatal cranial ultrasound. Furthermore, the quality of postnatal cranial ultrasound depends on the size of the fontanelle.

Experience from Brazil indicates that many neonates with suspected congenital Zika virus syndrome have a very small or closed fontanelle at birth and cranial ultrasound may not be feasible or reliable in providing useful clinical information in these cases.
3.3.3 Recommendations

5. In neonates with head circumference < -2 SD and ≥ -3 SD, or where the head is disproportionately small relative to face or body size, (and no strong indication from clinical examination of a genetic or environmental cause of microcephaly) neuroimaging should be performed if:
   a. Zika virus infection is suspected in the mother during pregnancy; or
   b. Any neurological signs or symptoms are present;
6. In neonates with head circumference < -3 SD neuroimaging should be performed if there is no strong indication from clinical examination of a genetic or environmental cause of microcephaly.
7. When neuroimaging is indicated:
   a. Either CT or MRI can be used.
      - CT is satisfactory to identify neuroimaging findings suggestive of congenital Zika virus syndrome.
      - MRI is satisfactory to identify neuroimaging findings suggestive of congenital Zika virus syndrome, and may also provide further detail and diagnose other conditions.
   b. If CT or MRI are not available, cranial ultrasound can be performed if the fontanelle is of adequate size.

Remarks

• Currently there are no known pathognomonic neurological findings for congenital Zika virus syndrome. Findings reported in neonates with congenital Zika virus syndrome include: cerebral calcification, brain atrophy and ventriculomegaly, cerebellar and brainstem anomalies, cortical gyral abnormalities and callosal abnormalities.

• Cerebral calcification is commonly seen in congenital infections. Some genetic disorders, such as Aicardi–Goutières syndrome and mutations in the OCLN gene, whose brain manifestations may mimic congenital infections.

Operational considerations

• In addition to availability, radiation exposure in CT and higher cost and potential need for sedation in MRI should be considered when selecting an imaging modality.
• Neuroradiological findings should be interpreted in the context of other clinical and laboratory information.
• When indicated, cranial ultrasound should be performed by an ultrasonographer experienced in neonatal cranial ultrasound.

3.4 Laboratory investigations of neonates for congenital Zika virus syndrome

3.4.1 Identifying other causes of microcephaly including congenital infections

The clinical and neuroimaging findings of neonates born with microcephaly due to congenital infections or some genetic disorders can be similar. In order to provide the most appropriate care and to counsel families of children with congenital microcephaly, it is important to establish the underlying etiology. A clinical history, including immunization status, past and recent infections, and exposures can ascertain risk factors or characteristics of one etiology or another. A careful physical examination of the neonate may also identify signs that may point toward a specific diagnosis.

Additional laboratory testing can help to make a diagnosis of other congenital infections such as CMV or rule out genetic disorders such as Aicardi-Goutières syndrome and mutations in the OCLN gene, whose brain manifestations may mimic congenital infections.

In order to attribute microcephaly or other neurological findings to in utero Zika virus exposure, other causes of congenital abnormalities must be excluded.

However, there is currently no validated laboratory diagnostic test or commercial assay to confirm congenital Zika virus infection or exposure in neonates. Zika virus ribonucleic acid (RNA) has been detected by reverse transcription polymerase chain reaction (RT-PCR) from the serum of neonates with perinatal transmission of Zika virus from the mother. Zika virus IgM has also been detected from the CSF of infants with congenital Zika virus syndrome. As the sensitivity and specificity of RT-PCR and serological testing for Zika virus in neonates with suspected congenital Zika virus infection is being established, it is recommended that both RT-PCR and serology be performed to determine congenital infection.

3.4.2 Recommendations

8. Serological testing for TORCH infections should be performed (unless excluded in the mother in pregnancy):
   a. in neonates with congenital microcephaly; or
   b. where the head is disproportionately small relative to face or body size;
   And
   c. where Zika virus infection is suspected in the mother during pregnancy; or
   d. any neurological signs or symptoms are present
9. The role of serological and virological testing for Zika virus in neonates should be assessed based on further data on sensitivity and specificity and understanding of cross-reactivity with other flaviviruses.
Operational considerations

• Positive CMV serology in a neonate is not a reliable indicator of in utero CMV infection. Diagnosis requires detection of CMV in urine, saliva, blood or other tissues within 2-3 weeks of birth.

