

Congenital cytomegalovirus: a systematic review of newborn screening

External review against programme
appraisal criteria for the UK National
Screening Committee

Version: Publication

Author: York Evidence Synthesis Group

Date: August 2026

**The UK National Screening Committee secretariat is hosted by the
Department of Health and Social Care**

Contents

Congenital cytomegalovirus: a systematic review of newborn screening	1
Contents.....	2
About the UK National Screening Committee (UK NSC)	5
Abbreviations.....	6
Plain English summary	7
Executive summary	9
Purpose of the review	9
Background.....	9
Focus of the review.....	9
Recommendation under review.....	10
Findings of this review and gaps in the evidence	10
Recommendations on screening.....	12
Limitations	13
Evidence uncertainties	13
Introduction and approach	14
Background.....	14
The condition.....	14
Screening and treatment	15
Current global landscape of newborn screening	15
Current policy context and previous reviews	16
Objectives	16
Methods.....	17
Databases/sources searched.....	18
Eligibility for inclusion in the review	19
Inclusion criteria	19
Appraisal of studies for quality and risk of bias.....	22
Data extraction and methods of synthesis.....	22

Question level synthesis	24
Overall review summary.....	24
Criterion 4:	25
Previous work.....	25
Description of the current evidence	25
Additional material relating to Criterion 4.....	38
Urine testing	38
Other testing for cCMV infection	42
Summary of Findings Relevant to Criterion 4	44
Criterion 7:	46
Description of the current evidence	46
Combination tests.....	70
Summary of Findings Relevant to Criterion 7	71
Criteria 11 & 13:	75
Description of the current evidence	75
Summary of Findings Relevant to Criteria 11 and 13	91
Description of the current evidence	93
Summary of Findings Relevant to Criterion 12	94
Review summary.....	95
Conclusions and implications for policy.....	95
Evidence gaps and recommendations for research	95
Limitations	96
Appendix 1 — Search strategy	98
Electronic databases and resources	98
Database search strategies	99
Website searches.....	108
Appendix 2 — Included and excluded studies	110
PRISMA flowchart	110
Publications included after review of full text articles.....	111

Appendix 3 — Summary and appraisal of studies relating to Criterion 7	119
Data Extraction	119
Appraisal of study quality and risk of bias	174
Appendix 4 – Relevant ongoing studies and unpublished studies	185
Appendix 5 – UK NSC reporting checklist for evidence summaries	190
References	194

About the UK National Screening Committee (UK NSC)

The UK NSC advises ministers and the NHS in the 4 UK countries about all aspects of [population screening](#) and supports implementation of screening programmes.

Conditions are reviewed against [evidence review criteria](#) according to the UK NSC's [evidence review process](#).

Read a [complete list of UK NSC recommendations](#).

UK National Screening Committee, Southside, 39 Victoria Street, London, SW1H 0EU

www.gov.uk/uknsc

Blog: <https://nationalscreening.blog.gov.uk/>

For queries relating to this document, please contact: uknsc@dhsc.gov.uk.

© Crown copyright 2016

You may re-use this information (excluding logos) free of charge in any format or medium, under the terms of the Open Government Licence v3.0. To view this licence, visit OGL or email psi@nationalarchives.gsi.gov.uk. Where we have identified any third party copyright information you will need to obtain permission from the copyright holders concerned.

Published August 2026

Abbreviations

cCMV	Congenital cytomegalovirus
CI	Confidence interval
CMV	Cytomegalovirus
CNS	Central nervous system
CSF	Cerebrospinal fluid
CT	Computed tomography
DBS	Dried blood spots
ECCI	European Congenital Cytomegalovirus Initiative
FN	False negative
FP	False positive
HTA	Health Technology Assessment
LSV	Lenticulostriate vasculopathy
MRI	Magnetic resonance imaging
NHS	Newborn hearing screening
NICU	Neonatal intensive care unit
NPV	Negative predictive value
NSC	National Screening Committee
PCR	Polymerase chain reaction
PPV	Positive predictive value
PRISMA	Preferred Reporting Items for Systematic Reviews and Meta-Analyses
QUADAS	Quality Assessment of Diagnostic Accuracy Studies
RCT	Randomised controlled trial
SNHL	Sensorineural hearing loss
TN	True negative
TP	True positive
US	Ultrasound
WMA	White matter abnormalities

Plain English summary

Cytomegalovirus (CMV) is a common virus which usually causes mild flu-like symptoms, but sometimes causes problems in babies and people with a weakened immune system like elderly people and people undergoing transplants. If the virus is passed on to a fetus during pregnancy, the baby may be born with CMV infection, called congenital CMV (cCMV). Around 10% of babies with cCMV are born with symptoms and around half of these babies will develop long-term complications, such as hearing loss and developmental problems. Around 10-15% of those without early symptoms will develop problems later. Testing for cCMV at birth could enable treatment which could prevent or reduce long-term harm. Testing should ideally be done within three weeks of birth to avoid confusing cCMV with a CMV infection acquired after birth, which is less likely to cause long-term harm.

The UK National Screening Committee (UK NSC) does not currently recommend screening for cCMV. As part of its regular reviews on the evidence, the UK NSC commissioned York Evidence Synthesis Group to review existing studies to assess whether screening tests for newborn babies (defined as up to 28 days old in this review) can identify which babies are likely to develop long-term complications and would benefit from early treatment. We found 217 studies and extracted data from 171 studies that were reported in full.

We found 39 studies assessing how accurate saliva and dried blood spot testing were at detecting cCMV. These found that both tests were very accurate and will probably identify 90% or more of all cCMV cases. Hence both saliva and dried blood spot testing could be used to screen newborns for cCMV.

We found 97 studies that examined tests or characteristics of newborn babies with cCMV to identify who might experience long-term harm. The types of tests or characteristics varied, and evidence on most tests or characteristics was limited. The factor most clearly associated with long-term harm from cCMV was babies who showed symptoms of cCMV infection at birth, particularly hearing loss. There was some evidence that performing magnetic resonance imaging (MRI) scans on newborns can predict long-term developmental issues.

We found 37 studies that looked at newborn screening programmes to spot cCMV. None compared screening to current practice, so there is no evidence that screening for cCMV gives benefits when compared to what is currently done. Testing all newborns for cCMV ("universal screening") will identify almost all cCMV cases, but will identify lots of babies with no symptoms, most of whom will experience no long-term harms, but will undergo further examinations and possibly unnecessary treatment. Screening only babies who fail a newborn hearing test could identify most newborns who will have long-

term symptoms, but will miss some newborns with other types of symptoms, or where symptoms develop later.

Overall, although there is a lot of evidence on newborn screening for cCMV, it is not sufficient to conclusively recommend that it be used in practice. More studies comparing cCMV screening to standard medical practice without screening are needed to confirm exactly how children might benefit from screening.

Executive summary

Purpose of the review

This review aimed to assess whether there have been significant developments in the evidence base on screening for congenital cytomegalovirus (cCMV) since the previous full review of newborn screening for cCMV was conducted in 2017, and since the most recent evidence map in 2022. The review aimed to determine if sufficient evidence exists to support the establishment of a screening programme for cCMV in newborn infants.

Background

Cytomegalovirus (CMV) is a common virus which usually causes mild flu-like symptoms. However, if the virus is passed on to a fetus during pregnancy, the baby may be born with CMV infection, called cCMV. cCMV is the lead infectious cause of neurological disabilities in children in Europe. In the UK, roughly 650,000 to 695,000 babies are born each year. Around 2,400 babies are born with cCMV every year in the UK; around 10% have symptoms at birth and around half of these babies will experience long-term sequelae, primarily hearing and neurodevelopmental impairment. In addition, around 10% of asymptomatic newborns develop hearing impairment throughout early childhood. Overall, each year around 400 babies born in the UK with cCMV will experience long-term problems. In the newborn infant, cCMV may be diagnosed by urine, saliva or blood polymerase chain reaction (PCR) assay in the first 21 days of life, to distinguish cCMV with CMV acquired after birth. Antiviral treatment is reserved for newborns with severe disease, central nervous system (CNS) related symptoms or sensorineural hearing loss (SNHL).

Focus of the review

The aim of this project was to assess the value of initiating a screening programme for cCMV in newborns to reduce the long-term health burden caused by cCMV infection. A systematic review was undertaken to assess 4 key review questions:

1. What is the diagnostic accuracy of cCMV testing of saliva in newborns?
2. What is the diagnostic accuracy of cCMV testing of dried blood spots (DBS) in newborns?
3. What tests in newborns with cCMV identify which are likely to suffer long-term health consequences from cCMV, who would benefit from prompt treatment?

4. Can a screening programme for cCMV reduce the burden of cCMV-related morbidity and mortality?

Recommendation under review

The UK National Screening Committee (UK NSC) does not currently recommend population or targeted screening for cCMV in newborns.

Findings of this review and gaps in the evidence

A single overarching systematic search and review was conducted to answer all the key questions in this report, adapted to relate to the criteria set out by the UK NSC for assessing the suitability of a screening programme. These key questions will help answer the review questions above. A total of 171 published studies were included in the review. A summary of results for each criterion and key question is presented below.

Criterion 4: There should be a simple, safe, precise and validated screening test.

Key question 1 – What is the diagnostic accuracy of cCMV testing of saliva in newborns?

Key question 2 – What is the diagnostic accuracy of cCMV testing of dried blood spots (DBS) in newborns?

The review included 39 studies relevant to these questions; 21 studies assessed saliva tests, 9 assessed DBS tests, 2 assessed both saliva and DBS tests, and 10 assessed other tests (3 of these also assessed saliva tests), including urine and cord blood testing. Individual studies and reviews were generally of good quality. Results from 1 systematic review (9 studies included in the meta-analysis) and our meta-analysis of 12 studies agreed that saliva screening can accurately detect cCMV with a sensitivity of about 93% for a specificity of over 99%. DBS testing was assessed in 1 systematic review and 10 further studies; results of the review and our meta-analysis of 4 studies demonstrated a similarly high accuracy to detect cCMV, with a sensitivity of between 84% and 90% for a specificity of over 99%. The positive predictive value of both tests was less certain, but may be low because cCMV infection is rare in most populations. Therefore, confirmatory testing for cCMV after a positive saliva or DBS test is probably needed. The 7 studies of urine testing confirmed that it is accurate enough to be considered a reference standard test for cCMV.

Overall, there is a substantial body of high-quality evidence for the accuracy of tests for cCMV. We conclude that saliva, DBS or urine testing could all be used for newborn screening for cCMV, and that further research on the accuracy of these tests is not required.

Criterion 7: There should be an agreed policy on the further diagnostic investigation of individuals with a positive test result and on the choices available to those individuals.

Key question 3 – What tests in newborns with cCMV identify which are likely to suffer long-term health consequences from cCMV, who would benefit from prompt treatment?

The review included 97 studies relevant to this question; 7 systematic reviews, 19 prognostic accuracy studies and 71 studies assessing the association between test results and/or antenatal and/or newborn characteristics and long-term health consequences. A range of tests and characteristics were assessed, however, much of the evidence is of limited quality.

Overall, the results indicate that the most reliable characteristics for predicting long-term sequelae in infants with cCMV are the presence of symptoms at birth and timing of maternal CMV infection. This concurs with current guidelines where newborns with more severe cCMV symptoms are likely to be offered antiviral treatment, but those with milder symptoms are not. Hence how to manage newborns with asymptomatic cCMV remains unclear.

Our results also indicate that the most useful test for predicting long-term hearing and neurodevelopmental sequelae is likely to be magnetic resonance imaging (MRI). Results of studies assessing viral load to predict long-term hearing sequelae are promising, particularly in newborns with asymptomatic cCMV, however, further research is required to confirm these findings. As the risk of long-term sequelae may depend on many factors, development of a multifactorial risk model for long-term harm in newborns with cCMV would be useful.

Criteria 11 & 13: There should be evidence from high quality randomised controlled trials that the screening programme is effective in reducing mortality or morbidity. The benefit gained by individuals from the screening programme should outweigh any harms for example from overdiagnosis, overtreatment, false positives, false reassurance, uncertain findings and complications.

Key question 4 – Can a screening programme for cCMV reduce the burden of cCMV-related morbidity and mortality?

The review included 37 studies relevant to this question. No systematic reviews were identified and no studies compared newborn cCMV screening to current practice, or any management without screening. Therefore, it is not possible to conclude whether newborn screening leads to genuine benefits or harms for newborns, and the evidence available to address this question is of low quality.

Fifteen studies examined universal screening for all newborns. These found that universal screening would identify nearly all cases of cCMV, but most of these would be asymptomatic at birth and not likely to develop long-term sequelae. One study of a low-cCMV prevalence population in Finland found that the

rate of long-term sequelae in newborns with cCMV was not different to the rate in newborns without cCMV.

Twenty-two studies examined targeted screening, primarily screening newborns who fail a newborn hearing test. These studies found that targeted screening identifies newborns who will have long-term hearing loss, but will not detect cCMV in newborns with other types of symptoms, or where symptoms develop later.

Overall, there is insufficient evidence to demonstrate that newborns will benefit from cCMV screening when compared to current medical practice. Randomised or non-randomised studies comparing newborn screening to practice without screening are required in this area for both universal and targeted screening.

Criterion 12: There should be evidence that the complete screening programme (test, diagnostic procedures, treatment/intervention) is clinically, socially and ethically acceptable to health professionals and the public.

The review included 5 studies relevant to this question. All were small surveys on the acceptability and feasibility of saliva testing, rather than newborn cCMV screening as a whole. The surveys mostly conclude that saliva testing was feasible and acceptable. Some concerns over screening increasing parental anxiety were raised. The evidence base is too small and of too limited quality to draw any meaningful conclusions.

Recommendations on screening

There has been a substantial amount of new research on newborn screening for cCMV since the previous review conducted in 2017. In particular, there has been substantial new research on the accuracy of saliva, DBS and urine testing for cCMV. This new evidence appears to be of sufficient quantity and quality to show that all 3 tests could reasonably be used to test for cCMV as part of a screening programme.

While there have been many studies on tests and characteristics that may predict long-term harms in newborns with cCMV, the quality of the evidence is limited and is insufficient to draw definitive conclusions on their value within a screening programme. The only characteristic that is clearly associated with long-term harm is displaying cCMV-related symptoms at birth, such as hearing loss. This raises questions about how newborns who are asymptomatic might be managed, given their low rate of long-term harm.

A particular limitation of the evidence base is the lack of any randomised or non-randomised evidence comparing the outcomes of newborn cCMV screening to standard medical practice, or ad-hoc cCMV testing, without screening. All evidence on existing screening programmes is non-comparative and so of

low quality. Randomised trials may be difficult to conduct in this area, but trials or real-world evidence comparing screening to practice without screening are needed. Without such evidence, even if not randomised, it is not possible to determine to what extent newborns will benefit from cCMV screening, so universal or targeted cCMV screening cannot be recommended for implementation at this time.

Limitations

Much of the evidence included in this systematic review is limited by the study design, size and quality. In addition, many of the included studies may not be applicable to the UK, owing to differences in health systems and the incidence of primary and non-primary maternal CMV infection in different countries. The evidence is too limited to draw definitive conclusions on the value of initiating a screening programme for cCMV in newborns to reduce the long-term health burden caused by cCMV infection.

Evidence uncertainties

The main uncertainty in the evidence is the lack of evidence comparing cCMV screening to standard medical practice. Randomised trials, non-randomised studies, or other real-world studies comparing newborn cCMV screening to practice without screening are needed to properly demonstrate whether screening for cCMV will provide long-term benefits to children, as the current evidence is inconclusive.

In terms of further diagnostic investigation of newborns with cCMV to identify which infants are likely to benefit from prompt treatment, the results of studies assessing viral load to predict long-term sequelae appear promising, particularly in newborns with asymptomatic cCMV. Therefore, further research on the prognostic accuracy of viral load is recommended.

Introduction and approach

Background

The condition

Cytomegalovirus (CMV) is a common virus which can cause mild flu-like symptoms or can be asymptomatic in most immunocompetent adults. Around half of the UK population is infected by CMV at some point in their life.¹ However, if the virus is passed on to a fetus during pregnancy (through the placenta), the baby may be born with CMV infection, called congenital CMV (cCMV). In Europe, cCMV is the lead infectious cause of neurological disabilities in children.^{2, 3} It is estimated that in the UK around 3-5 in 1000 babies are born with cCMV.¹ This equates to around 2,400 babies every year from approximately 600,000 live births.⁴ A very recent study examining the use of dried blood spots (DBS) to determine the incidence of cCMV in the UK, in London, found a rate of 0.66%, equating to around 3,960 infants with cCMV each year.⁵ While the majority have no signs of illness at birth, around 10% of infected newborns will have clinically apparent signs of cCMV at birth, with symptoms such as jaundice, petechiae, hepatosplenomegaly, thrombocytopenia, chorioretinitis, intrauterine growth restriction, microcephaly, deafness, seizures, and brain imaging abnormalities. Around half of the symptomatic babies will experience long-term sequelae, primarily hearing and neurodevelopmental impairment. For those infected newborns who are asymptomatic at birth, outcomes are less clear, although around 10% are likely to develop hearing impairment throughout early childhood,⁶ and a proportion may go on to have learning, concentration and communication difficulties. Overall, each year around 400 babies born in the UK with cCMV will experience long-term problems.⁷ Other publications suggests a rate of long-term sequelae of around 17-20%,⁸ or potentially around 900 babies per year experiencing long-term problems.⁹

Primary maternal CMV infection occurs in women who acquire CMV for the first time in the periconceptual period or during pregnancy. Non-primary infection occurs in women who are already seropositive for IgG before pregnancy and either have reactivation of latent virus during pregnancy or are infected by a different strain of the virus. CMV is transmitted through bodily fluids, most commonly through the saliva or urine of young children; pregnant women in contact with young children are at heightened risk of having pregnancies complicated by cCMV. There is some evidence that implementation of hygiene-based interventions during pregnancy has the potential to prevent CMV transmission.^{10, 11} The risk of fetal transmission is 30-35% with primary maternal infection, compared to around 3% or less with re-infection/reactivation. Primary maternal infection and infection occurring earlier in gestation are associated with higher rates of fetal transmission and earlier infection is also associated with poorer fetal outcomes.¹²

Screening and treatment

In the newborn infant, cCMV may be diagnosed by urine, saliva or blood polymerase chain reaction (PCR) assay in the first 21 days of life; after 21 days it is not possible to distinguish cCMV from postnatal CMV infection. Previous research has shown that saliva testing may result in more false positive test results (from maternal CMV shed in breastmilk).¹² Research has also shown that newborn dried blood spot (DBS) testing may have lower sensitivity than saliva or urine;⁷ however, it can be used to diagnose cCMV retrospectively after 21 days of age.

The European Expert Consensus Statement on Diagnosis and Management of Congenital Cytomegalovirus recommends that antiviral treatment should be reserved for babies with evidence of central nervous system (CNS) disease or other 'severe' disease.¹³ Updated consensus recommendations from the European Congenital Cytomegalovirus Initiative (ECCI), published in May 2024, recommend treating newborns with isolated sensorineural hearing loss (SNHL), in addition to those with CNS related symptoms.⁸ The reason for treatment in this scenario is to try to prevent late-onset SNHL in the other ear. Oral valganciclovir is the drug of choice and intravenous ganciclovir should be used in babies unable to tolerate oral drugs or where gastrointestinal absorption is uncertain. No treatment is recommended for babies with 'mild' disease. Whilst the definition of mild disease is not clear, the 2017 European Expert Consensus Statement states that mild disease includes those with isolated (1 or 2 at most) and otherwise clinically insignificant or transient findings. These can include petechiae, mild hepatomegaly or splenomegaly or biochemical/haematologic abnormalities (such as thrombocytopenia, anaemia, leukopenia, borderline raised liver enzyme abnormalities or conjugated hyperbilirubinemia), and babies who are small for gestational age (weight for gestational age <2 standard deviations) without microcephaly.¹³ However, evidence on treatment effectiveness is limited, especially in babies who do not show signs of infection. Much of the debate around treatment relates to the potential adverse effects of antiviral drugs.¹³ The updated consensus recommendations list the following indications for testing for cCMV infection in newborns: evidence of maternal primary CMV infection during pregnancy; presence of indicative features on prenatal ultrasonography or MRI; newborn clinical manifestations consistent with cCMV.⁸

Current global landscape of newborn screening

In the USA, universal cCMV screening was introduced in 2023 in Minnesota and it is projected to be introduced in Connecticut in 2025. In addition, a National Institutes of Health study of universal screening in New York is underway and it is anticipated that screening will continue after this study has ended.¹⁴ In Canada, 4 provinces are implementing universal newborn screening for cCMV, and 3 provinces have targeted screening programmes that offer cCMV testing to newborns who fail newborn hearing tests.¹⁵ cCMV screening is also in use in Japan as part of wider newborn screening programmes.¹⁶

Current policy context and previous reviews

The UK National Screening Committee (UK NSC) does not currently recommend population screening for cCMV in newborns.¹⁷ This recommendation, based on an evidence review published in 2017,¹⁸ concluded that there was insufficient evidence to recommend the introduction of newborn screening for cCMV because:

- It is not clear how to identify newborns that will develop long-term sequelae.
- The treatment offered to babies with asymptomatic cCMV infection is unclear.
- Screening is likely to identify a greater number of infants with cCMV, the majority of which are likely to have minimal symptoms or no symptoms. The management and treatment approach for these children is unclear, and it is unknown whether screening improves their outcomes.

A UK NSC evidence map on newborn screening for CMV, published in June 2022, concluded that no further work on screening for CMV should be commissioned at that time, but that the UK NSC would review the decision on screening for CMV in 3-years' time.⁷

Objectives

This review aimed to assess whether there have been significant developments in the evidence base since the previous review was conducted in 2017, and if sufficient support exists for a screening programme for cCMV in newborns to reduce the long-term health burden caused by cCMV infection.

The overall objective was to undertake a systematic review to assess the clinical evidence on:

- newborn screening tests for identifying cCMV infection
- tests to determine which newborns are likely to develop long-term complications from cCMV
- whether a screening programme for cCMV can reduce the burden of cCMV-related complications

The review appraised evidence on the key questions in Table 1, adapted to relate to the criteria set out by the UK NSC for assessing the suitability of a screening programme. These key questions will help answer the review objectives above.

Table 1: Key questions for the evidence summary and relationship to the UK NSC screening criteria

Criterion	Key questions	Studies Included
The Test		
4	There should be a simple, safe, precise and validated screening test.	KQ1: What is the diagnostic accuracy of cCMV testing of saliva in newborns? KQ2: What is the diagnostic accuracy of cCMV testing of dried blood spots (DBS) in newborns? 10 diagnostic accuracy studies assessed other tests, including urine
7	There should be an agreed policy on the further diagnostic investigation of individuals with a positive test result and on the choices available to those individuals.	KQ3: What tests in newborns with cCMV identify which are likely to suffer long-term health consequences from cCMV, who would benefit from prompt treatment? 7 systematic reviews 19 prognostic accuracy studies 71 association studies
The screening programme		
11	There should be evidence from high quality randomised controlled trials that the screening programme is effective in reducing mortality or morbidity.	KQ4: Can a screening programme for cCMV reduce the burden of cCMV-related morbidity and mortality? 15 evaluations of universal screening programmes 22 evaluations of targeted screening programmes
12	There should be evidence that the complete screening programme (test, diagnostic procedures, treatment/ intervention) is clinically, socially and ethically acceptable to health professionals and the public.	KQ4: Can a screening programme for cCMV reduce the burden of cCMV-related morbidity and mortality? 5 studies with surveys reporting acceptability and feasibility of a screening programme
13	The benefit gained by individuals from the screening programme should outweigh any harms for example from overdiagnosis, overtreatment, false positives, false reassurance, uncertain findings and complications.	KQ4: Can a screening programme for cCMV reduce the burden of cCMV-related morbidity and mortality? 15 evaluations of universal screening programmes 22 evaluations of targeted screening programmes

Methods

The current review was conducted by York Evidence Synthesis Group, in keeping with the UK NSC evidence review process. Database searches were conducted in November 2024 to identify studies relevant to the questions detailed in Table 1.

The systematic review was conducted following the general principles recommended in the Centre for Reviews and Dissemination's guidance and reported in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement.^{19, 20} The systematic review was registered with PROSPERO, registration number CRD42024616064.

The York team were supported throughout the project, from inception to the writing of this report, by a broad advisory group. This consisted of representatives from UK NSC, experts in cCMV infection and newborn screening, representatives of the charity CMV Action, and families affected by cCMV.

Databases/sources searched

The aim of the search was to systematically identify published, unpublished and ongoing studies of screening tests or programmes to detect cCMV in newborns, and additional screening tests to triage newborns with cCMV for antiviral treatment. Comprehensive searches of databases and trial registers were undertaken, supplemented by reference checking of relevant systematic reviews, manual searches of abstracts from relevant CMV conferences, and requests for relevant research from advisory group members.

A search strategy was designed in Ovid MEDLINE by an information specialist in consultation with the review team. The strategy combined terms for newborns with terms for CMV. Searches of the title, abstract and author keyword fields of database records along with relevant subject headings were included in the strategy. The search was limited to records published from 2013 onwards in order to update previous reviews undertaken for the UK NSC. No further limits were applied. The MEDLINE strategy was peer reviewed using aspects of the Peer Review of Electronic Search Strategies (PRESS) checklist,²¹ and then translated to run on further databases and resources.

The following databases were searched in November 2024: MEDLINE ALL (Ovid), Embase (Ovid), Maternity and Infant Care Database (Ovid), Cochrane Central Register of Controlled Trials (CENTRAL, Wiley), the Science Citation Index (Web of Science), CINAHL (Ebsco), LILACS, Cochrane Database of Systematic Reviews (CDSR, Wiley), Database of Abstract of Reviews of Effects (DARE) and KSR Evidence (Ovid).

Unpublished, ongoing or grey literature was identified through searching the HTA database, International HTA database (INAHTA), websites of international HTA organisations, Conference Proceedings Citation index – Science (Web of Science), ClinicalTrials.gov, WHO International Clinical Trials Registry Platform, EU Clinical Trials Register and PROSPERO. All search results were imported into EndNote 21 reference management software and deduplicated.

Supplementary searching included manual searches of the abstracts from the European Congenital Cytomegalovirus Initiative (ECCI) congresses 2022 and 2024, consulting with advisory group members for any additional studies that they were aware of and checking the reference lists of all recent relevant systematic reviews.

The full search strategies can be found in Appendix 1.

Eligibility for inclusion in the review

Deduplicated references identified by the electronic searches were uploaded into EPPI-Reviewer. The machine learning and text mining tool in EPPI-Reviewer (priority screening) was used to prioritise titles and abstracts for screening.²² The first 40% of prioritised records were independently screened by 2 reviewers to ensure eligibility criteria were applied consistently; disagreements between reviewers were resolved by discussion or, if necessary, a third independent reviewer. The remaining titles and abstracts were assessed by 1 reviewer. It was initially planned to double screen at least 10% of records, however, owing to the complexity of the review question the level of agreement between reviewers was unacceptably low, therefore, double screening was continued until an acceptable level of agreement was reached. The full texts of potentially eligible studies were assessed independently by 2 reviewers, using the same process for resolution of disagreements as outlined above. Non-English language publications were eligible for inclusion and translated for data extraction. Eligible ongoing studies (e.g., reported in protocols and trial registers) without relevant outcome data reported at the time of data extraction were included and tabulated as 'Ongoing Studies'. Eligible studies published as conference abstracts were included and tabulated as 'Unpublished Studies'.

Inclusion criteria

Population

The population of interest for screening for cCMV infection was newborns, defined as up to 28 days old.

Index tests and interventions

This assessment included any test used to detect cCMV in newborns, such as PCR assay of saliva, urine or blood. This assessment also included tests conducted in newborns for identifying which newborns who test positive for cCMV are likely to develop long-term complications, and so would benefit from prompt treatment. The relevant tests include CMV viral load tests, genetic tests and specific clinical or medical findings (e.g., newborn hearing tests and MRI). In addition, retrospective assessment of tests undertaken during pregnancy for maternal prognostic factors (i.e., assessment of booking bloods to determine timing of CMV infection) were included.

Both standalone and multiplex tests were eligible for inclusion. Tests that were only conducted after 28 days of life, or after initiation of antiviral treatment, were excluded.

Comparators

Studies did not have to include any comparator group or test to be included. However, studies comparing a screening test or programme to no screening test for detecting cCMV were included, where available, as were studies comparing different screening tests.

Reference standard

The reference standard for detecting cCMV infection was PCR urine testing. For studies reporting accuracy of tests to detect newborns likely to develop long-term complications from cCMV, the reference standard was confirmed clinical diagnosis of any morbidity (e.g., hearing loss, neurological impairment) in childhood, or absence of diagnosed morbidity in childhood. As newborns may be treated to prevent complications, which would interfere with this reference standard, alternatives were accepted, such as poor performance on newborn hearing tests, or other indicators of long-term morbidity.

Outcomes

When considering the accuracy of screening tests to detect long-term health consequences the key outcomes were standard diagnostic accuracy measures (including sensitivity, specificity, positive and negative predictive values).

When considering the overall clinical value and effectiveness of screening the outcomes of interest were:

- cCMV-related morbidities
 - hearing loss, both at birth and later onset or deterioration
 - neurological impairments in infants, including brain imaging results
 - neurodevelopmental outcomes
 - any other long-term negative outcomes associated with cCMV infection
- mortality
- use of additional tests and procedures
- incidental findings, however defined
- potential harms from screening or treatment
- health-related quality of life (of children and their families)
- acceptability of screening

Study designs

Clinical trials (including randomised and other controlled trials), prospective and retrospective cohort studies and case-control studies were sought. In addition, good quality, up to date, systematic reviews were included.

Table 2: Inclusion and exclusion criteria for the key questions

Key question	Inclusion criteria						
	Population	Target Condition	Intervention	Reference standard	Comparator	Outcome	Study type
KQ1: What is the diagnostic accuracy of cCMV testing of saliva in newborns?	Newborns (<28 days)	cCMV	PCR testing of saliva for CMV	Urine PCR testing (or any other reasonable reference standard)	Not required (or comparison with any other cCMV test)	Detection of cCMV	Systematic reviews Prospective or retrospective cohort studies Case-control studies
KQ2: What is the diagnostic accuracy of cCMV testing of dried blood spots (DBS) in newborns?	Newborns (<28 days)	cCMV	DBS testing for CMV	Urine PCR testing (or any other reasonable reference standard)	Not required (or comparison with any other cCMV test)	Detection of cCMV	Systematic reviews Prospective or retrospective cohort studies Case-control studies
KQ3: What tests in newborns with cCMV identify which are likely to suffer long-term health consequences from cCMV, who would benefit from prompt treatment?	Newborns (<28 days)	Any long-term sequelae related to cCMV	Any test of newborns with cCMV to predict long-term sequelae	Clinical confirmation of symptoms	None	Any long-term sequelae (hearing loss, neurodevelopmental issues etc)	Systematic reviews Prospective or retrospective cohort studies Case-control studies
KQ4: Can a screening programme for cCMV reduce the burden of cCMV-related morbidity and mortality?	Newborns (<28 days) being screened for cCMV	cCMV and its long-term sequelae	Any screening test for cCMV (saliva, DBS or urine)	None	None, or outcomes without screening	Presence of cCMV Antiviral treatment Any long-term sequelae (hearing loss, neurodevelopmental issues etc)	Systematic reviews RCTs Other controlled trials Prospective or retrospective cohort studies

Abbreviations: cCMV, congenital cytomegalovirus; CMV, cytomegalovirus; DBS, dried blood spots; KQ, key question; PCR, polymerase chain reaction; RCTs, randomised controlled trials.

Appraisal of studies for quality and risk of bias

Risk of bias and applicability of prognostic and diagnostic accuracy studies were assessed using the QUADAS-2 (Quality Assessment of Diagnostic Accuracy Studies) checklist²³ by 1 reviewer and independently checked by a second reviewer (3 reviewers shared this work). Risk of bias in systematic reviews was assessed by 1 reviewer and independently checked by a second reviewer using the ROBIS tool.²³ Disagreements were resolved through discussion.

For studies that only assessed an association between test results/characteristics and health outcomes, study quality was briefly summarised, but not formally assessed. For non-comparative studies of screening programmes study quality was not formally assessed, but general quality issues across the studies were evaluated.

Data extraction and methods of synthesis

A standardised data extraction form for study characteristics was developed and piloted. Data on the study design, patient characteristics, intervention(s), comparator(s) and relevant outcomes were extracted from included studies by 1 reviewer and independently checked by a second reviewer (4 reviewers shared this work). Disagreements were resolved through discussion.

The following diagnostic accuracy values were extracted or calculated for each test:

- the number of true positives (TP), which is the number of newborns with a positive cCMV test result who do have cCMV, i.e., the number of newborn babies correctly identified by the cCMV test as having cCMV
- the number of false positives (FP), which is the number of newborns with a positive cCMV test result who do not have cCMV, i.e., the number of newborn babies incorrectly identified by the cCMV test as having cCMV
- the number of true negatives (TN), which is the number of newborns with a negative cCMV test result who do not have cCMV, i.e., the number of newborn babies correctly identified by the cCMV test as not having cCMV
- the number of false negatives (FN), which is the number of newborns with a negative cCMV test result who do have cCMV, i.e., the number of newborn babies incorrectly identified by the cCMV test as not having cCMV

Measures of test accuracy are as follows:

- Sensitivity = $TP / (TP + FN)$

This is the true positive rate, which is the probability that a newborn with cCMV receives a positive cCMV test result; in other words, the ability of a test to correctly classify a newborn baby with cCMV.

- Specificity = $TN / (TN + FP)$

This is the true negative rate, which is the probability that a newborn without cCMV receives a negative cCMV test result; in other words, the ability of a test to correctly classify a newborn baby without cCMV.

- Positive predictive value (PPV) = $TP / (TP + FP)$

This is the probability that a newborn who receives a positive cCMV test result has cCMV.

- Negative predictive value (NPV) = $TN / (TN + FN)$

This is the probability that a newborn who receives a negative cCMV test result does not have cCMV.

These test accuracy measures were extracted or, where not directly reported, calculated from other reported data, where appropriate. Calculated measures, as opposed to directly reported measures, are identified as such in all results tables.

For ongoing studies or unpublished studies (conference abstracts or clinical trial/PROSPERO register records) that appeared to meet the review inclusion criteria, basic study details were extracted and are tabulated in Appendix 4.

Study characteristics and results are presented in structured tables (see Appendix 3) and as a narrative summary (see sections below). For diagnostic accuracy studies, where feasible, a joint bivariate meta-analysis of sensitivity and specificity was performed.²⁴ In cases where meta-analysis was performed the DerSimonian-Laird random-effects method was used, including assessment of heterogeneity using the I^2 statistic.^{25, 26}

Question level synthesis

Overall review summary

A single overarching systematic search and review was conducted to answer all the key questions in this report. The overall review results are summarised briefly here, with discussion of findings for each key question in the sections below.

The electronic searches identified a total of 6,258 records after deduplication between databases. The full texts of 422 records were ordered for further review; 224 were excluded at full paper stage and are listed in Appendix 2, along with the reasons for their exclusion. Eighteen additional records were identified from screening abstract books of ECCI congresses (held in 2022 and 2024). One additional record was identified through consulting with advisory group members. No additional records were found from screening reference lists of relevant systematic reviews. A flow diagram of the study selection process is presented in Appendix 2.

A total of 217 records met the review inclusion criteria. There were 171 studies published in full; study characteristics are presented in Appendix 3. There were 8 ongoing studies and 38 studies only reported as conference abstracts; tabulated in Appendix 4.

Criterion 4: There should be a simple, safe, precise and validated screening test.

Previous work

The previous UK NSC evidence review by Bazian Ltd¹⁸ included 1 systematic review and 2 prospective cohort studies that assessed diagnostic tests for the detection of cCMV. The systematic review (Wang et al., 2015)²⁷ was conducted on the screening of cCMV through the PCR assay of DBS and also included a meta-analysis. It included 14 studies and reported a sensitivity of 84% (95% confidence interval [CI]: 81 to 87) and a specificity of 100% (95% CI: 100 to 100) for detecting cCMV. The previous review concluded that while the specificity of the PCR assay of DBS was very high, the sensitivity was low, particularly when considering only the prospective studies (62%). The review was high-quality with generally a low risk of bias.

The 2 cohort studies assessed the usefulness of real-time PCR assays of both dried or liquid saliva²⁸ or PCR assays of liquid saliva.²⁹ Both cohort studies were found to have a high potential for verification bias as confirmatory testing was not conducted in the entire cohort. The previous review concluded that the value of saliva testing as a screening tool needed to be investigated further.

Bazian Ltd (2017)¹⁸ also pointed out that even if DBS and saliva testing were able to detect cCMV reliably, it was not possible to correlate the likelihood of adverse outcomes as none of the studies reported longer-term disease outcomes.

The UK NSC evidence map of 2022⁷ did not address this criterion.

Description of the current evidence

There were 39 studies identified as relevant for this research question and included in the review. Systematic reviews were given priority, followed by diagnostic accuracy studies. Studies were categorised according to the sample-type being tested. Where possible a diagnostic bivariate meta-analysis was conducted for studies that had not been included in previous systematic reviews. If a 2x2 contingency table was not provided, it was calculated using the sensitivity and specificity reported. Meta-analyses were conducted using the web-based software MetaDTA.^{30, 31}

Question 1 – What is the diagnostic accuracy of cCMV testing of saliva in newborns?

This question included any prospective or retrospective cohort studies, or case-control studies, that reported the diagnostic accuracy of saliva testing for cCMV in newborns when compared to urine testing as a reference standard. Existing systematic reviews of relevant studies were also included. Full details of eligibility criteria are presented in Table 2.

