Congenital Cytomegalovirus

A European Expert Consensus Statement on Diagnosis and Management

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Congenital cytomegalovirus (cCMV) is the most common congenital infection in the developed world. Reported prevalence varies between cohorts but is approximately 7 per 1000 births.1 About half of cytomegalovirus (CMV)-infected babies with clinically detectable disease at birth are destined to have significant impairments in their development, and cCMV infection is implicated in approximately 25% of all children with sensorineural hearing loss (SNHL).1,2 Meta-analysis shows that although long-term sequelae, especially SNHL, are more common in those with clinically detectable disease at birth, they are also found in 13% of those without clinical features attributable to CMV on initial examination.3

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Despite the significant long-term impact of cCMV infection, there is limited evidence on which to base many treatment decisions in clinical practice. In an era of enhanced perinatal screening, fetuses and newborns are increasingly tested for CMV after abnormalities were detected during routine ultrasonography or maternal serology. Furthermore, otherwise “asymptomatic”, congenitally CMV-infected, newborns are being identified after detection of SNHL through newborn hearing screening programs. Because of earlier diagnosis, babies with cCMV now presenting to pediatricians differ from those primarily included in clinical trials of treatment reported in the literature.

A symposium was convened during the 2015 conference of the European Society of Paediatric Infectious Diseases to discuss the current management of cCMV. In attendance were clinicians from throughout Europe, many of whom are involved in policy for cCMV for their region/country.

This article summarizes the discussions at this meeting alongside the evidence informing them. A balanced perspective of the controversies in this area is presented and areas of consensus highlighted. Finally, where evidence is lacking, suggestions are made for future research efforts to address areas of unmet medical need.

The authors acknowledge the coexisting need for studies on the management of babies with symptoms consistent with cCMV, but in whom this diagnosis cannot be firmly established, and of those with symptomatic postnatal CMV infection; this article does not, however, address these groups.

The internationally accepted GRADE system for evaluating evidence has been used to illustrate points where relevant (Table 1).3

### DEFINITIONS OF SYMPTOMATIC DISEASE

Classically, cCMV infection is categorized as “symptomatic” or “asymptomatic” at birth. Differing definitions and opinions on what constitutes “symptomatic” CMV infection, however, makes interpreting the literature challenging. Indeed, some of the largest cohort studies include babies with SNHL at birth in the group described as being “asymptomatic” because no “clinically apparent disease” was detectable during newborn examination.4 In modern healthcare systems, whereby cCMV is increasingly detected through screening for other conditions, alongside increased accessibility of investigations, such as magnetic resonance imaging (MRI), the traditional dichotomy between clinically “apparent” and “inapparent” disease is becoming less meaningful. Table 2 summarizes the accepted clinical features of what constitutes “symptomatic” CMV infection, including need for studies on the management of cCMV for their region/country.

#### TABLE 1. Grade System of Evaluating Evidence3

<table>
<thead>
<tr>
<th>Quality Rating</th>
<th>Definition</th>
<th>Example Methodology</th>
<th>Depiction in Text</th>
</tr>
</thead>
<tbody>
<tr>
<td>High</td>
<td>Further research is very unlikely to change our confidence in the estimate of effect</td>
<td>Randomized trials or double-upgraded observational studies</td>
<td>A</td>
</tr>
<tr>
<td>Moderate</td>
<td>Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate</td>
<td>Downgraded randomized trials or upgraded observational studies</td>
<td>B</td>
</tr>
<tr>
<td>Low</td>
<td>Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate</td>
<td>Double-downgraded randomized trials or observational studies</td>
<td>C</td>
</tr>
<tr>
<td>Very low</td>
<td>Any estimate of effect is very uncertain</td>
<td>Triple-downgraded randomized trials, or downgraded observational studies, or case series/case reports</td>
<td>D</td>
</tr>
</tbody>
</table>

#### TABLE 2. Possible Signs and Symptoms in Children With Congenital CMV5–8

<table>
<thead>
<tr>
<th>Clinically detectable symptoms/signs</th>
<th>Physical Examination</th>
<th>Neuroimaging</th>
<th>Laboratory results</th>
<th>Neurologic signs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Small for gestational age (birth weight &lt;−2 SD for gestational age)</td>
<td>Microcephaly (head circumference &lt;−2 SD for gestational age)</td>
<td>Petechiae or purpura (usually found within hours of birth and persist for several weeks)</td>
<td>Blueberry muffin rash (intra dermal hematoipoiesis)</td>
<td>Jaundice</td>
</tr>
<tr>
<td>Hepatomegaly</td>
<td>Splenomegaly</td>
<td>Neurologic physical examination</td>
<td>Microcephaly (head circumference &lt;−2 SD for gestational age)</td>
<td>Neurologic signs (lethargy, hypotonia, seizures, poor sucking reflex)</td>
</tr>
<tr>
<td>Elevated liver enzymes (ALT/AST)</td>
<td>Conjugated hyperbilirubinemia</td>
<td>Cerebrospinal fluid</td>
<td>Abnormal cerebral fluid indices, positive CMV DNA</td>
<td></td>
</tr>
</tbody>
</table>

*CMV-associated jaundice can be present at the first day after birth and usually persists longer than physiologic jaundice.