3.5 Management of neonates with congenital Zika virus syndrome

3.5.1 Early complications

Limited clinical data available from Brazil show that infants with congenital Zika virus syndrome are at high risk for a spectrum of complications including developmental delays, seizures, hearing and visual impairment, excessive irritability, early-onset spasticity, swallowing difficulties, arthrogryposis and hip dysplasia. Although the course of congenital Zika virus syndrome is unknown, children with congenital infections and/or microcephaly are at high risk for developmental delays and auditory and visual impairment and the risk of these are higher in the setting of an abnormal exam and/or neuroimaging. Existing WHO guidelines for screening and management of the sequelae associated with congenital Zika virus syndrome should be utilized for a comprehensive treatment approach, including guidelines for epilepsy, spasticity, hearing and vision. It is also essential to support parents and families of affected infants to deal with the anxiety and psychosocial distress experienced at these times. For all infants and families, support, care and treatment should follow a multidisciplinary approach.

It is anticipated that recommendations on the management of children with congenital Zika virus syndrome will be revised based on new evidence in late 2016.

3.5.2 Recommendations

10. Families of neonates with congenital Zika virus syndrome should be informed about the diagnosis and advised regarding management and prognosis.

11. Psychosocial support and advice should be provided to families of neonates with congenital Zika virus syndrome as described in WHO interim guidance on ‘Psychosocial support for pregnant women and for families with microcephaly and other neurological complications in the context of Zika virus’.

12. Infants with congenital Zika syndrome should receive a comprehensive neurodevelopmental assessment, and supportive therapy should be put in place for any difficulties noted including irritability, seizures, swallowing difficulties, early onset spasticity and hip dysplasia.

13. Multidisciplinary approaches should be adopted to provide early interventions and support to promote neurodevelopment, prevent contractures and manage early complications as outlined in WHO mhGAP and community-based rehabilitation guidelines.

Operational considerations

• Health care practitioners need to be trained and provided with resources to recognize and manage the reported neurological complications associated with congenital Zika virus syndrome.
• Community-based rehabilitation and support may be relevant especially in resource limited settings.

3.6 Follow-up of children in areas of Zika virus transmission

3.6.1 Short and long term follow-up

Little follow-up data is available to date regarding children affected by congenital Zika virus syndrome, the description of which is still relatively preliminary. Children identified in Brazil, where the largest number of cases has been reported, are still generally less than 12 months of age. Even though retrospective data from French Polynesia has revealed an increased incidence of microcephaly associated with a Zika virus outbreak in 2013-2015, little follow-up data of affected infants from that period are available.

However, follow-up management can be informed by existing experience and guidelines for other congenital infections and microcephaly-associated neurodevelopmental conditions. For example, presence of microcephaly and its severity are strongly associated with a variety of neurodevelopmental sequelae that become evident in early to late childhood from intellectual disability, epilepsy, cerebral palsy, visual impairment, and hearing loss. Head circumference monitoring may provide an indication of brain growth and the likelihood of neurodevelopmental abnormalities; repeat neurological examinations may identify signs and symptoms of abnormalities as they become evident. Findings may also be helpful for counselling parents and families. For example, children with congenital CMV infection who do not have neurologic symptoms within the first year of life are unlikely to develop neurodevelopmental or intellectual impairment later. It is also generally agreed that neurological sequelae of congenital CMV infection do not emerge after two years of age.

In developing recommendations for follow-up of children in areas of Zika virus transmission, the guideline development group considered other standard recommendations for neurodevelopmental follow-up of at-risk children and existing WHO guidelines for auditory and ophthalmologic screening.

Note: It is anticipated that recommendations on the follow-up of children with congenital Zika virus syndrome will be revised based on new evidence in late 2016.
3.6.2 Recommendations

14. Infants with congenital Zika virus syndrome should be followed up at 1 month, 3 months, 6 months, 9 months, 12 months, 18 months and 24 months of age. Additional follow-up should be provided if there are other complications. Further follow-up beyond 24 months of age will be required depending on the child’s condition and needs.

15. At each visit, head circumference should be measured in order to monitor postnatal brain growth. For term neonates, WCGS for attained head circumference should be used to interpret the head circumference measurement. For preterm neonates, Intergrowth-21 postnatal growth standards for attained head circumference should be used to interpret postnatal changes of head circumference until 64 weeks postmenstrual age. After this, WCGS for attained head circumference should be used to interpret the head circumference measurement.