Description of the evidence

Database searches yielded 1 systematic review, and 22 diagnostic accuracy studies. Three identified studies were included in the previous UK NSC evidence review by Bazian Ltd.¹⁸

Systematic Reviews

A systematic review assessed the diagnostic accuracy of PCR using saliva samples for newborns with cCMV (Zheng et al., 2022).³² Nine studies,^{28, 29, 33-39} published between 2006 and 2021, were included for quantitative analysis in this systematic review. The 2 studies included in the previous UK NSC evidence review¹⁸ were included in the Zheng et al. (2022) review.³² The index test was saliva PCR. Seven of the included studies used urine, assessed either by PCR or rapid culture, as the reference test. Bopanna et al. (2011)²⁸ used saliva rapid culture as the reference test and Alkhawaja et al. (2012)³³ used a culture of either saliva or urine.

The systematic review conducted by Zheng (2022) was of high quality overall with a low risk of bias. Of the studies included in the review, 4 were judged to be at high risk of bias, and the rest at moderate to low risk of bias.

The summary sensitivity and specificity of PCR saliva testing to detect cCMV infection were 91% (95% CI: 70 to 98) and 100% (95% CI: 100 to 100), respectively. A subgroup analysis was conducted where saliva PCR was compared with only urine culture. In this subgroup analysis the summary sensitivity and specificity were 97% (95% CI: 61 to 100) and 100% (95% CI: 99 to 100), respectively. Meta-analyses of PPV and NPV were not performed.

Diagnostic accuracy studies

Twenty-two studies were identified on the diagnostic accuracy of saliva testing. Details of these studies are presented in Table 3. Of these, 3 studies^{35, 38, 39} had been included in the Zheng et al. (2022) systematic review.³²

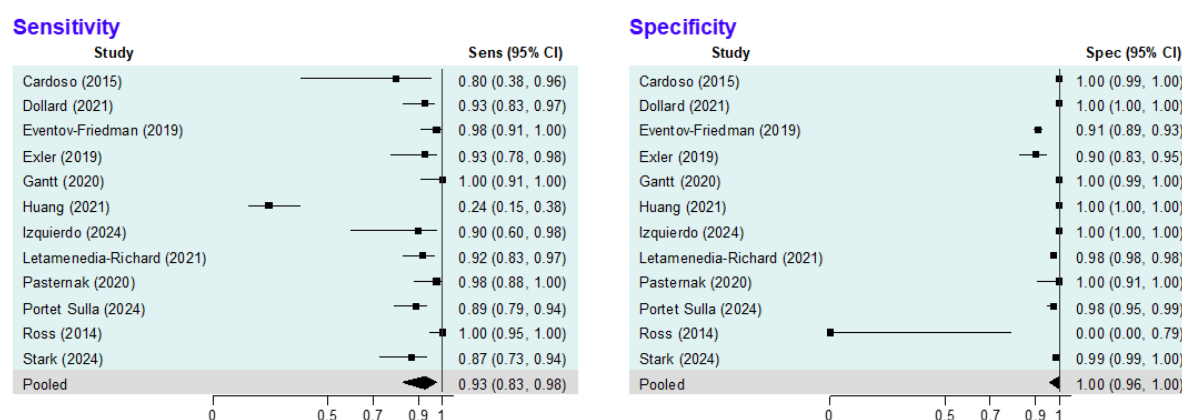
The studies were conducted in a wide variety of locations (7 in USA, 3 in France, 3 in Israel, 2 in China, 1 in Brazil, 1 in Germany, 1 in Turkey, 1 in Chile, 1 in USA and Italy, and 1 in Canada, USA, Italy and Australia), with only 1 from the UK. Most were prospective in design.

Most studies used urine PCR as the reference standard test, but some used repeated saliva testing. Studies varied substantially in size from 125 newborns to over 70,000. Diagnostic accuracy to detect cCMV was high, with reported sensitivities over 90% in most studies, and specificity generally close to 100%. There was considerable heterogeneity in PPV however, with estimates ranging from 12.9% to 100%.

Ten of the identified studies were not included in our meta-analysis. Eight⁴⁰⁻⁴⁷ were not included as they used saliva as the reference standard, either through culture testing or repeated testing. Two^{48, 49} were excluded from the meta-analysis as they did not report sufficient data for a bivariate meta-analysis.

The meta-analysis included 12 studies;^{35, 38, 39, 50-58} the forest plots showing the results are presented as Figure 1. The pooled estimates from bivariate meta-analyses for sensitivity and specificity were consistent with those in Zheng (2022).³² The pooled sensitivity was 93.2% (95% CI: 82.9 to 97.5) and the specificity was 99.5% (95% CI: 96.4 to 99.9). Meta-analyses of PPV and NPV gave results of 87% (95% CI: 63 to 96) and 100% (95% CI: 100 to 100), respectively. There was substantial heterogeneity in PPV values across studies, with some larger studies having positive predictive values below 50%.

Figure 1 Bivariate meta-analysis of diagnostic accuracy of saliva testing



Abbreviations: CI, confidence interval; Sens, sensitivity; Spec, specificity.

The quality of the studies was assessed using QUADAS-2 (Table 4). All studies generally had a low risk of bias across all domains. There was a high risk of bias in patient selection in Exler (2019)⁵⁰ and Stark (2024)⁵⁸ as both studies selected newborns using targeted screening based either on positive maternal cCMV tests during pregnancy (Exler 2019) or a pre-specified criteria for clinical symptoms (Stark 2024).

Table 3: Results of the saliva studies

Study details	Study Type	Reference standard	Patient numbers		Diagnostic accuracy, % (95% CI)			
			Total Screened	With cCMV	Sensitivity	Specificity	PPV	NPV
Studies included in meta-analysis								
Cardoso (2015) ³⁸ <i>Location:</i> Brazil	Prospective	Urine	333	5	80	100	100	99.7
Dollard (2021) ³⁹ <i>Location:</i> USA	Prospective	Urine	12554	56	92.9 (83.0 to 97.2)	99.9 (99.9 to 100)	86.7 (75.8 to 93.1)	100 (99.9 to 100)
Eventov-Friedman (2019) ³⁵ <i>Location:</i> Israel	Prospective	Urine and Saliva	856	58	98.3 (90.8 to 99.9)	91.5 (89.3 to 93.3)	45.6 (36.7 to 54.7)	99.9 (99.2 to 99.9)
Exler (2019) ⁵⁰ <i>Location:</i> Germany	Retrospective	Urine	133	29	93 (77 to 99)	90 (83 to 95)	73 (56 to 86)	98 (93 to 100)
Gantt (2020) ⁵¹ <i>Location:</i> Canada/USA/Italy/Australia	Prospective and Retrospective	Saliva (composite)	1480	39	100.0 [†] (100.0 to 100.0)	99.8 [†] (99.4 to 100.0)	Not reported	Not reported
Huang (2021) ⁵² <i>Location:</i> China	Prospective	Urine	6234	49	24.5 (14.6 to 38.1)	100 (99.9 to 100)	100 (75.7 to 100)	99.4 (99.2 to 99.6)
Izquierdo (2024) ⁵³ <i>Location:</i> Chile	Prospective	Urine	2186	10	90 [†] (60 to 98)	100 [†] (100 to 100)	99.15	100
Letamendia-Richard (2022) ⁵⁴ <i>Location:</i> France	Retrospective	Urine	15341	63	92 (84 to 97)	98 (95 to 100)	90.5 (84.9 to 100)	Not reported
Pasternak (2020) ⁵⁵ <i>Location:</i> Israel	Prospective	Urine	83	42	97.6 (87.4 to 100)	100 (91.4 to 100)	Not reported	Not reported

Study details	Study Type	Reference standard	Patient numbers		Diagnostic accuracy, % (95% CI)			
			Total Screened	With cCMV	Sensitivity	Specificity	PPV	NPV
Portet Sulla (2024) ⁵⁶ <u>Location:</u> France	Retrospective	Urine	275	70	88	98	21.8	Not reported
Ross (2014) ⁵⁷ <u>Location:</u> USA	Prospective	Saliva	80	Not reported	Not reported	Not reported	Not reported	Not reported
Stark (2024) ⁵⁸ <u>Location:</u> USA	Retrospective	Urine	908	38	86.8 (71.9 to 95.6)	99.4 (98.8 to 99.9)	89.2 (75.5 to 95.7)	99.4 (98.7 to 99.8)
Studies not included in meta-analysis								
Atwood (2023) ⁴⁰ <u>Location:</u> USA	Prospective (part of screening)	Repeated testing	696	5	Not reported	95.5 (93.7 to 96.9)	13.9 (4.7 to 29.5)	Not reported
Dunn (2023) ⁴¹ <u>Location:</u> USA/Italy	Prospective and Retrospective	Saliva (composite)	1853	17	94.1 [†] (71.3 to 99.9)	100.0 [†] (99.7 to 100.0)	Not reported	Not reported
Gunlemez (2023) ⁴⁸ <u>Location:</u> Turkey	Prospective	Urine or blood	545	Not reported	Not reported	99.1 (97.9 to 99.7) [†]	Not reported	Not reported
Leruez-Ville (2017) ⁴² <u>Location:</u> France	Prospective	Saliva (repeated)	11740	51	Not reported	Not reported	58.6 (47.5 to 69.1)	Not reported
Pinninti (2015) ⁴³ <u>Location:</u> USA	Prospective	Saliva	72239	266	Not reported	Not reported	Not reported	Not reported
Ross (2018) ⁴⁴ <u>Location:</u> USA	Prospective	Saliva/DBS	307	284	Not reported	Not reported	Not reported	Not reported

Study details	Study Type	Reference standard	Patient numbers		Diagnostic accuracy, % (95% CI)			
			Total Screened	With cCMV	Sensitivity	Specificity	PPV	NPV
Shlonsky (2021) ⁴⁹ <u>Location</u> : Israel	Prospective	Urine	1000	6	Not reported	98.1	24	100
Tan (2023) ⁴⁵ <u>Location</u> : UK	Prospective	Saliva	100	70	93.3 (85.3 to 97.1)	100 (86.7 to 100)	100 (94.9 to 100)	83.3 (68.2 to 92.1)
Wang (2017) ⁴⁶ <u>Location</u> : China	Prospective	Saliva	7720	46	93.9 (82.1 to 98.4)	99.9 (99.8 to 100)	88.5 (75.9 to 95.2)	100 (99.9 to 100)
Wunderlich (2023) ⁴⁷ <u>Location</u> : USA	Prospective	Repeated testing	94	Not reported	Not reported	Not reported	Not reported	Not reported
Abbreviations: CI, confidence interval; NPV, negative predictive value; PPV, positive predictive value. † Calculated using the 2 x 2 data provided in relevant publications.								

Table 4: Quality assessment results for the saliva studies

Paper	Domain 1: Patient selection		Domain 2: Index tests		Domain 3: Reference Standard		Domain 4: Flow and timing
		Applicability		Applicability		Applicability	
	Risk of bias from selection	Concerns about matching of patients to review question?	Risk of bias relating to index test	Concerns about matching of index tests to review question?	Risk of bias relating to reference test	Concerns about matching of reference test to review question?	Risk of bias from patient flow?
First author							
Cardoso (2015) ³⁸	Low	Low	Low	Low	Low	Low	Unclear
Dollard (2021) ³⁹	Low	Low	Low	Low	Low	Low	Unclear
Eventov-Friedman (2019) ³⁵	Low	Low	Low	Low	Low	Low	Low
Exler (2019) ⁵⁰	High	High	Low	Low	Low	Low	Low
Gantt (2020) ⁵¹	Low	Low	Low	Low	Low	Low	Low
Huang (2021) ⁵²	Low	Low	Low	Low	Low	Low	Low
Izquierdo (2024) ⁵³	Low	Low	Low	Low	Low	Low	Low
Letamendia-Richard (2022) ⁵⁴	Low	Low	Low	Low	Low	Low	Low
Pasternak (2020) ⁵⁵	Unclear	Low	Low	Low	Low	Low	Low
Portet Sulla (2024) ⁵⁶	High	High	Low	Low	Low	Low	Unclear
Ross (2014) ⁵⁷	High	High	Unclear	Unclear	Unclear	Unclear	High
Stark (2024) ⁵⁸	High	Low	Low	Low	Low	Low	Low
Atwood (2023) ⁴⁰	Unclear	Unclear	Low	Low	Low	Low	Unclear
Dunn (2023) ⁴¹	High	Unclear	Low	Low	Low	Low	Low
Gunlemez (2023) ⁴⁸	Low	Low	Unclear	Unclear	Unclear	Unclear	Unclear

Paper	Domain 1: Patient selection		Domain 2: Index tests		Domain 3: Reference Standard		Domain 4: Flow and timing
		Applicability		Applicability		Applicability	
First author	Risk of bias from selection	Concerns about matching of patients to review question?	Risk of bias relating to index test	Concerns about matching of index tests to review question?	Risk of bias relating to reference test	Concerns about matching of reference test to review question?	Risk of bias from patient flow?
Leruez-Ville (2017) ⁴²	Low	Low	Low	Low	Low	Low	Unclear
Pinninti (2015) ⁴³	Unclear	Low	Low	Low	Low	Low	Unclear
Ross (2018) ⁵⁹	Low	Low	Unclear	Unclear	Unclear	Unclear	Unclear
Shlonsky (2021) ⁴⁹	Low	Low	Low	Low	Low	Low	Low
Tan (2023) ⁴⁵	Unclear	High	Low	Low	Unclear	Low	High
Wang (2017) ⁴⁶	Low	Low	Low	Low	Low	Low	Unclear
Wunderlich (2023) ⁴⁷	Low	Low	Low	Low	Low	Low	Unclear

Question 2 – What is the diagnostic accuracy of cCMV testing of dried blood spots (DBS) in newborns?

This question included any prospective or retrospective cohort studies, or case-control studies, that reported the diagnostic accuracy of DBS testing for cCMV in newborns when compared to urine testing as a reference standard. Existing systematic reviews of relevant studies were also included.

Description of the evidence

The review included 1 systematic review, and 10 diagnostic accuracy studies. The systematic review was included in the previous UK NSC review in 2017.¹⁸

Systematic Reviews

No systematic reviews were identified after Wang (2015),²⁷ which was included in the previous UK NSC review.

As noted earlier, the Wang review was high-quality with generally a low risk of bias. It included 14 studies, all of which were published in or before 2012, so were not identified by our new review. Nine of the included studies were assessed as having a high risk of patient selection bias. The meta-analysis reported a sensitivity of 84% (95% CI: 81 to 87) and a specificity of 100% (95% CI: 100 to 100) for detecting cCMV. Sensitivity may have been lower in the prospective studies (62%) than in the retrospective studies (95%). The pooled positive predictive value (PPV) and negative predictive value (NPV) were 91% (95% CI: 84 to 95) and 99% (95% CI: 97 to 100), respectively.

Diagnostic accuracy studies

Ten studies^{39, 46, 60-67} were identified that were published after the Wang (2015) review.²⁷ A summary of these studies is provided in Table 5. Six of the 10 included studies were prospective in design, and 4 were retrospective or case-control studies. Studies were conducted in a range of international locations (3 in USA, 1 in France, 1 in Japan, 1 in Iran, 1 in Belgium, 1 in Spain and 1 in China), with only 1 from the UK. Studies varied substantially in size from 60 newborns to over 12,000. Diagnostic accuracy to detect cCMV was variable but generally high, with most studies reporting a sensitivity over 80% and specificity of nearly 100%.

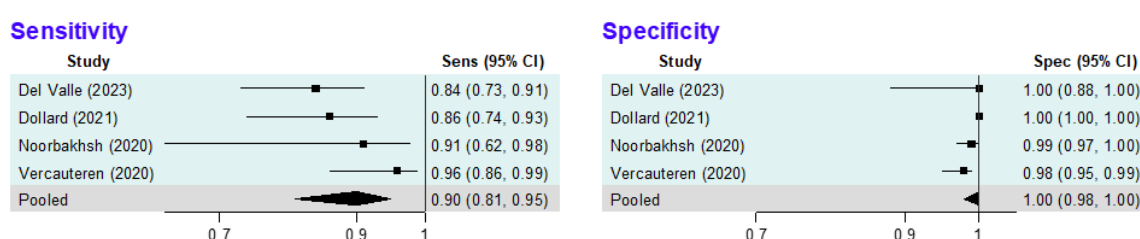
A meta-analysis was conducted to compare the diagnostic accuracy of testing DBS, using urine testing as a reference standard, as the latter is considered the gold standard. This meta-analysis included 4 of the included studies.^{39, 61, 64, 66} Five studies^{46, 60, 62, 63, 65, 67} were

not included in the meta-analysis as these studies compared the diagnostic accuracy of testing DBS to non-urine samples.

The results of the meta-analysis are shown in forest plots for sensitivity and specificity (Figure 2). The pooled sensitivity and specificity from the bivariate meta-analysis were 89.9% (95% CI: 81.3 to 94.8) and 99.9% (95% CI: 98.1 to 100).

These values were consistent with the estimates reported in the Wang (2015) systematic review. Meta-analyses for PPV and NPV gave results of 96% (95% CI: 86 to 99) and 99% (95% CI: 91 to 100), respectively. This is also broadly consistent with the Wang 2015 review.

Figure 2 Bivariate meta-analysis of diagnostic accuracy of DBS



Abbreviations: CI, confidence interval; DBS, dried blood spot; Sens, sensitivity; Spec, specificity.

The quality of the studies published after the Wang et al. (2015) systematic review was assessed using QUADAS-2 (Table 6). Generally, the included studies were at a low risk of bias across all domains.

Table 5: Results of the dried blood spot studies

Study details	Study Type	Reference standard	Patient numbers		Diagnostic accuracy, % (95% CI)			
			Total Screened	With cCMV	Sensitivity	Specificity	PPV	NPV
Studies included in meta-analysis								
Del Valle (2023) ⁶¹ <u>Location:</u> USA	Retrospective	Urine or Saliva	89	62	83.9	100	100	73
Dollard (2021) ³⁹ <u>Location:</u> USA	Prospective	Urine	12554	56	85.7 (74.3 to 92.6)	100 (100 to 100)	98 (89.3 to 99.6)	99.9 (99.9 to 100)
Noorbakhsh (2020) ⁶⁴ <u>Location:</u> Iran	Prospective	Urine	224	11	90.9 (58.7 to 99.7)	99 (96.6 to 99.8)	83.3 (51.5 to 97.9)	99.5 (97.4 to 99.9)
Vercauteren (2020) ⁶⁶ <u>Location:</u> Belgium	Prospective	Urine	276	48	95.8 (85.8 to 99.5)	97.8 (95.0 to 99.3)	90.2 (78.6 to 96.7)	99.1 (97.0 to 99.9)
Studies not included in meta-analysis								
Atkinson (2014) ⁶⁰ <u>Location:</u> UK	Retrospective / Case-Control	Not urine	Not reported	26	81 (60.6 to 93.4)	100	Not reported	Not reported
Fourgeaud (2023) ⁶² <u>Location:</u> France	Prospective	Saliva	256	Not reported	95	33	21	97
Moteki (2018) ⁶³ <u>Location:</u> Japan	Case-control	Cord blood	48	12	83.3	97.2	Not reported	Not reported
Ross (2017) ⁶⁵ <u>Location:</u> USA	Prospective	Saliva	313	313	Not reported	Not reported	Not reported	Not reported

Study details	Study Type	Reference standard	Patient numbers		Diagnostic accuracy, % (95% CI)			
			Total Screened	With cCMV	Sensitivity	Specificity	PPV	NPV
Vives-Onos (2019) ⁶⁷ <u>Location</u> : Spain	Case-control	Any	184	103	56 (47 to 65)	98 (91 to 99)	Not reported	Not reported
Wang (2017) ⁴⁶ <u>Location</u> : China	Prospective	Saliva	3953	11	39.3 (22.1 to 59.3)	99.9 (99.8 to 100)	78.6 (48.8 to 94.3)	99.6 (99.4 to 99.8)
Abbreviations: CI, confidence interval; NPV, negative predictive value; PPV, positive predictive value. † Calculated using the 2 x 2 data provided in the publications.								

Table 6: Quality assessment results for the dried blood spot studies

Paper	Domain 1: Patient selection		Domain 2: Index tests		Domain 3: Reference Standard		Domain 4: Flow and timing
		Applicability		Applicability		Applicability	
	Risk of bias from selection	Concerns about matching of patients to review question?	Risk of bias relating to index test	Concerns about matching of index tests to review question?	Risk of bias relating to reference test	Concerns about matching of reference test to review question?	Risk of bias from patient flow?
First author							
Del Valle (2023) ⁶¹	Low	Low	Low	Low	Low	Unclear	Low
Dollard (2021) ³⁹	Low	Low	Low	Low	Low	Low	Unclear
Noorbakhsh (2020) ⁶⁴	Low	Low	Low	Low	Low	Low	Low
Vercauteren (2020) ⁶⁶	Low	Low	Low	Low	Low	Low	Unclear
Atkinson (2014) ⁶⁰	High	High	Low	High	Low	High	High
Fourgeaud (2023) ⁶²	Low	Low	Low	Low	Low	Low	Unclear
Moteki (2018) ⁶³	Low	Low	Low	Low	Low	Low	Low
Ross (2017) ⁶⁵	Low	Low	Low	Low	Unclear	Unclear	High
Vives-Onos (2019) ⁶⁷	Unclear	Low	Unclear	Low	Low	Low	High
Wang (2017) ⁴⁶	Low	Low	High	Unclear	High	Unclear	High

Additional material relating to Criterion 4

Urine testing

The previous UK NCS evidence review by Bazian Ltd (2017)¹⁸ did not include urine testing, even though it was considered to be the gold standard. This was stated to be due to the practical difficulties in collecting urine from newborns, limiting the role of urine testing in universal screening.

Systematic Reviews

No systematic reviews of urine testing were identified.

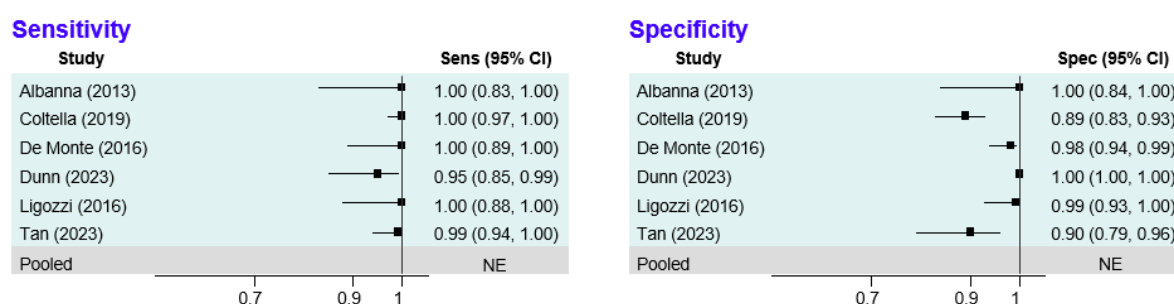
Diagnostic accuracy studies

Seven studies were identified (Table 7).^{41, 45, 52, 68-71} The studies were conducted in a wide variety of locations (2 in Italy, 1 in USA, 1 in France, 1 in Egypt, 1 in China), with only 1 from the UK. Three were prospective in design and 4 were retrospective or case-control studies. Four studies used urine viral cultures as the reference standard test. Studies were mostly small in size but varied from 58 newborns to just over 6,000. Diagnostic accuracy to detect cCMV was very high, with reported sensitivities and specificities at or close to 100% in most studies.

A meta-analysis was conducted on the studies that compared the diagnostic accuracy of urine PCR to urine viral cultures. Huang (2021)⁵² was not included as it compared urine PCR to saliva PCR and had already been included in the meta-analysis for saliva testing.

The bivariate meta-analysis could not reliably estimate pooled values for sensitivity and specificity due to the very high sensitivities and specificities in all included studies, and consequent lack of convergence of the models (see Figure 3). However, separate meta-analyses of sensitivity and specificity suggested that the sensitivity was over 99% (95% CI: 96 to 100) and specificity was also over 99% (95% CI: 92 to 100).

Figure 3 Diagnostic accuracy in studies of urine testing



Abbreviations: CI, confidence interval; NE, not evaluable; Sens, sensitivity; Spec, specificity.

The quality of the studies was assessed using QUADAS-2 (Table 8). As De Monte (2016)⁷⁰ was a French language publication it was not possible to assess its quality. Generally, all included studies were at low risk of bias across all domains.

Table 7: Results of the urine studies

Study details	Study Type	Reference standard	Patient numbers		Diagnostic accuracy, % (95% CI)			
			Total Screened	With cCMV	Sensitivity	Specificity	PPV	NPV
Studies included in meta-analysis								
Albanna (2013) ⁶⁸ <i>Location:</i> Egypt	Case-control	Urine culture	39	19	100	100	100	100
Coltella (2019) ⁶⁹ <i>Location:</i> Italy	Retrospective	Urine viral culture	255	107	100	89.19	Not reported	Not reported
De Monte (2016) ⁷⁰ <i>Location:</i> France	Retrospective	Urine viral culture	141	31	100 (86.9 to 100)	98.2 (93.2 to 99.2)	Not reported	Not reported
Dunn (2023) ⁴¹ <i>Location:</i> USA	Prospective	Urine composite	1624	43	95.4† (84.2 to 99.4)	100.0† (99.8 to 100.0)	Not reported	Not reported
Ligozzi (2016) ⁷¹ <i>Location:</i> Italy	Retrospective	Shell viral culture	110	29	100† (88.1 to 100)	98.8† (93.3 to 100.0)	Not reported	Not reported
Tan (2023) ⁴⁵ <i>Location:</i> UK	Prospective	Urine LDT P CR	148	96	99 (94.3 to 99.8)	90.4 (79.4 to 95.8)	95 (89.2 to 97.8)	97.9 (87.0 to 99.7)
Studies not included in meta-analysis								
Huang (2021) ⁵² <i>Location:</i> China	Prospective	Saliva	6234	49	28.6 (17.9 to 42.4)	100 (99.9 to 100)	100 (78.5 to 100)	99.4 (99.2 to 99.6)
Abbreviations: CI, confidence interval; NPV, negative predictive value; PPV, positive predictive value. † Calculated using the 2 x 2 data provided in the publications.								

Table 8: Quality assessment results for the urine studies

Paper	Domain 1: Patient selection		Domain 2: Index tests		Domain 3: Reference Standard		Domain 4: Flow and timing
		Applicability		Applicability		Applicability	
First author	Risk of bias from selection	Concerns about matching of patients to review question?	Risk of bias relating to index test	Concerns about matching of index tests to review question?	Risk of bias relating to reference test	Concerns about matching of reference test to review question?	Risk of bias from patient flow?
Albanna (2013) ⁶⁸	Low	Low	Low	Low	Low	Low	Low
Coltella (2019) ⁶⁹	Low	Low	Low	Low	Low	Low	Low
De Monte (2016) ⁷⁰	Unclear	Unclear	Unclear	Unclear	Unclear	Unclear	Unclear
Dunn (2023) ⁴¹	High	Unclear	Low	Low	Low	Low	Low
Ligozzi (2016) ⁷¹	Unclear	Unclear	Low	Low	Low	Low	Low
Tan (2023) ⁴⁵	Unclear	Unclear	Low	Low	Low	Low	Unclear
Huang (2021) ⁵²	Low	Low	Unclear	Unclear	Unclear	Unclear	Unclear

Other testing for cCMV infection

Systematic Reviews

No systematic reviews were identified.

Diagnostic accuracy studies

An additional 3 studies⁷²⁻⁷⁴ were identified that tested for cCMV using uncommon testing methods (Table 9). Wang (2013)⁷² compared the accuracy of diagnosing cCMV using throat swabs to umbilical cord blood. Barkai (2013)⁷³ compared the diagnostic accuracy of umbilical cord blood (as an alternative to DBS) to urine testing. The sensitivity and specificity reported in Barkai (2013) was quite high. Reyes (2020)⁷⁴ also looked at umbilical cord blood but did not report diagnostic accuracy results.

As these 3 studies examine different tests, and are not tests likely to be used in routine screening, no quality assessment or synthesis of these studies was performed.

Table 9: Results of the studies of other testing methods

Study details	Study Type	Testing Method	Reference standard	Patient numbers		Diagnostic accuracy, % (95% CI)			
				Total Screened	With cCMV	Sensitivity	Specificity	PPV	NPV
Wang (2013) ⁷² <u>Location</u> : China	Not reported	Throat swab	Cord blood (PCR)	Not reported	Not reported	67	75	57	82
Barkai (2013) ⁷³ <u>Location</u> : Israel	Retrospective (screening programme)	Cord blood	Urine	8105	24	100 (84.6 to 100.0)	99.1 (95.2 to 99.9)	Not reported	Not reported
Reyes (2020) ⁷⁴ <u>Location</u> : Spain	Retrospective	Cord blood (dried)	Mixed	16	16	Not reported	Not reported	Not reported	Not reported
Abbreviations: cCMV, congenital cytomegalovirus; CI, confidence interval; NPV, negative predictive value; PCR, polymerase chain reaction; PPV, positive predictive value.									

Summary of Findings Relevant to Criterion 4

Criterion Met

There is now a large body of clinical evidence, including 2 systematic reviews and 39 diagnostic accuracy studies published since 2012, on the accuracy of newborn testing for cCMV. This evidence, both the systematic reviews and the primary studies, is generally of good quality. It seems unlikely that any future studies or reviews will substantially differ from the conclusions reached in this review.

This review identified 1 existing systematic review (Zheng, et al., 2022) and 22 diagnostic accuracy studies of newborn saliva PCR testing to detect cCMV infection. Both the review and the primary studies were generally of good quality and at low risk of bias. The meta-analyses of diagnostic accuracy conducted in the Zheng (2022) review and in this review were consistent in their results. Both found that saliva testing will detect over 90% of all cCMV cases, with a very low rate of false-positive results (around 0.2%). This very high accuracy suggests that saliva PCR testing is a sufficiently accurate test that could reasonably be used in a cCMV screening programme. It should be noted that this high detection rate refers to all cCMV cases, including those which are asymptomatic at birth and/or do not go on to develop long-term symptoms.

Despite the high accuracy of saliva testing the positive predictive value is predicted to be around 87%, so some newborns with a positive saliva test will not have cCMV infection. The exact proportion is currently uncertain, as it was highly variable across studies. This suggests that confirmatory testing after a positive saliva test may be required.

This review identified 1 existing systematic review (Wang, et al., 2015; included in the previous UK NSC review of this topic) and 10 diagnostic accuracy studies published after that review, of newborn DBS PCR testing to detect cCMV infection. The review and the primary studies were generally of good quality and at low risk of bias. The meta-analyses of diagnostic accuracy conducted in the Wang et al. (2015) review²⁷ and in this review were consistent in their results. Both found that DBS testing will detect most cCMV cases (around 80% to 90% of cases), with a very low rate of false-positive results (less than 1%). This high accuracy suggests that DBS PCR testing is a sufficiently accurate test that could reasonably be used in a cCMV screening programme. However, it does appear to be less accurate than saliva PCR testing.

This review did not find any studies comparing saliva and DBS testing, so it is uncertain which is preferable in practice. Although DBS seems to have a lower diagnostic accuracy

this would need confirming in studies directly comparing the 2 methods. Also, it is possible that, while DBS is less accurate overall, the cCMV cases detected may be the more serious cases requiring treatment, but that requires further research. DBS may also have some practical benefits over saliva testing, as DBS testing is already in use for other conditions.

This review found only 7 studies on the diagnostic accuracy of urine testing. This is probably because urine testing was already well established as a reference standard test for cCMV infection before this review was conducted. The results from the included studies confirm that urine testing is highly accurate, and so would be an appropriate confirmatory test after a positive saliva or DBS test result in any cCMV screening programme.

Criterion 7: There should be an agreed policy on the further diagnostic investigation of individuals with a positive test result and on the choices available to those individuals.

What tests in newborns with cCMV identify which are likely to suffer long-term health consequences from cCMV, who would benefit from prompt treatment?

Previous work

The previous review included 3 cohort studies: Alarcon et al. (2013),⁷⁵ Bilavsky et al. (2015)⁷⁶ and Forner et al. (2015),⁷⁷ that assessed whether specific signs or symptoms were predictive of adverse long-term outcomes.¹⁸ All 3 studies were also identified by our searches. The previous review identified several limitations to these studies, including the small sample sizes. The previous review concluded that the available evidence provides only limited new information about treatment indications and there remains a lack of clarity about which newborns with cCMV will go on to develop adverse outcomes and might benefit from treatment.

The previous review included 1 systematic review and 5 cohort studies that reinforced the previously established observation that symptomatic newborns are more likely to develop adverse outcomes than asymptomatic newborns.¹⁸ One of these studies, was also identified by our searches.⁷⁸

The Evidence Map of 2022⁷ identified 14 studies, including 1 systematic review, relevant to this topic. Most examined the impact of viral load on long-term outcomes. The conclusion was that although some studies identified possible markers to distinguish which cCMV infected newborns will suffer long-term negative outcomes, there was no consensus about the predictive value of such markers or their reliability.

Description of the current evidence

Overall, 97 studies were identified as being relevant for this research question and were included in the review. Priority was given to systematic reviews, followed by “prognostic accuracy” studies, which we define as studies that formally assess the ability of a test to predict later health outcomes in terms of sensitivity and specificity or similar metrics. Other studies reported whether a test result, antenatal characteristic or newborn characteristic was associated with long-term health consequences, but without formal assessment of prognostic value. These are termed “association studies” in this report. These association

studies are only briefly summarised because of their large numbers, and because they are less informative as to the true clinical value of tests and their consequent lesser relevance to this assessment.

Owing to the large number of eligible studies, the evidence has been split according to the type of test or characteristic investigated.

Neuroimaging (MRI and ultrasound)

Systematic review evidence

One systematic review by Van de Walle et al.(2024)⁷⁹ assessed the value of MRI to predict clinical outcome in children with cCMV. This review included 20 studies. However, only 7 studies evaluated the associations between postnatal MRI and later clinical outcomes, and the review did not consider prognostic accuracy for detection of sequelae as an outcome. Critical appraisal of the systematic review revealed a high risk of bias. This was due to an unclear description of the prespecified study inclusion criteria in the absence of a published protocol, a limited search strategy, poor reporting of study characteristics and no attempt to narratively or quantitatively synthesise study findings.

No synthesis was undertaken, and associations between postnatal MRI and later outcomes were presented separately for each study. Therefore, studies included in this review are discussed in the sections below.

Prognostic accuracy studies

Sixteen studies assessing the accuracy of neuroimaging (either used alone or in combination with non-imaging tests) to predict neurodevelopmental delay, epilepsy or hearing loss were identified.^{75, 80-94} Full results from these studies are summarised in Table 10.

In general, neuroimaging tests demonstrated reasonable accuracy in predicting neurodevelopmental delay, epilepsy and hearing outcomes. The most accurate tests were MRI or ultrasound (and sometimes computed tomography) using the 'Alarcon' scoring. Using a threshold of 2 or more to predict neurodevelopmental delay, these tests yielded sensitivities from 77% to 94%, with specificities of 69% to 100%, in 2 studies.^{80,81} MRI had good accuracy to detect epilepsy, with a sensitivity of 92% (95% CI: 62 to 99) at a specificity of 68% (95% CI: 44 to 87).⁹¹

Combining neuroimaging tests with other tests sometimes improved accuracy. A sensitivity of 87% at a specificity of 100%⁷⁵ for predicting neurodevelopmental delay was yielded by a test result involving a score of ≥ 2 on the Noyola scale or a cerebrospinal fluid (CSF) beta 2-

m concentration above a threshold of 7.9mg/L. In another study,⁸⁹ a composite test result of normal cerebral ultrasound, no hearing loss, and normal platelet count at birth predicted the absence of neurological or auditory sequelae with a sensitivity of 92% at a specificity of 83%.

In general, studies were small, and most studies had either a high or unclear overall risk of bias (see Appendix 3).