ALT indicates alanine aminotransferase; AST, aspartate aminotransferase; SD, standard deviations.
cCMV disease with those symptoms detectable on newborn examination listed separately to those detectable only if specific investigations are conducted, for example, when cCMV is already suspected.5–8

Full Consensus Within This Expert Group Was That

1. For the purposes of research and publication, newborns identified as having cCMV disease after abnormal clinical examination at birth (such as microcephaly, small for gestational age (SGA), widespread petechiae, hepatosplenomegaly) should be differentiated from those babies identified through screening or investigation for other disorders, for example, those tested for CMV after known/likely maternal infection or abnormal newborn hearing screening. This differentiation would allow for more accurate assessment of the prognostic value of individual manifestations of “symptomatic” disease on longer-term outcomes as already shown in other publications.9

2. “Symptomatic” cCMV should be considered as “severe,” “moderate” or “mild” disease.

a. “Mild” disease includes those with isolated (1 or 2 at most), otherwise, clinically insignificant or transient findings, such as petechiae, mild hepatomegaly or splenomegaly or biochemical/hematologic abnormalities (such as thrombocytopenia, anemia, leukopenia, borderline raised liver enzyme abnormalities or conjugated hyperbilirubinemia) or SGA (defined as weight for gestational age <−2 standard deviations) without microcephaly.

b. “Severe” disease includes those with central nervous system (CNS) involvement (abnormal neurologic or ophthalmologic examination, microcephaly or neuroimaging consistent with cCMV disease [such as calcifications, moderate to severe ventriculomegaly, cysts, white matter changes, cerebral or cerebellar hypoplasia, hippocampal dysplasia, neuronal migration abnormalities])10 or with life-threatening disease.

The Majority Agreed That

2b. “Severe” disease also includes babies with evidence of severe single-organ disease (including those with clinically significant liver enzyme abnormalities [liver “failure”] and marked hepatosplenomegaly) or those with significant multiorgan involvement. Babies with transient or otherwise clinically insignificant abnormalities (ie, the babies are not “sick”) that resolve spontaneously over a few weeks are not included in this group even if these abnormalities are multiple.

2c. A further group exists that may be considered to have “moderate” disease. This group is heterogeneous and includes, for example, those with persistent (eg, more than 2 weeks duration) abnormalities of hematologic/biochemical indices or more than 2 “mild” disease manifestations (as listed earlier). Because of lack of evidence, full consensus could not be reached on how to approach this group, and treatment decisions are currently made on a case by case basis. Development of a validated clinical scoring system for disease severity at presentation and risk of sequelae would be beneficial for both counseling parents and informing treatment decisions.

3. Defining CNS involvement

a. It remains uncertain whether some, nonspecific findings detected on cranial ultrasound (CrUSS) and MRI (particularly isolated lenticulostrial vasculopathy [LSV]) constitute clinically significant CNS disease. LSV has been detected in 0.4%–5.8% of all neonates undergoing an ultrasound, and only 5% has been associated with cCMV.11–12 Some have suggested isolated LSV as a marker of risk for SNHL.11 Others have found only more extensive neuroimaging abnormalities to be of prognostic value.11,14 The majority at this meeting would not consider LSV in isolation to be a notable CNS manifestation of disease. It is suggested that neuroradiologic abnormalities not known to be clearly associated with CMV disease and adverse outcomes are discussed with a suitably experienced neuroradiologist, particularly, if the results of these discussions might influence treatment decisions.

b. The exact pathophysiology of SNHL is not clear but is likely secondary to infection and degradation of sensory structures within the inner ear.15,16 It is therefore debated whether isolated SNHL should truly be considered a CNS manifestation of infection and, as a consequence, whether such children should be considered comparable to those with CNS disease included in published clinical trials. No studies have addressed this specific population, but a nonrandomized cohort study observing the effects of valganciclovir in isolated SNHL is in progress (clinicaltrials.gov NCT02005822). The majority of experts at this meeting would categorize babies with isolated, confirmed SNHL in the “severe”/CNS group because bilateral SNHL is not only associated with likely long-term impairments but was also included in the criteria for recruitment in the only randomized controlled trials (RCTs) in cCMV. However, consensus was not reached because the spectrum of hearing loss is wide, and treatment of isolated SNHL has not been evaluated in any RCTs.