16. Developmental and neurological assessments should be performed with the full engagement of caregivers to identify developmental delays and other neurological abnormalities including epilepsy and disorders of movement, posture and swallowing.

17. Hearing should be screened in the first month of life as early as possible before discharge from hospital and further audiological evaluation and services should be provided as per the WHO guiding principles for newborn and infant hearing screening and the Position Statement from the Joint Committee on Infant Hearing.

18. There should be comprehensive ophthalmological assessment.

19. The health and well-being of the families and caregivers, including their psychological well-being should be assessed. Families and caregivers should be provided psychosocial support and parenting advice.

20. Infants born to mothers with suspected, probable or confirmed Zika virus infection during pregnancy, even without microcephaly or disproportionately small head relative to the face or body, should be followed up to detect, manage and investigate signs of neurodevelopmental abnormality including feeding difficulties, hearing or vision problem and poor head growth. Follow-up visits should occur at 3 months, 9 months and 24 months of age as a minimum.

21. Families and caregivers should be provided with psychosocial support and parenting advice.

Remarks

- Timing of the neuroassessment after birth needs to balance the sensitivity of the examination to detect impairments (which increases with age) against the need to identify impairments early to maximize the efforts of interventions.
- Undiagnosed hearing loss could hinder language development.

Operational considerations

- It is important to have a well-established protocol for assessment and training of caregivers and health workers, especially in the absence of physiological tests for hearing.
- Hearing assessment can be undertaken through behavioural measures and evaluation for age-appropriate language milestones. A follow up regime is needed whereby the children can be repeatedly evaluated by trained health professionals. In case of any delay in these milestones or in case of parental/caregiver suspicion, the child should undergo full audiological evaluation to establish the degree and nature of hearing loss.
- Once a diagnosis of hearing loss is established, suitable interventions based on the hearing loss and co-morbidities must be made available to the child and family.
Table 2. Summary of recommendations