Table 10: Summary of evidence evaluating the accuracy of neuroimaging tests to predict sequelae of cCMV infection

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Alarcon et al. 2013 ⁷⁵ Prospective Spain Follow-up: mean 8.7 years	With cCMV: 26 Symptoms at birth: 26 Long-term outcomes: 15	Mixed (CSF, microcephaly, neuroimaging)	Neurodevelopmental delay (in terms of motor function, cognition, behaviour, hearing, vision, and epilepsy) at mean 8.7 years	Neuroimaging (Noyola score of ≥ 2) OR Adjusted microcephaly (threshold unclear) Sens: 66% Spec: 100% CSF concentration ($>7.9\text{mg/L}$) OR Neuroimaging (Noyola score of ≥ 2)^A Sens: 87% Spec: 100%
Alarcon et al. 2016 ⁸⁰ Prospective Spain Follow-up: mean 8.7 years	With cCMV: 26 Symptoms at birth: 26 Long-term outcomes: 18	MRI/ Ultrasound/CT	Neurodevelopmental delay (in terms of motor function, cognition, behaviour, hearing, vision, and epilepsy) at mean 8.7 years	Neuroimaging with MRI/US/CT (Noyola scale ≥ 2) Sens: 61% Spec: 100% Neuroimaging with MRI/US/CT (Alarcon scale ≥ 2) Sens: 77% Spec: 100%
Alarcon et al. 2024 ⁸¹ Retrospective Spain Follow-up: median 4 years	With cCMV: 160 Symptoms at birth: 103 Long-term outcomes: 53	MRI/ US	Moderate/severe disabilities: neurodevelopmental impairments (including hearing, CP, cognition, behavioral, epilepsy, visual) at median 4 years	Neuroimaging with MRI/US (Alarcon scale ≥ 2) Sens: 90% (95% Ct 78.2 to 96.7) Spec: 69% (95% Ct 58.1 to 78.5) TPWMA on MRI/US (1 or more present) Sens: 77.8% (95% Ct 40 to 97.2) Spec: 69.1% (95% Ct 52.9 to 82.4)
Blazquez-Gamero et al. 2019 ⁸² Prospective Spain Follow-up: 12 months	With cCMV: 107 Symptoms at birth: 77 Long-term outcomes: 37	MRI/ symptoms at birth	Any sequelae (hearing or neurological) at 12 months	Abnormal US and abnormal MRI (any category of symptoms at birth) Sens: 83.8% (95% Ct 68.0 to 93.8) Spec: 79.4% (95% Ct 62.1 to 91.3) Abnormal US and abnormal MRI (with symptoms at birth) Sens: 81.1% (95% Ct 64.8 to 92.0) Spec: 85.3% (95% Ct 68.9 to 95.1) Abnormal US and abnormal MRI (with NO symptoms at birth) Sens: 2.7% (95% Ct 0.7 to 14.2) Spec: 94.1% (95% Ct 80.3 to 99.3)
Capretti et al. 2014 ⁸³ Prospective Italy Follow-up: 6 years	With cCMV: 40 Symptoms at birth: 11 Long-term outcomes: 7	MRI/ Ultrasound	Psychomotor development at 6 years	US (pathological) Sens: 57% (95% Ct 20 to 88) Spec: 94% (95% Ct 78 to 96) MRI (pathological) Sens: 100% (95% Ct 56 to 100) Spec: 94% (95% Ct 78 to 98)

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Castellanos et al. 2022 ⁸⁴ Retrospective Spain Follow-up: 4 years	With cCMV: 36 Symptoms at birth: 20 Long-term outcomes: 18	MRI/ Ultrasound	Neurological sequelae (WMA and other neuroimaging findings, delayed development, hearing loss) at 4 years	US(pathological) Sens: 83.3% (95% Ct 58 to 100) Spec: 44.4% (95% Ct 18.7 to 70.2) MRI(Noyola score >=2) Sens: 94% (95% Ct 80 to 100) Spec: 66.6% (95% Ct 36 to 97.5)
Chebib et al. 2022 ⁸⁵ Retrospective France Follow-up: 21 months	With cCMV: 130 Symptoms at birth: 94 Long-term outcomes: 83	Antenatal infection point / fetal imaging	Inner ear impairment at median 21 months	Infection in first trimester OR Imaging anomaly of fetus Sens: 91% (95% Ct 83 to 98) Spec: 48% (95% Ct 36 to 60) Infection in first trimester AND Imaging anomaly of fetus Sens: 38% (95% Ct 28 to 47) Spec: 92% (95% Ct 86-98)
Craeghs et al. 2021 ⁸⁶ Prospective Belgium Follow-up: mean around 2.5 years	With cCMV: 411 Symptoms at birth: 164 Long-term outcomes: 75	MRI/ Ultrasound	Hearing loss at mean 2.5 years	US(pathological) Sens: 43% (95% Ct 31 to 54) Spec: 84% (95% Ct 79 to 88) MRI(pathological) Sens: 39% (95% Ct 28 to 50) Spec: 78% (95% Ct 72 to 83)
De Juan et al. 2020 ⁸⁷ Retrospective Spain Follow-up: mean 19 months	With cCMV: 60 Symptoms at birth: unclear Long-term outcomes: 32	US/MRI	Neurological sequelae at mean of 19 months	US or MRI(score of 1 or more on Alarcon scale) Sens: 71.9% (95% Ct 53.3 to 86.3) Spec: 71.4% (95% Ct 51.3 to 86.8)
Foulon et al. 2019 ⁸⁸ Prospective Belgium Follow-up: median 31 months	With cCMV: 173 Symptoms at birth: 9 Long-term outcomes: 20	Trimester of infection / US/MRI	Hearing loss (mild or worse) at median 31 months	US(abnormal) Sens: 47.1% (95% Ct 23 to 72.2) Spec: 92.5% (95% Ct 86.2 to 96.5) MRI(abnormal) Sens: 57.1% (95% Ct 18.4 to 90.1) Spec: 66.7% (95% Ct 43.0 to 85.4)
Fourgeaud et al. 2024 ⁸⁹ Prospective France Follow-up: 24 months	With cCMV: 250 Symptoms at birth: 24 Long-term outcomes: 19	<i>Lack of</i> risk factors for SNHL / psychomotor problems, as aimed to predict NO sequelae	<i>Lack of</i> SNHL (bilateral or unilateral) OR psychomotor sequelae (motor delay, psychomotor delay, autism, nystagmus) at 24 months	No hearing loss at birth, normal cerebral ultrasound, and normal platelet count (for detection of NO sequelae) Sens: 92% Spec: 83%

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Gianattasio et al. 2018 ⁹⁰ Retrospective Italy Follow-up: around 6 years	With cCMV: 170 Symptoms at birth: 112 Long-term outcomes: Unclear	US/MRI/CT	An impaired neurologic outcome was defined in the presence of 1 or more of the following problems: developmental/cognitive impairment, motor delay requiring rehabilitation, epilepsy and behavioural/emotional problems. Follow up 1-6 years.	US(Noyola >=1) Sens: 75.6% (95% Ct 62.2 to 86.7) Spec: 43.3% (95% Ct 31.7 to 56.7) CT(Noyola scale >=1) Sens: 66.7% (95% Ct 53.3 to 80) Spec: 85.7% (95% Ct 76.8 to 94.6) MRI(Noyola scale >=1) Sens: 42.6% (95% Ct 29.8 to 57.5) Spec: 89.1% (95% Ct 80 to 96.4) US(Alarcon >=1) Sens: 88.9% (95% Ct 80 to 97.8) Spec: 40% (95% Ct 28.3 to 51.7) CT(Alarcon >=1) Sens: 88.9% (95% Ct 80 to 97.8) Spec: 62.5% (95% Ct 50 to 75) MRI(Alarcon >=1) Sens: 87.2% (95% Ct 76.6 to 95.7) Spec: 50.9% (95% Ct 38.2 to 63.6)
Kwak et al. 2018 ⁹¹ Retrospective South Korea Follow-up: mean 3.2 years	With cCMV: 31 Symptoms at birth: 31 Long-term outcomes: 11	MRI	Epilepsy at mean 3.2 years	MRI(>=2.5 points on un-named scale) Sens: 91.7% (95% Ct 61.5 to 99.8) Spec: 68.4% (95% Ct 43.5 to 87.4)
Lucignani et al. 2021 ⁹² Retrospective Italy Follow-up: 2 years	With cCMV: 44 Symptoms at birth: 27 Long-term outcomes: 30	MRI	Development of adverse neurological outcomes [including hearing loss, severe neurological delay or CP] by end of follow up at minimum of 2 years	MRI(score >0 'old system') Sens: 93.3% (95% Ct 77.9 to 99.2) Spec: 50% (95% Ct 23.0 to 77.0) MRI(score >2 'new system') Sens: 70% (95% Ct 50.6 to 85.3) Spec: 80%
Nishida et al. 2020 ⁹³ Prospective Japan Follow-up: 18 months	With cCMV: 42 Symptoms at birth: 23 Long-term outcomes: 19	MRI/ clinical symptoms	Neurodevelopmental impairment (NDI) was defined as a developmental quotient <80, hearing dysfunction, blindness, or epilepsy requiring anti-epileptic drugs at 18 months	MRI(any brain abnormality) Sens: 89.5% (95% Ct 66.9 to 98.7) Spec: 52.6% (95% Ct 28.9 to 75.6) MRI(>=1 of the defined 3 MRI abnormalities of V, PVC or WMA) Sens: 89.5% (95% Ct 66.9 to 98.7) Spec: 52.6% (95% Ct 28.9 to 75.6) MRI(>=2 of the defined 3 MRI abnormalities) Sens: 89.5% (95% Ct 66.9 to 98.7) Spec: 89.5% (95% Ct 66.9 to 98.7)

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Vande Walle et al. 2024 ⁹⁴ Retrospective Belgium Follow-up: up to 4 years	With cCMV: 255 Symptoms at birth: unclear Long-term outcomes: 53	MRI	Cognitive development (Bayley Scales of Infant and Toddler Development) Severe impairment < 50. Motor development (AIMS). AIMS < P10 abnormal. Hearing evaluation (ABR). ABR thresholds ≤ 40 dB nHL considered normal. Follow up median 12 months	MRI (Apparent Diffusion Coefficient of white matter [ADC] > 0.0497) – optimum threshold for predicting hearing impairment Sens: 86% Spec: 62% MRI (ADC > 0.0671) – optimum threshold for predicting cognitive impairment Sens: 83% Spec: 58% MRI (ADC > 0.159) – optimum threshold for predicting motor development Sens: 57% Spec: 72%
Abbreviations: ABR, Auditory Brainstem Response; ADC, apparent diffusion co-efficient; AIMS, Alberta Infant Motor Scale; cCMV, congenital cytomegalovirus; CMV, Cytomegalovirus; CP, cerebral palsy; CSF, cerebrospinal fluid; CT, computed tomography; dB, decibels; DNA, deoxyribonucleic acid; MRI, magnetic resonance imaging; NDI, neurodevelopmental impairment; PCR, polymerase chain reaction; PVC, periventricular cyst; Sens, sensitivity; SNHL, sensorineural hearing loss; Spec, specificity; TPWMA, temporal pole white matter abnormalities; US, ultrasound; V, ventriculomegaly; WM, white matter; WMA, white matter abnormalities.				

Association studies

Seven studies that assessed the association between neuroimaging and long-term outcomes, but did not report prognostic accuracy data, were identified.⁹⁵⁻¹⁰¹ Studies evaluated US⁹⁵⁻⁹⁹ and/or MRI.^{96-98, 100, 101} Five studies were prospective^{95, 96, 99-101} and 2 were retrospective.^{97, 98} Sample sizes ranged from 11 to 211 infants with cCMV, and median follow-up ranged from 12 to 68 months. Studies were conducted in Italy, Switzerland, Netherlands, Belgium and Spain.

MRI

Vande Walle et al. (2023)¹⁰⁰ showed that white matter abnormalities (WMA) on MRI are associated with developing adverse motor sequelae and cognitive delay, but not late-onset hearing loss. Other studies demonstrated associations between abnormal MRI findings in the newborn period and subsequent hearing,⁹⁷ neurodevelopmental^{96, 101} or neurodevelopmental and hearing⁹⁸ deficits, but these findings have either not been statistically tested^{96, 98, 101} or have not demonstrated a significant effect.⁹⁷

Ultrasound

Three studies⁹⁷⁻⁹⁹ demonstrated that ultrasound abnormalities are predictors of hearing and neurodevelopmental sequelae. However, 4 studies^{95-97, 99} found that ultrasound abnormalities were not significantly associated with hearing and neurodevelopmental sequelae, particularly after adjustment for confounding. These inconsistent findings suggest that ultrasound may be less useful than MRI for predicting sequelae.

Antenatal characteristics – timing of maternal infection

Systematic review evidence

Two relevant systematic reviews were found. The systematic review by Chatzakis et al. (2020)¹⁰² included 17 studies. Of those, 6 studies (772 participants) evaluated the associations between timing of infection and sensorineural hearing loss (SNHL) or neurodevelopmental impairment at follow-up, and were included in meta-analyses.

The review had a low risk of bias. The protocol was published on PROSPERO and set appropriate inclusion criteria. Searches, data extraction and synthesis were all conducted appropriately.

Primary maternal infection in the first trimester led to a prevalence of SNHL or neurodevelopmental impairment of 22.8% (95% CI: 15.4 to 30.2; $I^2=27.6\%$) at follow up. Primary maternal infection in the second and third trimesters led to smaller rates of 0.1% (95% CI: 0 to 0.8; $I^2=24\%$) and 0% (95% CI: 0 to 2.1; $I^2=0\%$) at follow up, respectively.

The systematic review by Buca et al. (2021)¹⁰³ included 26 studies, but only a subset were used in the relevant analyses. Two of the 26 included studies were included in our review.^{104,}

¹⁰⁵ This review had a low risk of bias, with clear inclusion criteria, and good methods of searching, data extraction and synthesis. The pooled percentage of children with abnormal neurodevelopmental outcomes was 5.44% (95% CI: 2.3 to 9.8) in 7 studies evaluating first trimester infection, 0% (95% CI: 0 to 7) in 6 studies evaluating second trimester infection, and 0% (95% CI: 0 to 7.7) in 4 studies evaluating third trimester infection. The pooled percentage of children with hearing problems was 11.35% (95% CI: 6.7 to 17.1) in 7 studies evaluating first trimester infection, 7% (95% CI: 2.1 to 14.5) in 6 studies evaluating second trimester infection, and 0% (95% CI: 0 to 7.7) in 4 studies evaluating third trimester infection.

Prognostic accuracy studies

Two studies were identified. Foulon et al. (2019)⁸⁸ (see Table 11) was included in the Chatzakis et al. (2020) review. It was a relatively large study (72 for this analysis) but had an unclear overall risk of bias (see Appendix 3). It found that primary infection during the first trimester had a sensitivity of 56% (95% CI: 21 to 86) at a specificity of 81% (95% CI: 69 to 90) for predicting hearing loss. Infection during the first *or* second trimester had a sensitivity of 89% (95% CI: 52 to 100) at a specificity of 33% (95% CI: 22 to 46) for predicting hearing loss. Neither of these thresholds are likely to be adequate in terms of avoiding unacceptably high levels of either false negative or false positive results.

Chebib et al. (2022)⁸⁵ was rated at serious risk of bias and so should be interpreted with caution (see Appendix 3). It showed primary infection in the first trimester *or* having an imaging anomaly of the fetus had a sensitivity of 91% (95% CI: 83 to 98) and specificity of 48% (95% CI: 36 to 60) for detecting inner ear dysfunction at 21 months. Similarly, a positive test defined by infection in the first trimester and having an imaging anomaly of the fetus had a sensitivity of 38% (95% CI: 28 to 47) and specificity of 92% (95% CI: 86 to 98) (Table 10).

Table 11: Summary of evidence evaluating the accuracy of timing of infection to predict sequelae of cCMV infection

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Foulon et al. 2019 ⁸⁸ Prospective Belgium Follow-up: median 31 months	With cCMV: 173 Symptoms at birth: 9 Long-term outcomes: 20	Trimester of infection	Hearing loss (mild or worse) at median 31 months	Trimester of infection (1st) Sens: 55.6% (95% Ct 21.2 to 86.3) Spec: 81% (95% Ct 69.1 to 89.8) Trimester of infection (1st or 2nd) Sens: 88.9% (95% Ct 51.8 to 99.7) Spec: 33.3% (95% Ct 22 to 46.3)
Abbreviations: cCMV, congenital cytomegalovirus; CI, confidence interval; MRI, magnetic resonance imaging; Sens, sensitivity; Spec, specificity; US, ultrasound.				

Association studies

Sixteen studies that assessed the association between timing of maternal infection and long-term outcomes were identified.^{98, 104-118} Three of these studies were included in the systematic review by Chatzakis et al. (2020).^{107, 108, 111} Therefore, we have summarised the 13 additional association studies, not included in the Chatzakis review.

Six of these studies were prospective^{104, 106, 115-118} and 7 were retrospective.^{98, 105, 109, 110, 112-114} Sample sizes ranged from 27 to 753 infants with cCMV and follow-up ranged from 12 to 185 months (mean). Studies were conducted in Finland, Italy, Netherlands, Belgium, France, USA and Israel.

Five of these studies showed that primary infection in the first trimester significantly increased risk of long-term neurological and/or hearing sequelae.^{105, 110, 112, 115, 116} Seven studies^{98, 104, 106, 109, 113, 114, 117} demonstrated the same direction of effect, reporting a greater prevalence of long-term neurological and/or hearing sequelae with earlier infection. However, these effects were either not statistically tested,^{98, 106, 109, 113, 117} or yielded a non-significant result.^{104, 114} In contrast, 1 study¹¹⁸ showed a greater prevalence of hearing sequelae in those with later infection, but this very small study did not conduct a statistical analysis.

Antenatal characteristics – type of maternal infection

Systematic review evidence

One systematic review by Maltezou et al. (2020)¹¹⁹ compared long-term sequelae in children born following maternal primary or non-primary infection. The review reported data from 9 studies. Eight of these studies (835 persons) presented evidence assessing the association of the type of maternal infection and later outcomes. The review was at low risk of bias, with a protocol published on PROSPERO, and appropriate methods of searching, data extraction and synthesis.

Meta-analyses showed that maternal infection status (primary versus non-primary infection) did not influence the risk of developing unilateral SNHL (OR 0.80 (95% CI: 0.54 to 1.20)), bilateral SNHL (OR 0.66 (95% CI: 0.30 to 1.42)) or other neurologic outcomes (OR 0.73 (95% CI: 0.43–1.25)).

Prognostic accuracy studies

No prognostic accuracy studies assessing type of maternal infection were identified.

Association studies

Nine studies were identified.^{111-113, 117, 118, 120-123} Three of these studies^{111, 122, 123} were included in the systematic review by Maltezou et al. (2020).¹¹⁹ We have summarised the other 6 studies.^{112, 113, 117, 118, 120, 121} Four studies were prospective^{117, 118, 120, 121} and 2 were retrospective.^{112, 113} Sample sizes ranged from 22 to 186 infants with cCMV and follow-up ranged from 12 to 185 months (mean). The studies were from Finland, Italy, the USA and Iran.

Two studies^{112, 113} found that a significantly higher proportion of infants born to mothers with non-primary infection (compared to primary infection) had long-term hearing or neurodevelopmental sequelae. Another study¹¹⁸ found no numerical difference in the incidence of hearing sequelae between primary and non-primary maternal infection groups. Two studies^{117, 120} found that prevalence of hearing sequelae was slightly higher in the primary infection group. Finally, 1 study¹²¹ stated that there was no correlation between the type of infection and long-term visual sequelae, but no data were clearly presented to support this.

Gestational age at birth

Systematic review evidence

No systematic reviews were identified.

Prognostic accuracy studies

No prognostic accuracy studies were identified.

Association studies

Two studies^{101, 124} evaluated the association between gestational age and hearing and neurodevelopmental sequelae. One study was prospective¹⁰¹ and the other was retrospective.¹²⁴ Follow up was between 6 and 72 months. Sample sizes ranged from 11 to 102 infants with cCMV. The studies were from Italy and the Netherlands.

One study¹²⁴ showed that significantly more infants who developed long-term impairment were born premature, compared with infants who did not develop long-term impairment ($p=0.024$). The other study¹⁰¹ suggested that neurodevelopmental sequelae were numerically more likely in infants with a lower gestational age.

Symptoms at birth

Systematic review evidence

Three relevant systematic reviews were found.¹²⁵⁻¹²⁷ Smyrli et al. (2024)¹²⁵ assessed neurodevelopmental outcomes of children who had presented with asymptomatic cCMV at

birth. The review had a low risk of bias overall. The protocol was published on PROSPERO, and appropriate methods of data extraction and synthesis were used. The only limitation was the limited number of databases used in the search. All 28 studies in the systematic review evaluated the long-term outcomes for children with asymptomatic cCMV. The review included 28 studies, 19 of which reported analyses of outcomes in children who were asymptomatic at birth versus children who were symptomatic at birth. Sixteen of those 19 studies showed that adverse neurodevelopmental outcomes were less common among children with asymptomatic cCMV at birth. No formal meta-analyses were undertaken.

Vos et al. (2021)¹²⁶ assessed the association between symptomatic status at birth and childhood and late-onset hearing loss. The review included 65 studies, but only 17 were relevant to this assessment. The review had a low risk of bias overall, with a published protocol, and appropriate methods of searching, data collection and synthesis. The narrative review did not form any firm conclusions about an association between symptomatic status at birth and subsequent hearing loss in children with cCMV, although they found that hearing loss was associated with both symptomatic and asymptomatic disease.

Riga et al. (2018)¹²⁷ also evaluated the relationship between symptoms at birth and hearing loss, at up to 10 years of age. Eleven studies were included. The review was deemed to be at high risk of bias, principally because of limitations in the search strategy and synthesis methodology. Among asymptomatic newborns, SNHL was diagnosed in 8.9%, while for symptomatic newborns the percentage raised to 44.2%. This synthesis appeared to be a simple summation of study results rather than a proper meta-analysis.

Prognostic accuracy studies

Nishida et al. (2020)⁹³ evaluated the accuracy of the existence of symptoms and found a sensitivity of 84% (95% CI: 60 to 97) at a specificity of 73.7% (95% CI: 49 to 91) (Table 12). The combination of 2 or more clinical symptoms and/or ventriculopathy, periventricular cysts and white matter abnormalities on imaging yielded a sensitivity of 90% (95%CI: 67 to 99) at a specificity of 75% (95% CI: 49 to 91) (Table 10). The combination of 3 or more clinical symptoms and/or ventriculopathy, periventricular cysts and white matter abnormalities on imaging yielded a sensitivity of 79% (95%CI: 54 to 94) at a specificity of 95% (95% CI: 74 to 100) (Table 10). More results are available in Appendix 3. This was a small (n=32) study, with a relatively short follow up of 18 months, and with unclear risk of bias (see Appendix 3).

Blazquez-Gamero et al. (2019)⁸² evaluated symptoms at birth combined with neuroimaging to predict any hearing or neurological sequelae. Neuroimaging in symptomatic newborns had a sensitivity of 81% (95% CI: 65 to 92) at a specificity of 85% (95% CI: 69 to 95).

Neuroimaging in asymptomatic newborns was not prognostic, with a sensitivity of 2.7% (95%

CI: 0.07 to 14) at a specificity of 94% (95% CI: 80 to 99). These results are summarised in Table 10. This moderately sized (n=71) study was at high risk of bias and there were concerns about applicability (see Appendix 3).

Alarcon et al. (2013)⁷⁵ assessed the accuracy of adjusted microcephaly for predicting neurodevelopmental delay, yielding a sensitivity of 44% and specificity of 100% (Table 12). The study also evaluated the accuracy of adjusted microcephaly in combination with positive neuroimaging findings (sensitivity of 66% at a specificity of 100%) and in combination with C SF beta2-m concentration (sensitivity of 82% at a specificity of 100) (Table 10). This small (n=26) study was also at high risk of bias and had concerns about applicability (see Appendix 3).

Table 12: Summary of evidence evaluating the accuracy of symptoms at birth to predict sequelae of cCMV infection

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Alarcon et al. 2013 ⁷⁵ Prospective Spain Follow-up: mean 8.7 years	With cCMV: 26 Symptoms at birth: 26 Long-term outcomes: 15	Mixed (CSF, microcephaly, neuroimaging)	Neurodevelopmental delay (in terms of motor function, cognition, behaviour, hearing, vision, and epilepsy) at mean 8.7 years	Adjusted microcephaly (threshold unclear) Sens: 44% Spec: 100%
Nishida et al. 2020 ⁹³ Prospective Japan Follow-up: 18 months	With cCMV: 42 Symptoms at birth: 23 Long-term outcomes: 19	Clinical symptoms	Neurodevelopmental impairment (NDI) was defined as a developmental quotient <80, hearing dysfunction, blindness, or epilepsy requiring anti-epileptic drugs at 18 months	Clinical symptoms (present) Sens: 84.2% (95% CI: 60.4 to 96.6) Spec: 73.7% (95% CI: 48.8 to 90.9)
Abbreviations: cCMV, congenital cytomegalovirus; CI, confidence interval; CSF, cerebrospinal fluid; NDI, neurodevelopmental impairment; Sens, sensitivity; Spec, specificity.				

Association studies

Thirty-six studies^{78, 97, 99, 105, 114-118, 120-122, 124, 128-150} were identified. However, 8 of these studies^{78, 136, 145-150} were included in the systematic reviews.¹²⁵⁻¹²⁷ Therefore, we have summarised the 28 additional association studies. There were 16 prospective cohort studies and 12 retrospective cohort studies. Sample sizes ranged from 22 to 753 infants with cCMV, and follow-up ranged from 3 months to 18 years, where reported (most studies followed children up for over 2 years). Studies were conducted in Italy, Belgium, the Netherlands, the USA, Australia, Indonesia, China, Taiwan, Switzerland, Spain, Iran, Finland, and UK/Sweden.

Twenty-six of the studies compared the outcomes of children with symptomatic cCMV at birth with those with asymptomatic cCMV at birth.^{97, 99, 105, 114-118, 120-122, 124, 128-135, 137-139, 142-144} Nine of these 26 studies showed that symptoms at birth were significantly associated with an increased risk of long-term visual,¹²¹ hearing,^{97, 114, 132} neurodevelopmental and hearing¹²⁸ and unspecified^{116, 122, 99, 105} long-term sequelae. In the other 17 studies statistical analyses were not carried out but the same direction of effect was evident, with a larger numerical prevalence of hearing,^{115, 117, 118, 120, 131, 134, 135, 142} visual,^{133, 137} language,¹¹⁴ or neurodevelopmental/hearing^{124, 129, 130, 138, 139, 143, 144} sequelae in the symptomatic groups.

Two studies^{140, 141} did not compare outcomes in generally symptomatic and asymptomatic infants, instead focussing on the effects of specific baseline symptoms. One study¹⁴⁰ compared infants with CNS involvement against those without CNS involvement and those with only petechial rash. Significantly more infants in the CNS group had SNHL at birth, but the frequency of late-onset SNHL did not differ between groups. Significantly more children in the CNS group had an IQ below 70. The other study¹⁴¹ compared infants who were small for gestational age with those appropriate for gestational age; long-term sequelae were only reported for the 14 symptomatic infants (all of whom had long-term sequelae). The study concluded that long-term outcomes do not seem to be affected by isolated impaired fetal growth, however, the study is severely limited by the very small sample size.

Audiological screening result at birth

Systematic review evidence

No systematic reviews were identified.

Prognostic accuracy studies

Two studies evaluated the prognostic accuracy of audiological screening at birth. Dhondt et al. (2023)¹⁵¹ assessed the accuracy of any hearing loss at birth for predicting vestibular function at 4 years. Although children who did not go on to develop vestibular dysfunction

rarely had hearing loss at birth, yielding a specificity 91% (95% CI: 86 to 95), only a small proportion of those who did develop vestibular function had baseline hearing loss, yielding a sensitivity 35% (95% CI: 15 to 59) (Table 13). This was a relatively large study (n=169), with good applicability but a high risk of overall bias (see Appendix 3).

Forgeaud et al. (2024)⁸⁹ also evaluated hearing loss as part of a composite test involving platelet count and US measurements at birth. The results (sensitivity 92% at a specificity of 83%) are discussed in more depth in the neuroimaging section and are presented in Table 10.

Table 13: Summary of evidence evaluating the accuracy of audiological tests to predict sequelae of cCMV infection

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Dhondt et al. 2023 ¹⁵¹ Prospective Belgium Follow-up: 4 years	With cCMV: 169 Symptoms at birth: 76 Long-term outcomes: 20	Hearing loss at birth	Vestibular and motor function at 4 years	Hearing loss at birth (present) Sens: 35% (95% Ct 15.4 to 59.2) Spec: 91.3% (95% Ct 85.5 to 95.3)
Abbreviations: cCMV, congenital cytomegalovirus; CI, confidence interval; Sens, sensitivity; Spec, specificity.				

Association studies

Two studies^{152, 153} were identified. Both were retrospective cohort studies. Sample sizes ranged from 44 to 61 infants with cCMV and follow-up ranged from 46 to 54 months (mean). Studies were conducted in Italy and the USA. Both studies^{152, 153} found that a higher proportion of infants with a 'refer' result on newborn hearing screening developed SNHL than those with a bilateral 'pass' result. One study¹⁵² found that worse hearing threshold at birth was a significant risk factor for progression of hearing loss ($p=0.02$).

Viral load

Systematic review evidence

No systematic reviews were identified.

Prognostic accuracy studies

Yamaguchi et al. (2022)¹⁵⁴ assessed the accuracy of viral load (at a threshold of >2950 copies/ml) for detecting developmental delay at 36 months. The test was poor at predicting later developmental delay, with a sensitivity of 57% (95% CI: 18 to 90). However, the test was able to accurately predict a lack of developmental problems, with a specificity of 84% (95% CI: 60 to 97) (Table 14). This was a small study ($n=26$) which was at an overall high risk of bias (see Appendix 3).

Table 14: Summary of evidence evaluating the accuracy of viral load to predict sequelae of cCMV infection

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Yamaguchi et al. 2022 ¹⁵⁴ Retrospective Japan Follow-up: 36 months	With cCMV: 39 Symptoms at birth: 21 Long-term outcomes: 7	Viral load	Developmental delay at 36 months	CMV DNA load (droplet digital PCR $\geq$ 2950 copies/ml) Sens: 57.1% (95% CI 18.4 to 90.1) Spec: 84.2% (95% CI 60.4 to 96.6)
Abbreviations: cCMV, congenital cytomegalovirus; CI, confidence interval; CMV, cytomegalovirus; DNA, deoxyribonucleic acid; PCR, polymerase chain reaction; Sens, sensitivity; Spec, specificity.				

Association studies

Eleven studies were identified.^{56, 77, 105, 115, 155-161} There were 8 prospective cohort studies, 2 retrospective cohort studies and 1 *post hoc* analysis. Sample sizes ranged from 9 to 387 infants with cCMV and follow-up ranged from 6 months to 6 years, where reported. Studies were conducted in Italy, Belgium, France, the USA, China and Japan. Four studies included children with symptomatic cCMV,^{155-157, 160} 3 studies included children with asymptomatic cCMV,^{77, 158, 159} and 4 studies included a mixed population.^{56, 105, 115, 161}

All 4 studies of infants with symptomatic cCMV found no significant association between viral load and SNHL severity. Of the 4 studies that included a mixed population, 3^{56,161,115} found an association between higher viral load and the increased occurrence of SNHL and neurodevelopmental sequelae. The fourth reported no significant association between viraemia and neurodevelopmental sequelae, although no raw data were presented.

All 3 studies^{77, 158, 159} that included only infants with asymptomatic cCMV found a statistically significant association between higher viral load at baseline and the development of late-onset sequelae.

Genotypes and immune cell markers

Systematic review evidence

No systematic reviews were identified.

Prognostic accuracy studies

One study¹⁶² showed that the sensitivity and specificity of a 16 gene signature to predict SNHL occurring at any point in a 3 year follow up period were 96% and 89%, respectively (Table 15). This study was deemed at a high risk of bias.

Table 15: Summary of evidence evaluating the accuracy of genetic tests to predict sequelae of cCMV infection

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Ouellette et al. 2020 ¹⁶² Prospective USA Follow-up: 36 months	With cCMV: 80 Symptoms at birth: 49 Long-term outcomes: 24	Genes	Late-onset SNHL at 3 years	16 gene classifier signature (present) Sens: 95.8% Spec: 89.3%
Abbreviations: cCMV, congenital cytomegalovirus; Sens, sensitivity; SNHL, sensorineural hearing loss; Spec, specificity.				

Association studies

Nine studies were identified.^{99, 143, 144, 163-168} There were 6 prospective cohort studies and 3 retrospective cohort studies. Sample sizes ranged from 12 to 131 infants with cCMV and follow-up ranged from 6 months to 6 years of age. Studies were conducted in the Netherlands, Spain, Poland, the USA and China. Two studies included children with symptomatic cCMV,^{163, 164} and 7 studies included a mixed population.^{99, 143, 144, 165-168} A range of different genotypes were assessed in these studies and 3 of the studies assessed immune cell responses. Most of the genotype studies were limited by low patient numbers and missing data. Results were mixed, with many studies reporting no significant association between the genotypes assessed and long-term sequelae; those that reported an association stated that further studies are required. Studies that assessed immune cell markers reported an association between lower levels of T cells^{99, 166} or B cells¹⁴⁴ and long-term sequelae.

CNS involvement (CSF)

Systematic review evidence

No systematic reviews were identified.

Prognostic accuracy studies

Alarcon et al. (2013)⁷⁵ evaluated the prognostic accuracy of cerebrospinal fluid (CSF) beta 2-m concentration above a threshold of 7.9mg/L. Used alone, this test had a sensitivity of 69% at a specificity of 100%. When used in combination with microcephaly, accuracy improved to a sensitivity of 82% at a specificity of 100% (see Table 16). This was a small study (n=26) with concerns about applicability and a high overall risk of bias (see Appendix 3).

Results for a combination of CSF beta2-m concentration and neuroimaging have been described in the neuroimaging section and are summarised in Table 10.

Table 16: Summary of evidence evaluating the accuracy of CSF tests to predict sequelae of cCMV infection

Study properties	Sample size	Baseline tests performed	Long-term outcomes reported	Baseline test accuracy (threshold for positive test)
Alarcon et al. 2013 ⁷⁵ Prospective Spain Follow-up: mean 8.7 years	With cCMV: 26 Symptoms at birth: 26 Long-term outcomes: 15	Mixed (CSF, microcephaly, neuroimaging)	Neurodevelopmental delay (in terms of motor function, cognition, behaviour, hearing, vision, and epilepsy) at mean 8.7 years	CSF concentration (>7.9mg/L) Sens: 69% Spec: 100% CSF concentration (>7.9 mg/L) OR adjusted microcephaly (threshold unclear) Sens: 82% Spec: 100%
Abbreviations: cCMV, congenital cytomegalovirus; CSF, cerebrospinal fluid; Sens, sensitivity; Spec, specificity.				

Association studies

Two studies that assessed the association between CNS involvement (detection of CMV DNA in cerebrospinal fluid by PCR as a marker of CNS involvement) were identified.^{169, 170} One was a prospective cohort study and 1 a retrospective cohort study. Sample sizes ranged from 136 to 168 infants with cCMV and follow-up ranged from 4-8 weeks to 6 months. Studies were conducted in Poland and Spain. One study¹⁶⁹ found that CNS involvement was associated with neurological abnormalities such as seizures at 4-8 weeks follow up, but the study with longer-term follow-up¹⁷⁰ found no association between a positive result and neurological sequelae or SNHL at 6 months of age. Both studies concluded that the test may not be a useful prognostic marker in clinical practice.

Combination tests

Systematic review evidence

No systematic reviews were identified.

Prognostic accuracy studies

Four prognostic accuracy studies were identified that used combination tests.^{75, 82, 85, 89} In 3 of these studies,^{82, 85, 89} all the combination tests involved neuroimaging, and in the other study⁷⁵ 1 combination test involved neuroimaging, whilst the other covered CSF and symptoms. The combination tests involving neuroimaging have been presented in Table 10, and discussed in the appropriate sections. The CSF/symptom combination test has been presented in Table 16.

Association studies

Many studies looked at the associations of more than 1 prognostic variable with sequelae, but analyses were usually carried out for each prognostic variable independently, and not in combination. Only 2 studies looked at the likelihood of sequelae in terms of the interactions between more than 1 variable.^{76, 117}

Bilavsky et al. (2015)⁷⁶ evaluated the association between hearing sequelae and the combination of symptomatic status and presence of lenticulostriate vasculopathy (LSV). Broadly, the study found that having LSV with no treatment leads to more long-term hearing loss than having LSV *with* treatment ($p < 0.001$) or being asymptomatic ($p = 0.008$). Although the study was moderately sized, with a reasonably long follow-up, the study was limited by the unknown LSV status of the asymptomatic group, who may not have had LSV status measured.

Demmler-Harrison et al. (2020)¹¹⁷ looked at the association of hearing sequelae with the combination of symptomatic status and type of maternal CMV infection. Newborns were

more likely to develop hearing loss if symptomatic and the mother had a primary infection than if they were symptomatic and the mother had a non-primary infection. They were also more likely to develop hearing loss if asymptomatic and the mother had a primary infection than asymptomatic and the mother had a non-primary infection. However, these findings were not statistically analysed.

Summary of Findings Relevant to Criterion 7

Criterion Not Met

The review included 94 studies relevant to this question; 4 systematic reviews, 19 prognostic accuracy studies and 71 studies assessing the association between test results and/or antenatal and/or newborn characteristics and long-term health consequences. A range of tests and characteristics were assessed; however, much of the evidence is of limited quality.

Overall, the results indicate that the most reliable characteristics for predicting long-term sequelae in infants with cCMV are the presence of symptoms at birth and timing of maternal CMV infection. Newborns with more severe cCMV symptoms are likely to be offered antiviral treatment under current guidelines.⁸ Our results indicate that the most useful test for predicting long-term hearing and neurodevelopmental sequelae is likely to be MRI. Results of studies assessing viral load to predict long-term hearing sequelae are promising, particularly in newborns with asymptomatic cCMV, however, further research is required to confirm these findings. As there appear to be multiple factors that affect the risk of long-term sequelae, research to generate a multifactorial risk model for long-term harms in newborns with cCMV may be useful.

A summary is presented below for each test/characteristic assessed.

Neuroimaging (MRI and ultrasound)

Fifteen prognostic accuracy studies and 7 other cohort studies assessed the accuracy of neuroimaging to predict long-term sequelae in infants with cCMV. In general, neuroimaging tests demonstrated reasonable accuracy in predicting neurodevelopmental delay, epilepsy and hearing outcomes. The most accurate tests were MRI or US (and sometimes CT) using 'Alarcon' scoring. In general, studies were small, and most studies had either a high or unclear risk of bias.

Studies assessing the association between US abnormalities and long-term sequelae had conflicting results; 3 studies reported an association between US abnormalities and hearing and/or neurodevelopmental sequelae, whilst 2 studies reported a numerically higher

prevalence of adverse outcomes in infants without abnormalities detected on US. Overall, MRI appears to be a more useful modality than US for predicting long-term sequelae.

Antenatal characteristics (timing of maternal infection)

One good quality systematic review pooled evidence from 6 studies and found that primary maternal infection in the first trimester led to a much higher prevalence of adverse long-term sequelae than primary maternal infection in the second or third trimesters. However, we identified an additional 2 prognostic accuracy studies (of low quality) that did not demonstrate adequate accuracy of timing of maternal infection for predicting adverse hearing outcomes. Our review also included 13 cohort studies. Twelve of the studies demonstrated that infection in the first trimester significantly increased the risk of long-term neurological or hearing sequelae (5 of which reported a statistically significant association). Only 1 small study showed a greater prevalence of hearing sequelae in those with later infection.

Whilst some of these studies were limited by low patient numbers, missing data and relatively short-term follow-up, the finding that first trimester infection is associated with more severe long-term hearing or neurodevelopmental sequelae was generally consistent between studies and is likely to be valid.

Antenatal characteristics (type of maternal infection)

One good quality systematic review pooled evidence from 8 studies and found that type of maternal CMV infection (primary or non-primary infection) did not influence the risk of developing SNHL or other neurological outcomes at follow-up. We identified an additional 6 cohort studies. Their results were inconsistent; 2 studies reported a statistically significant higher proportion of infants born to mothers with non-primary infection having long-term hearing and/or neurodevelopmental sequelae, 2 studies suggested the opposite direction of effect, and 2 studies reported no correlation between type of maternal infection and long-term sequelae. Confidence in these findings is limited by the small sample size in most studies and conflicting results. Overall, type of maternal CMV infection has not been shown to be a reliable predictor of long-term sequelae.