WHEN SHOULD TESTING FOR CONGENITAL CMV BE CONSIDERED?

Indications for testing for cCMV are based on the presence of one or more of the most frequently observed clinical features (Table 3).17 Unfortunately, predictive values for each of these features are not available.

Full Consensus Within This Expert Group Was That Testing for cCMV Should Be Performed in

1. Fetuses with ultrasound/MRI imaging consistent with cCMV disease (by appropriately timed antenatal testing of amniotic fluid).18 (Quality C, Level 1)

2. Newborns where there is a maternal history of suspected primary CMV infection during pregnancy. If antenatal testing of amniotic fluid has been conducted, it is suggested that CMV infection should still be confirmed at birth because both false-positive and -negative results have been reported.19 (Quality C, Level 1)

3. Newborns with signs/symptoms consistent with cCMV disease (see Table 2; including those with findings consistent with cCMV on antenatal imaging). (Quality B, Strength 1)

4. Children with confirmed SNHL.16 Systems need to be established to ensure testing for cCMV occurs, where possible, in the first 21 days of life because dried blood spot (DBS) are not always readily available for testing (see below). (Quality B, Strength 1)

The Majority Agreed That

5. Newborns who are SGA should not routinely be tested. Studies in SGA newborns have shown the prevalence of cCMV to be 0%–5.2%.19,22 However, the majority of


**TABLE 3. Clinical Features That Should Lead to Testing for Congenital CMV**

<table>
<thead>
<tr>
<th>Feature</th>
<th>Notes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Neonates</strong></td>
<td></td>
</tr>
<tr>
<td>Physical examination</td>
<td></td>
</tr>
<tr>
<td>Hepatosplenomegaly</td>
<td></td>
</tr>
<tr>
<td>Petechiae, purpura or blueberry muffin rash in a newborn</td>
<td></td>
</tr>
<tr>
<td>Jaundice (prolonged or conjugated hyperbilirubinemia)</td>
<td></td>
</tr>
<tr>
<td>Microcephaly (head circumference &lt;−2 SD for gestational age)</td>
<td>Consider if symmetrically small for gestational age (&lt;−2 SD for gestational age)</td>
</tr>
<tr>
<td><strong>Neurology</strong></td>
<td></td>
</tr>
<tr>
<td>Seizures with no other explanation</td>
<td></td>
</tr>
<tr>
<td>Laboratory parameters</td>
<td></td>
</tr>
<tr>
<td>Prolonged jaundice with transaminitis</td>
<td></td>
</tr>
<tr>
<td>Conjugated hyperbilirubinemia</td>
<td></td>
</tr>
<tr>
<td>Unexplained thrombocytopenia, consider if leukopenia or anemia</td>
<td></td>
</tr>
<tr>
<td><strong>Neuroimaging</strong></td>
<td></td>
</tr>
<tr>
<td>Intraocular calcification (often periventricular)</td>
<td></td>
</tr>
<tr>
<td>Intraocular ventriculomegaly without other explanation</td>
<td></td>
</tr>
<tr>
<td>Consider in the case of periventricular cysts, subependymal pseudocysts, germinal cysts, white matter abnormalities, cortical atrophy, migration disorders, cerebellar hypoplasia, lenticulostriate vasculopathy</td>
<td></td>
</tr>
<tr>
<td><strong>Visual examination</strong></td>
<td></td>
</tr>
<tr>
<td>Abnormal findings on ophthalmologic examination consistent with congenital CMV (eg, chorioretinitis)</td>
<td>Consider if congenital cataracts</td>
</tr>
<tr>
<td><strong>Failed neonatal hearing screen</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Maternal serology</strong></td>
<td></td>
</tr>
<tr>
<td>Evidence of maternal seroconversion*</td>
<td></td>
</tr>
<tr>
<td>Consider in women with known CMV infection (known IgG seropositive at start of pregnancy), particularly, if symptoms or virologic examination consistent with suspected CMV reactivation/reinfection*</td>
<td></td>
</tr>
<tr>
<td>Prematurity†</td>
<td></td>
</tr>
<tr>
<td>Older children</td>
<td></td>
</tr>
<tr>
<td>Sensorineural hearing loss: new diagnosis</td>
<td></td>
</tr>
</tbody>
</table>

Features in bold are those where there is consensus for testing. Features in italics are those that might lead to testing in individual circumstances and depending on local practice.

*Seek expert clinical virology advice for interpretation of virologic investigations in pregnancy.
†Baseline screening to differentiate between congenital and postnatal CMV infection is helpful for extremely premature infants (<28 weeks gestational age) who are at increased risk of symptomatic postnatal infection. SD indicates SD indicates standard deviations.