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| 1 | Neonates should have their head circumference measured in the first 24 hours of life:  
|   | a. For term neonates (37-42 weeks), WHO Child Growth Standards for size at birth should be used to interpret measurements. If accurate gestational age is known, Intergrowth-21 Size at Birth Standards are preferred.  
|   | b. For preterm neonates, Intergrowth-21 Size at Birth Standards for gestational age and sex should be used to interpret measurements. |
| 2 | All mothers should be asked about clinical signs and symptoms suggestive of Zika virus infection and/or laboratory confirmation of Zika virus infection during pregnancy, including when the possible infection occurred (first, mid or final trimester). |
| 3 | Neonates should be examined to assess whether the head appears disproportionately small relative to the face or body (craniofacial disproportion). |
| 4 | In neonates with congenital microcephaly or in whom the head appears disproportionately small relative to the face or body, a full history and physical and neurological examination, including assessment of hearing and vision, should be performed in order to detect additional malformations potentially associated with Zika virus infection. |
| 5 | In neonates with head circumference < -2 SD and ≥ -3 SD, or where the head is disproportionately small relative to face or body size, (and no strong indication from clinical examination of a genetic or environmental cause of microcephaly) neuroimaging should be performed if:  
|   | a. Zika virus infection is suspected in the mother during pregnancy; or  
|   | b. Any neurological signs or symptoms are present. |
| 6 | In neonates with head circumference < -3 SD neuroimaging should be performed if there is no strong indication from clinical examination of a genetic or environmental cause of microcephaly. |
| 7 | When neuroimaging is indicated:  
|   | a. Either CT or MRI can be used.  
|   | - CT is satisfactory to identify neuroimaging findings suggestive of congenital Zika virus syndrome.  
|   | - MRI is satisfactory to identify neuroimaging findings suggestive of congenital Zika virus syndrome, and may also provide further detail and diagnose other conditions.  
|   | b. If CT or MRI are not available, cranial ultrasound can be performed if the fontanelle is of adequate size. |
| 8 | Serological testing for TORCH infections should be performed (unless excluded in the mother in pregnancy):  
|   | a. in neonates with congenital microcephaly, or  
|   | b. where the head is disproportionately small relative to face or body size,  
|   | And  
|   | c. where Zika virus infection is suspected in the mother during pregnancy, or  
|   | d. any neurological signs or symptoms are present |
| 9 | The role of serological and virological testing for Zika virus in neonates should be assessed based on further data on sensitivity and specificity and understanding of cross-reactivity with other flaviviruses. |
| 10 | Families of neonates with congenital Zika syndrome should be informed about the diagnosis, and advised regarding management and prognosis. |
| 11 | Psychosocial support and advice should be provided to families of neonates with congenital Zika virus syndrome as described in WHO interim guidance on ‘Psychosocial support for pregnant women and for families with microcephaly and other neurological complications in the context of Zika virus’. |
| 12 | Infants with congenital Zika virus syndrome should receive a comprehensive neurodevelopmental assessment, and supportive therapy should be put in place for any difficulties noted, including irritability, seizures, swallowing difficulties, early onset spasticity and hip dysplasia. |
| 13 | Multidisciplinary approaches should be adopted to provide early interventions and support to promote neurodevelopment, prevent contractures and other neurological abnormalities including epilepsy and disorders of movement, posture and swallowing. |
| 14 | Infants with congenital Zika virus syndrome should be followed up at 1 month, 3 months, 6 months, 9 months, 12 months, 18 months and 24 months of age. Additional follow-up should be provided if there are other complications. Further follow-up beyond 24 months of age will be required depending on the child’s condition and needs. |
| 15 | At each visit, head circumference should be measured in order to monitor postnatal brain growth. For term newborns, WCGS for attained head circumference should be used to interpret the head circumference measurement. For preterm newborns, Intergrowth-21 preterm postnatal growth standards for attained head circumference should be used to interpret postnatal changes of head circumference until 64 weeks postmenstrual age. After this WCGS for attained head circumference should be used to interpret the head circumference measurement. |
| 16 | Developmental and neurological assessments should be performed with the full engagement of caregivers to identify developmental delays and other neurological abnormalities including epilepsy and disorders of movement, posture and swallowing. |
| 17 | Hearing should be screened in the first month of life as early as possible before discharge from hospital and further audiological evaluation and services should be provided as per the WHO guiding principles for newborn and infant hearing screening and the Position Statement from the Joint Committee on Infant Hearing. |
| 18 | There should be comprehensive ophthalmological assessment. |
| 19 | The health and well-being of the families and caregivers, including their psychological well-being should be assessed. Families and caregivers should be provided psychosocial support and parenting advice. |
| 20 | Infants born to mothers with suspected, probable or confirmed Zika virus infection during pregnancy, even without microcephaly or disproportionately small head relative to the face or body, should be followed up to detect, manage and investigate signs of neurodevelopmental abnormality including feeding difficulties, hearing or vision problem and poor head growth. Follow-up visits should occur at 3 months, 9 months and 24 months of age as a minimum. |
| 21 | Families and caregivers should be provided with psychosocial support and parenting advice. |
4. Guidance development methods

4.1 Methods

4.1.1 Evidence retrieval, assessment and synthesis

A systematic search of evidence was undertaken between February and March 2016 with search terms reflecting the scope of the guidelines (see below). No time or language limits were implemented.

The term “microcephaly” was used in PubMed to search for relevant literature while the term “congenital microcephaly” was used in the Embase. Excluding animal studies and case reports, 5895 and 4489 articles were identified through each database respectively.

The titles of these 10 384 publications were reviewed for relevance and duplication and articles that focused on single etiology or mechanisms causing microcephaly were also excluded, leaving 855 articles for further assessment. Abstracts of these publications were screened and 139 articles considered directly relevant, reporting outcomes among children with microcephaly with a focus on neurodevelopmental outcomes or being associated with congenital infections. An additional 83 articles focused on neuroimaging findings. Additional articles suggested by experts in the field were also reviewed even if not captured by the original search strategy. The reference lists of articles identified through this search were also reviewed and papers deemed relevant also included.

Guidelines developed by national and international organizations within the last five years were additionally identified as primary sources of information.

Population characteristics and major findings of manuscripts were captured and summarized. Evidence summaries were developed as per the scope of the guideline.

Additional searches in PubMed searching “Zika virus” and restricted to data published after 2015 were performed on 25 April 2016 to reflect the most up to date published evidence.

4.1.2 Guideline Development Group

A guideline development group (GDG) was established that provided clinical experience or technical expertise in the areas of microcephaly, paediatric neurology, paediatric neuro-imaging, neonatology and epidemiology/surveillance of birth defects, including experts from affected countries. Participants were identified based on searches of the published literature, as known experts in the field/from affected countries and for geographic representation.