Antenatal characteristics (gestational age)

Two cohort studies assessed the association between gestational age and long-term sequelae. As might be expected, both studies reported that more infants who were born premature developed adverse outcomes.

Symptoms at birth

One good quality systematic review assessed neurodevelopmental outcomes of children who had presented with asymptomatic cCMV at birth. Most of the included studies that compared symptomatic and asymptomatic infants found that adverse neurodevelopmental outcomes were less common among children with asymptomatic cCMV at birth.

We identified an additional 30 cohort studies. Twenty-eight of the studies compared the outcomes of children with symptomatic cCMV at birth with those with asymptomatic cCMV at birth; all 28 studies showed that symptoms at birth were associated with an increased risk of long-term sequelae (9 of which reported a statistically significant association). Whilst some studies were limited by small sample sizes and/or short-term follow-up, the evidence was largely consistent, suggesting that there is an association between symptomatic cCMV at birth and poorer long-term outcomes, as might be expected.

Audiological screening result at birth

Two prognostic accuracy studies assessed the accuracy of audiological screening at birth to predict hearing sequelae in infants with cCMV. One study (which was at a high risk of bias) did not demonstrate adequate accuracy of audiological screening at birth for predicting adverse hearing outcomes. The other study demonstrated a high level of accuracy for predicting SNHL or psychomotor sequelae. Our review also included 2 other small cohort studies, both studies found that a higher proportion of infants with a 'refer' result on newborn hearing screening developed SNHL than those with a bilateral 'pass' result.

Viral load

One very small prognostic accuracy study (at a high risk of bias) assessed the accuracy of viral load for detecting developmental delay; the test was poor at predicting later developmental delay.

Our review also included 11 other cohort studies. Four of those studies included infants with symptomatic cCMV; all 4 studies found a trend towards lower baseline viral load and better hearing outcomes, but the findings did not reach statistical significance. Three studies included infants with asymptomatic cCMV; all 3 studies found a statistically significant association between higher viral load at baseline and the development of late-onset sequelae (primarily SNHL). Four studies included a mixed population of symptomatic and asymptomatic infants; 3 of the studies found an association between higher viral load and the development of hearing and/or neurodevelopmental sequelae.

Overall, viral load appears to be a promising test for predicting long-term hearing (and possibly neurodevelopmental) sequelae, particularly amongst asymptomatic infants.

Genotypes and immune cell markers

One prognostic accuracy study (at a high risk of bias) reported a good level of accuracy of a 16 gene signature to predict SNHL. Our review also included 9 other cohort studies; 6 studies assessed a range of different genotypes and 3 studies assessed immune cell markers. Most of the genotype studies were limited by low patient numbers and missing data. Results were mixed, with many studies reporting no significant association between the genotypes assessed and long-term sequelae. Studies that assessed immune cell markers reported an association between lower levels of T cells or B cells and long-term sequelae. Study authors drew cautious conclusions and/or recommended further research. No firm conclusions can be drawn from any of the studies.

CNS involvement (cerebrospinal fluid)

One very small prognostic accuracy study (at a high risk of bias) assessed the accuracy of CNS involvement, assessed using cerebrospinal fluid, for detecting neurodevelopmental delay; results indicated reasonable predictive accuracy, particularly when used in combination with microcephaly. Our review also included 2 cohort studies; both studies concluded that the test may not be a useful prognostic marker in clinical practice.

Criteria 11 & 13: There should be evidence from high quality randomised controlled trials that the screening programme is effective in reducing mortality or morbidity. The benefit gained by individuals from the screening programme should outweigh any harms for example from overdiagnosis, overtreatment, false positives, false reassurance, uncertain findings and complications.

Can a newborn screening programme for cCMV reduce the burden of cCMV-related morbidity and mortality?

Previous work

The most recent UK NSC evidence map⁷ identified only 2 cohort studies that investigated the impact of screening on morbidity.^{148, 171} Both studies were identified by our searches and are included in our assessment. The previous review concluded that the available evidence provides only limited new information and there was close to no evidence on the benefits of early treatment or interventions for cCMV compared to late treatment after the presentation of symptoms.

The earlier UK NSC evidence review from 2017¹⁸ identified 1 study it judged to be relevant.¹⁷² This study examined 178 newborns with symptomatic cCMV and found that newborns identified by screening were less likely to have long-term disease sequelae than newborns identified through routine referral (28% versus 51%, $p=0.003$). However, the review concluded that this study could not demonstrate whether this difference was due to the direct effect of screening. This study did not meet the inclusion criteria for the current review and so is not considered further.

Description of the current evidence

The aim was to identify all systematic reviews and randomised or non-randomised studies that reported evidence on an evaluation of screening for cCMV in practice. To be eligible for inclusion, studies had to report outcomes in line with the protocol, specifically: symptoms at birth (e.g. outcome of hearing test or results of neuroimaging); use of antiviral therapy; or long-term sequelae (e.g. long-term hearing loss or neurodevelopmental issues). Studies reporting acceptability of screening to physicians or parents were also included.

Systematic reviews

The review identified 2 systematic reviews that investigated the value of newborn cCMV screening.

Hilditch et al (2018)¹⁷³ included 2 studies of universal screening and 4 of targeted screening. All 6 studies were eligible for our review and were found in our searches. The Hilditch review did not perform any quantitative or qualitative synthesis of the studies, so is of limited value for this assessment. We therefore chose to examine the included studies as part of our review in the sections below. The Hilditch review concluded that newborn screening will identify asymptomatic newborns with cCMV that would otherwise have gone undiagnosed. They also stated that those newborns could benefit from antiviral therapy, but that treatment remains controversial. There may be other benefits of early diagnosis, including early intervention and hearing augmentation.

Pollick et al (2024)¹⁷⁴ included 12 studies: 3 of universal screening, 8 of targeted screening and 1 of pre-natal screening. Only 4 of the studies met the inclusion criteria for this review (and are included in our evaluation below), limiting the relevance of the Pollick review. The Pollick review found that universal screening will identify all cCMV cases, but is at risk of more false positives, and may be more costly and logistically burdensome. Hearing-targeted screening may be more cost-efficient but will miss asymptomatic cases and newborns with late-onset hearing loss. It concluded that continued research that examines the potential benefits and disadvantages of screening and how this may impact detection of hearing and other neurodevelopmental outcomes is required.

Individual studies

Of the 422 papers considered at full text screening 36 studies met the inclusion criteria for this question. None of the included studies were randomised controlled trials (RCTs) and none were non-randomised studies comparing a screened population with a non-screened population. Therefore, strictly, there is no evidence to meet this criterion. However, the non-comparative studies which were identified are reported here.

There were 14 studies that evaluated universal screening (i.e. screening of all newborns regardless of symptoms, see Table 17), 12 studies evaluated targeted screening in newborns who had failed a newborn hearing screen (see Table 19), and 10 studies evaluated targeted screening in newborns who showed signs of cCMV at birth (including hearing loss) (see Table 21).

Presentation of results was limited in all studies, largely restricted to presenting numbers of cCMV diagnoses, and numbers with and without symptoms at birth. In the subset of studies

that reported long-term follow-up, reporting was mostly limited to numbers receiving antiviral therapy and numbers with long-term sequelae. Given the absence of comparative studies and the limitations of the results presented it was decided not to perform a quality assessment or risk of bias assessment of the included studies. However, we note the following general quality and risk of bias concerns across the studies:

- as there is no comparative evidence it is not possible to determine whether symptoms at birth, or long-term sequelae, are caused by CMV infection, or are unrelated to infection;
- it is not possible to evaluate which symptoms or long-term sequelae would have been identified, managed and treated in the absence of cCMV screening;
- as management protocols were not usually reported, there may have been differences in how symptomatic infection, asymptomatic infection and uninfected newborns were managed and followed-up;
- as use of antiviral therapy was not randomised or controlled it is not possible to determine whether treatment itself prevented or moderated long-term sequelae, or whether sequelae would have occurred without treatment;
- follow-up duration, where reported, was variable across studies; and
- studies were performed in a wide variety of locations with differing levels of cCMV prevalence. Therefore, results of the studies may not be applicable to the UK context.

Given the limitations in the evidence base, results presented in this report focus on summarising the overall conclusions and findings of the studies, and summarising event rates (such as frequency of long-term sequelae) using random-effects meta-analysis.

Universal screening

The review identified 15 studies evaluating the performance of universal newborn screening for cCMV, using either saliva, DBS or urine testing. The characteristics and general conclusions of these studies are summarised in Table 17. Eight studies used primarily saliva testing to screen for cCMV, 3 used urine testing, 3 used DBS and 1 used a mix of tests.

Only 1 of these studies (Puhakka 2019)¹⁷⁵ included any comparative evidence. This study compared long-term outcomes in children with symptomatic infection, asymptomatic infection and no infection after saliva-based screening. It compared 40 cases of cCMV in Finland with 54 matched controls without cCMV. It found no difference in neurodevelopmental outcomes or in hearing between cases and controls after 18 months of follow-up (for example, 66% of cases and 76% of controls passed an audiological exam, $p=0.572$). It concluded that screening was unwarranted in Finland because of the low prevalence of cCMV infection.

Based on the reported conclusions of the 15 included studies, 4 were broadly in favour of implementing universal screening,^{54, 148, 176, 177} 2 did not favour screening,^{175, 178} 2 suggested more research is required or were neutral,^{36, 179} 3 concluded that screening was feasible without stating if it should be used^{171, 180, 181} and the 4 remaining studies did not conclude directly on whether screening should be used.¹⁸²⁻¹⁸⁵ Therefore, there seems to be little consistency across the included studies as to whether universal screening should be implemented.

Arguments made in favour of universal screening included (see Table 17):

- most cCMV cases would not be diagnosed without universal screening;
- it identifies newborns with late neurological sequelae that would otherwise be missed;
- it identifies newborns who would benefit from ongoing audiologic surveillance;
- antiviral treatment of symptomatic cCMV leads to normal development or mild long-term sequelae; and
- it predicts late-onset SNHL and neurodevelopmental disorders.

Arguments made against current implementation of universal screening included (see Table 17):

- outcomes at 18 months of age in Finland did not differ between infected infants and healthy control infants;
- targeted cCMV screening among neonates who fail a hearing screen was preferable to universal screening;
- most newborns with cCMV had a normal cranial ultrasound; and
- universal urine screening likely identifies subclinical symptomatic cCMV.

Most studies concluded that universal screening, whichever test was used, was feasible and practical to implement. One study¹⁸³ reported that pooled saliva screening (in pools of 10 newborns) was beneficial and reasonable when compared to individual sample screening.

The included studies reported numbers of newborns who were diagnosed cCMV, numbers who had symptoms (including hearing loss and neuroimaging abnormalities), numbers given antiviral therapy, and numbers with long-term sequelae (hearing loss or neurodevelopmental issues). Table 18 summarises these data across the studies by presenting results from a meta-analysis of all studies reporting data on each outcome. This gives an “average” number of events that might be observed within a universal screening programme. Given the rarity of most events, results are presented as expected numbers of events per 10,000

babies screened, per 10,000 with cCMV, and, for antiviral therapy and long-term sequelae, per 10,000 with and without symptoms at birth. As this meta-analysis pools results across 3 types of screening (saliva, DBS and urine) and across a wide range of countries and settings, results are very heterogeneous, so should only be interpreted as a broad view of the consequences of universal screening.

The meta-analysis results in Table 18 demonstrate that, as expected, cCMV infection is rare, occurring in only 0.4% of newborns across the included universal screening studies. Symptomatic cCMV, and hence long-term sequelae, is very rare, at less than 0.1% of all newborns. Only 23% of newborns with cCMV displayed symptoms at birth, and cranial abnormalities may be more common than hearing loss. This concurs with claims made by the studies that universal screening will detect cCMV cases that will be missed by any other form of management. Antiviral therapy was only given to a minority of newborns with confirmed cCMV (around 11%), and around 18% of those with cCMV developed long-term sequelae.

Antiviral therapy was never given to children who were classified as being asymptomatic at birth, and only 6% of newborns with asymptomatic cCMV went on to develop long-term sequelae. This concurs with the findings of Puhakka (2019)¹⁷⁵ that it is unclear whether these findings can be distinguished from hearing loss and neurodevelopmental issues that are unrelated to cCMV. Among newborns with symptomatic cCMV, 69% received antiviral therapy, so there will still be some children with symptoms who do not receive therapy. Around 56% of these newborns will develop long-term sequelae.

Table 17: Details of 15 studies relating to evaluation of universal cCMV screening performance

Study details	Screening test	Confirmation test	Follow-up beyond infancy	Number screened, Number with cCMV (Prevalence)	Results presented	Study conclusions
Universal screening (15 studies)						
Dunn 2025 ¹⁷⁷ Canada	DBS	Urine	No	551,034 601 with cCMV	Symptoms at birth Antiviral treatment	A lower than expected prevalence of cCMV. Screening identified many children who would otherwise not have been diagnosed and who would benefit from ongoing audiologic surveillance
Kaye 2024 ¹⁷⁹ USA	DBS	Urine	No	60115 184 with cCMV	Symptoms at birth Antiviral treatment	Further evaluation of the newborn screening using dried blood spot sensitivity is warranted
Papaevangelou 2017 ¹⁸⁴ Greece	DBS	Not stated	Yes (5 years)	2149 10 with cCMV	Symptoms up to 5 years	<i>No conclusions related to screening reported</i>
Barkai 2014 ³⁶ Israel	Saliva	Urine	Yes	9845 48 with cCMV	Hearing tests at birth and up to 18 months Antiviral treatment	Saliva is a feasible and effective means of universal neonatal CMV screening that can detect affected infants who might benefit from treatment and follow-up. The long-term clinical significance of screening and its cost-effectiveness are yet to be determined
Chiereghin 2022 ¹⁷⁶ Italy	Saliva	Urine or Saliva	Yes	3151 21 with cCMV	Viral load Symptoms at birth Antiviral treatment	Without universal neonatal CMV screening, some infected infants who develop late neurological sequelae may not be recognized

Study details	Screening test	Confirmation test	Follow-up beyond infancy	Number screened, Number with cCMV (Prevalence)	Results presented	Study conclusions
Demortier 2021 ¹⁸² France (Reunion/Mayotte)	Saliva	Urine	No	1042 16 with cCMV	Symptoms at birth Survey of healthcare workers (with limited response)	Parents found screening to be acceptable
Letamendia-Richard 2022 ⁵⁴ France	Saliva	Urine	No	15356 63 with cCMV	Viral load Symptoms at birth Survey of healthcare workers	Universal screening for cCMV on saliva samples is feasible and highly acceptable 62% would probably not have been diagnosed without universal screening
Merav 2024 ¹⁸³ Israel	Saliva (pooled samples)	Saliva then urine	No	15805 54 with cCMV	Performance of pooled versus individual screening	Wide feasibility and benefits of pooled saliva testing
Puhakka 2019 ¹⁷⁵ Finland	Saliva	Urine	Yes (up to 18 months)	19868 40 with cCMV	Viral load Neurological and hearing symptoms at 18 months Comparison with uninfected control children	Outcomes at 18 months of age did not differ between the infected infants and healthy control infants. With such a low burden in Finland, universal newborn screening for cCMV seems unwarranted
Yamamoto 2020 ¹⁷⁸ Brazil	Saliva	Urine	Yes (median 36 months)	11900 68 with cCMV	Hearing loss at birth Hearing loss at long-term follow-up	Integrating targeted cCMV screening among neonates who fail a hearing screen could be a reasonable, cost-effective strategy to identify newborns with early-onset cCMV-related hearing loss
Zeytinoglu 2018 ¹⁸⁵ Turkey (paper in Turkish)	Saliva	Urine or blood	No	1000 2 with cCMV	Symptoms at birth	<i>No conclusions related to screening reported</i>
Kruc 2024 ¹⁸⁰ USA	Saliva or DBS	Urine	Yes (up to 5 years)	23644 87 with cCMV	Symptoms at birth	Among infants with cCMV identified in a universal screening study, the majority

Study details	Screening test	Confirmation test	Follow-up beyond infancy	Number screened, Number with cCMV (Prevalence)	Results presented	Study conclusions
						had a normal cranial ultrasound
Koyano 2017 ¹⁴⁸ Japan	Urine	Not stated	Yes (Over 2 years)	23405 72 with cCMV	Viral load Symptoms at birth Antiviral treatment Symptoms at follow-up	Newborn urine–filter paper screening of congenital CMV infection is warranted
Yamada 2020 ¹⁷¹ Japan	Urine	Urine	Yes (duration unreported)	11736 56 with cCMV	Symptoms at birth Neurological assessments on long-term follow-up	Universal urine screening and antiviral therapy for neonates with symptomatic cCMV may decrease neurological impairments, because 58% of the treated infants had normal development or mild sequelae. Universal urine screening likely identifies subclinical symptomatic cCMV
Yamaguchi 2017 ¹⁸¹ Japan	Urine	Not stated	Yes (6 months)	23368 60 with cCMV	Viral load MRI at birth MRI at follow-up	Urinary CMV load may effectively predict the incidence of late-onset SNHL and neurodevelopmental disorders
Abbreviations: cCMV, congenital cytomegalovirus; CMV, cytomegalovirus; DBS, dried blood spots; MRI, magnetic resonance imaging; SNHL, sensorineural hearing loss.						

Table 18: Meta-analysis of event rates in 15 studies of universal screening

Event	Number of studies with data	Predicted rate per 10,000 screened newborns (95% confidence interval)	Predicted rate per 10,000 newborns with cCMV (95% confidence interval)
Diagnosis of cCMV	14	37 (38 to 50)	x
Has symptoms at birth (all symptoms)	13	8 (4 to 15)	2293 (1379 to 3563)
Has hearing loss at birth	9	3 (1 to 5)	896 (601 to 1304)
Has cranial MRI/US abnormalities at birth	7	5 (2 to 10)	1676 (724 to 3419)
Receives antiviral treatment	10	4 (2 to 6)	1092 (682 to 1702)
Has long-term sequelae	8	7 (4 to 11)	1757 (970 to 2975)
Has long-term hearing loss	8	4 (3 to 6)	1225 (923 to 1609)
Has long-term neurological or developmental issues	4	6 (3 to 12)	1856 (694 to 4107)
	Number of studies with data	Predicted rate per 10,000 newborns with symptomatic cCMV (95% confidence interval)	Predicted rate per 10,000 newborns with asymptomatic cCMV (95% confidence interval)
Receives antiviral treatment	6	6920 (5278 to 8187)	Zero
Has long-term sequelae	5	5558 (2682 to 8101)	585 (335 to 1002)
Has long-term hearing loss	6	4085 (2340 to 6091)	550 (334 to 891)
Has long-term neurological or developmental issues	3	3495 (1788 to 5701)	380 (158 to 886)
Abbreviations: cCMV, congenital cytomegalovirus; MRI, magnetic resonance imaging; US, ultrasound.			

Targeted screening based on failure of newborn hearing screen

The review identified 12 studies evaluating the performance of targeted newborn screening for cCMV, among newborns who failed a newborn hearing screen (or equivalent hearing test), using either saliva, DBS or urine testing. The properties and general conclusions of these studies are summarised in Table 19. Six studies used saliva testing to screen for cCMV, 4 used saliva or urine testing, 1 used DBS and 1 used a mix of tests.

Based on the reported conclusions of the 12 included studies, 3 were broadly in favour of implementing targeted screening.¹⁸⁶⁻¹⁸⁸ Two did not favour screening,^{189, 190} although 1 of these was specifically in newborns in a neonatal intensive care unit.¹⁸⁹ Four suggested more research is required or were neutral.¹⁹¹⁻¹⁹⁴ Three concluded that targeted screening was feasible without stating if it should be used.¹⁹⁵⁻¹⁹⁷ Hence the studies seem broadly to suggest

that targeted screening may be valuable, but some further evaluation is needed, such as clearly demonstrating its cost-effectiveness.

Arguments made in favour of targeted screening included (see Table 19):

- it identifies the majority of infants with cCMV-related SNHL at birth;
- it contributes to the early detection of infants born with cCMV-related SNHL who could benefit from treatment; and
- it ensures appropriate clinical, neurodevelopmental and audiological follow-up.

Arguments made against current implementation of targeted screening included (see Table 19):

- Many infants with CMV-related SNHL will not be detected using a newborn hearing screen; and
- it is problematic to perform in NICUs due to timing difficulties.

Most studies concluded that targeted screening, whichever test was used, was feasible and practical to implement.

As for universal screening, the included studies reported numbers of newborns who were diagnosed with cCMV, numbers who had symptoms, and numbers receiving antiviral therapy. None of the studies, however, reported any long-term follow-up data. Table 20 summarises these data across the studies by presenting results from a meta-analysis of all studies, as in Table 18 for universal screening.

The meta-analysis results in Table 20 demonstrate that cCMV infection is uncommon even in newborns who fail a hearing screen, occurring in 1.5% of newborns across the included studies. About 75% of newborns with confirmed cCMV displayed symptoms at birth (beyond failure of the hearing test in itself), although it is possible that some symptoms were diagnosed as a consequence of testing positive for cCMV. Antiviral therapy was only given to about half of newborns with cCMV (54%), representing just under 1% of those screened. It is perhaps surprising that antiviral therapy was given so infrequently, despite both a diagnosis of cCMV and failure of a hearing test; this may indicate issues with delayed diagnosis meaning treatment cannot be offered. No studies reported any long-term outcome data.

Table 19: Details of 12 studies relating to evaluation of hearing loss-targeted cCMV screening performance

Study details	Screening test	Confirmation test	Follow-up beyond infancy	Number screened, Number with cCMV	Results presented	Study conclusions
Screening in infants who fail a newborn hearing test (12 studies)						
Chung 2022 ¹⁸⁷ Netherlands	DBS	Not stated	No	1374 59 with cCMV	Hearing loss and neuroimaging in infancy	Newborns who fail newborn hearing screening should be tested for CMV to ensure appropriate clinical, neurodevelopmental and audiological follow-up
Diener 2017 ¹⁹⁵ USA	Mixed	None	No	314 21 with cCMV	Hearing loss at birth Compliance with screening	Incorporating CMV screening into an established NHS program is a viable option
Ari-Even Roth 2017 ¹⁸⁶ Israel	Saliva	Urine	No	187 4 with cCMV	Hearing loss at birth	Targeted cCMV screening contributed to the early detection of infants born with cCMV-related isolated SNHL who could potentially benefit from antiviral treatment
Beswick 2021 ¹⁹¹ Australia	Saliva	Any	No	489 4 with cCMV	Stakeholder interviews	cCMV screening alongside hearing tests is achievable, a number of barriers need to be addressed
Fourgeaud 2022 ¹⁹⁶ France	Saliva	Saliva + blood	No	236 2 with cCMV	Hearing at birth Antiviral use	Targeted cCMV screening is feasible
Kadambari 2015 ¹⁹⁴ UK	Saliva	Not stated	No	203 2 with cCMV	Survey of screeners	Rapid diagnostic testing for cCMV needs a more detailed clinical and cost-effectiveness analysis
Ronner 2022 ¹⁸⁸ USA	Saliva	Not stated	Yes (duration not stated)	891 8 with cCMV	Hearing at birth Antiviral use Hearing at follow-up	Implementation of a targeted screening program for CMV in children who do not pass the NHS resulted in significantly higher rates of CMV testing and earlier referral to infectious disease
Fowler 2017 ¹⁹² USA	Saliva and D BS	Not stated	No	Unclear (31 with cCMV failed hearing test)	Hearing tests at birth	A targeted CMV approach that tests newborns who fail their NHS identified the majority of infants with CMV-related SNHL at birth. However,

Study details	Screening test	Confirmation test	Follow-up beyond infancy	Number screened, Number with cCMV	Results presented	Study conclusions
						43% of the infants with CMV-related SNHL in the neonatal period and cCMV infants who are at risk for late-onset SNHL were not identified by NHS
Medoro 2021 ¹⁸⁹ USA (NICUs only)	Urine or saliva	Urine	No	1498 3 with cCMV	Hearing tests at birth	Targeted CMV testing for failed hearing screen in the NICU is problematic
Rawlinson 2018 ¹⁹⁷ Australia	Urine or saliva	Not stated	No	336 19 with cCMV	Hearing test and MRI at birth	Timely CMV testing for newborns who fail NHS and who have permanent hearing loss confirmed, has been shown to be clinically practical
Vancor 2019 ¹⁹⁰ USA	Urine or saliva	Not stated	No	171 2 with cCMV	Symptoms at birth	Consideration should be given to other strategies for identifying children at risk of hearing loss as a result of cCMV infection
Williams 2014 ¹⁹⁸ UK	Urine or saliva	Not stated	No	407 6 with cCMV	Symptoms and neuroimaging at birth	Targeted salivary screening for cCMV within the NHS is feasible, acceptable and detects infants with cCMV-related SNHL who could benefit from early treatment
Abbreviations: cCMV, congenital cytomegalovirus; CMV, cytomegalovirus; DBS, dried blood spots; MRI, magnetic resonance imaging; NHS, newborn hearing screening; NICU, neonatal intensive care unit; SNHL, sensorineural hearing loss.						

Table 20: Meta-analysis of event rates in 12 studies of hearing-loss targeted screening

Event	Number of studies with data	Predicted rate per 10,000 screened newborns (95% confidence interval)	Predicted rate per 10,000 newborns with cCMV (95% confidence interval)
Diagnosis of cCMV	12	147 (80 to 269)	x
Has symptoms at birth (all symptoms)	10	118 (69 to 202)	7548 (6825 to 8400)
Has cranial MRI/US abnormalities at birth	Insufficient data		
Receives antiviral treatment	9	80 (53 to 118)	5360 (3450 to 7170)
Has long-term sequelae	Insufficient data		
Abbreviations: cCMV, congenital cytomegalovirus; MRI, magnetic resonance imaging; US, ultrasound.			

Targeted screening based on signs suggestive of cCMV infection

The review identified 10 studies evaluating the performance of newborn screening for cCMV, where screening was based on signs consistent with possible cCMV infection. Included signs varied across studies. Failure of hearing tests was consistently included, but other possible signs included known maternal CMV infection, microcephaly, baby being small for gestational age, among others. The properties and general conclusions of these studies are summarised in Table 21. One study used saliva testing to screen for cCMV, 5 used urine testing, 3 used saliva or urine testing, 1 used DBS testing.

Based on the reported conclusions of the 10 included studies, 3 were broadly in favour of implementing targeted screening.¹⁹⁹⁻²⁰¹ One did not favour targeted screening.¹⁵² Two were more neutral.^{202, 203} Two concluded that it was feasible without stating if it should be used.^{101, 204} Two did not report conclusions on the value of screening.^{205, 206} Hence the studies seem broadly to suggest that targeted screening may be valuable, but findings were not consistent.

Arguments made in favour of targeted screening included (see Table 21):

- it helps lead to the diagnosis of cCMV infection that may otherwise be overlooked; and
- it allows for early recognition of asymptomatic cases with hearing loss.

Arguments made against current implementation of targeted screening included (see Table 21):

- it may overlook cCMV infection that is asymptomatic at birth; and

- a universal newborn screening programme should be adopted instead of targeted screening.

Most studies concluded that targeted screening, whichever test was used, was feasible and practical to implement.

As for universal screening, the included studies reported numbers of newborns who were diagnosed with cCMV, numbers who had symptoms, numbers receiving antiviral therapy, and numbers with long-term sequelae. Table 22 summarises these data across the studies by presenting results from a meta-analysis of all studies, as in Table 18 for universal screening.

The meta-analysis results in Table 22 demonstrate that cCMV occurred in 1.5% of newborns across the included studies. About 54% of newborns with confirmed cCMV (and 0.7% of newborns screened) were judged to be symptomatic at birth, and only 28% of newborns with confirmed cCMV received antiviral therapy, despite those babies having signs and symptoms consistent with infection. This is notably lower than the proportion receiving therapy when screening is based on failure of a hearing test only (Table 20), suggesting that expanding targeted screening beyond failure of a hearing test may be picking up more asymptomatic cases who do not receive antiviral therapy. About 18% of newborns with cCMV went on to have long-term sequelae. This is very similar to the proportion in the universal screening studies (Table 18).

Table 21: Details of 10 studies relating to evaluation of symptom targeted cCMV screening performance

Study details	Screening test	Confirmation test	Follow-up beyond infancy	Number screened, Number with cCMV	Results presented	Study conclusions
Screening in infants with symptoms suggestive of cCMV infection (10 studies)						
Lu 2018 ²⁰⁴ Taiwan	DBS	Urine or saliva	Yes (3 months)	1716 3 with cCMV	Hearing tests in infancy	Hearing, genetic, and CMV screenings are feasible
Webb 2024 ²⁰⁶ Australia	Saliva	Urine or saliva	Yes (up to 6 months)	96 1 with cCMV	Parent survey	Parent experiences of targeted cCMV screening were positive
Forli 2024 ¹⁵² Sweden	Urine	Not stated	Yes (3-12 years)	1615 25 with cCMV	Hearing loss at birth Symptoms at birth Hearing at follow-up	cCMV screening in newborns allows the detection of many cases; but many are still missed. Important to adopt a universal newborn screening program or a program extended to newborns at higher risk
Fujii 2017 ²⁰² Japan	Urine	Urine	None	575 20 with cCMV	Comparison with universal screening Symptoms at birth	Urine test identifies cCMV cases efficiently. Risk-based and universal newborn screening strategies for cCMV infection each have advantages
Palma 2021 ²⁰¹ Italy	Urine	Not stated	None	Not stated 20 with cCMV	Hearing loss at birth	Coupling cCMV screening and hearing screening is worthwhile as it allows early recognition of asymptomatic cases with hearing loss
Tan 2023 ²⁰⁵ Singapore	Urine	Not stated	Yes (2-3 months)	1220 11 with cCMV	Hearing loss	<i>No conclusions related to screening reported</i>
Zhang 2023 ²⁰³ Japan	Urine	Not stated	Unclear	102 3 with cCMV	Symptoms at birth Antiviral treatment	May overlook cCMV infection that is asymptomatic at birth; however, it is feasible and

Study details	Screening test	Confirmation test	Follow-up beyond infancy	Number screened, Number with cCMV	Results presented	Study conclusions
						helps lead to the diagnosis of cCMV infection that may otherwise be overlooked
Masarweh 2021 ¹⁹⁹ Israel	Urine or saliva	Not stated	Unclear	693 18 with cCMV	Hearing loss at birth Antiviral treatment	This examination policy can detect nearly all newborns with congenital CMV infection in whom therapy is indicated
Poletti 2024 ¹⁰¹ Italy	Urine or saliva	Not stated	Yes (up to 36 months)	940 11 with cCMV	Hearing loss at birth Antiviral treatment Hearing loss at follow-up Neurodevelopment at follow-up	Implementation of an expanded cCMV screening program appears feasible and of clinical value
McCrary 2020 ²⁰⁰ USA	Urine, saliva, DBS	Not stated	Unclear	754 21 with cCMV	Symptoms at birth	An expanded targeted early CMV testing program has the potential to identify symptomatic cCMV infants who would not be identified otherwise
Abbreviations: cCMV, congenital cytomegalovirus; CMV, cytomegalovirus; DBS, dried blood spots.						

Table 22: Meta-analysis of event rates in 10 studies of symptom targeted screening

Event	Number of studies with data	Predicted rate per 10,000 screened newborns (95% confidence interval)	Predicted rate per 10,000 newborns with cCMV (95% confidence interval)
Diagnosis of cCMV	8	147 (82 to 262)	x
Has symptoms at birth (all symptoms)	8	71 (29 to 172)	5438 (3795 to 6991)
Has hearing loss at birth	5	13 (3 to 69)	1983 (704 to 4471)
Has cranial MRI/US abnormalities at birth	Insufficient data		
Receives antiviral treatment	5	43 (20 to 93)	2828 (1472 to 4741)
Has long-term sequelae	4	15 (6 to 38)	1800 (964 to 3111)
Abbreviations: cCMV, congenital cytomegalovirus; MRI, magnetic resonance imaging; US, ultrasound.			

Summary of Findings Relevant to Criteria 11 and 13

Criterion Not Met

The review identified 14 studies on universal cCMV screening and 22 studies of targeted cCMV screening relevant to this question. None of the studies were randomised, and only 1 study reported any form of non-randomised comparative evidence. The lack of comparative evidence inevitably limits the quality and value of the available evidence because it is not possible to determine whether symptoms at birth, or long-term sequelae, are caused by CMV infection, or are unrelated to infection. Nor is it possible to evaluate which symptoms or long-term sequelae would have been identified, managed and treated in the absence of cCMV screening. Given this fundamental limitation, there is currently no evidence to demonstrate that either universal or targeted screening of newborns for cCMV will provide long-term benefits to children with cCMV.

Key potential benefits of universal screening are that most cCMV cases would not be diagnosed without universal screening, because most newborns with cCMV do not have clear symptoms and because of potential issues with lack of awareness of cCMV among clinicians. It will also identify newborns with later development of neurological sequelae that would otherwise be missed. However, key concerns with universal screening are that most newborns with cCMV are asymptomatic at birth, without hearing loss and with a normal cranial ultrasound. Most of these babies will not develop long-term sequelae caused by cCMV. Universal screening also may identify many cases of subclinical symptomatic cCMV.

As asymptomatic newborns with cCMV are not currently offered antiviral therapy, the management pathway for most children diagnosed with cCMV is uncertain. The evidence suggests that only a minority of children diagnosed with cCMV through universal screening (perhaps around 11%) will receive antiviral therapy, and so the potential long-term clinical significance of universal screening and its cost-effectiveness are unknown. A further concern is that the only comparative study found that rates of long-term harms were similar between children with and without cCMV infection. This was in Finland, where primary maternal CMV infection rates are low, so the exact prevalence of cCMV could be an important factor in determining the clinical and cost-effectiveness of cCMV screening.

Key potential benefits of targeted screening after failure of a newborn hearing test are that it identifies the majority of infants with CMV-related SNHL at birth who could benefit from treatment. It allows for early recognition of asymptomatic cases with hearing loss. Key difficulties are that it will miss cases that are asymptomatic at birth, or who have symptoms that are not hearing loss. There may also be difficulties with timing, as both the hearing test and the cCMV test must be conducted quickly after birth if antiviral treatment is to be instigated promptly.

Unlike universal screening, most newborns who test positive for cCMV after targeted screening are symptomatic, and potentially go on to receive antiviral therapy. Hence targeted screening has a clearer treatment and management pathway than universal screening. However, it remains to be shown that targeted screening and subsequent antiviral therapy leads to better clinical outcomes than informal or ad-hoc testing for cCMV, or treating and managing sequelae caused by cCMV as symptoms emerge.

Adding additional signs or symptoms over hearing loss to targeted screening does not seem to have any clear benefits. The 12 studies that considered this were inconsistent in which symptoms were added to hearing loss. The overall impact appeared to be increasing the number of asymptomatic cases of cCMV that were detected, with a consequent reduction in the proportion of newborns who received antiviral treatment.

Criterion 12: There should be evidence that the complete screening programme (test, diagnostic procedures, treatment/ intervention) is clinically, socially and ethically acceptable to health professionals and the public.

Previous work

The acceptability of cCMV screening has not been assessed in any previous UK NSC evidence review on this topic.

Description of the current evidence

The studies of universal or targeted screening described under Criteria 11 and 13 generally concluded that screening was feasible and practical. However, only 5 of those studies included any kind of formal survey of parents, doctors, nurses or other stakeholders. Demortier (2021)¹⁸² and Letamendia-Richard (2022)⁵⁴ surveyed healthcare workers as part of a universal screening programme using saliva testing (see Table 17). Beswick (2021)¹⁹¹, Kadambari (2015)¹⁹⁴ and Webb (2024)²⁰⁶ surveyed stakeholders, screeners and parents, respectively, as part of a targeted screening programme using saliva testing (see Table 19 and Table 21).

As these were simple surveys, no quality assessment of these studies was undertaken.

Demortier (2021)¹⁸² surveyed 9 healthcare workers in La Reunion and Mayotte. They agreed that screening was generally acceptable, although with some limitations due to time taken to obtain parental consent. Responses were mixed as to whether universal screening for cCMV was relevant and useful.

Letamendia-Richard (2022)⁵⁴ surveyed 72 healthcare workers, of which 92.7% found the salivary test easy to perform; 91.4% reported that time devoted to sample collection was acceptable. Time spent obtaining informed consent was also acceptable for 78.6% of the respondents.

Beswick (2021)¹⁹¹ analysed feedback from an unstated number of stakeholders. They identified problems with limited funding for screening and training burdens. Nurses found that a cCMV diagnosis increased parental anxiety and found a lack of awareness of cCMV. The rarity of cCMV and its symptoms meant there was “study fatigue” and a reduced recruitment rate.

Kadambari (2015)¹⁹⁴ surveyed 24 screeners who generally felt that screening was a good idea and was feasible. They also generally felt confident explaining the screening process to parents.

Webb (2024)²⁰⁶ surveyed 65 parents, most of whom thought that saliva testing for cCMV was easy to do (92%); that testing their baby for cCMV was a good idea (92.0%) and that they were glad their baby was screened for cCMV (96.9%).

Summary of Findings Relevant to Criterion 12

Criterion Not Met

Evidence on the acceptability of cCMV screening is extremely limited, covering only 5 studies, mostly of small size. The evidence base is therefore too small, and of too limited quality, to draw any reasonable conclusions.

Evidence was mostly from screeners and medical professionals who generally concluded that screening was feasible and acceptable. Some concerns over screening increasing parental anxiety were raised. Evidence from parents was available from only 1 small study, where parents were very supportive of screening.

The current evidence relates only to the acceptability of saliva testing, and there is no evidence on the acceptability of cCMV screening more broadly. For example, there is no evidence on the acceptability of screening where newborns are asymptomatic, or on the acceptability of antiviral treatment, general management of cCMV, or long-term follow-up.

Review summary

Conclusions and implications for policy

This report assesses newborn screening for congenital cytomegalovirus (cCMV) infection against select UK National Screening Committee (UK NSC) criteria for appraising the viability, effectiveness and appropriateness of a screening programme.