Studies report a prevalence of 1.4%–1.8%, which is not significantly higher than the prevalence of cCMV in the general population. Therefore, evidence is insufficient to justify screening all newborns with isolated SGA for cCMV. None of these studies distinguish between asymmetrical (with normal head circumference) and symmetrical SGA, but when head circumference was mentioned, most SGA babies with cCMV had microcephaly (head circumference <−2 SD standard deviations).23,24 Because of this, and the poor prognostic outcome of children with CMV and microcephaly, many present at this meeting test those babies with symmetrical SGA but not those with preserved head growth.14 (Quality C, Strength 2)

6. Prematurity. Evidence that premature babies have a higher incidence of cCMV is limited.20,22 Testing extremely premature babies (<28 weeks gestational age) at birth does, however, assist in differentiating between congenital and postnatal infection. This may be very helpful in guiding the management of these babies that are particularly vulnerable to symptomatic postnatal infection. However, consensus was not reached regarding this area, with cost being a factor among other considerations.24 (Quality C, Strength 2)

7. Testing of babies born to mothers who are known to be CMV seropositive at the establishment of pregnancy. Although maternal nonprimary CMV infection is known to be important when considering the overall burden of cCMV disease, testing all babies born to these women, particularly in populations with high maternal seroprevalence, is tantamount to universal neonatal screening.23,26 Identifying women with nonprimary CMV who are at highest risk of transmitting infection to their fetus remains elusive. It was agreed that individual case discussion and local policy should therefore dictate practice in this area. Further research is clearly needed.

**LABORATORY DIAGNOSIS OF CONGENITAL CMV INFECTION**

Testing for cCMV using CMV polymerase chain reaction (PCR) in urine is highly reliable: sensitivity is 100% and specificity 99%.27 One negative urine specimen in a neonate is therefore sufficient to exclude infection, and repeat sampling is not necessary. After 21 days, a urine positive for CMV could be because of CMV acquired postnatally from, for example, passage through the birth canal or through breast milk. As CMV PCR techniques are becoming more sensitive, earlier testing, before the age of 14 days, is recommended.23

CMV PCR testing of saliva is an alternative and is easy to perform. Samples should be taken immediately before feeding in breastfed newborns, and confirmed with urine, as false-positive results have been reported.21,23

PCR assay of neonatal DBS can be performed retrospectively in an attempt to diagnose cCMV after the first 21 days of life. Sensitivity is around 84% in meta-analysis but is highly variable depending on the laboratory techniques used and the population being tested; a negative DBS PCR cannot, therefore, be used to definitively exclude a diagnosis of cCMV.22

**Full Consensus Within This Expert Group Was That**

1. Testing for cCMV should be performed using a single CMV PCR of urine obtained within 21 days of birth but ideally within 14 days of birth (Quality B, Strength 1).
2. Saliva PCR testing can be an alternative, but a positive result should be confirmed using urine (Quality B, Strength 1).
3. After the age of 21 days, CMV DNA PCR of stored DBS can be used to diagnose cCMV retrospectively; sensitivity is relatively low, and a negative test cannot be used to definitively exclude a diagnosis of cCMV (Quality B, Strength 1).

**RECOMMENDED INVESTIGATIONS AFTER CONFIRMING A DIAGNOSIS OF CONGENITAL CMV INFECTION**

After a virologic diagnosis of cCMV infection has been made, additional investigations are necessary to evaluate the extent of disease and to assist with discussions regarding prognosis and treatment.

**Full Consensus Within This Expert Group Was That**

1. The investigations below are conducted in any baby in whom a diagnosis of cCMV is confirmed, looking specifically for the manifestations of disease (Table 2):
   - Complete blood count, liver enzymes, (conjugated) bilirubin
   - Renal function (before initiating therapy)
5. Babies with “mild” cCMV disease (as defined earlier) should not receive treatment. No studies have clearly addressed treatment in this group. Most present at this meeting would not, therefore, treat babies with 1 or 2 isolated or transient, clinically insignificant, manifestations of disease (Quality C, Strength 2).

6. Babies with “moderate” cCMV disease (as defined earlier). Evidence for treating babies with multiple, but not severe, manifestations of disease (including jaundice, hepatosplenomegaly without significantly raised liver enzymes, SGA) is limited. It is, therefore, recommended that these cases are discussed on a case-by-case basis with a clinician with experience of managing babies with cCMV (such as a pediatric infectious disease specialist) (Quality B, Strength 2).