The guideline meeting was held on 17-19 March 2016 at the WHO headquarters in Geneva, Switzerland. This meeting was jointly organized by the World Health Organization (WHO) headquarters Departments of Maternal, Newborn, Child and Adolescent Health (MCA), Mental Health and Substance Abuse (MSD), Nutrition for Health and Development (NHD) and Reproductive Health and Research (RHR).

4.1.3 Finalization of recommendations

Draft recommendations were prepared by a WHO secretariat. A chairperson with expertise in managing group processes and interpreting evidence was nominated at the opening of the consultation and the nomination approved by the GDG. The GDG were asked to consider the draft recommendations in light of the evidence presented.

A decision-making framework was presented to guide the discussions. It included considerations such as (i) the desirable and undesirable effects of these recommendations; (ii) the available evidence; (iii) likely values and preferences of health workers and communities related to the recommended interventions in different settings; and (iv) feasibility and resource implications for programme managers in different settings. The GDG discussed the evidence and considered these issues to reach a consensus and to finalize the recommendations.

The draft recommendations were shared with an external peer-review group to ensure that there were no important omissions, contradictions or inconsistencies with scientific evidence or programmatic feasibility; and to assist with clarifying language, especially in relation to implementation and interpretation by policymakers and programme staff.

4.2 Declaration of interests

All GDG members completed a standard WHO Declaration of Interests (DOI) form before participating in the technical consultation or any activities related to the development of the guidance. Participants of the technical consultation also made verbal declarations of their DOI statements prior to the consultation and no conflicts were identified.

4.3 Acknowledgements

This guideline process was coordinated by the WHO Departments of Maternal, Newborn, Child and Adolescent Health (MCA) and Mental Health and Substance Abuse (MSD). The WHO secretariat included Rajiv Bahl, Anthony Costello, Tarun Dua, Nigel Rollins and Shekhar Saxena from WHO Geneva and Pablo Duran from the Centro Latinoamericano de Perinatología, Department of Women’s and Reproductive Health, WHO Regional Office for the Americas and Pilar Ramón-Pardo from the WHO Regional Office of the Americas.

Support was provided by staff from the WHO Incident Management System, WHO Geneva, including Ian Clarke, Qiu Yi Khut, Anaïs Legand and William Perea.
Dr Ganeshwaran H Mochida (Boston Children’s Hospital and Harvard Medical School, Boston, United States of America) and Dr Archana A. Patel (Boston Children’s Hospital and Harvard Medical School) were engaged as consultants in order to review and synthesize the evidence and draft the guideline. Dylan Vaughan (Boston Children’s Hospital, Boston, United States) assisted in literature retrieval. Michelle Griffin and Steven Morris from Public Health England, United Kingdom, assisted with preparations and documentation of the guideline meeting.

The guideline development group members were: Satinder Aneja (Lady Hardinge Medical College, New Delhi, India); James Barkovich (University of California, San Francisco, United States of America); Marianne Besnard (Ta’aone Hospital, Tahiti); Helen Cross (Institute of Child Health, London, United Kingdom); Richard Leventer (Murdoch Children’s Research Institute, University of Melbourne, Australia); Amira Masri (University of Jordan, Jordan); Cynthia Moore (Division of Birth Defects and Developmental Disabilities, Centers for Disease Control and Prevention, Atlanta, United States of America); Charles Newton, (Kenya Medical Research Institute (KEMRI) - Wellcome Trust Research Programme, Nairobi, Kenya); Alessandra Augusta Barroso Penna e Costa (Instituto Fernandes Figueira, Fundação Oswaldo Cruz, Rio de Janeiro, Brazil); Tania R.D. Saad Salles (University of Rio de Janeiro, Brazil); Vanessa van der Linden (Recife, Brazil), and Khalid Yunis (American University of Beirut, Lebanon).

4.4 Review date

These recommendations have been produced under WHO emergency procedures and will remain valid until December 2016. Literature will be reviewed routinely to determine whether updates need to be made to the guideline, specifically with regards to the spectrum of disease. The Departments of Maternal, Newborn, Child and Adolescent Health and Mental Health and Substance Abuse at WHO Geneva will be responsible for reviewing this guidance at that time, and updating it as appropriate. WHO welcomes queries and suggestions regarding the content of this guidance. Please email suggestions to mncahi@who.int.

5. References


48. McCarthy M. Severe eye damage in infants with microcephaly is presumed to be due to Zika virus. BMJ (Clinical research ed). 2016;352:i855.