There has been considerable new research in this area since the previous evidence review in 2017 and the evidence map in 2022. This new evidence particularly demonstrates the high accuracy and practicality of saliva or DBS testing to detect cCMV infection in newborns. There has been an expansion in the evidence on tests or characteristics that will predict long-term sequelae of cCMV infection. Whilst this evidence is of limited quality, the results indicate that presence of cCMV symptoms at birth, and timing of maternal CMV infection, are both associated with long-term sequelae. There is also promising, but not yet conclusive, evidence that the use of MRI scans and assessment of CMV viral load may be useful in identifying newborns at higher risk of long-term sequelae. There have also been numerous studies reporting on the use of cCMV screening in practice, demonstrating its general feasibility and practicality.

However, despite the now substantial evidence base on cCMV screening, we conclude that further evidence is required before a change to the current recommendation can be made. This is primarily because there have been no studies, either randomised or non-randomised, that formally compare the outcomes of a cCMV screening programme to no screening or current clinical management, so there is, as yet, no conclusive evidence that newborn screening for cCMV will be beneficial. Therefore, newborn screening for cCMV is still not recommended.

Evidence gaps and recommendations for research

The most substantial gap in the current evidence, where there is clear need for future clinical studies, is to properly compare the clinical impact of cCMV screening to current clinical practice without screening. Ideally this should consist of cluster randomised trials, where locations or medical centres are randomised to: universal cCMV screening, targeted screening after failing a newborn hearing screen, or standard practice without screening. Newborns should be tested using saliva, DBS or urine PCR, with a confirmatory urine test if required. Appropriate tests should be performed to identify cCMV symptoms and antiviral therapy offered to newborns with symptoms. Follow-up should be sufficient to identify long-term sequelae, as far as is possible, and confirm the impact of screening and antiviral treatment. Ideally,

trials should also survey parents and caregivers, to evaluate the acceptability of screening, treatment and long-term management.

As randomised trials may be impractical, non-randomised comparative studies may also be useful. We note that, with the introduction of cCMV screening in some Canadian provinces and US states, studies comparing locations that have adopted screening to those that have not may be useful.

This review has also identified some other areas where further clinical research would be useful. As MRI appears to be a promising means of predicting long-term sequelae, further studies evaluating the prognostic accuracy of MRI to detect long-term hearing, developmental and neurological outcomes would be useful, as well as studies evaluating the practical implications of using MRI scans as part of cCMV screening, such as identifying which newborns would most benefit from an MRI scan. Similarly, further studies assessing the predictive value of CMV viral load to identify newborns at risk of long-term sequelae would be useful. These could form part of the randomised trials of newborn cCMV screening.

Development of a tool to predict long-term sequelae that incorporates the full range of plausible predictors (such as symptoms at birth, trimester of infection, specific imaging findings, or viral load) would also be useful. This is because the causes of long-term sequelae are almost certainly multifactorial, and so an accurate test will need to cover a range of predictive variables. The design of such a tool could initially be informed by a large cohort study in a neonatal population with cCMV, investigating the magnitude of association of plausible predictors with sequelae using multivariable logistic regression. Such a risk tool could then be validated in a separate cohort, to establish how well it can predict sequelae.

We note that there is now a substantial evidence base on the accuracy of both saliva and DBS testing for cCMV, with both covered by extensive systematic reviews (including ongoing reviews, listed in Appendix 4). We therefore suggest that further diagnostic accuracy studies are not required. However, there may still be value in studies comparing the accuracy and practicality of saliva screening with DBS screening.

Limitations

Many of the included studies were limited in terms of their design, size and quality. In addition, very few studies were conducted in the UK, limiting their applicability to UK practice, owing to differences in health systems and the prevalence of maternal CMV in different countries.

Acknowledgments

The York Evidence Synthesis group would like to thank all those who supported and provided advice for this review. Specifically, all those at UK NSC, all clinical advisers, CMV Action and all parent representatives.

Appendix 1 — Search strategy

Electronic databases and resources

The search strategy included searches of the databases shown in Table 1. All searches were restricted to literature published from 2013 onwards.

Table 23 Databases and resources searched

Database/resource	Platform	Date of search	Date range/segment
MEDLINE ALL	Ovid	19/11/2024	1946 to Nov 18, 2024
Embase	Ovid	19/11/2024	1974 to 2024 Nov 18
Maternity and Infant Care	Ovid	19/11/2024	1971 to Nov 05, 2024
CINAHL	Ebsco	19/11/2024	Inception to 20241119
LILACS	https://pesquisa.bvsalud.org/portal/advanced/?lang=en	19/11/2024	Inception to last update (date unavailable)
Cochrane Central Register of Controlled Trials (CENTRAL)	Cochrane Library, via Wiley	19/11/2024	Issue 10 of 12, Oct 2024
Cochrane Database of Systematic Reviews (CDSR)	Cochrane Library, Wiley	19/11/2024	Issue 11 of 12, Nov 2024
Science Citation Index (SCI)	Web of Science, Clarivate Analytics	19/11/2024	1900 – 2024-11-16
Conference Proceedings Citation Index – Science (CP-SCI)	Web of Science, Clarivate Analytics	19/11/2024	1990 – 2024-11-16
KSR Evidence	Ovid	19/11/2024	2015 to 2024 Week 45
Database of Abstracts of Reviews of Effects (DARE)	CRD databases	19/11/2024	Inception – 31 st March 2015
Health Technology Assessment (HTA) database	CRD databases	19/11/2024	Inception – 31 st March 2018
International Health Technology Assessment (HTA) database	https://database.inahta.org/	19/11/2024	Inception – last update (date unavailable)
PROSPERO	https://www.crd.york.ac.uk/prospero/	19/11/2024	Inception – last update (date unavailable)
ClinicalTrials.gov	https://clinicaltrials.gov/ct2/	19/11/2024	Inception – last update (date unavailable)
WHO International Clinical Trials Registry Platform (ICTRP)	https://trialsearch.who.int/Default.aspx	19/11/2024	Inception – last update (date unavailable)
EU Clinical Trials Register	https://www.clinicaltrialsregister.eu/ctr-search/search	20/11/2024	Inception – last update (date unavailable)
HTA Agency websites	Various (see below for individual HTA agency URLs accessed)	19/11/2024	Inception – last update (date unavailable)

Search Terms and structure

Search terms included combinations of free text and subject headings (e.g. Medical Subject Headings [MeSH] for MEDLINE, and Emtree terms for Embase). The search strategy was structured using aspects of the population, combined using the Boolean operator AND:

- Population: age – newborns (0-28 days)

AND

- Population: health condition – CMV

Individual search strategies for each database and resource can be viewed below.

Database search strategies

MEDLINE ALL

via Ovid <http://ovidsp.ovid.com/>

Date range: 1946 to November 18, 2024

Date searched: 19th November 2024

Records retrieved: 2638

```
1  exp Infant, Newborn/ (696555)
2  Neonatology/ (3278)
3  (newborn$ or new born$ or newly born$).ti,ab,kf. (212956)
4  (neonat$ or neo nat$).ti,ab,kf. (334932)
5  (infant$ or infancy).ti,ab,kf. (579331)
6  (baby$ or babies).ti,ab,kf. (88964)
7  at birth.ti,ab,kf. (61025)
8  (preterm or preterms or pre term or pre terms).ti,ab,kf. (104600)
9  (preemie$ or premie or premies).ti,ab,kf. (239)
10 (low adj2 (birthweight$ or birth weight$)).ti,ab,kf. (43582)
11 (lbw or vlbw or elbw).ti,ab,kf. (11240)
12 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 (1297214)
13 Cytomegalovirus/ (23937)
14 exp Cytomegalovirus Infections/ (29425)
15 cytomegalovir$.ti,ab,kf. (49521)
16 ((cytomegal$ or cyto megal$) adj6 (virus$ or viral$ or vir?emi$)).ti,ab,kf. (10032)
17 cytomegaloinfection$.ti,ab,kf. (1)
18 ((cytomegal$ or cyto megal$) adj infection$).ti,ab,kf. (11208)
19 ((cytomegal$ or cyto megal$) and (virus$ or viral$ or vir?emi$)).ti,ab,kf. (156)
20 (CMV or HCMV).ti,ab,kf. (37691)
21 cCMV.ti,ab,kf. (739)
22 human herpesvirus 5.ti,ab,kf. (37)
23 human herpes virus 5.ti,ab,kf. (14)
24 (HHV 5 or HHV5).ti,ab,kf. (66)
25 salivary gland virus$.ti,ab,kf. (59)
26 13 or 14 or 15 or 16 or 17 or 18 or 19 or 20 or 21 or 22 or 23 or 24 or 25 (65657)
27 12 and 26 (7380)
28 exp animals/ not humans.sh. (5278373)
29 27 not 28 (7098)
30 limit 29 to yr="2013 -Current" (2638)
```

Key:

/ = subject heading (MeSH heading)

sh = subject heading (MeSH heading)

exp = exploded subject heading (MeSH heading)
\$ = truncation
? = optional wildcard – one or no characters
ti,ab,kf = terms in title or abstract or author keyword fields
adj = terms next to each other in order specified
adj2 = terms within two words of each other (any order)

Embase

via Ovid <http://ovidsp.ovid.com/>

Date range: 1974 to 2024 November 18

Date searched: 19th November 2024

Records retrieved: 4804

- 1 newborn/ (632221)
- 2 baby/ (14822)
- 3 neonatology/ (6041)
- 4 (newborn\$ or new born\$ or newly born\$).ti,ab,kw. (243697)
- 5 (neonat\$ or neo nat\$).ti,ab,kw. (442847)
- 6 (infant\$ or infancy).ti,ab,kw. (624640)
- 7 (baby\$ or babies).ti,ab,kw. (127181)
- 8 at birth.ti,ab,kw. (81954)
- 9 exp low birth weight/ (65734)
- 10 exp prematurity/ (138950)
- 11 (preterm or preterms or pre term or pre terms).ti,ab,kw. (145796)
- 12 (preemie\$ or premie or premies).ti,ab,kw. (393)
- 13 (low adj2 (birthweight\$ or birth weight\$)).ti,ab,kw. (55243)
- 14 (lbw or vlbw or elbw).ti,ab,kw. (15481)
- 15 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14 (1435440)
- 16 Cytomegalovirus/ (41472)
- 17 exp human cytomegalovirus/ (8760)
- 18 exp cytomegalovirus infection/ (45608)
- 19 cytomegalovir\$.ti,ab,kw. (63173)
- 20 ((cytomegal\$ or cyto megal\$) adj6 (virus\$ or viral\$ or vir?emi\$)).ti,ab,kw. (13634)
- 21 cytomegaloinfection\$.ti,ab,kw. (1)
- 22 ((cytomegal\$ or cyto megal\$) adj infection\$).ti,ab,kw. (13895)
- 23 ((cytomegal\$ or cyto megal\$) and (virus\$ or viral\$ or vir?emi\$)).ti,ab,kw. (181)
- 24 (CMV or HCMV).ti,ab,kw. (61625)
- 25 cCMV.ti,ab,kw. (863)
- 26 human herpesvirus 5.ti,ab,kw. (51)
- 27 human herpes virus 5.ti,ab,kw. (16)
- 28 (HHV 5 or HHV5).ti,ab,kw. (99)
- 29 salivary gland virus\$.ti,ab,kw. (13)
- 30 16 or 17 or 18 or 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26 or 27 or 28 or 29 (113281)
- 31 15 and 30 (9938)
- 32 animal/ or animal experiment/ (4904537)
- 33 (animal or animals or rat or rats or mouse or mice or murine or pig or pigs or piglet or piglets or porcine or rabbit or rabbits or hamster or hamsters or sheep or monkey or monkeys or macaque or macaques).ti,ab,hw. (7805171)
- 34 32 or 33 (7805171)
- 35 exp human/ or exp human experiment/ (27309330)
- 36 34 not (34 and 35) (5829658)
- 37 31 not 36 (9328)
- 38 limit 37 to yr="2013 -Current" (4804)

Key:

/ = subject heading (Emtree heading)

exp = exploded subject heading (Emtree heading)

hw = subject heading word

\$ = truncation

? = optional wildcard – one or no characters

ti,ab,kw = terms in title or abstract or author keyword fields

adj = terms next to each other in order specified

adj2 = terms within two words of each other (any order)

Maternity and Infant Care

via Ovid <http://ovidsp.ovid.com/>

Date range: 1971 to November 05, 2024

Date searched: 19th November 2024

Records retrieved: 404

- 1 (newborn\$ or new born\$ or newly born\$).af. (48922)
- 2 (neonat\$ or neo nat\$).af. (77600)
- 3 (infant\$ or infancy).af. (116339)
- 4 (baby\$ or babies).af. (42172)
- 5 at birth.af. (11025)
- 6 (preterm or preterms or pre term or pre terms).af. (36985)
- 7 (preemie\$ or premie or premies).af. (72)
- 8 (low adj2 (birthweight\$ or birth weight\$)).af. (17959)
- 9 (lbw or vlbw or elbw).af. (4041)
- 10 or/1-9 (178316)
- 11 cytomegalovir\$.af. (1060)
- 12 ((cytomegal\$ or cyto megal\$) adj6 (virus\$ or viral\$ or vir?emi\$)).af. (188)
- 13 cytomegaloinfection\$.af. (0)
- 14 ((cytomegal\$ or cyto megal\$) adj infection\$).af. (579)
- 15 ((cytomegaly\$ or cyto megaly\$) and (virus\$ or viral\$ or vir?emi\$)).af. (0)
- 16 (CMV or HCMV).af. (681)
- 17 cCMV.af. (99)
- 18 (human herpesvirus 5 or human herpes virus 5).af. (0)
- 19 (HHV 5 or HHV5).af. (1)
- 20 salivary gland virus\$.af. (0)
- 21 or/11-20 (1118)
- 22 10 and 21 (795)
- 23 limit 22 to yr="2013 -Current" (404)

Key:

\$ = truncation

? = optional wildcard – one or no characters

af = terms in any database fields

adj = terms next to each other in order specified

adj2 = terms within two words of each other (any order)

CINAHL

via Ebsco <https://www.ebsco.com/>

Date range: Inception to 20241119

Date searched: 19th November 2024

Records retrieved: 783

S1	(MH "Infant, Newborn+")	162,261	
S2	(MH "Neonatology")	1,542	
S3	TI ((newborn* or new-born* or (newly N0 born*)) OR AB ((newborn* or new-born* or (newly N0 born*))		39,452
S4	TI (neonat* or neo-nat*) OR AB (neonat* or neo-nat*)	86,610	
S5	TI (infant* or infancy) OR AB (infant* or infancy)	139,479	
S6	TI (baby* or babies) OR AB (baby* or babies)	38,698	

S7	TI (preterm or preterms or pre-term or pre-terms) OR AB (preterm or preterms or pre-term or pre-terms)	43,268
S8	TI (preemie* or premie or premies) OR AB (preemie* or premie or premies)	355
S9	TI (low N2 (birthweight* or birth-weight*)) OR AB (low N2 (birthweight* or birth-weight*))	15,037
S10	TI (lbw or vlbw or elbw) OR AB (lbw or vlbw or elbw)	4,174
S11	S1 OR S2 OR S3 OR S4 OR S5 OR S6 OR S7 OR S8 OR S9 OR S10	315,539
S12	(MH "Cytomegaloviruses")	1,701
S13	(MH "Cytomegalovirus Infections+")	3,874
S14	TI cytomegalovir* OR AB cytomegalovir*	5,216
S15	TI ((cytomegal* or cyto-megal*) N6 (virus* or viral* or vir#emi*)) OR AB ((cytomegal* or cyto-megal*) N6 (virus* or viral* or vir#emi*))	1,092
S16	TI cytomegaloinfection* OR AB cytomegaloinfection*	0
S17	TI ((cytomegal* or cyto-megal*) N0 infection*) OR AB ((cytomegal* or cyto-megal*) N0 infection*)	1,544
S18	TI ((cytomegal* or cyto-megal*) and (virus* or viral* or vir#emi*)) OR AB ((cytomegal* or cyto-megal*) and (virus* or viral* or vir#emi*))	2
S19	TI (CMV or HCMV) OR AB (CMV or HCMV)	3,174
S20	TI cCMV OR AB cCMV	152
S21	TI ("human herpesvirus 5" or "human herpes virus 5") OR AB ("human herpesvirus 5" or "human herpes virus 5")	0
S22	TI ("HHV 5" or "HHV5") OR AB ("HHV 5" or "HHV5")	0
S23	TI ("salivary gland virus" or "salivary gland viruses") OR AB ("salivary gland virus" or "salivary gland viruses")	1
S24	S12 OR S13 OR S14 OR S15 OR S16 OR S17 OR S18 OR S19 OR S20 OR S21 OR S22 OR S23	7,044
S25	S11 AND S24	1,308
S26	MH animals+	102,528
S27	MH (animal studies)	158,518
S28	TI (animal model*)	4,039
S29	S26 OR S27 OR S28	252,096
S30	MH (human)	2,840,690
S31	S29 NOT S30	217,272
S32	S25 NOT S31	1,289
S33	S25 NOT S31 Limiters - Publication Date: 20130101-20241231	783

Key:

MH = CINAHL subject heading

MH with + = exploded CINAHL subject heading

* = truncation

= wildcard - one or no characters

TI = title field

AB = abstract field

N2 = terms within two words of each other (any order)

Latin American and Caribbean Health Sciences Literature (LILACS)

via <https://pesquisa.bvsalud.org/portal/advanced/?lang=en>

Date range: Inception to current

Date searched: 19th November 2024

Records retrieved: 233

Advanced search screen used, terms searched in the title, abstract and subject fields. Filters applied: results from LILACS databases, publication year range: 2013-2024

1. newborn* OR new-born* OR "newly born" OR neonat* OR neo-nat* OR infant* OR infancy OR baby* OR babies OR "at birth" OR preterm OR preterms OR pre-term OR pre-terms OR preemie* OR premie OR premies OR lbw OR vlbw OR elbw OR "low birthweight" OR "low birthweights" OR "low birth-weight" OR "low birth-weights"

AND

cytomegalovir* OR cytomegaloinfection* OR CMV OR HCMV OR cCMV OR "human herpesvirus 5" OR "human herpes virus 5" OR "HHV 5" OR "HHV5" OR "salivary gland virus" OR "salivary gland viruses"

123 records

2. newborn* OR new-born* OR "newly born" OR neonat* OR neo-nat* OR infant* OR infancy OR baby* OR babies OR "at birth" OR preterm OR preterms OR pre-term OR pre-terms OR preemie* OR premie OR premies OR lbw OR vlbw OR elbw OR "low birthweight" or "low birthweights" or "low birth-weight" or "low birth-weights"

AND

(cytomegal* OR cyto-megal*) AND (infection* OR virus* OR viral* OR viremi* OR viraemi*)

110 records

Key:

* = truncation

Cochrane Library

via Wiley <http://onlinelibrary.wiley.com/>

Cochrane Central Register of Controlled Trials (CENTRAL): Issue 10 of 12, Oct 2024

Records retrieved: 192

Cochrane Database of Systematic Reviews (CDSR): Issue 11 of 12, Nov 2024

Records retrieved: 6

Date searched: 19th November 2024

#1	MeSH descriptor: [Infant, Newborn] explode all trees	24460	
#2	MeSH descriptor: [Neonatology] this term only	63	
#3	(newborn* or (new next born*) or (newly next born*)):ti,ab,kw	37964	
#4	(neonat* or (neo next nat*)):ti,ab,kw	30453	
#5	(infant* or infancy):ti,ab,kw	81631	
#6	(baby or babies):ti,ab,kw	12265	
#7	(at next birth):ti,ab,kw	39722	
#8	(preterm or preterms or (pre next term) or (pre next terms)):ti,ab,kw	18139	
#9	(preemie* or premie or premies):ti,ab,kw	57	
#10	(low near/2 (birthweight* or birth weight*)):ti,ab,kw	12915	
#11	(lbw or vlbw or elbw):ti,ab,kw	2022	
#12	#1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11	121007	
#13	MeSH descriptor: [Cytomegalovirus] this term only	461	
#14	MeSH descriptor: [Cytomegalovirus Infections] explode all trees	1061	
#15	cytomegalovir*:ti,ab,kw	2682	
#16	((cytomegal* or cyto-megal*) near/6 (virus* or viral* or viremi* or viraemi*)):ti,ab,kw	370	
#17	cytomegaloinfection*:ti,ab,kw	0	
#18	((cytomegal* or cyto-megal*) next infection*):ti,ab,kw	1803	
#19	((cytomegaly* or cyto-megaly*) and (virus* or viral* or viremi* or viraemi*)):ti,ab,kw	3	
#20	(CMV or HCMV):ti,ab,kw	2368	
#21	cCMV:ti,ab,kw	12	
#22	"human herpesvirus 5":ti,ab,kw	0	
#23	"human herpes virus 5":ti,ab,kw	0	
#24	("HHV 5" or "HHV5"):ti,ab,kw	0	
#25	(salivary next gland next virus*):ti,ab,kw	0	
#26	#13 or #14 or #15 or #16 or #17 or #18 or #19 or #20 or #21 or #22 or #23 or #24 or #25	3438	
#27	#12 and #26 with Cochrane Library publication date Between Jan 2013 and Nov 2024, in Cochrane Reviews, Cochrane Protocols	6	
#28	#12 and #26 with Publication Year from 2013 to 2024, in Trials	192	

Key:

MeSH descriptor = subject heading (MeSH heading)

* = truncation

? = wildcard - zero or one characters

ti,ab,kw = terms in title, abstract or keyword fields

next = terms are next to each other

near/2 = terms within two words of each other (any order)

Science Citation Index (SCI)

Conference Proceedings Citation Index – Science (CP-SCI)

via Web of Science, Clarivate Analytics <https://clarivate.com/>

Date searched: 19th November 2024

Date range SCI: 1900 – 2024-11-16

Date range CP-SCI: 1990 – 2024-11-16

Records retrieved: 2606

- | | | |
|--|---|-----------------|
| 1: TS=(cytomegalovir*) | Editions: WOS.SCI,WOS.ISTP | Results: 57207 |
| 2: TS=((cytomegal* or cyto-megal*) NEAR/6 (virus* or viral* or vir\$emi*)) | Editions: WOS.SCI,WOS.ISTP | Results: 12259 |
| 3: TS=(cytomegaloinfection*) | Editions: WOS.SCI,WOS.ISTP | Results: 0 |
| 4: TS=((cytomegal* or cyto-megal*) NEAR/1 infection*) | Editions: WOS.SCI,WOS.ISTP | Results: 19533 |
| 5: TS=((cytomegaly* or cyto-megaly*) and (virus* or viral* or vir\$emi*)) | Editions: WOS.SCI,WOS.ISTP | Results: 118 |
| 6: TS=(CMV or HCMV) | Editions: WOS.SCI,WOS.ISTP | Results: 40762 |
| 7: TS=(cCMV) | Editions: WOS.SCI,WOS.ISTP | Results: 748 |
| 8: TS=("human herpesvirus 5" or "human herpes virus 5") | Editions: WOS.SCI,WOS.ISTP | Results: 43 |
| 9: TS=("HHV 5" or "HHV5") | Editions: WOS.SCI,WOS.ISTP | Results: 60 |
| 10: TS=("salivary gland virus") | Editions: WOS.SCI,WOS.ISTP | Results: 74 |
| 11: #1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 | Editions: WOS.SCI,WOS.ISTP | Results: 73539 |
| 12: TS=(newborn* or new-born* or "newly born") | Editions: WOS.SCI,WOS.ISTP | Results: 196051 |
| 13: TS=(neonat* or neo-nat*) | Editions: WOS.SCI,WOS.ISTP | Results: 341407 |
| 14: TS=(infant* or infancy) | Editions: WOS.SCI,WOS.ISTP | Results: 550643 |
| 15: TS=(baby* or babies) | Editions: WOS.SCI,WOS.ISTP | Results: 76840 |
| 16: TS=("at birth") | Editions: WOS.SCI,WOS.ISTP | Results: 56577 |
| 17: TS=(preterm or preterms or pre-term or pre-terms) | Editions: WOS.SCI,WOS.ISTP | Results: 126050 |
| 18: TS=(preemie* or premie or premies) | Editions: WOS.SCI,WOS.ISTP | Results: 229 |
| 19: TS=(low NEAR/2 (birthweight* or birth-weight*)) | Editions: WOS.SCI,WOS.ISTP | Results: 52040 |
| 20: TS=(lbw or vlbw or elbw) | Editions: WOS.SCI,WOS.ISTP | Results: 11324 |
| 21: #12 OR #13 OR #14 OR #15 OR #16 OR #17 OR #18 OR #19 OR #20 | Editions: WOS.SCI,WOS.ISTP | Results: 991736 |
| 22: #11 and #21 | Editions: WOS.SCI,WOS.ISTP Timespan: All years (Publication Date) | Results: 5492 |

Key:

TS = topic tag; searches in title, abstract, author keywords and keywords plus fields

* = truncation

\$ = represents zero or one character

NEAR/2 = terms within two words of each other (any order)

EB Health - KSR Evidence

via Ovid <http://ovidsp.ovid.com/>

Date range: 2015 to 2024 Week 45

Date searched: 19th November 2024

Records retrieved: 60

- 1 (newborn\$ or new born\$ or newly born\$).af. (2142)
- 2 (neonat\$ or neo nat\$).af. (5888)
- 3 (infant\$ or infancy).af. (7028)
- 4 (baby\$ or babies).af. (1475)
- 5 at birth.af. (772)
- 6 (preterm or preterms or pre term or pre terms).af. (3846)
- 7 (preemie\$ or premie or premies).af. (2)
- 8 (low adj2 (birthweight\$ or birth weight\$)).af. (1490)
- 9 (lbw or vlbw or elbw).af. (426)
- 10 or/1-9 (13134)
- 11 cytomegalovir\$.af. (311)
- 12 ((cytomegal\$ or cyto megal\$) adj6 (virus\$ or viral\$ or vir?emi\$)).af. (71)
- 13 cytomegaloinfection\$.af. (0)
- 14 ((cytomegal\$ or cyto megal\$) adj infection\$).af. (119)
- 15 ((cytomegaly\$ or cyto megaly\$) and (virus\$ or viral\$ or vir?emi\$)).af. (0)
- 16 (CMV or HCMV).af. (274)
- 17 cCMV.af. (22)
- 18 (human herpesvirus 5 or human herpes virus 5).af. (0)
- 19 (HHV 5 or HHV5).af. (2)
- 20 salivary gland virus\$.af. (0)
- 21 or/11-20 (380)
- 22 10 and 21 (61)
- 23 limit 22 to yr="2013 -Current" (60)

Key:

\$ = truncation

? = optional wildcard – one or no characters

af = terms in any field

adj = terms next to each other in order specified

adj2 = terms within two words of each other (any order)

Database of Abstracts of Reviews of Effects (DARE)**Health Technology Assessment (HTA) database**

via <http://www.crd.york.ac.uk/CRDWeb/>

Date range DARE: Inception – 31st March 2015

Date range HTA database: Inception – 31st March 2018

Date searched: 19th November 2024

Records retrieved: 1

- 1 MeSH DESCRIPTOR Infant, Newborn EXPLODE ALL TREES IN DARE,HTA 1057
- 2 MeSH DESCRIPTOR Neonatology IN DARE,HTA 5

3	(newborn* or new-born* or "newly born") IN DARE, HTA	1332
4	(neonat* or neo-nat*) IN DARE, HTA	1352
5	(infant* or infancy) IN DARE, HTA	2740
6	(baby* or babies) IN DARE, HTA	428
7	("at birth") IN DARE, HTA	128
8	(preterm or preterms or pre-term or pre-terms) IN DARE, HTA	828
9	(preemie* or premie or premies) IN DARE, HTA	0
10	(low adj2 (birthweight* or birth-weight*)) IN DARE, HTA	317
11	((birthweight* or birth-weight*) adj2 low) IN DARE, HTA	22
12	(lbw or vlbw or elbw) IN DARE, HTA	43
13	#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9 OR #10 OR #11 OR #12	3498
14	MeSH DESCRIPTOR Cytomegalovirus IN DARE,HTA7	
15	MeSH DESCRIPTOR Cytomegalovirus Infections EXPLODE ALL TREES IN DARE,HTA	31
16	(cytomegalovir*) IN DARE, HTA	61
17	((cytomegal* or cyto-megal*) adj6 (virus* or viral* or viremi* or viraemi*)) IN DARE, HTA	7
18	((virus* or viral* or viremi* or viraemi*) adj6 (cytomegal* or cyto-megal*)) IN DARE, HTA	9
19	(cytomegaloinfection*) IN DARE, HTA	0
20	((cytomegal* or cyto-megal*) adj1 infection*) IN DARE, HTA	46
21	(infection* adj1 (cytomegal* or cyto-megal*)) IN DARE, HTA	8
22	((cytomegal* or cyto-megal*) and (virus* or viral* or viremi* or viraemi*)) IN DARE, HTA	0
23	(CMV or HCMV) IN DARE, HTA	33
24	(cCMV) IN DARE, HTA	0
25	("human herpesvirus 5" or "human herpes virus 5") IN DARE, HTA	0
26	("HHV 5" or HHV5) IN DARE, HTA	0
27	("salivary gland virus" or "salivary gland viruses") IN DARE, HTA	0
28	#14 OR #15 OR #16 OR #17 OR #18 OR #19 OR #20 OR #21 OR #22 OR #23 OR #24 OR #25 OR #26 OR #27	71
29	#13 AND #28	9
30	(*) IN DARE, HTA FROM 2013 TO 2015	20407
31	#29 AND #30	1

Key:

MeSH DESCRIPTOR = subject heading (MeSH heading)

* = truncation

adj2 = terms within two words of each other (order specified)

International Health Technology Assessment (HTA) database

via <https://database.inahta.org/>

Date range: Inception – 18th November 2024

Date searched: 19th November 2024

Records retrieved: 1

((("Cytomegalovirus Infections"[mhe]) OR ("Cytomegalovirus"[mh]) OR (("human herpesvirus 5" OR "human herpes virus 5" OR "HHV 5" OR "HHV5" OR "salivary gland virus" OR "salivary gland viruses") [Title] OR ("human herpesvirus 5" OR "human herpes virus 5" OR "HHV 5" OR "HHV5" OR "salivary gland virus" OR "salivary gland viruses") [abs] OR ("human herpesvirus 5" OR "human herpes virus 5" OR "HHV 5" OR "HHV5" OR "salivary gland virus" OR "salivary gland viruses") [Keywords]) OR ((CMV or HCMV or cCMV) [Title] OR (CMV or HCMV or cCMV) [abs] OR (CMV or HCMV or cCMV) [Keywords]) OR ((cytomegaloinfection*) [Title] OR (cytomegaloinfection*) [abs] OR (cytomegaloinfection*) [Keywords]) OR (((cytomegal* or cyto-megal*) [Title] OR (cytomegal* or cyto-megal*) [abs] OR (cytomegal* or cyto-megal*) [Keywords]) AND ((infection* or virus* or viral* or viremi* or viraemi*) [Title] OR (infection* or virus* or viral* or viremi* or viraemi*) [abs] OR (infection* or virus* or viral* or viremi* or viraemi*) [Keywords])) OR ((cytomegalovir*) [Title] OR (cytomegalovir*) [abs] OR (cytomegalovir*) [Keywords])) AND (((("low birthweight" or "low birthweights" or "low birth-weight" or "low birth-weights") [Title] OR ("low birthweight" or "low birthweights" or "low birth-weight" or "low birth-weights") [abs] OR ("low birthweight" or "low birthweights" or "low birth-weight" or "low birth-weights") [Keywords]) OR ((preterm or preterms or pre-term or pre-terms or preemie* or premie or premies or lbw or vlbw or elbw) [Title] OR (preterm or preterms or pre-term or pre-terms or preemie* or premie or premies or lbw or vlbw or elbw) [abs] OR (preterm or preterms or pre-term or pre-terms or preemie* or premie or premies or lbw or vlbw or elbw) [Keywords])

elbw)[Keywords]) OR (("at birth")[Title] OR ("at birth")[abs] OR ("at birth")[Keywords]) OR ((newborn* or new-born* or "newly born" or neonat* or neo-nat* or infant* or infancy or baby* or babies)[Title] OR (newborn* or new-born* or "newly born" or neonat* or neo-nat* or infant* or infancy or baby* or babies)[abs] OR (newborn* or new-born* or "newly born" or neonat* or neo-nat* or infant* or infancy or baby* or babies)[Keywords]) OR ("Neonatology"[mh] OR ("Infant, Newborn"[mhe]))

Date limit 2013 onwards applied – 1 record retrieved

Key:

[Keywords] = search of keyword field

[abs] = search of abstract field

[Title] = search of title field

[mh] = subject heading search

[mhe] = exploded subject heading search

* = truncation

PROSPERO

via <https://www.crd.york.ac.uk/prosperto/>

Date searched: 19th November 2024

Records retrieved: 106

#1	MeSH DESCRIPTOR Infant, Newborn EXPLODE ALL TREES	1301
#2	MeSH DESCRIPTOR Neonatology	2
#3	newborn* or new-born* or (newly adj1 born*)	4750
#4	neonat* or neo-nat*	9274
#5	infant* or infancy or baby* or babies	14476
#6	"at birth"	1923
#7	preterm or preterms or pre-term or pre-terms or preemie* or premie or premies	5738
#8	low adj2 (birthweight* or birth-weight*)	2359
#9	lbw or vlbw or elbw	480
#10	#1 OR #2 OR #3 OR #4 OR #5 OR #6 OR #7 OR #8 OR #9	21414
#11	MeSH DESCRIPTOR Cytomegalovirus	23
#12	MeSH DESCRIPTOR Cytomegalovirus Infections EXPLODE ALL TREES	31
#13	cytomegalovir*	290
#14	(cytomegal* or cyto megal*) adj6 (virus* or viral* or viremi* or viraemi*)	89
#15	cytomegaloinfection*	0
#16	(cytomegal* or cyto megal*) adj infection*	150
#17	(cytomegal* or cyto megal*) and (virus* or viral* or viremi* or viraemi*)	3
#18	CMV or HCMV or cCMV	289
#19	"human herpesvirus 5" or "human herpes virus 5" or HHV5 or "HHV 5"	4
#20	"salivary gland virus" or "salivary gland viruses"	3
#21	#11 OR #12 OR #13 OR #14 OR #15 OR #16 OR #17 OR #18 OR #19 OR #20	398
#22	#10 AND #21	106

Key:

MeSH DESCRIPTOR = subject heading (MeSH heading)

* = truncation

TI,KW,RQ = terms in title, keyword or research question field

adj2 = terms within two words of each other

ClinicalTrials.gov

<https://clinicaltrials.gov/ct2/>

Date searched: 19th November 2024

Records retrieved: 504

1. cytomegalovirus OR CMV OR cCMV OR HCMV OR "cytomegalovirus infection" OR "congenital cytomegalovirus" | 0 Days to 28 Days old =114 studies

2. cytomegalovirus OR CMV OR cCMV OR HCMV OR "cytomegalovirus infection" OR "congenital cytomegalovirus" | 0 Months to 1 Months old =121 studies

3. cytomegalovirus OR CMV OR cCMV OR HCMV OR "cytomegalovirus infection" OR "congenital cytomegalovirus" | 0 Weeks to 4 Weeks old =114 studies

4. cytomegalovirus OR CMV OR cCMV OR HCMV OR "cytomegalovirus infection" OR "congenital cytomegalovirus" | Other terms: newborn* OR new-born* OR "newly born" OR neonat* OR neo-nat* OR infant* OR infancy OR baby* OR babies OR "at birth" =109 studies

5. cytomegalovirus OR CMV OR cCMV OR HCMV OR "cytomegalovirus infection" OR "congenital cytomegalovirus" | Other terms: preterm OR preterms OR pre-term OR pre-terms OR preemie* OR premie OR premies OR lbw OR vlbw OR elbw OR "low birthweight" OR "low birthweights" OR "low birth-weight" OR "low birth-weights" = 46 studies

WHO International Clinical Trials Registry Platform (ICTRP)

<https://trialsearch.who.int/Default.aspx>

Date searched: 19th November 2024

Records retrieved: 66

Basic search interface used. No date limits available in basic search interface, therefore results from all years downloaded and records pre-2013 removed in EndNote.

1. 65 records for 58 trials found for: (newborn* OR new-born* OR "newly born" OR neonat* OR neo-nat* OR infant* OR infancy OR baby* OR babies) AND (cytomegalovir* OR CMV OR cCMV OR HCMV)

2. 8 records for 8 trials found for: ("at birth" OR preterm OR preterms OR pre-term OR pre-terms OR preemie* OR premie OR premies OR lbw OR vlbw OR elbw OR "low birthweight" OR "low birthweights" OR "low birth-weight" OR "low birth-weights") AND (cytomegalovir* OR CMV OR cCMV OR HCMV)

EU Clinical Trials Register

<https://www.clinicaltrialsregister.eu/ctr-search/search>

Date of search: 20th November 2024

Records retrieved: 38

Advanced search

1. 38 result(s) found for: cytomegalovirus OR CMV OR HCMV OR cCMV filter applied: infant, newborn, preterm

Also searched for EU clinical trials via the new portal Clinical Trials Information System (CTIS) <https://euclinicaltrials.eu/search-for-clinical-trials/?lang=en>

1. cytomegalovirus – scanned 18 results – 0 with a population of newborns

2. CMV – scanned 32 results – 0 with a population of newborns

3. HCMV – 0 results

4. cCMV – 0 results

Website searches

HTA Agencies

Date searched: 19th November 2024

Records retrieved: 3

Browsed or searched the following HTA Agency websites to check for additional reports not found through database searches. A date limit of 2013 was applied.