TREATMENT

No antiviral drugs are currently licensed for the treatment of cCMV. Although many case reports and cohort studies have reported on treatment for cCMV, there are results from only 2 RCTs.7,41–44 The first of these studies evaluated 6 weeks’ intravenous ganciclovir treatment in neonates (<1 month of age), gestational age ≥32 weeks and clinically apparent disease in the newborn period with evidence of CNS disease (including microcephaly, intracranial calcification, abnormal CSF indices for age, hearing deficit and chorioretinitis).1 Improved hearing and neurodevelopmental outcomes were shown, but there was significant loss to follow-up.7,44 A more recent trial compared 6-week to 6-month treatment with oral valganciclovir and included babies with any evidence of symptomatic (including non-CNS) cCMV disease.45 Few babies enrolled, however, had isolated, mild clinical features, and none in the 6-month treatment group had isolated SNHL (D Kimberlin 2015, personal email correspondence, 28 April). A modest benefit on both 2-year hearing and neurodevelopmental outcomes was shown with the 6-month treatment course. The longer treatment course improved likelihood of better hearing outcomes most notably in those with preexisting CNS involvement. Longer duration of therapy was only statistically significant, however, for “total ear” hearing as opposed to “best ear” hearing (which is of greater functional significance) and only once adjusted for baseline CNS involvement. Given the natural resolution of some features of cCMV disease in published cohorts, alongside the delayed onset of hearing loss and fluctuations in SNHL reported in cCMV, it is even more challenging to draw any conclusions regarding treatment effect from uncontrolled studies.7,16,45

Clinical trials to date do not, therefore, provide good evidence on which to base treatment decisions for many of the infants presenting to clinicians in everyday clinical practice.

Table 4 provides guidance on which infants should be offered treatment after a risk versus benefit discussion with the family. This table and associated text indicate areas where consensus was reached. Much discussion focused around the treatment of babies with less severe cCMV disease and whether the minimal additional benefit shown in the 6-month treatment course was sufficient to justify such a prolonged course of treatment. Although clinical findings such as SGA and petechiae have been shown in historical cohorts to predict risk for SNHL, more recent reanalysis of data indicates that these findings in isolation are generally associated with disease-free outcomes in babies presenting without other manifestations of symptomatic disease.5,6,6 Opinion on the severity, or number, of symptoms justifying antiviral treatment remains divided, and it is therefore strongly recommended that clinicians discuss treatment initiation and duration with an expert in this area.

Full Consensus Within This Expert Group Was That

1. Babies with evidence of CNS disease should receive antiviral treatment (Quality A, Strength 1). Treatment should be preferably for 6-months duration (Quality B, Strength 2).

2. Babies with no clinical/laboratory findings consistent with CMV disease should not receive treatment because no evidence exists to support treatment in this group (Quality D, Strength 1 [not to treat]).

3. Babies with evidence of life-threatening disease or severe single-organ disease or multiorgan involvement should receive treatment. Although evidence is limited, particularly for life-threatening disease, consensus was that treatment should be considered in this group (Quality B, Strength 1). Consensus could not be reached on duration of treatment in this group.

4. Oral valganciclovir is now the drug of choice. Intravenous ganciclovir should be used in babies unable to tolerate oral drug or where gastrointestinal absorption is uncertain (Quality A, Strength 1).

The Majority Agreed That

1. Babies with evidence of life-threatening disease or severe single-organ disease or multiorgan involvement should receive treatment. Although evidence is limited, particularly for life-threatening disease, consensus was that treatment should be considered in this group (Quality B, Strength 1). Consensus could not be reached on duration of treatment in this group.

The Majority Agreed That

5. Babies with “mild” cCMV disease (as defined earlier) should not receive treatment. No studies have clearly addressed treatment in this group. Most present at this meeting would not, therefore, treat babies with 1 or 2 isolated or transient, clinically insignificant, manifestations of disease (Quality C, Strength 2).

6. Babies with “moderate” cCMV disease (as defined earlier). Evidence for treating babies with multiple, but not severe, manifestations of disease (including jaundice, hepatosplenomegaly without significantly raised liver enzymes, SGA) is limited. It is, therefore, recommended that these cases are discussed on a case-by-case basis with a clinician with experience of managing babies with cCMV (such as a pediatric infectious disease specialist) (Quality B, Strength 2).
TABLE 4. Summary of Treatment Recommendations