Agency for Healthcare Research and Quality (AHRQ), USA

1. <https://www.ahrq.gov/research/findings/ta/index.html>

Browsed list of technology assessments, topic refinements and archive of technology assessments – 0 relevant reports found
108

2. <https://www.ahrq.gov/research/findings/evidence-based-reports/search.html>

Searched by keyword cytomegalovirus, CMV, cCMV, HCMV – 0 relevant reports found

Adelaide Health Technology Assessment (AHTA), AUSTRALIA

<https://health.adelaide.edu.au/adelaide-health-technology-assessment/research-services/publications/>

Browsed following lists (2013 onwards):

reports and monographs – none relevant

protocols – none relevant

Technology Briefs and Prioritising summaries – non relevant

Presentations and abstracts – none relevant

Agency for Care Effectiveness – Singapore

<https://www.ace-hta.gov.sg/>

Browsed lists of technology guidance, horizon scanning reports, scientific publications – 0 relevant reports found

Austrian Institute for Health Technology Assessment

<https://eprints.aihta.at/>

cytomegalovirus – 32 hits browsed – 1 relevant

Canadian Agency for Drugs and Technologies in Health (CADTH), CANADA

<https://www.cadth.ca/>

General search https://www.cadth.ca/search?s=&facets_query=&page=0

1. cytomegalovirus – 21 results, 2 potentially relevant

Browsed projects in progress page and topics under consideration page – 0 relevant

Health Information and Quality Authority, IRELAND HIQA

<https://www.hiqa.ie/reports-and-publications/health-technology-assessments>

Browsed all 152 health technology assessments – none relevant

Scottish Health Technologies Group

<https://shtg.scot/our-advice/>

cytomegalovirus – 0 results

in progress: <https://shtg.scot/what-we-do/work-programme/>

Browsed all – none relevant

Health Technology Wales

<https://healthtechnology.wales/reports-guidance/>

cytomegalovirus – 0 results

National Institute for Health and Care Excellence

<https://www.nice.org.uk/>

General search box:

1. cytomegalovirus – 8 results, none relevant.

4. <https://www.nice.org.uk/guidance/published> Searched for cytomegaloviurs - 2 results, none relevant

in development – none relevant

awaiting development – none relevant

National Institute for Health Research Journals Library

<https://www.journalslibrary.nihr.ac.uk/search/#/>

1. cytomegalovirus – 7 results browsed, none relevant

Belgian Health Care Knowledge Centre

<https://www.kce.fgov.be/en/all-reports-0>

Filtered to HTA reports – browsed 2013-2024 – none

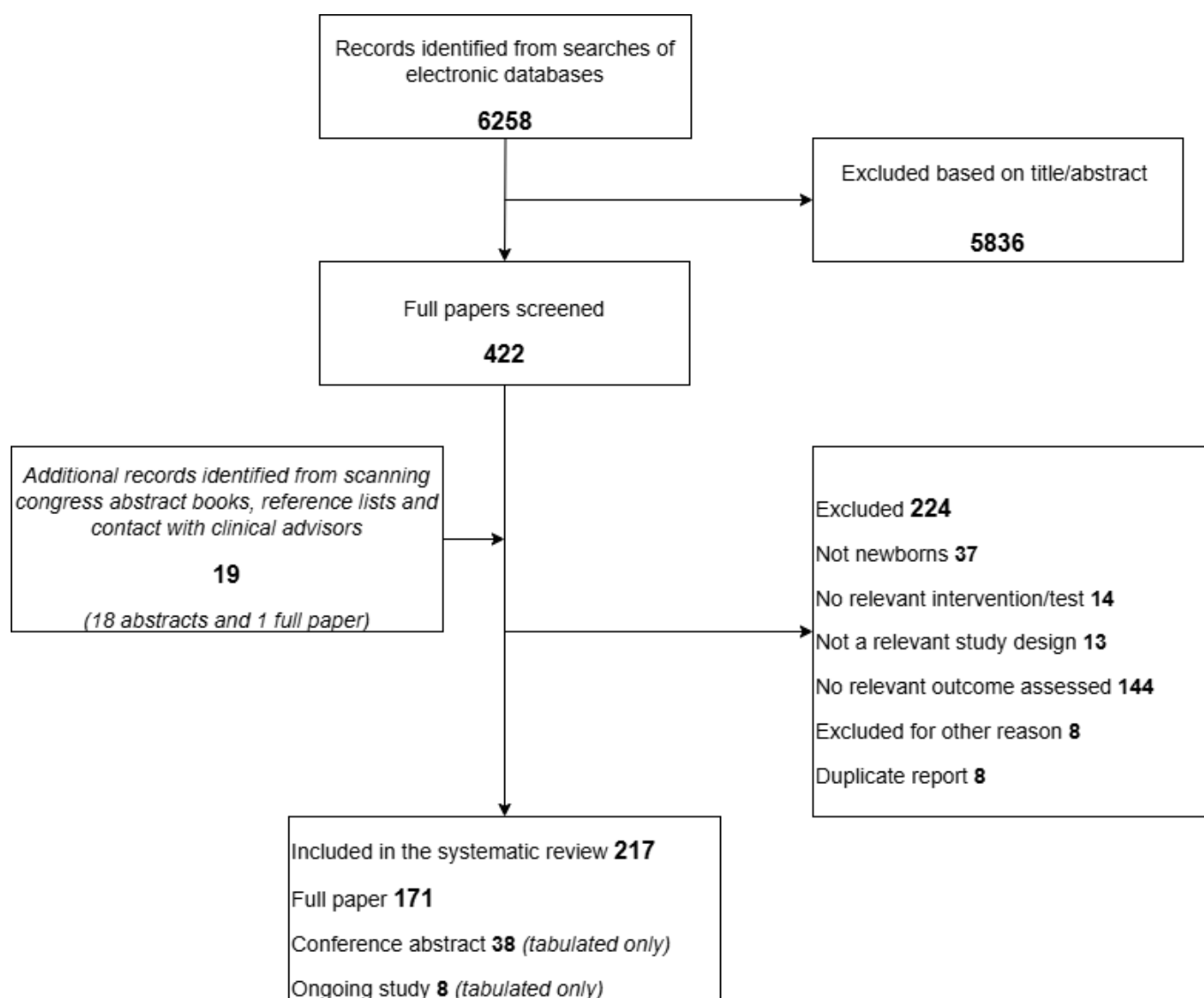
Results from the above searches were imported into Endnote 21 and duplicates removed.

Appendix 2 — Included and excluded studies

PRISMA flowchart

Figure 4 summarises the volume of studies included and excluded at each stage of the review. A total of 217 studies were ultimately judged to be relevant to 1 or more review questions; 171 studies were published in full and were data extracted. There were 8 ongoing studies and 38 studies only reported as conference abstracts; these are tabulated in Appendix 4. Publications that were included or excluded after the review of full text articles are detailed in the tables below.

Figure 4: Flow diagram of the study selection process



Publications included after review of full text articles

The 171 publications included after review of full texts are summarised in Table 24 below.

Table 24: Summary of publications included after review of full text articles, and the question(s) each publication relates to

Study	The test	The screening programme	Implementation criteria	Comments
Atwood et al., 2023 ⁴⁰	Q1		4	Diagnostic accuracy study
Cardoso et al., 2015 ³⁸	Q1		4	Diagnostic accuracy study
Dollard et al., 2021 ³⁹	Q1 & Q2		4	Diagnostic accuracy study
Dunn et al., 2023 ⁴¹	Q1		4	Diagnostic accuracy study
Eventov-Friedman et al., 2019 ³⁵	Q1		4	Diagnostic accuracy study
Exler et al., 2019 ⁵⁰	Q1		4	Diagnostic accuracy study
Gantt et al., 2020 ⁵¹	Q1		4	Diagnostic accuracy study
Gunlemez et al., 2023 ⁴⁸	Q1		4	Diagnostic accuracy study
Huang et al., 2021 ⁵²	Q1		4	Diagnostic accuracy study
Izquierdo et al., 2024 ⁵³	Q1		4	Diagnostic accuracy study
Leruez-Ville et al., 2017 ⁴²	Q1		4	Diagnostic accuracy study
Letamendia-Richard et al., 2022 ⁵⁴	Q1	Q4	4, 11, 12 & 13	Diagnostic accuracy study and Screening evaluation
Pasternak et al., 2020 ⁵⁵	Q1		4	Diagnostic accuracy study
Pinninti et al., 2015 ⁴³	Q1		4	Diagnostic accuracy study
Portet Sulla et al., 2024 ⁵⁶	Q1 & Q3		4 & 7	Association study and diagnostic accuracy study
Ross et al., 2014 ⁵⁷	Q1		4	Diagnostic accuracy study
Ross et al., 2018 ⁴⁴	Q1		4	Diagnostic accuracy study
Shlonsky et al., 2021 ⁴⁹	Q1		4	Diagnostic accuracy study
Stark et al., 2024 ⁵⁸	Q1		4	Diagnostic accuracy study
Tan et al., 2023 ⁴⁵	Q1		4	Diagnostic accuracy study
Wang et al., 2017 ⁴⁶	Q1 & Q2		4	Diagnostic accuracy study
Wunderlich et al., 2023 ⁴⁷	Q1		4	Diagnostic accuracy study
Zheng et al., 2022 ³²	Q1		4	Systematic review
Atkinson et al., 2014 ⁶⁰	Q2		4	Diagnostic accuracy study
Del Valle Penella et al., 2023 ⁶¹	Q2		4	Diagnostic accuracy study
Fourgeaud et al., 2023 ⁶²	Q2		4	Diagnostic accuracy study
Moteki et al., 2018 ⁶³	Q2		4	Diagnostic accuracy study
Noorbakhsh et al., 2020 ⁶⁴	Q2		4	Diagnostic accuracy study
Ross et al., 2017 ⁶⁵	Q2		4	Diagnostic accuracy study
Vercauteren et al., 2020 ⁶⁶	Q2		4	Diagnostic accuracy study
Vives-Onos et al., 2019 ⁶⁷	Q2		4	Diagnostic accuracy study
Wang et al., 2015 ²⁷	Q2		4	Systematic review
Albanna et al., 2013 ⁶⁸			4	Diagnostic accuracy study
Barkai et al., 2013 ⁷³			4	Diagnostic accuracy study
Coltella et al., 2019 ⁶⁹			4	Diagnostic accuracy study
De Monte et al., 2016 ⁷⁰			4	Diagnostic accuracy study
Ligozzi et al., 2016 ⁷¹			4	Diagnostic accuracy study
Reyes et al., 2020 ⁷⁴			4	Diagnostic accuracy study
Wang et al., 2013 ⁷²			4	Diagnostic accuracy study
Airlangga et al., 2024 ¹²⁸	Q3		7	Association study
Alarcon et al., 2013 ⁷⁵	Q3		7	Prognostic accuracy study
Alarcon et al., 2016 ⁸⁰	Q3		7	Prognostic accuracy study
Alarcon et al., 2024 ⁸¹	Q3		7	Prognostic accuracy study
Amir et al., 2016 ¹⁰⁴	Q3		7	Association study
Auriti et al., 2022 ¹²⁹	Q3		7	Association study
Bartlett et al., 2018 ¹⁴⁷	Q3		7	Association study
Bilavsky et al., 2015 ⁷⁶	Q3		7	Association study

Study	The test	The screening programme	Implementation criteria	Comments
Blazquez-Gamero et al., 2019 ⁸²	Q3		7	Prognostic accuracy study
Buca et al., 2021 ¹⁰³	Q3		7	Systematic review
Capretti et al., 2014 ⁸³	Q3		7	Prognostic accuracy study
Capretti et al., 2017 ¹²¹	Q3		7	Association study
Capretti et al., 2021 ¹³⁰	Q3		7	Association study
Castellanos et al., 2022 ⁸⁴	Q3		7	Prognostic accuracy study
Chatzakis et al., 2020 ¹⁰²	Q3		7	Systematic review
Chebib et al., 2022 ⁸⁵	Q3		7	Prognostic accuracy study
Chen et al., 2019 ¹⁶³	Q3		7	Association study
Coscia et al., 2020 ¹¹²	Q3		7	Association study
Craeghs et al., 2021 ⁸⁶	Q3		7	Prognostic accuracy study
Czech-Kowalska et al., 2021 ¹⁶⁹	Q3		7	Association study
De Cuyper et al., 2024 ¹¹⁵	Q3		7	Association study
De Juan et al., 2020 ⁸⁷	Q3		7	Prognostic accuracy study
Demmler-Harrison et al., 2020 ¹¹⁷	Q3		7	Association study
Dhondt et al., 2021 ¹³¹	Q3		7	Association study
Dhondt, 2023 ¹⁵¹	Q3		7	Prognostic accuracy study
Dong et al., 2023 ¹⁶⁴	Q3		7	Association study
Elkan Miller et al., 2021 ¹⁰⁶	Q3		7	Association study
Faure-Bardon et al., 2019 ¹⁰⁷	Q3		7	Association study
Forli et al., 2024 ¹⁵²	Q3		7	Association study
Forner et al., 2015 ⁷⁷	Q3		7	Association study
Foulon et al., 2019 ⁸⁸	Q3		7	Prognostic accuracy study
Fourgeaud et al., 2024 ⁸⁹	Q3		7	Prognostic accuracy study
Giannattasio et al., 2017 ¹²³	Q3		7	Association study
Giannattasio et al., 2017 ⁹⁵	Q3		7	Association study
Giannattasio et al., 2018 ⁹⁰	Q3		7	Prognostic accuracy study
Goderis et al., 2016 ⁷⁸	Q3		7	Association study
Goycochea-Valdivia et al., 2017 ¹⁷⁰	Q3		7	Association study
Hu et al., 2024 ¹³²	Q3		7	Association study
Jin et al., 2019 ¹³³	Q3		7	Association study
Kabani et al., 2023 ¹⁶¹	Q3		7	Association study
Takei et al., 2024 ¹⁵⁷	Q3		7	Association study
Kasztelewicz et al., 2017 ¹⁶⁵	Q3		7	Association study
Kawada et al., 2015 ¹⁶⁰	Q3		7	Association study
Korndewal et al., 2017 ¹⁴⁶	Q3		7	Association study
Keymeulen et al., 2023 ¹¹⁶	Q3		7	Association study
Koyano et al., 2018 ¹⁴⁸	Q3		7	Association study
Kwak et al., 2018 ⁹¹	Q3		7	Prognostic accuracy study
Lanzieri et al., 2018 ¹³⁴	Q3		7	Association study
Lipitz et al., 2013 ¹⁰⁸	Q3		7	Association study
Lo et al., 2022 ¹³⁵	Q3		7	Association study
Lucignani et al., 2021 ⁹²	Q3		7	Prognostic accuracy study
Maes et al., 2017 ¹⁵⁰	Q3		7	Association study
Maltezou et al., 2020 ¹¹⁹	Q3		7	Systematic review
Mardegan et al., 2024 ⁹⁷	Q3		7	Association study
Marsico et al., 2019 ¹⁵⁵	Q3		7	Association study
Massoud et al., 2023 ¹⁰⁹	Q3		7	Association study
Medoro et al., 2024 ¹⁶⁶	Q3		7	Association study
Meyer et al., 2017 ¹⁵⁶	Q3		7	Association study
Minsart et al., 2020 ¹⁴⁵	Q3		7	Association study
Nigro et al., 2020 ¹¹⁰	Q3		7	Association study

Study	The test	The screening programme	Implementation criteria	Comments
Nishida et al., 2020 ⁹³	Q3		7	Prognostic accuracy study
Noorbakhsh et al., 2022 ¹²⁰	Q3		7	Association study
Oosterom et al., 2015 ⁹⁸	Q3		7	Association study
Ouellette et al., 2020 ¹⁶²	Q3		7	Prognostic accuracy study
Pati et al., 2013 ¹⁶⁷	Q3		7	Association study
Pinninti et al., 2016 ¹⁴⁰	Q3		7	Association study
Poletti de Chaurand et al., 2024 ¹⁰¹	Q3	Q4	7, 11 & 13	Association study and screening evaluation
Puhakka et al., 2017 ¹¹¹	Q3		7	Association study
Puhakka et al., 2022 ¹¹⁸	Q3		7	Association study
Riga et al., 2018 ¹²⁷	Q3		7	Systematic review
Rohren et al., 2024 ¹⁵³	Q3		7	Association study
Rovito et al., 2017a ¹⁴³	Q3		7	Association study
Rovito et al., 2017b ¹⁴⁴	Q3		7	Association study
Rovito et al., 2017c ¹²⁴	Q3		7	Association study
Rovito et al., 2018 ¹⁶⁸	Q3		7	Association study
Royackers et al., 2013 ¹³⁶	Q3		7	Association study
Salome et al., 2020 ¹⁵⁸	Q3		7	Association study
Salome et al., 2023 ¹³⁷	Q3		7	Association study
Scaramuzzino et al., 2022 ¹¹³	Q3		7	Association study
Seneviratne et al., 2022 ¹³⁸	Q3		7	Association study
Simonazzi et al., 2014 ¹⁴¹	Q3		7	Association study
Smyrli et al., 2024 ¹²⁵	Q3		7	Systematic review
Soriano-Ramos et al., 2024 ⁹⁹	Q3		7	Association study
Tapasak et al., 2022 ¹⁴²	Q3		7	Association study
Topham et al., 2019 ¹⁴⁹	Q3		7	Association study
Townsend et al., 2013 ¹²²	Q3		7	Association study
Turiziani Colonna et al., 2020 ¹¹⁴	Q3		7	Association study
Vande Walle et al., 2023 ¹⁰⁰	Q3		7	Association study
Vande Walle, 2024 ⁷⁹	Q3		7	Systematic review
Vande Walle, 2024 ⁹⁴	Q3		7	Prognostic accuracy study
Vila-Bedmar et al., 2024 ⁹⁶	Q3		7	Association study
Vos et al., 2021 ¹²⁶	Q3		7	Systematic review
Wang et al., 2021 ¹⁵⁹	Q3		7	Association study
Yamaguchi et al., 2022 ¹⁵⁴	Q3		7	Prognostic accuracy study
Zavattoni et al., 2014 ¹⁰⁵	Q3		7	Association study
Zavattoni et al., 2016 ¹³⁹	Q3		7	Association study
Ari-Even Roth et al., 2017 ¹⁸⁶		Q4	11 & 13	Screening evaluation
Barkai et al., 2014 ³⁶		Q4	11 & 13	Screening evaluation
Beswick et al., 2021 ¹⁹¹		Q4	11, 12 & 13	Screening evaluation
Chiereghin et al., 2022 ¹⁷⁶		Q4	11 & 13	Screening evaluation
Chung et al., 2023 ¹⁸⁷		Q4	11 & 13	Screening evaluation
Demortier et al., 2021 ¹⁸²		Q4	11, 12 & 13	Screening evaluation
Diener et al., 2017 ¹⁹⁵		Q4	11 & 13	Screening evaluation
Dunn et al., 2025 ¹⁷⁷		Q4	11, & 13	Screening evaluation
Forli et al., 2024 ²⁰⁷		Q4	11 & 13	Screening evaluation
Fourgeaud et al., 2022 ¹⁹⁶		Q4	11 & 13	Screening evaluation
Fowler et al., 2017 ¹⁹²		Q4	11 & 13	Screening evaluation
Fujii et al., 2017 ²⁰²		Q4	11 & 13	Screening evaluation
Hilditch et al., 2018 ¹⁷³		Q4	11 & 13	Systematic review
Kadambari et al., 2015 ¹⁹⁴		Q4	11, 12 & 13	Screening evaluation
Kaye et al., 2024 ¹⁷⁹		Q4	11 & 13	Screening evaluation
Kruc et al., 2024 ¹⁸⁰		Q4	11 & 13	Screening evaluation
Lu et al., 2018 ²⁰⁴		Q4	11 & 13	Screening evaluation

Study	The test	The screening programme	Implementation criteria	Comments
Masarweh et al., 2021 ¹⁹⁹		Q4	11 & 13	Screening evaluation
McCrary et al., 2020 ²⁰⁰		Q4	11 & 13	Screening evaluation
Medoro et al., 2021 ¹⁸⁹		Q4	11 & 13	Screening evaluation
Merav et al., 2024 ¹⁸³		Q4	11 & 13	Screening evaluation
Palma et al., 2021 ²⁰¹		Q4	11 & 13	Screening evaluation
Papaevangelou et al., 2019 ¹⁸⁴		Q4	11 & 13	Screening evaluation
Pollick et al., 2024 ¹⁷⁴		Q4	11 & 13	Systematic review
Puhakka et al., 2019 ¹⁷⁵		Q4	11 & 13	Screening evaluation
Rawlinson et al., 2018 ¹⁹⁷		Q4	11 & 13	Screening evaluation
Ronner et al., 2022 ¹⁸⁸		Q4	11 & 13	Screening evaluation
Tan et al., 2023 ²⁰⁵		Q4	11 & 13	Screening evaluation
Vancor et al., 2019 ¹⁹⁰		Q4	11 & 13	Screening evaluation
Webb et al., 2024 ²⁰⁶		Q4	11, 12 & 13	Screening evaluation
Williams et al., 2014 ¹⁹⁸		Q4	11 & 13	Screening evaluation
Yamada et al., 2020 ¹⁷¹		Q4	11 & 13	Screening evaluation
Yamaguchi et al., 2017 ¹⁸¹		Q4	11 & 13	Screening evaluation
Yamamoto et al., 2020 ¹⁷⁸		Q4	11 & 13	Screening evaluation
Zeytinoglu et al., 2019 ¹⁸⁵		Q4	11 & 13	Screening evaluation
Zhang et al., 2023 ²⁰³		Q4	11 & 13	Screening evaluation

Publications excluded after review of full text articles

Of the 422 publications included after the review of titles and abstracts, 224 were ultimately judged not to be relevant to this review. These publications, along with reasons for exclusion, are listed in Table 25.

Table 25: Publications excluded after review of full text articles

Reference	Reason for exclusion
Akiko et al., 2021 ²⁰⁸	Population
Akiva et al., 2023 ²⁰⁹	Outcome
Albano et al., 2014 ²¹⁰	No eligible test
Albano et al., 2016 ²¹¹	Outcome
Albano et al., 2017 ²¹²	Outcome
Alifieraki et al., 2022 ²¹³	Outcome
Almeida et al., 2023 ²¹⁴	Outcome
Alwan et al., 2019 ²¹⁵	Outcome
Amin et al., 2021 ²¹⁶	Population
Annabelle et al., 2023 ²¹⁷	Outcome
Anonymous, 2016 ²¹⁸	Other (erratum for other study)
Arancibia et al., 2023 ²¹⁹	No eligible test
Arellano-Galindo et al., 2014 ²²⁰	Outcome
Banjac et al., 2022 ²²¹	Study design
Barriga et al., 2024 ²²²	Outcome
Ben Shoham et al., 2023 ²²³	Outcome
Berg et al., 2019 ²²⁴	Population
Bergevin et al., 2015 ²²⁵	No eligible test
Beswick et al., 2019 ²²⁶	No eligible test
Bilavsky et al., 2016 ²²⁷	Population
Bilavsky et al., 2017 ²²⁸	Outcome
Blazquez-Gamero et al., 2020 ²²⁹	Outcome
Blazquez-Gamero et al., 2021 ²³⁰	Duplicate report
Boppana et al., 2016 ²³¹	Population
Boscarino et al., 2024 ²³²	No eligible test

Reference	Reason for exclusion
Branas et al., 2015 ²³³	Outcome
Brunet, 2024 ²³⁴	Study design
Campione et al., 2022 ²³⁵	No eligible test
Canadian Agency for Drugs and Technologies in Health, 2024 ²³⁶	Outcome
Central Hospital Nancy France, 2023 ²³⁷	Outcome
Chasqueira et al., 2023 ²³⁸	Outcome
Chauhan et al., 2021 ²³⁹	Outcome
Chiereghin et al., 2017 ²⁴⁰	Outcome
Chisholm et al., 2013 ²⁴¹	Outcome
Chisholm et al., 2014 ²⁴²	Population
Choudhary et al., 2015 ²⁴³	Population
Correa et al., 2016 ²⁴⁴	Outcome
Cushing et al., 2022 ²⁴⁵	Outcome
Czech-Kowalska et al., 2021 ²⁴⁶	Outcome
Dallaire et al., 2021 ²⁴⁷	Outcome
Dallaire et al., 2022 ²⁴⁸	Population
Dar et al., 2017 ²⁴⁹	Outcome
De Cuyper et al., 2023 ²⁵⁰	Outcome
de Vries et al., 2013 ²⁵¹	Population
Del Valle Penella et al., 2023 ²⁵²	Outcome
Demmler-Harrison G J, 2021 ²⁵³	Study design
Dharnidhar et al., 2019 ²⁵⁴	Outcome
Dimopoulou et al., 2020 ²⁵⁵	Outcome
Dobbins et al., 2019 ²⁵⁶	Outcome
Dougherty et al., 2022 ²⁵⁷	Population
Dreher et al., 2013 ²⁵⁸	Outcome
Dreher et al., 2014 ¹⁷²	No eligible test
Ebrahimi-Rad et al., 2017 ²⁵⁹	Outcome
Eres Saritas et al., 2023 ²⁶⁰	Outcome
Eventov-Friedman et al., 2014 ²⁶¹	Outcome
Farnan et al., 2023 ²⁶²	Outcome
Faure-Bardon et al., 2020 ²⁶³	Population
Fernandes et al., 2021 ²⁶⁴	No eligible test
Folkins et al., 2013 ²⁶⁵	Outcome
Gandhoke et al., 2013 ²⁶⁶	Population
Gantt et al., 2018 ²⁶⁷	Study design
Garnham et al., 2023 ²⁶⁸	Outcome
Gillespie et al., 2023 ²⁶⁹	Study design
Goderis et al., 2013 ²⁷⁰	Duplicate report
Goel et al., 2018 ²⁷¹	Outcome
Goetzl et al., 2019 ²⁷²	Population
Gonzalez Lopez et al., 2015 ²⁷³	Outcome
Gorzer et al., 2015 ²⁷⁴	Outcome
Gunes et al., 2024 ²⁷⁵	Outcome
Gunkel et al., 2014 ³⁷	Population
Gupta et al., 2023 ²⁷⁶	Population
Gutierrez Posso et al., 2023 ²⁷⁷	Outcome
Gwee et al., 2014 ²⁷⁸	Outcome
Hadar et al., 2016 ²⁷⁹	Outcome
Hadar et al., 2017 ²⁸⁰	Outcome
Hamid et al., 2022 ²⁸¹	Outcome
Hamilton et al., 2014 ²⁸²	Population
Hasan et al., 2021 ²⁸³	Outcome
Hernandez-Alvarado et al., 2023 ²⁸⁴	No eligible test
Hirth et al., 2018 ²⁸⁵	Other (insufficient information to screen against inclusion criteria)
Hirth et al., 2021 ²⁸⁶	Other (no data relating to review questions)
Hranilovich et al., 2016 ²⁸⁷	Outcome

Reference	Reason for exclusion
Hranilovich et al., 2020 ²⁸⁸	Population
Hu et al., 2021 ²⁸⁹	Outcome
Huang et al., 2024 ²⁹⁰	Other (no relevance to review questions)
Ikuta et al., 2013 ²⁹¹	Other (no relevance to review questions)
Imafuku et al., 2020 ²⁹²	Outcome
Ishida et al., 2015 ²⁹³	Study design
Ismail et al., 2013 ²⁹⁴	Outcome
Izquierdo et al., 2023 ²⁹⁵	Duplicate report
Javid et al., 2016 ²⁹⁶	Outcome
Jedlinska-Pijanowska et al., 2020 ²⁹⁷	Outcome
Jedlinska-Pijanowska et al., 2021 ²⁹⁸	Outcome
Kachramanoglou et al., 2021 ²⁹⁹	Outcome
Kakkar et al., 2015 ³⁰⁰	Other (insufficient information to screen against inclusion criteria)
Karamchandani et al., 2022 ³⁰¹	Outcome
Karimi et al., 2023 ³⁰²	Outcome
Kawano et al., 2016 ³⁰³	Outcome
Keymeulen et al., 2022 ³⁰⁴	Outcome
Khalil et al., 2016 ³⁰⁵	Outcome
Kim et al., 2023 ³⁰⁶	Population
Kipfmüller et al., 2018 ³⁰⁷	Outcome
Kitamura et al., 2022 ³⁰⁸	Population
Kobayashi et al., 2015 ³⁰⁹	Outcome
Kohmer et al., 2019 ³¹⁰	Population
Koontz et al., 2015 ³¹¹	Population
Korndewal et al., 2016 ³¹²	Outcome
Kravchenko et al., 2015 ³¹³	Population
Kravchenko et al., 2021 ³¹⁴	Outcome
Krishna et al., 2020 ³¹⁵	No eligible test
Kruc et al., 2022 ³¹⁶	Outcome
Kyriakopoulou et al., 2023 ³¹⁷	Outcome
Lakhan et al., 2023 ³¹⁸	Outcome
Lam et al., 2017 ³¹⁹	Outcome
Lee et al., 2013 ³²⁰	Population
Leruez-Ville et al., 2013 ³²¹	Population
Leruez-Ville et al., 2020 ³²²	Outcome
Levit et al., 2020 ³²³	Outcome
Lewald et al., 2023 ³²⁴	Outcome
Liao et al., 2023 ³²⁵	Outcome
Lilleri et al., 2023 ³²⁶	Outcome
Lim et al., 2013 ³²⁷	Population
Lipitz et al., 2020 ³²⁸	Outcome
Lopez et al., 2017 ³²⁹	Outcome
Luck et al., 2016 ³³⁰	Outcome
Mack et al., 2017 ³³¹	Study design
Mahajan et al., 2018 ³³²	Outcome
Maltezou Panagiota et al., 2019 ³³³	Duplicate report
Mappa et al., 2023 ³³⁴	Population
Mareri et al., 2013 ³³⁵	Outcome
Martinez et al., 2021 ³³⁶	Outcome
Mastutik et al., 2022 ³³⁷	Population
Matsuo et al., 2014 ³³⁸	Outcome
McCrary et al., 2019 ³³⁹	Outcome
Medoro et al., 2017 ³⁴⁰	Duplicate report
Medoro et al., 2019 ³⁴¹	Other (insufficient information to screen against inclusion criteria)
Medoro et al., 2019 ³⁴²	Outcome
Medoro et al., 2020 ³⁴³	Outcome
Melamed et al., 2020 ³⁴⁴	Outcome

Reference	Reason for exclusion
Meogrossi et al., 2023 ³⁴⁵	Population
Merino-Hernandez et al., 2023 ³⁴⁶	Outcome
Minami et al., 2021 ³⁴⁷	No eligible test
Mirsalehi et al., 2024 ³⁴⁸	Outcome
Morimoto et al., 2023 ³⁴⁹	Study design
Mu et al., 2023 ³⁵⁰	Population
Mujtaba et al., 2016 ³⁵¹	Outcome
Nagel et al., 2020 ³⁵²	Outcome
Namdeo et al., 2013 ³⁵³	Outcome
Nishida et al., 2016 ³⁵⁴	Outcome
Nishida et al., 2016 ³⁵⁵	Outcome
Niz Xavier et al., 2015 ³⁵⁶	Outcome
Noorbakhsh et al., 2020 ³⁵⁷	Outcome
Noorbakhsh et al., 2023 ³⁵⁸	Outcome
Norton et al., 2020 ³⁵⁹	Study design
Ohyama et al., 2019 ³⁶⁰	Outcome
Oneko et al., 2020 ³⁶¹	Outcome
Orb et al., 2024 ³⁶²	Outcome
Palma et al., 2019 ³⁶³	Outcome
Palma et al., 2023 ³⁶⁴	Outcome
Paradowska et al., 2013 ³⁶⁵	Outcome
Paradowska et al., 2014 ³⁶⁶	Outcome
Paradowska et al., 2019 ³⁶⁷	Outcome
Parsons et al., 2021 ³⁶⁸	Other (insufficient information to screen against inclusion criteria)
Pathirana et al., 2019 ³⁶⁹	Outcome
Pellegrinelli et al., 2019 ³⁷⁰	Population
Peterson et al., 2020 ³⁷¹	Outcome
Pitlick et al., 2015 ³⁷²	Outcome
Rahniayu et al., 2022 ³⁷³	Population
Raynor et al., 2021 ³⁷⁴	Outcome
Reina et al., 2014 ³⁷⁵	Outcome
Renaud et al., 2018 ³⁷⁶	Outcome
Reynders et al., 2024 ³⁷⁷	Outcome
Ronchi et al., 2020 ³⁷⁸	Outcome
Roy et al., 2024 ³⁷⁹	Population
Rymer-Haskel et al., 2019 ³⁸⁰	Outcome
Sahiner et al., 2015 ³⁸¹	Outcome
Sakamoto et al., 2015 ³⁸²	Outcome
Sarkar et al., 2019 ³⁸³	Outcome
Schleiss et al., 2023 ³⁸⁴	Population
Schleiss et al., 2023 ³⁸⁵	Population
Shahar-Nissan et al., 2022 ³⁸⁶	Outcome
Shahrokhi et al., 2020 ³⁸⁷	Outcome
Shears et al., 2022 ³⁸⁸	Outcome
Smiechura et al., 2014 ³⁸⁹	Outcome
Smiljkovic et al., 2017 ³⁹⁰	Outcome
Smiljkovic et al., 2019 ³⁹¹	Outcome
Smiljkovic et al., 2020 ³⁹²	Outcome
Smyrli et al., 2022 ³⁹³	Duplicate report
Southern Illinois University, 2016 ³⁹⁴	Outcome
Spalletti-Cernia et al., 2016 ³⁹⁵	No eligible test
Ssentongo et al., 2021 ³⁹⁶	Outcome
Stoyell et al., 2024 ³⁹⁷	Outcome
Stoykova et al., 2022 ³⁹⁸	Outcome
Suarez et al., 2023 ³⁹⁹	Outcome
Thompson et al., 2020 ⁴⁰⁰	Outcome
Torii et al., 2024 ⁴⁰¹	Outcome
Torre et al., 2016 ⁴⁰²	Outcome

Reference	Reason for exclusion
Turner et al., 2014 ⁴⁰³	Outcome
Uematsu et al., 2016 ⁴⁰⁴	Outcome
Vande Walle et al., 2021 ⁴⁰⁵	Outcome
Vargas et al., 2022 ⁴⁰⁶	No eligible test
Viswanathan et al., 2019 ⁴⁰⁷	Outcome
Vives-Onos et al., 2014 ⁴⁰⁸	Outcome
Wang et al., 2017 ⁴⁰⁹	Population
Wang et al., 2022 ⁴¹⁰	Outcome
Waters et al., 2014 ²⁹	Outcome
Webb et al., 2022 ⁴¹¹	Duplicate report
Wei et al., 2014 ¹⁹³	Outcome
Williams et al., 2015 ⁴¹²	Study design
Wissel et al., 2017 ⁴¹³	Study design
Wolf, 2024 ⁴¹⁴	Study design
Wu et al., 2022 ⁴¹⁵	Outcome
Wu et al., 2022 ⁴¹⁶	Population
Wujcicka et al., 2017 ⁴¹⁷	Outcome
Yamaguchi et al., 2023 ⁴¹⁸	Outcome
Yamaguchi et al., 2023 ⁴¹⁹	Outcome
Yang et al., 2021 ⁴²⁰	Outcome
Yao, 2018 ⁴²¹	Outcome
Zavattoni et al., 2014 ⁴²²	Study design
Zavattoni et al., 2018 ⁴²³	Outcome
Zeng et al., 2017 ⁴²⁴	Population
Zhang et al., 2013 ⁴²⁵	No eligible test
Zhang et al., 2016 ⁴²⁶	Outcome
Zheng et al., 2022 ⁴²⁷	Duplicate report

Appendix 3 — Summary and appraisal of studies relating to Criterion 7

Data Extraction

Detailed characteristics of studies relevant to criterion 7 are tabulated below.

Table 26: Systematic reviews relevant to criterion 7

Study details	Test/characteristic assessed (of relevance to this review)	Population (of relevance to this review)	Outcome (and timeframe)	Results/conclusions and comments
Vande Walle, 2024 ⁷⁹	Postnatal brain MRI A 1.5 or 3T scanner was used in the axial, coronal and/or sagittal planes. Sequences included T2 ($\pm$ fat saturation), T1, DWI (with ADC) and FLAIR. Mild sedation, anaesthesia or a vacuum mattress were sometimes used to reduce motion artefacts. Postnatal scoring systems described in some studies. Prenatal MRI was included in the paper, but details of prenatal MRI have not been	Newborns and children with cCM V. No definition of upper age limit for baseline MRI	Neurodevelopmental impairments (cognitive and motor), sensorineural hearing loss, epilepsy, behavioural problems, visual impairments. Timeframes not consistently described.	An analysis of the associations between prenatal MRI and later outcomes was included in the paper, but details of these associations have not been included in this review as they are out of scope. Out of the 20 studies included in the SR, only 7 studies covered the relevant analyses between post-natal MRI and the outcomes below. No quantitative or narrative synthesis were undertaken, and associations between postnatal MRI and later outcome were merely given per study. These results have been summarised below. The numbers of participants in each included study were not presented in the SR [Vande Walle (2024)]. Other reporting deficits described below relate to perceived limitations in the SR, but some may be related to similar reporting limitations in the original sources. <u>Neurodevelopmental outcomes</u> <i>Kwak et al:</i> MRI score was associated with epilepsy ($p=0.004$). Polymicrogyria on MRI was associated with epilepsy ($p=0.012$) and neurodevelopmental impairment (NDI) ($p=0.01$). Ventriculomegaly ($p=0.045$), calcification ($p=0.006$), and white matter abnormalities (no p value given) on MRI each associated with epilepsy. For all outcomes, outcome follow up length was not presented in the SR, and presence of any statistical adjustment was unclear.