<table>
<thead>
<tr>
<th>Disease Manifestation</th>
<th>Treatment Recommendation</th>
<th>Level of Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>CNS disease</td>
<td>Ganciclovir/valganciclovir: duration 6 months*</td>
<td>Treatment: Quality A, Strength 1 (to treat)</td>
</tr>
<tr>
<td>Microcephaly, CNS calcification, chorioretinitis</td>
<td>Ganciclovir/valganciclovir: minimum of 6 weeks, up to 6 months*†</td>
<td>Treatment: Quality B, Strength 2</td>
</tr>
<tr>
<td>White matter changes (or other abnormalities on MRI consistent with CMV disease)*</td>
<td>No treatment</td>
<td>Treatment: Quality B, Strength 2</td>
</tr>
<tr>
<td>Other “severe” disease (includes life-threatening or severe single-organ or multiorgan non-CNS disease)</td>
<td>No treatment</td>
<td>Treatment: Quality C, Strength 2 (for no treatment)</td>
</tr>
<tr>
<td>“Mild” disease: isolated or transient disease (eg, jaundice, Petechiae, SGA in isolation; max 2 abnormalities)</td>
<td>Ganciclovir/valganciclovir: Duration 6 months*</td>
<td>Treatment: Quality C, Strength 1</td>
</tr>
<tr>
<td>No clinical or biochemical findings of disease (a detectable CMV viremia)</td>
<td>Consider treatment after discussion with specialist</td>
<td>Treatment: Quality C, Strength 2</td>
</tr>
<tr>
<td>Majority opinion: but no consensus Isolated hearing deficit§</td>
<td>Duration: Minimum of 6 weeks and up to 6 months*</td>
<td>Treatment: Quality B, Strength 2</td>
</tr>
<tr>
<td>“Moderate” disease (see text for definition; eg, multiple minor findings consistent with CMV disease)⁹</td>
<td>Treatment: Quality D, Strength 1 (for no treatment)</td>
<td></td>
</tr>
</tbody>
</table>

There is currently only evidence for starting treatment in the first month of life.

*Limited evidence without full consensus: see text for further description.

†It was suggested (without consensus) that treatment might continue in this group until the underlying clinical manifestation of disease (eg, hepatitis) resolved because benefit of 6 months treatment is unclear.

§No studies address this particular group, although they were included in eligibility criteria for treatment in both published RCTs of treatment.

7. Treatment of isolated SNHL: The majority at this meeting would include SNHL at birth in their indications for treatment because this was in the inclusion criteria for treatment in previous RCTs. Furthermore, the main benefit of treatment is in preserving hearing rather than improving hearing once damage exists, with good outcomes reported in observational studies (with likely bias).⁷,⁴¹,⁴⁷ There was no, however, consensus, and it is acknowledged that no RCTs have specifically addressed treatment effect in this group of babies who are usually now identified through newborn hearing screening programs (Grade C, Strength 1).

8. Drug dose and formulation: Although oral valganciclovir is now first-line treatment in most cases, it is currently unknown whether valganciclovir reaches target areas as effectively as ganciclovir or, indeed, where drug should be targeted (eg, CNS or inner ear) because no studies have directly compared the 2 drugs. In those with severe disease, particularly if absorption is uncertain, intravenous ganciclovir is, therefore, preferred by some in early stages of treatment until oral therapy can be reliably tolerated (Quality C, Strength 1).

9. Treatment duration in cases without CNS involvement: In those infants in whom the decision is taken to give antiviral treatment, the majority would treat for 6 months. However, there was no consensus on this point in light of the modest benefit shown for longer treatment courses in the only RCT (Quality B, Strength 2).

10. Treating babies older than 28 days: Treatment of older children has not been addressed in any RCTs, although it is acknowledged that the 28-day cutoff is also not evidence based. Retrospective case series of small numbers of babies treated outside the newborn period have reported good outcomes.⁴⁸,⁴⁹ Babies found to have SNHL after hearing screening at birth often do not have a diagnosis of CMV confirmed until outside the 1-month “window of evidence” for treatment. No consensus was reached on how late it might be acceptable to start treatment in this scenario, or in the eventuality of hearing deterioration. Two RCTs are currently evaluating the use of treatment in older children with cCMV and SNHL (clinicaltrials.gov NCT01649869 and NCT02606266), which may clarify this debate. (Evidence for treating outside the newborn period Quality D, Strength 2.)

SIDE EFFECTS OF ANTIVIRAL TREATMENT

Much of the debate around treating less severely affected babies relates to the potential side effects of currently available antiviral drugs. Significant neutropenia is frequently observed during antiviral treatment in infants. This is reported less commonly with valganciclovir than with ganciclovir (21% compared with 65%).⁷,⁴¹,⁴⁵,⁵⁰ Neutropenia generally occurs during the first month of treatment, with no increased toxicity observed after 6 weeks in those randomized to receive 6-month treatment compared with placebo in the only RCT evaluating this.⁴¹ The oral administration of ganciclovir also removes the burden of hospitalization and risk of nosocomial infections and central line complications observed during treatment with ganciclovir. Hepatotoxicity has been reported in up to 30% of those treated with ganciclovir and thrombocytopenia in a similar proportion.⁵¹ In the most recent study of treatment with valganciclovir, deranged liver function was observed, but this was neither clinically nor statistically significant when compared with placebo. In all studies, abnormal biochemical and hematologic parameters resolved after drug discontinuation.