Study details	Test/characteristic assessed (of relevance to this review)	Population (of relevance to this review)	Outcome (and timeframe)	Results/conclusions and comments
	included in this review as they are out of scope.			<i>Nishida et al</i>: Ventriculomegaly ($p<0.001$), periventricular cysts ($p<0.005$), and white matter abnormalities ($p=0.02$) were associated with NDI. NDI could be predicted (Youden index of 0.78) by presence of 2 out of the 3 MRI abnormalities. For all outcomes, follow up length was not presented in the SR, and presence of statistical adjustment was unclear. <i>Vande Walle</i>: Isolated white matter abnormalities were associated with lower motor scores ($p=0.01$), and impaired mental development ($p=0.07$). For all outcomes, follow up length was not presented in the SR, and presence of statistical adjustment was unclear. <u>Hearing outcomes</u> <i>Alarcon</i>: Temporal pole lesions (but not white matter lesions) were significantly associated with sensorineural hearing loss (SNHL), but follow up length was not presented in the SR. Results appear to be based on an adjusted regression analysis. <i>Craeghs</i>: There was a [presumably univariate] significant association between abnormal MRI and hearing loss ($p=0.007$), but follow up length was not presented in the SR. On multivariate analysis only calcifications were associated with hearing loss. <i>De Cuyper</i>: Periventricular cysts were independently and significantly associated with hearing loss, but follow up length was not presented in the SR. <i>Kwak</i>: No associations between MRI abnormalities and SNHL were observed, but follow up length was not presented in the SR. <i>Vande Walle</i>: Associations between white matter lesions and hearing loss ($p=0.008$) were observed, but follow up length was not presented in the SR. <u>Combined neurodevelopmental/hearing loss outcomes</u> <i>Blazquez-Gamero</i>: Abnormal MRI was strongly associated with NDI/hearing loss sequelae ($p=0.028$) at 12 months. This was an adjusted association, based on multivariable logistic regression. <u>Other information</u> MRI was also compared to US, but only in terms of the anomalies detected by each. No analysis was made of the differential ability to predict later outcomes.

Study details	Test/characteristic assessed (of relevance to this review)	Population (of relevance to this review)	Outcome (and timeframe)	Results/conclusions and comments
				No data on predictive accuracy of post-natal MRI were presented. Protocol not published as far as is known.
Chatzakis, 2020 ¹⁰²	Timing of infection: first vs second vs third trimester	Neonates of pregnant women with confirmed primary CMV infection in the immediate preconception period (up to 12 weeks preconception) or during pregnancy were included.	The only outcome relevant was severe sensorineural hearing loss (SNHL) or neurodevelopmental impairment at follow-up (from 6 months to 5 years).	Although 17 studies were included in the review, only 6 studies^{88, 107, 108, 111, 428, 429} contained the relevant outcome of sensorineural hearing loss (SNHL) and/or neurodevelopmental impairment at follow-up, and so only results relating to these are included here. Data from these 6 studies (comprising 738 participants) were synthesised using meta-analysis to evaluate the association of time of maternal infection and severe SNHL or neurodevelopmental impairment at follow-up. Maternal infection in the first trimester resulted led to a pooled prevalence of SNHL or neurodevelopmental impairment of 22.8% (95% Ct 15.4-30.2; $I^2=27.6\%$) at follow up. Maternal infections in the second and third trimesters led to smaller rates of 0.1% (95% Ct 0-0.8; $I^2=24\%$) and 0% (95% Ct 0-2.1; $I^2=0\%$) at follow up, respectively. No evidence of sub-grouping or meta-regression by potential outcome modifiers was presented. The meta-analyses were flawed by not evaluating the between-group effects (for example a more valid approach might have analysed the odds ratios for later impairment of each of second and third trimester groups with reference to the first trimester group), and instead provided the within-group prevalences. Although this approach prevents a rigorous analysis of group difference, the large magnitude of effect in this analysis provides some evidence that timing of infection may be an important factor determining the likelihood of neurodevelopmental and hearing sequelae. A further (sensitivity) meta-analysis restricted to the 4 studies that were conducted prospectively, and had used IgG avidity assays for the timing of maternal infection produced similar results. No data on predictive accuracy of timing of infection were presented. Protocol published on PROSPERO.

Study details	Test/characteristic assessed (of relevance to this review)	Population (of relevance to this review)	Outcome (and timeframe)	Results/conclusions and comments
Maltezou, 2020 ¹¹⁹	Type of maternal infection: primary infection (PI; where the mother is initially infected during the index pregnancy) or non-primary infection (NPI; where the mother is initially infected before the index pregnancy).	Congenitally CMV infected children born following maternal PI or NPI	Relevant outcomes were unilateral SNHL, bilateral SNHL, or other neurologic outcomes. Follow up ranged from 1-5 years (set at a minimum of 12 months in SR protocol).	The paper reported some results for analyses from 9 studies (n=879) that are not reported here as they are outside the scope of the current review. However, 8 of these studies (n=835) produced data that were relevant to the meta-analyses of 'later outcomes'. Meta-analyses showed that maternal infection status (primary versus non-primary infection) did not influence the risk of developing unilateral SNHL [pooled OR of 8 studies = 0.80 (95 % CI 0.54–1.20, p = 0.282) I²=0%], bilateral SNHL [pooled OR of 5 studies = 0.66 (95 % CI 0.30–1.42, p = 0.283) I²=36.2%] or other neurologic outcomes [pooled OR of 4 studies = 0.73 (95 % CI 0.43–1.25, p = 0.518) I²=0%]. This paper suggests that type of maternal infection is not an important factor determining hearing or neurological outcomes at follow up. No data on predictive accuracy of type of maternal infection were presented. Protocol published on PROSPERO.
Smyrli, 2024 ¹²⁵	Children with asymptomatic cCMV at birth.	Children with asymptomatic and symptomatic cCMV	Neurodevelopmental outcomes. Age at last follow up ranged from 1 month to 18 years.	All 28 studies (number of participants unclear) in the SR evaluated the outcomes for children with asymptomatic cCMV. For the 19/28 studies (number of participants unclear) that included an analysis of asymptomatic versus <i>symptomatic cCMV</i>, 16/19 studies showed that adverse neurodevelopmental outcomes were less common among children with asymptomatic cCMV at birth. Where reported, antivirals were received by children with asymptomatic cCMV in 6 studies and by children with symptomatic cCMV in 11 studies. Although this might create bias, the bias would favour the symptomatic group, which would be expected to reduce rather than augment the overall finding that asymptomatic children have better long-term outcomes. No formal meta-analyses were undertaken, as a narrative approach had been decided pre-hoc based on the anticipated heterogeneity between studies. No data on predictive accuracy of type of maternal infection were presented.

Study details	Test/characteristic assessed (of relevance to this review)	Population (of relevance to this review)	Outcome (and timeframe)	Results/conclusions and comments
				Protocol published on PROSPERO.
Buca,2021 ¹⁰³	Trimester of infection	Children with cCMV and normal prenatal USat diagnosis	Neurodevelopment/ hearing. Age at last follow up ranged from 1 month to 10 years (median)	26 studies were included in the systematic review, but only a proportion of these were relevant to the subgroup analysis of the association between trimester of infection and neurodevelopmental/hearing sequelae. It is unclear which (or how many) of these eligible studies there were, as the number of studies with data on trimester of infection in the study characteristics table does not tally with the number of studies with relevant data in the results section. The pooled proportion of children with abnormal neurodevelopmental outcome was 5.4% (95% Ct 2.3 to 9.8) in 7 studies evaluating first trimester infection, 0% (95% Ct 0 to 7) in 6 studies evaluating second trimester infection, and 0% (95% Ct 0 to 7.7) in 4 studies evaluating third trimester infection. The pooled proportion of children with hearing problems was 11.4% (95% Ct 6.7 to 17.1) in 7 studies evaluating first trimester infection, 7% (95% Ct 2.1 to 14.5) in 6 studies evaluating second trimester infection, and 0% (95% Ct 0 to 7.7) in 4 studies evaluating third trimester infection. No data on predictive accuracy of timing of maternal infection were presented. No evidence that the protocol was published.
Vos, 2021 ¹²⁶	Symptoms at birth	Children with cCMV	Late-onset hearing loss, up to 18 years	65 articles were included, but only 17 were relevant for assessing the association between cCMV symptoms at birth and late-onset hearing loss. Other studies were not relevant as they covered post-natal CMV, hearing loss at birth, or did not analyse the association between symptoms at birth and outcomes. This narrative review did not form any firm conclusions about an association between symptomatic status at birth and subsequent hearing loss in children with cCMV; late-onset permanent childhood hearing loss occurred in children in all cCMV subgroups (symptomatic, asymptomatic and unspecified). No data on predictive accuracy of symptoms at birth were presented.

Study details	Test/characteristic assessed (of relevance to this review)	Population (of relevance to this review)	Outcome (and timeframe)	Results/conclusions and comments
				Protocol published on PROSPERO.
Riga, 2018 ¹²⁷	Symptoms at birth	Children with cCM V	CMV-induced hearing loss, up to 6-10 years of age	11 articles were included. It is unclear how many of these were included in the synthesis of results. A synthesis was carried out on a pooled population of 1089 children with cCM V. Among asymptomatic participants SNHL was diagnosed in 8.9% (70/784), while for symptomatic children the percentage raised to 44.2% (107/242). This synthesis appears to be a simple summation of study results rather than a proper meta-analysis. No data on predictive accuracy of symptoms at birth were presented. No evidence that the protocol was published.

Table 27: Prognostic accuracy studies relevant to criterion 7

Study	Details of neonatal sample, cCMV diagnosis method/timing and timing of baseline predictive tests	Specific sequelae predicted with follow up	Baseline test(s) to predict the defined sequelae, with thresholds defining a positive test	n	TP	FN	FP	TN	Sen (%)	Spec (%)	PPV (%)	NPV (%)	AUC
Alarcon et al. 2013 ⁷⁵	Symptomatic neonates only. 3 deaths during follow-up. 15 with at least 1 sequel of neurodevelopmental delay 17/26 patients received antivirals Identification of CMV or viral DNA in urine or blood, or detection of CMV IgM or viral antigen in blood during first 2 weeks. CSF baseline tests performed at mean 8.5 days. Timing of other baseline tests unclear.	Neurodevelopmental delay (in terms of motor function, cognition, behaviour, hearing, vision, and epilepsy) at mean 8.7 years	adjusted microcephaly (threshold unclear)	26	-	-	-	-	44	100	100	44	0.72
			CSF conc. >7.9mg/L	26	-	-	-	-	69	100	100	63	0.84
			Neuroimaging (on any of MRI, US or CT) score of ≥ 2 on Noyola scale [#]	26	-	-	-	-	61	100	100	53	0.80
			CSF conc. >7.9mg/L OR adjusted microcephaly (threshold unclear)	26	-	-	-	-	82	100	100	70	0.91
			adjusted microcephaly (threshold unclear) OR neuroimaging (on any of MRI, US or CT) score of ≥ 2 on Noyola scale	26	-	-	-	-	66	100	100	57	0.83
			CSF >7.9mg/L OR neuroimaging (on any of MRI, US or CT) score of ≥ 2 on Noyola scale	26	-	-	-	-	87	100	100	77	0.92
Alarcon et al. 2016 ⁸⁰	Symptomatic neonates only. 3 deaths during follow-up. 15 with at least 1 sequel of neurodevelopmental delay 17/26 patients received antivirals Identification of CMV or viral DNA in urine or blood, or detection of CMV IgM or viral antigen in blood during first 2 weeks. Timing of baseline tests unclear.	Neurodevelopmental delay (in terms of motor function, cognition, behaviour, hearing, vision, and epilepsy) at mean 8.7 years	Neuroimaging (on any of MRI, US or CT) $\geq$ score of 2 on 'new scale' ^{**} (contains additional characteristics in each scoring category, allowing more scores of 2 or above)	26	-	-	-	-	77	100	100	66	0.88
			Neuroimaging (on any of MRI, US or CT) score of ≥ 2 on Noyola scale [#]	26	-	-	-	-	61	100	100	53	0.80

Alarcon et al. 2024 ⁸¹	103 symptomatic, 57 asymptomatic 41 with moderate/severe disability at follow up 96/160 received antivirals Diagnosis method of cCMV not reported. Baseline neuroimaging in first 3 months of life	Moderate/severe disabilities: neurodevelopmental impairments (including hearing, CP, cognition, behavioral, epilepsy, visual) at median 4 years	Neuroimaging (US or MRI) composite score (as 'new' score in Alarcon, 2016**) $\geq$ score of 2	160	-	-	-	-	90	69	63	92	0.89
			One or more temporal pole white matter abnormalities (TPWMAs) observed	160	-	-	-	-	78	69	35	94	0.73
Blazquez-Gamero et al. 2019 ⁸²	77 symptomatic, 30 asymptomatic 37 with sequelae during follow up. 96/107 received antivirals Only children diagnosed with cCMV during the fetal period or at birth (first 14 days of life) were included in this study. Retrospective DBS after 14 days Excluded. MRI done in first 3 months of life.	Any sequelae (hearing or neurological) at 12 months	Abnormal US AND Abnormal MRI (with any category of symptoms at birth). ##	71	31	6	7	27	84	79	82	82	-
			Abnormal US AND Abnormal MRI (WITH symptoms at birth). ##	71	30	7	5	29	81	85	86	81	-
			Abnormal US AND Abnormal MRI (with NO symptoms at birth). ##	71	1	36	2	32	03	94	33	47	-
Capretti et al. 2014 ⁸³	11 symptomatic and 29 asymptomatic 7 with pathological psychomotor development at follow up 7/40 had antivirals Shell-vial procedure for CMV identification from infant	Psychomotor development at 6 years	cUS: cUS scans were defined pathological if they showed ventriculomegaly, pseudocysts, cerebral or cerebellar hypoplasia, hippocampal dysplasia, neuronal migration abnormalities, white matter abnormalities or calcifications.	40	4	3	2	31	57	94	66	91	-

	urine, within first 2 weeks of life. US performed in first 2 weeks and MRI at week 4.		cMRI: Cerebral MRI scans were defined as pathological if they showed intracranial calcification, ventriculomegaly, pseudocysts or cerebellar hypoplasia	40	7	0	2	31	100	94	78	100	-
Castellanos et al. 2022 ⁸⁴	Patients being followed up in a pediatric neurology department. 20 symptomatic, 16 asymptomatic 18 with neurological sequelae at follow up 20/36 had antivirals Congenital CMV infection was confirmed by positive PCR or urine culture findings for CMV in the first 15 days of life or positive D BS in first 48 hours of life. Timing of MRI and US tests not reported.	Neurological sequelae (WMA and other neuroimaging findings, delayed development, hearing loss) at 4 years	Brain MRI categorized by modified Noyola score. Threshold of score used is unclear, but appears to be ≥ 2	29	15	1	4	9	94	67	80	88	-
			Transfontanellar US. Pathological brain ultrasound findings were defined as intracranial calcifications, ventriculomegaly, germinolytic cysts, and/or lenticulostriate vasculopathy	30	10	2	10	8	83	44	50	80	-
Chebib et al. 2022 ⁸⁵	94 symptomatic, 36 asymptomatic 83 with inner ear impairment at 21 months No data on numbers using anti-virals No clear information on cCMV test or timing Index tests were both antenatal	Inner ear impairment at median 21 months	Infection in first trimester OR Imaging anomaly of fetus						91	48			
			Infection in first trimester AND imaging anomaly of fetus						38	92			

Craeghs et al. 2021 ⁸⁶	164 symptomatic, 247 asymptomatic 75 with hearing loss at follow up. No data on number using antivirals Diagnosis of cCMV infection made by PCR on a urine sample in first 2 weeks of life. MRI and US done at neonatal stage but precise timing not reported.	Hearing loss at mean 2.5 years	MRI Pathological if 1 or more of: WM injury, WM injury with cyst formation, periventricular pseudocysts, intraventricular adhesions, gyration disorders, calcifications, vermis hypoplasia, cortical atrophy, ventriculomegaly, or other abnormalities.	275	29	46	45	155	39	78	39	77	-
			US Pathological if 1 or more of: intraventricular adhesions, periventricular, WM injury with or without cyst formation, lenticulostriatal vasculopathy, calcifications, vermis or even cerebellar hypoplasia, hyperechogenic caudal pit, ventriculomegaly, or other abnormalities.	342	32	43	44	223	43	84	11	92	-

De Juan et al. 2020 ⁸⁷	Not reported how many were symptomatic 32/60 with neurological sequelae Number using antivirals not reported Patients diagnosed with CMV infection within the first 3 weeks of birth with positive urine or blood PCR tests. Patients older than 3 weeks of age with positive serology (CMV-specific IgM) or viral load in plasma or a blotting paper blood sample from newborn screening were also included Unclear when the US and MRI tests were used.	Neurological sequelae at a mean of 19 months	Ultrasonography and 1.5 Tesla MRI were used. Noyola et al. classification used (score of 1 or more used as threshold by reviewers)	60	23	9	8	20	72	71	74	69	
Dhondt et al. 2023 ¹⁵¹	76 symptomatic, 93 asymptomatic 20 with vestibular impairment at follow up. No data on numbers using anti-virals No information on cCMV test or timing Hearing test conducted at birth	Vestibular function at 4 years	Hearing loss at birth	169	7	13	13	136	35	91	35	91	-
Foulon et al. 2019 ⁸⁸	9 symptomatic and 146 asymptomatic 20 with mild or worse SNHL at follow up	Hearing loss (mild or worse) at median 31 months	Trimester of infection: positive was first trimester	72	5	4	12	51	56	81	29	93	-
			Trimester of infection: positive was first or second trimester	72	8	1	42	21	89	33	16	95	-

	Number on antivirals not provided Method of cCMV diagnosis not described. Timing of US/MRI not reported		US: 'abnormal' was: intracranial calcifications, migrational abnormalities, white matter disease, periventricular cysts, cerebral atrophy, ventriculomegaly, ventricular adhesions, and lenticulostriate vasculopathy.	137	8	9	9	111	47	93	47	93	--
			MRI: 'abnormal' was: intracranial calcifications, migrational abnormalities, white matter disease, periventricular cysts, cerebral atrophy, ventriculomegaly, ventricular adhesions, and lenticulostriate vasculopathy.	28	4	3	7	14	57	67	36	82	-

Fourgeaud et al. 2024 ⁸⁹	Prediction tool developed in a cohort of 227 neonates with cCMV, but the accuracy of the developed tool was tested in a separate validation cohort of n=102. 24 symptomatic and 78 asymptomatic in validation cohort 19 in validation cohort had sequelae during follow up Number on antivirals from validation cohort not reported. Newborns <1 month of age with cCMV diagnosed in the first 10 days of life were recruited during routine care diagnosis. Imaging and tests appear to have been assessed at birth.	SNHL (bilateral or unilateral) OR psychomotor sequelae (motor delay, psychomotor delay, autism, nystagmus) at 24 months	Risk factors for SNHL/ psychomotor problems: Hearing, US, platelets). This DA analyses the detection of NO sequelae, so the index test is also detecting normality; thus the threshold is no hearing loss at birth, normal cerebral ultrasound, and normal platelet count	102	-	-	-	-	92	83	63	97	0.94
Giannattasi o et al. 2018 ⁹⁰	112 symptomatic and 58 asymptomatic, but only symptomatic included in the diagnostic accuracy analysis Unclear how many had impaired neurologic outcomes.	An impaired neurologic outcome was defined in the presence of 1 or more of the following problems: developmental/co gnitive impairment, motor	Noyola Brain MRt Noyola score of 1 or more.	102	-	-	-	-	43	89	-	-	-
			Noyola US: threshold was a Noyola score of 1 or more.	105	-	-	-	-	76	43	-	-	-
			Noyola CT: threshold was a Noyola score of 1 or more	101	-	-	-	-	67	86	-	-	-
			Alorcon Brain MRt 'new' Alorcon** score of 1 or more	102	-	-	-	-	87	51	40	91	-
			Alorcon US Threshold was an Alorcon score of 1 or more.	105	-	-	-	-	89	40	34	90	-

	19 ganciclovir, 28 valganciclovir and 29 with both treatments Diagnosis of cCMV infection was based on virus detection by polymerase chain reaction assay in urine samples collected within 2 weeks of birth. Timing of neuroimaging unclear.	delay requiring rehabilitation, epilepsy and behavioral/emotional problems. Follow up 1-6 years.	Alorcon CT: Threshold was an Alorcon score of 1 or more	101	-	-	-	-	89	63	48	95	-
Kwak et al. 2018 ⁹¹	31 symptomatic only. 12 developed epilepsy during follow up. 17 using antivirals. Diagnosis confirmed by urine CMV culture or DNA polymerase chain reaction within 4 weeks after birth. MRI at mean of 33 days (but up to 375 days)	Epilepsy at mean 3.2 years	Brain MRI One point given to each of the following findings: ventriculomegaly, white matter abnormality, polymicrogyria, and calcification. Threshold for a positive test was ≥ 2.5 MRI points	31	11	1	6	13	92	68	65	93	0.87
Lucignani et al. 2021 ⁹²	27 symptomatic and 17 asymptomatic 30 with adverse neurological outcomes 22 received antivirals Congenital CMV infection was defined by PCR urine/blood and IgM CMV in the serum in first 2 weeks of life in neonates born to CMV infected mothers. Brain MRI at mean of 1.2 years.	Development of adverse neurological outcomes [including hearing loss, severe neurological delay or CP] by end of follow up at minimum of 2 years	Brain MRI Scores assigned based on presence of lesions involving 3 different locations (cerebral cortex, white matter, cerebellum), presence of calcifications and ventricular dilatation, and/or presence of ventriculomegaly. A positive test is >0 points	44	28	2	7	7	93	50	80	78	-
			Brain MRI 'New score', based on weight at birth, new maternal infection or recurrent infection, trimester of infection and symptoms at birth. Threshold for positive test ≥ 2 .	44	21	9	3	11	70	80	88	55	-

Nishida et al. 2020 ⁹³	19 symptomatic, 23 asymptomatic 19 with neurodevelopmental impairment outcomes 20 with antiviral therapy Diagnosis of cCMV in urine within 3 weeks of birth. MRI within 3 months of birth (median 24 days)	Neurodevelopmental outcomes (Neurodevelopmental impairment (NDI) was defined as a developmental quotient <80, hearing dysfunction, blindness, or epilepsy requiring anti-epileptic drugs at 18 months)	MRI— any of the 3 brain MRI abnormalities (V, PVC, WMA)	38	17	2	9	10	89	53	65	83	-
			MRI— ventriculomegaly (V)	38	16	3	3	16	84	84	84	84	-
			MRI— periventricular cyst (PVC)	38	11	8	1	18	58	95	92	69	-
			MRI— white matter abnormality (WMA)	38	16	3	8	11	84	58	67	79	-
			MRI— V and/or PVC	38	17	2	3	16	89	84	85	89	-
			MRI— V and/or WMA	38	17	2	9	10	89	53	65	83	-
			MRI— PVC and/or WMA	38	17	2	8	11	89	58	68	85	-
			MRI— >=1 of the MRI abnormalities	38	17	2	9	10	89	53	65	83	-
			MRI— >=2 of the MRI abnormalities	38	17	2	2	17	89	89	89	89	-
			Clinical symptoms (CS) - hepatosplenomegaly, liver dysfunction, thrombocytopenia, eye complications, and hearing dysfunction	38	16	3	5	14	84	74	76	82	-
			CS and/or V	38	18	1	6	13	95	68	75	93	-
			CS and/or PVC	38	16	3	5	14	84	74	76	82	-
			CS and/or WMA	38	18	1	10	9	95	47	64	90	-
			CS and/or V and/or PVC and/or WMA	38	18	1	1	18	95	47	95	95	-
			>=2 of CS and/or MRI abnormalities	38	17	2	5	14	89	74	77	88	-
			>=3 of CS and/or MRI abnormalities	38	15	4	1	18	79	95	94	82	-

Ouellette et al. 2020 ¹⁶²	49 symptomatic and 31 asymptomatic 24 with vestibular impairment at follow up 19 on antivirals cCMV infection was defined as a positive culture or PCR test of urine (88%) and/or a positive PCR from saliva (12%) samples obtained in the first 21 days of age. Positive saliva samples were all confirmed by CMV urine culture or PCR Biomarkers appear to be collected soon after birth, but unclear	Late-onset SNHL at 3 years	Biomarkers: 16-gene classifier	52	-	-	-	-	96	89	-	-	0.98
Vande Walle et al. 2024 ⁹⁴	No data on number symptomatic. 18 known to have developed cognitive impairment by follow up. 35 known to have developed a motor impairment by follow up. Unclear how many with hearing impairment at follow up.	Cognitive development: assessed using the Bayley Scales of Infant and Toddler Development. Severe impairment below 50. Follow up median 12 months.	Apparent Diffusion Coefficient (ADC) values of the white matter in MRI at threshold of ≥ 0.0671	242	-	-	-	-	83	58	-	-	0.74
	No report of number on antivirals Viral isolation and/or PCR on urine or saliva within the first 3 weeks. Retrospective diagnosis (after age of 21 days) made by PCR on DBS	Motor development. assessed using AIMS. AIMS below P10 was considered abnormal. Follow up median 12 months.	Apparent Diffusion Coefficient (ADC) values of the white matter in MRI at threshold of ≥ 0.159	239	-	-	-	-	57	72	-	-	0.64

	MRI evaluated at a median 21 days.	Hearing evaluation using ABR. ABR thresholds ≤ 40 dB nHL were considered normal. Follow up median 12 months	Apparent Diffusion Coefficient (ADC) values of the white matter in MRI at threshold of ≥ 0.0497	250	-	-	-	-	86	62	-	-	0.79
Yamaguchi et al. 2022 ¹⁵⁴	21 symptomatic and 18 asymptomatic 7 with developmental delay at 36 months. Number using antivirals not reported All patients were diagnosed with cCMV by qPCR of their blood or urine within 21 days of age. Viral loads measured at diagnosis (i.e. at or before 21 days)	Developmental delay at 36 months	CMV DNA load using droplet digital PCR ≥ 2950 copies/ml	26	4	3	3	16	57	84	57	84	0.74

Data in italics have been calculated or inferred by the reviewer from data provided in the paper.

Abbreviations: ABR, Auditory Brainstem Response; ADC, apparent diffusion co-efficient; AIMS, Alberta Infant Motor Scale; AUC, Area under the curve; cCMV, congenital cytomegalovirus; cMRI, cranial MRI; CMV, Cytomegalovirus; CP, cerebral palsy; CS, clinical symptoms; CSF, cerebrospinal fluid; CT, computed tomography; cUS, cranial US; dB, decibels; DBS, dried blood spot; DNA, deoxyribonucleic acid; FN, false negatives; FP, false positives; Ig, immunoglobulin; MRI, magnetic resonance imaging; NDI, neurodevelopmental impairment; NPV, negative predictive value; PCR, polymerase chain reaction; PPV, positive predictive value; PVC, periventricular cyst; sen, sensitivity; SNHL, sensorineural hearing loss; spec, specificity; TN, true negatives; TP, true positive; TPWMA, Temporal pole white matter abnormalities; US, ultrasound; V, ventriculomegaly; WM, white matter; WMA, white matter abnormalities.

Table 28: Association studies relevant to criterion 7

Study details	Test/characteristic assessed	Population	Outcome (and timeframe)	Results/conclusions and comments
Neuroimaging (n=3 studies)				
Giannattasio et al., 2017 ⁹⁵ Prospective cohort study Italy	Head ultrasound (lenticulostriated vasculopathy; LSV) within the first month of life	Infants with cCMV N=161 (105 symptomatic)	Neurodevelopment and hearing outcomes Follow-up: mean 4.5 years	Head ultrasound (HUS) was abnormal in 77/105 (73%) symptomatic infants and 18/56 (32%) asymptomatic infants. Of those with abnormal HUS (n=95), neurodevelopmental impairment was observed in 25/64 (39%) patients with LSV and in 17/31 (55%) patients without LSV ($p>0.05$). Hearing impairment was observed in 24/64 (38%) patients with LSV and in 14/31 (45%) patients without LSV ($p>0.05$); indicating that presence of LSV (alone or with other findings) is not a risk factor for unfavourable outcome. There was also no correlation between presence of LSV and outcomes when restricting the analysis to symptomatic patients, or to asymptomatic patients. The authors concluded that “Although LSV is a common HUS finding in infants with cCMV infection, its presence is not predictive of an adverse outcome. Our data suggest that HUS as a single neuroimaging investigation is unreliable in selecting candidates for antiviral therapy, mainly in the presence of LSV as isolated finding”. This conclusion appears appropriate and is based on a reasonable size cohort of infants with cCMV and abnormal HUS. The study methods and length of follow-up were acceptable.
Vande Walle et al., 2023 ¹⁰⁰ Prospective cohort study Belgium	MRI performed within the first 45 days of life (median age at MRI was 21 days)	Newborns with cCMV N=211 (13/207 patients [6.3%] were clinically symptomatic)	Hearing and neuromotor evaluation Follow-up: median 12 months	White matter was normal in 121 patients, doubtful in 62 and abnormal in 28. Neonatal hearing loss occurred in 4/27 (14.8%) patients with abnormal, 1/118 (0.8%) patients with normal and 1/62 (1.6%) patients with doubtful white matter ($p<0.01$). Nine patients had late-onset hearing loss; all of whom had normal white matter on neonatal brain MRI. Impaired cognitive development was seen in 3/27 (11.1%) patients with abnormal, 3/114 (2.6%) patients with normal and 1/59 (1.7%) patients with doubtful white matter (no significant difference). Alberta Infant Motor Score (AIMS) was below P75 in 21/26 (80.8%) patients with abnormal, 73/114 (64%) patients with normal and 36/57 (63.2%) patients with doubtful white matter. In a subgroup of 77 patients with ≥ 18 months follow-up, AIMS score was below P75 in 10/13 (76.9%) patients with abnormal, 13/34 (38.2%) patients with normal and 7/20 (35%) patients with doubtful white matter ($p<0.05$).

				The authors concluded that “Abnormal white matter was associated with neonatal hearing loss and mild, lower motor scores. A tendency towards impaired cognitive development was seen. Patients with doubtful white matter did not show worse clinical outcome”. This conclusion appears appropriate and is based on a relatively large cohort of infants with cCMV, although only 28 infants had white matter abnormalities. The length of follow-up was relatively short, with only 77 patients having follow-up ≥18 months (of which, only 13 had white matter abnormalities). This study is included in the Vande Walle 2024 systematic review of MRI
Vila-Bedmar et al., 2024⁹⁶ Prospective cohort study Spain	Ultrasound (US) and MRI performed within the first 2 weeks of life	Newborns with cCMV diagnosed through a screening programme N=15 (2 symptomatic)	SNHL, neuro-developmental disorders, chorioretinitis and epilepsy Follow-up: median 68 months	Brain abnormalities were found on 4/15 US and 10/15 MRI; white matter abnormalities (WMA) were the most common finding on MRI. Eight children were treated with antivirals. Two children were diagnosed with psychomotor delay and 2 with attention-deficit hyperactivity disorder; all of them presented brain abnormalities on MRI, especially severe and extensive WMA, that were not detected by US (2 children were symptomatic at birth). All 5 children with normal MRI developed normally without long-term sequelae. The authors concluded that “Children with isolated WMA without other clinical symptoms showed overall good outcomes, although more studies with a larger sample and a control group should be performed. Interobserver agreement about the presence of abnormalities in MRI was good”. The methods and length of follow-up appear appropriate. However, this was a very small study, therefore, the authors’ conclusion that further research is required appears appropriate.
Antenatal characteristics (n=10 studies)				
Amir et al., 2016¹⁰⁴ Retrospective cohort study Israel	First trimester infection vs second trimester infection (diagnosed by positive amniotic fluid findings)	Infants infected with CMV in 1st or 2nd trimester with normal fetal imaging N=98 (60 symptomatic)	Late-onset SNHL (and motor and cognitive delay) Follow-up: median 32 months (27 months for infants infected during the first trimester and 45 months for those infected in the second trimester)	Fifty-two infants were treated with an antiviral agent, the main indication for treatment was abnormal cranial ultrasound finding. Fifty-two infants were infected in the first trimester and 46 in the second trimester. Moderate and severe hearing losses were associated more often with first-trimester infection than second-trimester infection ($p=0.053$); 6/104 ears vs 0/92 ears at the time of the last test. 95/104 (91.4%) infants had normal hearing in the first trimester group vs 91/92 (98.9%) in the second trimester group. The authors concluded that “First trimester infection is associated with a trend towards more severe hearing loss than second trimester infection.

				Overall, the outcome of these infants is good, with a low rate of sequelae on short-term follow-up and intermediate-term follow-up". This conclusion appears appropriate and is based on a reasonable size cohort of infants with cCMV and acceptable methods and length of follow-up.
Elkan Miller et al., 2021¹⁰⁶ Prospective cohort study Israel	Second trimester infection vs third trimester infection (mostly diagnosed by amniocentesis, according to patient preference)	Newborns with cCMV N=127 (proportion symptomatic at birth not reported)	Hearing and neurodevelopmental outcome Follow-up: median 4 years	Maternal infection was during the second trimester in 100/127 infants and third trimester in 27/127 infants. 7/100 (7%) infants had long-term sequelae (hearing loss or neurodevelopmental impairment) following second trimester infection vs 1/27 (3.7%) following third trimester infection. Twenty-one (21%) second trimester infection infants received antiviral therapy vs 3 (11%) third trimester infection infants. The authors concluded that "Second trimester infection is associated with a slight risk of developing mild childhood sequelae, mostly partial unilateral hearing loss, which may develop late in childhood." This conclusion appears appropriate and is based on a relatively large cohort of infants with second trimester infection. The study methods and length of follow-up were acceptable.
Faure-Bardon et al., 2019¹⁰⁷ Prospective cohort study France	Trimester of infection (based upon serial measurements of immunoglobulin (Ig) M and IgG and on IgG avidity in sera collected at each trimester)	Newborns with cCMV N=234; 19 lost to follow-up (70 symptomatic)	SNHL and neurological sequelae Follow-up: median 24 months	126 newborns were infected in the first trimester, 72 in the second trimester and 36 in the third trimester. SNHL and/or neurological sequelae were only seen in children infected following a primary infection in the first trimester of pregnancy (32.4% vs 0% infants infected in the second or third trimester; $p<0.0001$). The authors concluded that "These results suggest that a CMV infection can be severe only when the virus hits the fetus in the embryonic or early fetal period. Recent guidelines recommend auditory follow-ups for at least 5 years for all infected children. This raises parental anxiety and generates significant costs. We suggest that auditory and specialized neurologic follow-ups may be recommended only in cases of a maternal infection in the first trimester". This conclusion appears appropriate and is based on a relatively large cohort of infants with first or second trimester infection (only 36 infants had third trimester infection). The study methods and length of follow-up were acceptable. This study is included in the Chatzakis 2020 systematic review of time of maternal infection.

Lipitz et al., 2013¹⁰⁸ Prospective cohort study Israel	First trimester infection vs second trimester infection (based upon seroconversion or presence of IgG and IgM antibodies associated with low IgG avidity)	cCMV (diagnosed as fetus) N=137 (proportion symptomatic at birth not reported)	Hearing loss and neurodevelopmental delay Follow-up: median 27 months of age	Maternal infection was during the first trimester in 66 infants and second trimester in 71 infants. Mothers with first-trimester infection were significantly more likely to have infants with auditory or neurodevelopmental sequelae than patients with second-trimester infection (13/66 infants [19.7%] vs 4/71 infants [5.6%]; $p=0.01$). The authors concluded that “The risk of sequelae is higher following first- than second-trimester CMV infection”. This conclusion appears appropriate and is based on a relatively large cohort of infants. The study methods and length of follow-up were acceptable. This study is included in the Chatzakis 2020 systematic review of time of maternal infection.
Massoud et al., 2023¹⁰⁹ Retrospective cohort study France	First trimester infection vs second trimester infection (mostly determined using maternal CMV IgG and IgM and IgG avidity test)	cCMV (diagnosed as fetus) N=27 surviving to infancy (out of the original sample of 49, 22 were terminated or had an intrauterine death) (proportion symptomatic at birth not reported)	Ophthalmological or sensorineural deficit, motor delay and/or cognitive impairment Follow-up: mean 51 months	Maternal infection was during the first trimester in 15 infants and second trimester in 12 infants. 9/15 first-trimester infection infants had a normal outcome, 5/15 had mild sequelae and 1/15 had severe sequelae. 11/12 second-trimester infection infants had a normal outcome and 1/12 had mild sequelae. Mild sequelae were mostly hearing loss. The authors summarised that “Fetuses infected with cytomegalovirus during the first trimester of pregnancy have significant risks of sequelae, which may persist into the second trimester, particularly for sensorineural disorders”. This was a small study investigating both prenatal imaging and trimester of infection in pregnant women with CMV infection. Most of the results reported related to prenatal imaging, with few results reported relating to trimester of infection.
Nigro et al., 2020¹¹⁰ Prospective cohort study Italy	Timing of maternal seroconversion (first trimester infection confirmed by positive amniotic fluid findings)	Infants of pregnant women diagnosed with primary CMV infection N=108 with cCMV (106 with long-term follow-up; proportion symptomatic at birth not reported)	Hearing loss, psychomotor retardation, cerebral palsies, visual impairment Follow-up: mean 4 years	Twenty-one infants had abnormalities and 85 were categorised as ‘normal’ at long-term follow-up. In a multivariate logistic regression, odds of later abnormalities were lower if maternal infection occurred in later gestation (mean 16.2 weeks) compared to earlier gestation (mean 10.1 weeks) [adjusted odds ratio 0.9 (95% CI: 0.83 to 0.98); $p=0.017$]. The authors concluded that “Maternal viremia predicts fetal infection and neonatal outcome. This may help patient counselling. High-dose HIG may prevent fetal infection and disease, and is associated with the resolution of DNAemia”.