Long-term side effects have not been evaluated in neonates treated with ganciclovir or valganciclovir. Animal studies raise the theoretical risk of gonadotoxicity and carcinogenicity.⁵²,⁵³ Although this has not been observed in humans to date, parents should be counseled about these potential risks, particularly when considering treatment in those groups in which benefit has not been clearly shown. No adverse long-term effects have been documented in a small cohort of babies treated in early neonatal studies and followed up to puberty (NCT00031421, unpublished data).

MONITORING OF BABIES DURING TREATMENT

Table 5 summarizes a proposed monitoring strategy for babies treated for cCMV. These recommendations are based on the safety monitoring and data obtained from the published RCTs.⁷,⁴¹
TABLE 5. Monitoring and Follow-Up According to Treatment Status

<table>
<thead>
<tr>
<th>No Treatment Given</th>
<th>Treatment Given</th>
</tr>
</thead>
<tbody>
<tr>
<td>—</td>
<td>Investigations whilst on treatment*</td>
</tr>
<tr>
<td>—</td>
<td>FBC,* LFT† and U&amp;E suggested weekly for first 4 weeks and then at least monthly</td>
</tr>
<tr>
<td>—</td>
<td>Weight measurement and drug dose review at time of blood sampling</td>
</tr>
<tr>
<td>—</td>
<td>Viral load at baseline (Quality C, Strength 2).</td>
</tr>
<tr>
<td>—</td>
<td>Consider Viral load 2–4 weekly whilst on antiviral therapy (not consensus)</td>
</tr>
<tr>
<td>—</td>
<td>Consider therapeutic drug monitoring if:</td>
</tr>
<tr>
<td>—</td>
<td>Viral load increase &gt;1.0 log₁₀ during treatment¶</td>
</tr>
</tbody>
</table>
|**Consider Viral load monitoring: Some centers report monitoring viral load to assist in decisions regarding adequate drug dosing and detection of potential drug resistance; however, most experts at this meeting do not conduct this routinely. Treatment duration is not altered by any viral parameters, and rebound of virus after treatment discontinuation is well documented with no demonstrable association with long-term outcomes.** (Quality D, Strength 2)

Follow up

Audiology assessment every 3–6 months in the first year, then every 6 months until 3 years of age and then every 12 months until 6 years old) (Quality C, Strength 1)

Pediatric infectious disease clinic review (or general pediatric clinic after consultation with a specialist) until at least 1 year, and ideally 2 years, of life. (Quality D, Strength 1)

Monitor development. (Quality D, Strength 1)

Ophthalmic assessment as directed by ophthalmologist, but baseline and annual review up to age 5 years in those with clinically detectable symptoms/signs at birth recommended.** (Quality D, Strength 2)

FBC indicates full blood count; LFT, liver function tests; U&E, urea, creatinine and electrolytes.

* Interrupt treatment or consider granulocyte colony stimulating factor (GCSF) if absolute neutrophil count <0.5 × 10⁹/L. Decreasing dose may be considered for less severe neutropenia.

† Increase frequency or seek advice if there is deterioration.

‡ Measuring viral load is not evidence based but offers some evaluation of virus response and enables detection of possible viral resistance.

§ Consider CMV resistance testing (sequencing) in unexplained elevations/breakthrough of viremia.

¶ Measuring viral load is not evidence based but offers some evaluation of virus response and enables detection of possible viral resistance.

** There is limited evidence on late ocular manifestations of cCMV. They are rare and include visual impairment and strabismus.

There are no data to support therapeutic drug monitoring.55,56 Therapeutic drug monitoring may, however, have a role when toxicity is a concern (eg, in those with impaired renal function) or where there are concerns about treatment response.

Full Consensus Within This Expert Group Was That

1. Where treatment is given, babies should have regular weight measurement and safety monitoring to enable appropriate dose adjustment of medication (see Table 5). (Quality A, Strength 1)

2. Where treatment is given, parents should be fully counselled about both the known and potential side effects of treatment with current antivirals. (Quality A, Strength 1 for short-term side effects; Long-term, no published studies)

3. Although there are theoretical risks of longer term treatment toxicity, no large cohorts have been followed up to enable this to be fully evaluated in humans treated during early life. Where possible, children receiving antiviral treatment should, therefore, be entered into a registry to enable ongoing pharmacovigilance.

Only a Minority Agreed That

1. Viral load monitoring: Some centers report monitoring viral load to assist in decisions regarding adequate drug dosing and detection of potential drug resistance; however, most experts at this meeting do not conduct this routinely. Treatment duration is not altered by any viral parameters, and rebound of virus after treatment discontinuation is well documented with no demonstrable association with long-term outcomes. (Quality D, Strength 2). If viral load is checked after discontinuing drug, it is suggested that parents are forewarned of the likelihood that virus will be detectable and that this is of unknown significance.

FOLLOW-UP

Table 5 summarizes recommended follow-up of babies with cCMV (both treated and untreated).

The recommendation for audiologic follow-up is based on long-term surveillance studies of SNHL in cCMV.54,56 Frequent follow-up is suggested during the first 2 years of life because this is the period of highest risk for development of cCMV-associated hearing loss and a critical period for language development. Early detection of SNHL during this period is also most likely to improve long-term outcomes.55 Monitoring should continue into early childhood, however, because deterioration in hearing continues throughout early life.55 (Quality B, Strength 1).

Neurodevelopmental follow-up is suggested at 1 and 2 years of age ideally with formal neurodevelopmental assessment. This is not, however, routinely conducted in all centers, and there is no evidence-based benefit in this particular group, although early detection of functional impairments is generally agreed to be beneficial.

Ophthalmic follow-up is recommended annually at least until children can talk in those with clinically detectable disease at birth, but not in those without, because deterioration in vision has been observed in this group (Quality C, Strength 1).6

Families should be given information for local/national support groups where these exist (see acknowledgements). Where cCMV parent groups are not easily accessible, parents of children with hearing loss may find support from groups for those with hearing impairment.
RECOMMENDATIONS FOR FUTURE PEDIATRIC RESEARCH

1. Clinical trials addressing treatment of those with more “minor” manifestations of disease/no clinically detectable disease at birth and those with isolated SNHL.
2. Clinical studies of antenatal therapies to decrease transmission of infection and cCMV disease once infection is established.
3. Publications relating to cCMV should make it clear how those included were identified (ie, babies presenting with clinically detected “symptoms” vs screened babies identified through existing antenatal or postnatal screening pathways including hearing screening programs), or after further investigation of abnormalities, such as thrombocytopenia, found incidentally when blood sampling is performed for other indications.
4. Development of clinical prediction models to better categorize severity of disease (CNS vs non-CNS and babies with single vs multiple findings of disease) and associated outcomes to assist counseling of parents.
5. Studies of neuroimaging, particularly MRI, and added value with regards to predicting long-term impairments particularly in those without clinically detectable disease at birth through studies involving unselected cCMV cohorts.
6. Clinical trials of alternative treatment durations and new anti-CMV therapies when available.
7. Biomarkers. It seems unlikely that a predefined duration of treatment will be similarly beneficial in babies with such varying clinical manifestations of disease and likely variable viral burden and host immune function. The development of both host and virologic biomarkers of long-term outcomes would greatly enhance design of future RCTs and enable more accurate counseling and resource allocation.
8. All children receiving treatment should be captured in a registry to enable ongoing pharmacovigilance for any long-term effects of antiviral medication.
9. Identification of risk factors for maternal virus transmission, particularly, in those mothers with previous known exposure to CMV (CMV IgG seropositive).

CONCLUSIONS

As stated at the outset, this article represents the consensus opinion of a group of professionals with a particular interest in cCMV. It highlights that much of our practice is based on limited data but identifies areas where there is nonetheless consensus among experts. Recent publications have shown potential cost-effectiveness of screening at birth for cCMV, although these calculations are constrained by the issues raised in this article regarding true quantification of benefits of treatment and agreed treatment duration in certain patient groups. It will be challenging to address many of the research questions raised through RCTs, given the significant resources and long-term follow-up required alongside potential difficulties in recruiting into such studies when treatment is anecdotally being offered more freely. Collecting accurate data on disease manifestations and treatment outcomes in different patient groups alongside maternal demographics can, however, inform treatment strategies as previously shown very effectively for the management of pediatric human immunodeficiency virus. This requires a unified approach to initial diagnostic tests, definitions of symptomatology and follow-up which is currently being addressed by a network of clinicians with an interest in this area through both national and European initiatives such as Paediatric European Network for Treatment of AIDS - Infectious Diseases, the European Congenital CMV Initiative and European Society of Paediatric Infectious Diseases and European Society for Clinical Virology (ESCV). It should also be reiterated that this article focuses on postnatal aspects of diagnosis and treatment. There is an associated and simultaneous need for work alongside obstetric and fetal medicine colleagues to address similar uncertainties in aspects of antenatal care. It is hoped that through such collaborations, progress will be made in decreasing infection and disease in fetuses, newborns and subsequently older children with cCMV.

ACKNOWLEDGMENTS

The CMV-infected children and their parents/caregivers who have cared for and who challenge us on a daily basis to provide evidence-based and consistent care including those parents who are involved with supporting other parents through local or national support groups including (but not exclusively) CMV Action (www.cmvaaction.org.uk); https://www.stopcito.megalovirus.org.

REFERENCES