				This study investigated several factors predictive of infant outcome, including HIG administration, fetal ultrasound and timing of maternal infection; few results were reported relating to timing of maternal infection.
Giannattasio et al., 2017 ¹²³ Prospective cohort study Italy	Primary maternal infection vs non-primary maternal infection	Children with cCMV N=158 (88 symptomatic)	Hearing and neurodevelopmental assessment Follow-up: Beyond 3 years of age	Ninety-three infants were born to mothers with primary CMV infection and 65 to mothers with non-primary infection. Fifty-one symptomatic infants received antiviral treatment; 25 in the primary infection group and 26 in the non-primary infection group. Frequency of impaired neurodevelopmental outcome (23.7% vs 24.6%) and hearing loss (25.8% vs 26.2%) did not differ between the primary infection and non-primary infection groups. The authors concluded that “Neurodevelopmental and hearing sequelae are not affected by the type of maternal CMV infection. Preventing strategies should be developed for both primary and non-primary infections”. This conclusion appears appropriate and is based on a relatively large cohort of infants. The study methods and length of follow-up were acceptable. This study is included in the Maltezou 2020 systematic review of type of maternal infection and the Smyrli 2024 systematic review of impact of symptoms.
Coscia et al., 2020 ¹¹² Retrospective cohort study Italy	Trimester of infection and primary vs non-primary maternal infection	Newborns with cCMV at Neonatal Care Unit N=91 (results presented for infants where timing (n=52) and type (n=49) of maternal infection could be determined)	Hearing and neurodevelopment Follow-up: 12 months (asymptomatic infants) and 24 months (symptomatic infants)	Data regarding the trimester of infection were missing for 39 patients. Determined cases accounted for 40% patients infected in the first trimester, 29% in the second trimester and 16% in the third. Data regarding type of infection were missing for 42 patients. Determined cases included 86% patients with primary infection and 14% with non-primary infection. A higher proportion of first-trimester infection infants had long-term sequelae, however, the difference between groups was only statistically significant for ‘need for cochlear implant’; 28% vs 0% (p=0.003). A higher proportion of infants born to mothers with non-primary infection had long-term sequelae, however, the difference between groups was only statistically significant for ‘need for cochlear implant’; 43% vs 10% (p=0.05). The authors concluded that “We underline the possibility of re-infection in previously immunized mothers (non-primary infection) with unfavorable neonatal and long-term outcomes”. Data on timing and type of maternal infection were missing for a large proportion of patients; therefore, the results are based on a small number of infants with relatively short-term follow-up.

Puhakka et al., 2017¹¹¹ Retrospective cohort study Finland	Timing of infection and primary vs non-primary maternal infection	Infants with symptomatic cCMV N=26 (all symptomatic)	Hearing, vision and neurology Follow-up: mean 21.7 months (vision), 44.6 months (hearing), 47.2 months (neurology)	Fourteen infants had non-primary maternal infection, 7 had primary infection in the first trimester or near conception and 5 during the second or third trimester. Long-term sequelae (SNHL, neurologic abnormality and/or visual impairment) occurred in 15 infants: 6/7 after primary infection in the first trimester, 0/5 after primary infection in the second or third trimester and 9/14 after non-primary infection. The authors concluded that “In this register-based cohort, non-primary infections caused the majority of symptomatic congenital CMV infections, and resulted in significant morbidity”. This was a very small study with missing data on long-term outcomes for some patients (particularly hearing and ophthalmologic examinations). Therefore, the results should be interpreted with caution. This study is included in the Chatzakis 2020 systematic review of time of maternal infection and the Maltezou 2020 systematic review of type of maternal infection.
Scaramuzzino et al., 2022¹¹³ Retrospective cohort study Italy	Primary maternal infection vs secondary maternal infection and trimester of infection	Babies with symptomatic cCMV N=53 (all symptomatic)	SNHL and neurodevelopmental impairment Follow-up: Mean 37.2 months (primary infection) and 55 months (secondary infection)	Forty babies were born to mothers with primary infection and 13 to mothers with secondary infection. Of those with primary infection, 18 were infected during the first trimester, 13 were infected during the second trimester and 8 were infected during the third trimester (1 unknown timing of infection). Treatment was given to 57.5% primary infection babies and 84.6% secondary infection babies. There was a higher rate of long-term sequelae (tetraparesis, epilepsy, motor and speech delay, and unilateral SNHL) in children of mothers with secondary infection, which was statistically significant for tetraparesis (p=0.002) and unilateral SNHL (p=0.009). Only children born to mothers with primary infection presented bilateral SNHL (6 vs 0; p=not significant). 66.7% first-trimester infected babies had neurodevelopmental and auditory sequelae vs 23% second-trimester infected and 12.5% third-trimester infected babies. The authors concluded that “Our data suggest that the risk of symptomatic cCMV and long-term sequelae is similar in children born to mother with primary and secondary CMV infection; it is important to pay appropriate attention to seropositive mothers in order to prevent reinfection and to detect and possibly treat infected babies”. The methods and length of follow-up appear appropriate. However, this was a small study and the number of children suffering some of the long-term

				sequelae reported (particularly tetraparesis, epilepsy and cognitive delay) was very small.
Symptoms/characteristics at birth (n=23 studies)				
Airlangga et al., 2024 ¹²⁸ Prospective cohort study Indonesia	Symptomatic vs asymptomatic cCMV	Newborns with cCMV N=26 (9 symptomatic)	Hearing function and central auditory neuroanatomy evaluation Follow-up: 3 months and 6 months	The proportion of children with global developmental delay was significantly higher in infants with symptomatic cCMV infection (88.9% vs 47.1%; $p<0.001$). The proportion of children with abnormal hearing function at 3 months old was significantly higher in infants with symptomatic cCMV infection (69.6% vs 30.4%; $p<0.001$). The authors concluded that "SNHL along with its fluctuation and progression are common in cCMV-infected infants. cCMV infection may induce structural changes in the central auditory pathway". This was a very small study with short-term follow-up.
Auriti et al., 2022 ¹²⁹ Retrospective cohort study Italy	Symptomatic vs asymptomatic cCMV	Infants with acquired infection (CMV, toxoplasma or syphilis) N=84 with cCMV (29 symptomatic)	Cognitive, motor, audiological, visual and language outcomes Follow-up: At least 24 months	Thirty-one (36.9%) newborns received antiviral treatment. At 1 year of age, 12/29 (41.4%) infants who were symptomatic at birth had 1 sequela and 4/29 (13.8%) had 2 or more sequelae, compared with 6/55 (10.9%) and 0/55 (0%) in asymptomatic infants, respectively. Of 69 infants with longer term follow-up (2-4 years of age), 14/27 (51.9%) symptomatic infants had 1 sequela and 5/27 (18.5%) had 2 or more sequelae, compared with 8/42 (19.0%) and 0/42 (0%) asymptomatic infants, respectively. The authors concluded that "Infected babies symptomatic at birth have a worse prognosis than asymptomatic ones. Long-term sequelae may occur in infected children asymptomatic at birth after the first year of life. Multidisciplinary follow-up until 4-6 years of age should be performed in all infected children, regardless of the presence of symptoms at birth". The methods and length of follow-up appear appropriate. However, this study was small, with only 29 symptomatic infants.
Bartlett et al., 2018 ¹⁴⁷ Retrospective cohort study Australia	Symptomatic vs asymptomatic cCMV	Infants with cCMV N=302 (214 symptomatic)	Hearing loss and developmental delay Follow-up: Not reported	Fifty-eight (19.2%) infants received antiviral treatment. Of 257 cCMV cases born in the era of universal neonatal hearing screening, hearing loss was reported in 75/178 (42.1%) symptomatic (49/75 cases reported ≤ 30 days of age, 17/75 reported >30 days of age and 9/75 had uncertain timing) and 21/79 (26.6%) asymptomatic cases. Developmental delay was reported in 30/178 (16.9%) symptomatic (6/30 cases reported ≤ 30 days of age, 18/30

				reported >30 days of age and 5/30 had uncertain timing) and 1/79 (1.3%) asymptomatic cases. The authors concluded that “There appears to be under-reporting and under-recognition of congenital CMV despite increasing use of antiviral therapy. Universal newborn CMV screening should be considered to facilitate follow-up of affected children and targeted linkage into hearing and developmental services, and to provide population-level infant CMV epidemiology to support research and evaluation of antiviral and adjunctive therapies”. Whilst this appears to be a large, well conducted study, it is unclear how long patients were followed up for and the conclusion does not relate to the association between symptomatic cCMV infection and long-term sequelae. This study is included in the Smyrli 2024 systematic review of impact of symptoms.
Capretti et al., 2021¹³⁰ Prospective cohort study Italy	Symptomatic vs asymptomatic cCMV	Newborns with cCMV N=30 (10 symptomatic)	SNHL and neurodevelopmental delay Follow-up: median 30 months	All 10 symptomatic infants received antiviral therapy. 6/10 symptomatic infants had SNHL and 1/10 had neurodevelopmental delay, whereas none of the 20 asymptomatic infants had SNHL or neurodevelopmental delay. The authors concluded that “A detectable CMV-specific CD8 + T-cell response, evaluated using the QuantiFERON-CMV assay, correlates with the lack of CMV-related symptoms and the control of CMV DNAemia”. This was a very small study investigating immune monitoring using QuantiFERON-CMV assay; few results were reported relating to the association between symptomatic cCMV infection and long-term sequelae.
Dhondt et al., 2021¹³¹ Prospective cohort study Belgium	Symptomatic vs asymptomatic cCMV	cCMV infected children N=93 (41 symptomatic)	SNHL and vestibular function Follow-up: average 10.2 months	Thirty-two symptomatic infants received antiviral therapy. Delayed onset SNHL occurred in 3/52 asymptomatic and 2/41 symptomatic patients (12 symptomatic patients had hearing loss at birth). The median age of diagnosis of delayed onset hearing loss was 13 months (range 2-20 months). The authors concluded that “cCMV can impair not only the auditory but also the vestibular function. Similar to the hearing loss, vestibular loss in cCMV can be highly variable. It can be unilateral or bilateral, limited or extensive, stable or progressive, and early or delayed in onset. As the vestibular function can deteriorate over time and even normal-hearing subjects can be affected, vestibular evaluation should be part of the standard otolaryngology follow-up in all children with cCMV”.

				Whilst this appears to be a reasonable sized study, the length of follow-up was relatively short and the conclusions do not relate to the association between symptomatic cCMV infection and long-term sequelae. All symptomatic infants received antiviral therapy, whilst none of the asymptomatic infants did, which could confound the finding relating to delayed onset SNHL
Goderis et al., 2016⁷⁸ Prospective cohort study (Flemish CMV registry) Belgium	Symptomatic vs asymptomatic cCMV	Children with cCMV N=379 (123 symptomatic)	Hearing loss Follow-up: not reported (mean age at diagnosis of delayed onset hearing loss was 18 months)	Fifty-six symptomatic infants received antiviral treatment. Delayed onset hearing loss occurred in 10.6% symptomatic patients and 7.8% asymptomatic patients; 63% symptomatic patients and 8% asymptomatic patients had hearing loss overall. Almost half of the symptomatic patients who developed delayed onset hearing loss had preexisting contralateral hearing loss. Mild hearing loss was more frequent in asymptomatic (52% ears) vs symptomatic (23.5% ears) infants, compared with moderate, severe or profound hearing loss. The authors concluded that "Symptomatic and asymptomatic cCMV infections are a major cause of hearing loss in childhood. Reliable estimates of the long-term outcome of cCMV infection are mandatory to increase vigilance, especially among pregnant women and to draw attention to preventive measures, vaccine development, and prenatal and postnatal therapy. Universal screening of newborns for cCMV infection should be initiated and combined with longitudinal audiometric follow-up". This study included a relatively large cohort of infants with cCMV, the methods and length of follow-up appear to have been acceptable. The conclusion that symptomatic and asymptomatic cCMV infections are a major cause of hearing loss appears appropriate.
Jin et al., 2019¹³³ Retrospective subanalysis of a prospective cohort study USA	Symptomatic vs asymptomatic cCMV	Infants with cCMV N=186 (77 symptomatic)	Visual impairment and SN HL Follow-up: mean 11-12 years	Eleven symptomatic patients developed cortical visual impairment (CVI) (11/77; 14.3%), compared to no asymptomatic patients (0/109; 0%). Comparisons were made between these 11 patients and the remaining 66 symptomatic patients who did not develop CVI; All 11 patients with CVI also had SNHL, compared with 69.7% patients without CVI. Microcephaly, white matter abnormality, seizure at birth, intracranial calcifications, encephalomalacia and dilatation of ventricles occurred more frequently in patients with CVI than without. Five predictive models based on 11 variables were created using logistic regression to predict which symptomatic patients would have developed CVI. Model 5 showed the variables of microcephaly, seizure at birth, optic atrophy, chorioretinitis/retinal scars, strabismus, intracranial calcifications, neonatal hearing loss and severe visual impairment

				were predictors of CVI; this model correctly identified 10/11 cases of CVI, giving the lowest false-positive rate (23.1%), overall accuracy 94.8%. The authors concluded that “CVI may result from symptomatic congenital CMV infection. The relationship of CVI and its risk factors in patients with CMV suggests the potential to predict the development of CVI through predictive modelling in future research. Early screening of CVI in children born with symptomatic congenital CMV can facilitate educational, social, and developmental interventions”. Whilst this was a reasonable sized study with long-term follow-up, it focussed on comparing symptomatic cCMV patients with (n=11) and without (n=66) CVI. The models proposed would need to be prospectively validated in a much larger study.
Korndewal et al., 2017¹⁴⁶ Retrospective cohort study Netherlands	Symptomatic vs asymptomatic cCMV	Infants with cCMV diagnosed retrospectively using dried blood spot PCR and healthy controls N=133 with cCMV (26 symptomatic)	SNHL, visual impairment and neurological impairment Follow-up: up to 6 years of age	Long-term impairment, diagnosed in the first 6 years of life, was more frequent in symptomatic than asymptomatic cCMV infection: multiple domains 19.2% vs 8.4%; SNHL 7.7% vs 2.8%; epilepsy 7.7% vs 0%; autism spectrum disorder 7.7% vs 1.9%; cognitive impairment 15.4% vs 3.7%; fine motor impairment 19.2% vs 7.5%; gross motor impairment 15.4% vs 8.4%; language disorder 23.1% vs 7.5% (other results presented). This study concluded that “Children with cCMV were twice as likely to have long-term impairment up to the age of 6 years, especially developmental delays and sensorineural hearing loss, than cCMV-negative comparison children, with a risk difference of 12.8%. These insights into the risk of cCMV-associated impairment can help optimize care and stimulate preventive measures”. This was a large study comparing long-term impairment between children with cCMV vs those without, as well as those with symptomatic cCMV vs asymptomatic cCMV. The authors’ conclusions appear appropriate. However, only 26 children had symptomatic cCMV, therefore, the number of children in the comparison of interest to this review was small. This study is included in the Smyrli 2024 systematic review of impact of symptoms.
Koyano et al., 2018¹⁴⁸	Symptomatic vs asymptomatic cCMV	Infants with cCMV N=60 (17 symptomatic)	Hearing, development and ‘other’ (including death, epilepsy, autism spectrum disorder and	Ten of the 17 symptomatic infants received antiviral treatment. All 7 untreated symptomatic infants had long-term sequelae (100%), including hearing loss (4), developmental delay (3), death (1), epilepsy (2) and autism spectrum disorder (1). Seven of the 10 treated symptomatic infants had long-

Prospective cohort study Japan			attention deficit-hyperactivity disorder) Follow-up: At least 2 years	term sequelae (70%), including hearing loss (7), developmental delay (3) and epilepsy (2). Five of the 43 asymptomatic infants had long-term sequelae (12%), including hearing loss (1), speech delay (2), autism spectrum disorder (1) and attention deficit-hyperactivity disorder (1). The authors concluded that “The rate of late-onset sequelae observed in Japan is similar to that reported in the USA and Europe. The treatment of symptomatic patients with antiviral agents results in favorable clinical outcomes. Thus, newborn urine-filter paper screening of congenital CMV infection is warranted”. The methods and length of follow-up appear appropriate (outcomes of symptomatic infants who received antiviral treatment were reported separately from those who did not receive treatment). However, this was a relatively small study including only 17 symptomatic infants. This study is included in the Smyrli 2024 systematic review of impact of symptoms.
Lanzieri et al., 2018 ¹³⁴ Prospective cohort study USA	Symptomatic vs asymptomatic cCMV	Patients with cCMV N=168 (76 symptomatic)	SNHL Follow-up: up to 18 years of age	Seventy-eight (51%) ears of symptomatic patients were diagnosed with congenital/early-onset SNHL, compared with 10 (5%) ears of asymptomatic patients. Twenty-five (17%) ears of symptomatic patients were diagnosed with delayed-onset SNHL, compared with 20 (11%) ears of asymptomatic patients. Forty-nine (32%) and 154 (84%) ears had normal hearing, respectively. In affected ears, all frequency-specific hearing thresholds worsened with age. At the end of follow-up, 56 (74%) symptomatic patients and 20 (22%) asymptomatic patients had SNHL, among whom 47 (84%) and 10 (50%) had bilateral SNHL, respectively. The authors concluded that “Congenital/early-onset SNHL frequently resulted in severe to profound loss in symptomatic and asymptomatic case patients. White matter lucency in asymptomatic case patients was significantly associated with SNHL by age 5 years”. This conclusion appears appropriate and is based on a relatively large cohort of infants. The study methods and length of follow-up were acceptable. However, it is unclear whether the results/conclusion relating to white matter lucency related to delayed-onset SNHL or congenital/early-onset SNHL
Lo et al., 2022 ¹³⁵	Symptomatic vs asymptomatic cCMV	Infants with cCMV N=39 (10 symptomatic, 20	SNHL	SNHL developed in 60% of children who were initially symptomatic and 34.5% of children who were initially asymptomatic (with normal hearing or isolated hearing loss). Of the 10 symptomatic infants, 4 had SNHL at birth (as

Prospective cohort study Taiwan		asymptomatic with normal hearing and 9 asymptomatic with isolated hearing loss)	Follow-up: median 4 years	well as other symptoms) and 2 patients subsequently developed late-onset S NHL. One asymptomatic infant with normal hearing subsequently developed late-onset SNHL. All 9 asymptomatic infants with isolated hearing loss had SNHL. All 13 infants who failed newborn hearing screening (NHS) developed SNHL, along with 3 infants who passed NHS. Initial viral loads were higher in children who were symptomatic at birth, who failed NHS, and who developed SNHL. The authors concluded that “The presence of cCMV-related symptoms at birth, failure in NHS, and blood viral load might be the prognostic factors for hearing outcomes. Regular audiologic examinations are necessary in all children with cCMV infection even after CMV DNAemia clearance”. The methods and length of follow-up appear appropriate. However, this was a small study including only 10 symptomatic infants.
Maes et al., 2017 ¹⁵⁰ Prospective cohort study Belgium	Symptomatic vs asymptomatic cCMV	Infants with cCMV, Connexin 26 mutation and controls N=40 (24 with cCMV; 8 symptomatic with impaired hearing, 8 symptomatic with normal hearing and 8 asymptomatic)	Motor performance and vestibular function Follow-up: 6 months of age	Six symptomatic children with normal hearing and 4 hearing-impaired symptomatic children received antiviral treatment. Mean values for gross and fine motor performance were lower for symptomatic children than asymptomatic children, particularly gross motor performance, where the mean score for the symptomatic hearing-impaired group was significantly lower than the asymptomatic cCMV group ($p=0.034$). The authors concluded that “The weakest gross motor performance was found in symptomatic hearing-impaired cCMV-infected children with absent cVEMP responses. These results suggest that abnormal saccular responses are a major factor for this delayed motor development, although more work is needed including comprehensive vestibular function testing to verify this”. This was a very small study with short-term follow-up, therefore, the results should be interpreted with caution. This study is included in the Smyrli 2024 systematic review of impact of symptoms.
Rovito et al., 2017 ^{c124} Retrospective cohort study Netherlands	Symptomatic vs asymptomatic cCMV	Children retrospectively diagnosed with cCMV from neonatal DBS	Long-term impairments (hearing, visual, neurological, motor, cognitive, speech-language) Follow-up: 6 years of age	Ten of the 18 (55.6%) symptomatic infants developed 1 or more long-term impairment vs 15/84 (17.9%) asymptomatic infants. Eleven infants were born premature and 91 were term. Significantly more infants who developed long-term impairment were born premature, compared with infants who did not develop long-term impairment (24% vs 6.5% respectively; $p=0.024$). All 6 premature infants with birthweight <2500g developed long-term impairment vs only 1 premature infant with birthweight ≥ 2500g.

		N=102 with cCMV (18 symptomatic)		The authors concluded that “Although these findings demonstrate limitations in the use of routine neonatal screening data as predictors for long-term cCMV outcome, a possible influence of cCMV in the amino acids metabolism of preterm neonates, and of higher viral loads in the metabolism of longer chain fatty acids were shown. Finally, by considering the use of routine neonatal screening data, this study represents a first step in identifying prognostic markers for cCMV outcome”. The conclusion does not specifically relate to the association between symptomatic cCMV infection and long-term sequelae. The finding of an association between symptomatic cCMV (including prematurity and birthweight) and the development of long-term impairment appears to be credible, although only 18 children in the study had symptomatic cCMV.
Royackers et al., 2013¹³⁶ Prospective cohort study Belgium	Symptomatic vs asymptomatic cCMV	Infants with cCMV N=98 (24 symptomatic)	Late-onset hearing loss Follow-up: average 5.4 years	Eight symptomatic children with hearing loss at birth were treated with ganciclovir. Two symptomatic children with normal hearing at birth developed late-onset hearing loss (detected within 5 months of birth). One asymptomatic child with normal hearing at birth developed late-onset hearing loss (detected at the age of 2 years and 9 months). The authors concluded that “Hearing loss caused by congenital cytomegalovirus infection cannot be defined unequivocally either with respect to the level of hearing loss or its evolution over time. Treating symptomatic children with ganciclovir leads to a better prognosis during the first year of life, after which progression or fluctuation again becomes more likely. However, overall, progression is more common in the untreated symptomatic group. Asymptomatic children with SNHL are more likely to have a stable hearing status”. The methods and length of follow-up appear appropriate. However, this was a small study including only 24 symptomatic infants and only 3 cases of late-onset hearing loss.
Salome et al., 2023¹³⁷ Prospective cohort study Italy	Symptomatic vs asymptomatic cCMV	Infants with cCMV N=250 (123 symptomatic)	Long-term visual impairment Follow-up: median 4.7 years (symptomatic infants) and 2.9 years (asymptomatic infants)	Fundoscopy abnormalities were identified in the neonatal period in 16/123 (13%) symptomatic infants and in none of the asymptomatic infants. Chorioretinitis was the most common finding (10/16 cases), with the others presenting with chorioretinal scars. Fifteen of the 16 infants with ocular involvement received antiviral therapy. Seven of the 16 showed normal psychomotor development, 8/16 developed severe psychomotor delay and 1/16 showed speech delay and mild psychomotor delay. No patient (symptomatic or asymptomatic) developed new/late-onset retinal lesions.

				Throughout follow-up, 5/16 infants with ocular involvement at birth were diagnosed with visual impairment. The authors concluded that "Chorioretinal lesions are a fairly common finding at birth in neonates with symptomatic cCMV, often associated with long-term visual impairment. Asymptomatic infants do not show ophthalmological abnormalities in the short or long term. This information is relevant both to parental counselling and to cost-effective patient management". Whilst this was a large study with relatively long-term follow-up, the results focussed on the 16 infants with ocular involvement in the neonatal period. No patients (symptomatic or asymptomatic) developed new/late-onset retinal lesions.
Seneviratne et al., 2022¹³⁸ Retrospective cohort study Australia	Symptomatic vs asymptomatic cCMV	cCMV cases N=45 (25 diagnosed within 3 weeks of birth and 20 diagnosed >3 weeks from birth; 35 symptomatic)	Developmental delay, SN HL, cerebral palsy, death Follow-up: 24 months	Thirteen symptomatic newborns received antiviral therapy. A degree of developmental delay during the first 24 months of follow-up was reported in 22 newborns: 9/13 (69.2%) symptomatic treated newborns; 10/22 (45.5%) symptomatic untreated newborns; and 3/10 (30%) asymptomatic newborns. SNHL: 3/13 (23.1%) symptomatic treated newborns, 5/22 (22.7%) symptomatic untreated newborns and 0 asymptomatic newborns. Cerebral palsy: 2/13 (15.4%) symptomatic treated newborns, 2/22 (9.1%) symptomatic untreated newborns and 0 asymptomatic newborns. One child died (symptomatic untreated). The authors concluded that "Up to half of the neonates with cCMV were at risk of developing a degree of developmental delay at our centre. Whether these outcomes are related to CMV infection or are confounded by the co-existence of prematurity is unclear and needs further evaluation in prospective studies". This was a small study, in which 20 of the newborns were diagnosed >3 weeks from birth, so may not have had congenital cytomegalovirus, therefore, the results should be interpreted with caution.
Topham et al., 2019¹⁴⁹ Prospective cohort study USA	Symptomatic vs asymptomatic cCMV	Children with symptomatic or asymptomatic cCMV (and controls) N=112 with cCMV (36 symptomatic) and 29 controls	Inattention and hyperactivity (also lifetime hearing loss) Follow-up: at least 6 years	Children with symptomatic cCMV had a higher proportion of elevated Attention Problems T-scores at follow up compared with the asymptomatic cCMV group (39% vs 17%; p=0.012). There were no differences in the proportions of children with elevated Hyperactivity T-scores at follow up between symptomatic cCMV and asymptomatic cCMV groups (28% vs 20%). Children with symptomatic cCMV had higher mean Attention Problems T-scores at follow up versus those with asymptomatic cCMV (scores of 56.0 vs 50.6; p=0.006) and higher mean Hyperactivity T-scores versus those with

				asymptomatic cCMV (scores of 54.3 vs 50.3; $p=0.031$). After adjusting for IQ, differences in mean Attention Problems or Hyperactivity T-scores were no longer significant. Significantly more symptomatic children had lifetime hearing loss: 28/36 (78%) vs 16/76 (21%) asymptomatic children; $p<0.001$. Mean IQ score was significantly lower in symptomatic vs asymptomatic children (90.5 vs 109.2; $p<0.001$). The authors concluded that “Children with asymptomatic cCMV are not at increased risk of inattention or hyperactivity compared with controls. However, our study suggests an increased prevalence of inattention and hyperactivity among children with symptomatic cCMV. Differences in IQ were confirmed to have a confounding effect. Evaluation for attention-deficit/hyperactivity disorder may be warranted in this population”. This conclusion appears appropriate and is based on a reasonable sized cohort of infants and relatively long-term follow-up. This study is included in the Smyrli 2024 systematic review of impact of symptoms.
Zavattoni et al., 2016¹³⁹ Retrospective cohort study Italy	Symptomatic vs asymptomatic cCMV	Newborns with cCMV N=46 (36 live births; 8 symptomatic)	Long-term sequelae Follow-up: Not reported	Long-term sequelae were reported in 3/7 (42.8%) symptomatic (1 lost to follow-up) and 7/22 (31.8%) asymptomatic (5 lost to follow-up, 1 died) newborns. In the symptomatic cCMV group, 1 had communicative disorders, 1 neurodevelopmental deficits and 1 motor and language disabilities. In the asymptomatic cCMV group, 2 had hearing loss, 3 had language disabilities, 1 had diabetes and 1 had cognitive impairment. The authors conclusions are unclear (there is no conclusions section). This was a very small study with only 8 symptomatic infants, unclear length of follow-up and no clear conclusions drawn.
Minsart et al., 2020¹⁴⁵ Retrospective cohort study Canada	Symptomatic (moderate to severe symptoms) vs asymptomatic cCMV	Mother-child pairs referred to Women and Children's Infectious Diseases Center with confirmed cCMV	Hearing function and neurodevelopmental symptoms Follow-up: median 2.1 years hearing, median 29.5 months neurodevelopmental symptoms	Fifty-one of the 77 infants received antiviral therapy. There were 3 neonatal deaths attributed to CMV disease. Among 74 children who survived the neonatal period, 50 had normal hearing initially; 5/50 (10%) developed late-onset SNHL after a median time of 1.2 years, all of whom had symptoms at birth. Nine out of 59 symptomatic children developed severe neurodevelopmental symptoms, all of which had moderate to severe symptoms at birth. No asymptomatic children had severe neurodevelopmental symptoms. The authors concluded that “Outcome of cCMV infection in a setting without routine CMV screening and third-trimester scan varies according to the

		N=77 with cCMV (59 symptomatic)		reason for screening and timing of diagnosis. Late detection of prenatal ultrasound anomalies and postnatal diagnosis following neonatal symptoms represented more than half of the series, and remain challenging situations. Awareness of this should help perinatologists to recognize cCMV patterns and ensure optimal management”. The methods and length of follow-up appear appropriate. However, this was a relatively small study including only 18 asymptomatic infants. This study is included in the Smyrli 2024 systematic review of impact of symptoms.
Hu et al., 2024¹³² Prospective cohort study China (non-English language publication)	Symptomatic vs asymptomatic cCMV (and severity of symptoms)	Newborns with cCMV N=176 (79 asymptomatic, 12 mildly symptomatic, 85 moderately to severely symptomatic)	SNHL Follow-up: not reported	Fifty cases had SNHL at birth, with different degrees of severity: 30 mild, 9 moderate, 5 severe and 6 extremely severe SNHL. Among the 50 infants with SNHL, 29 (58%) completed hearing follow-up; 19 (65.5%) had improved hearing thresholds, 7 (24.1%) remained stable and 3 (10.3%) progressed. Among 121 infants with normal hearing at birth, 52 (43%) completed hearing follow-up; 2 (1.7%) exhibited late-onset hearing loss. Univariate logistic regression analysis showed that disease grade at birth, SNHL, abnormal neuroimaging, and premature birth were all associated with poor hearing outcomes (all p<0.1). Multivariate logistic regression analysis found that SNHL and premature birth were independent risk factors for poor hearing outcomes (all p<0.05). The authors concluded that “SNHL resulting from cCMV infection presents symptoms at birth and can be improved by antiviral therapy. Poor hearing outcomes are associated with SNHL and prematurity”. This information has been extracted from the Chinese publication using the English abstract and a crude translation of the text (using Google Translate™). Therefore, the results and assessment of validity are very limited. Only 81/176 (46%) infants completed at least 1 hearing follow-up (at 3, 6, 12 and 24 months of age).
Tapasak et al., 2022¹⁴² Retrospective cohort study USA	Symptoms at birth, gender and race/ethnicity	cCMV patients N=170 (153 symptomatic)	SNHL Follow-up: 50.6 months (average follow-up for the 83 infants with SNHL eligible for evaluation)	In this cohort, 46% symptomatic children and 41% asymptomatic children developed SNHL. Of the 170 infants with cCMV, 83 were determined to have SNHL eligible for evaluation. Newborn hearing screening (NBHS) results were available for 69/83 infants, of which 26 passed. 22/26 (85%) eventually developed SNHL by their last visit. Over the monitoring period, 3/20 (15%) children with unilateral SNHL progressed to bilateral SNHL. After excluding those with profound hearing loss at detection, 32/68 (47%) had deterioration in hearing. Within our cohort, females with cCMV were significantly more likely

				to have SNHL than males with cCMV (48/77 [62.3%] vs 35/93 [37.6%]; $p<0.01$). Otherwise, there were no statistically significant differences in incidence of SNHL, CNS involvement, or radiographic evidence of disease between race/ethnicity or gender. The authors concluded that “Approximately one half of children with symptomatic cCMV in our cohort developed SNHL related to CMV. This hearing loss progression may manifest as a worsening in severity and/or progression from unilateral to bilateral hearing loss. The majority of children with symptomatic cCMV eventually end up with severe to profound range loss in at least one ear. Children with symptomatic cCMV are at high risk of delayed onset and progressive hearing loss and should be monitored closely”. This conclusion appears appropriate and is based on a relatively large cohort of infants, although only 17 infants had asymptomatic cCMV. The authors note that their results on the proportion of asymptomatic patients with hearing loss are higher than in previous research and attribute this discrepancy to the very low presence of asymptomatic patients in their study.
Pinninti et al., 2016¹⁴⁰ Prospective cohort study USA	CNS involvement vs no CNS involvement (transient findings) and a group with only petechial rash	Infants with symptomatic cCMV N=160	Hearing and neurodevelopment Follow-up: average 4.6 years	Infants with microcephaly, seizures, lethargy/hypotonia, poor suck and/or neuroimaging findings, with or without other findings, were categorised as the CNS group (n=84). Infants with jaundice, purpura, hepatosplenomegaly, elevated aspartate aminotransferase or thrombocytopenia with or without petechial rash and without CNS involvement were considered as the transient findings group (n=53). Infants with petechial rash without other findings comprised the third group (n=23). None of the included infants received antiviral therapy. Significantly more infants in the CNS group had SNHL at birth (58.2% vs 37.7% in the transient symptoms group and 21.7% in the petechiae group; $p=0.0007$). Of those with SNHL, bilateral SNHL and moderate-profound hearing loss were more frequent in the CNS group and transient symptom group than the petechiae group, but the differences were not statistically significant. The frequency of late-onset SNHL did not differ between groups: 16% in the CNS group vs 13.2% in the transient symptoms group and 8.7% in the petechiae group. Of those assessed for cognitive outcome, significantly more children in the CNS group had IQ<70 (23/39; 59%) than in the transient symptom group (8/39; 20%); $p<0.0001$. None of the 10 children in the petechiae group had IQ<70. The authors concluded that “In infants with symptomatic cCMV, those with evidence of CNS involvement at birth are at significantly higher risk for SNHL and cognitive deficits than those with transient symptoms or a petechial rash

				suggesting a direct relationship between the severity of disease at birth and the risk for adverse outcomes”. This conclusion appears appropriate and is based on a relatively large cohort of infants. However, only 88 children had cognitive outcome data, so the finding relating to cognitive deficit is less certain. The study methods and length of follow-up were acceptable.
Simonazzi et al., 2014¹⁴¹ Retrospective cohort study Italy	Small for gestational age vs not small for gestational age	Infants born to CMV infected mothers N=119 with cCMV (14 symptomatic)	Long-term sequelae (hearing impairment and neurodevelopmental delay) Follow-up: not reported	Eight of the 119 infants with cCMV were small for gestational age (SGA). SGA was defined as birth weight less than the 10th centile. Of the 14 symptomatic newborns, 2 were SGA and 12 were appropriate for gestational age (AGA). Both SGA newborns had long-term sequelae; 1 had monolateral deafness and 1 had monolateral deafness and speech disability. All 12 AGA newborns had long-term sequelae; 4 had hearing loss, 1 had monolateral deafness, 5 had bilateral deafness, 1 had learning disability and 1 had speech disability and learning disability. The authors concluded that “Congenital CMV infection does not seem to be associated with a higher incidence of SGA, and long-term outcomes do not seem to be affected by isolated impaired fetal growth”. The length of follow-up was not reported and long-term sequelae were only reported for the 14 symptomatic newborns, of which only 2 were SGA, therefore, the results should be interpreted with caution.
Audiological screening result at birth (n=2 studies)				
Forli et al., 2024¹⁵² Retrospective cohort study Italy	Newborn hearing screening result – refer vs pass	Children diagnosed with cCMV through newborn hearing screening or symptoms N=61 (35 symptomatic)	SNHL, ophthalmological and neurodevelopmental abnormalities Follow-up: mean 54 months	Nineteen symptomatic newborns received antiviral treatment. At newborn hearing screening, 39 newborns were ‘pass’ bilaterally, 8 were ‘refer’ unilaterally and 14 were ‘refer’ bilaterally. Twenty children (32.7%) developed SNHL (10 symptomatic and 10 asymptomatic at birth); 17 (85%) were ‘refer’ at newborn hearing screening and 3 (15%) were ‘pass’. In 5 patients (25%) a progression of the hearing threshold was observed (mean age at progression 26 months); risk factors for progression were worse final hearing threshold ($p=0.0001$) and worse hearing threshold at birth ($p=0.02$). 4/61 children (6.6%) presented with ophthalmological disorders compatible with cCMV and 16/40 (40%) were diagnosed with neurodevelopmental abnormalities. The authors concluded that “This study highlights the diverse and significant long-term sequelae of cCMV infection detected through early screening. With a significant proportion of cCMV children developing sensorineural hearing loss, ophthalmological and neurodevelopmental issues, the results

				emphasize the importance of continuous, multi-disciplinary follow-up. Early identification and tailored interventions are crucial for improving the long-term health and quality of life of children affected by cCMV. This conclusion appears appropriate and is based on a reasonable sized cohort of infants. The study methods and length of follow-up were acceptable. Whilst only 22 infants were given a 'refer' newborn hearing screening result, the results indicate that infants with a 'refer' result are more likely to develop SNHL than those with a 'pass' result.
Rohren et al., 2024¹⁵³ Retrospective cohort study USA	Newborn hearing screening result – refer (bilateral/unilateral) vs pass	Children with cCMV N=44	SNHL Follow-up: mean age 46 months	Twenty children received antiviral therapy. 33/44 (75%) children had SNHL of varying degree and age at onset (mean age at hearing loss detection 13 months). 17/44 (39%) received a bilateral 'refer' result on universal newborn hearing screen (UNHS). 17/44 (39%) children passed bilaterally; of which 6/17 (35%) acquired SNHL, detected at a mean age of 20 months. 10/44 children had a unilateral 'refer' result; 5/10 (50%) acquired SNHL in the contralateral ear, detected at a mean age of 24 months. The authors concluded that "Among children with cCMV included in this study who passed UNHS in both ears, 35% demonstrated delayed-onset SNHL. Notably, of those children who referred unilaterally, 50% later demonstrated SNHL in the contralateral ear. These findings have implications for audiological monitoring, and potentially antiviral therapy, of children with cCMV. As implementation of universal cCMV screening moves forward, a key aspect of follow-up will be appropriate long-term audiologic monitoring". This conclusion appears appropriate; the methods and length of follow-up were acceptable. However, this was a small study with only 17 and 10 children in the unilateral 'refer' and 'pass' groups, respectively.
Viral load (n=7 studies)				
Marsico et al., 2019¹⁵⁵ Post hoc analysis USA	Whole blood viral load (determined by real-time PCR before and during therapy)	Infants with symptomatic cCMV enrolled in 2 prospective studies of antiviral treatment N=120 (all symptomatic), of which baseline	Hearing and neurodevelopmental assessment Follow-up: up to 24 months	All patients received antiviral treatment (73 for 6 weeks and 47 for 6 months). In patients treated for 6 months, a lower baseline viral load was associated with better hearing outcome at 12 months; median baseline viral load was 3.56 log for children with improved/protected hearing vs 3.93 log for those with worsened hearing or maintenance of the same degree of hearing loss ($p \leq 0.02$). However, clinically meaningful viral load thresholds predictive of SNHL were not identified. No significant association was found between baseline viral load and 6 and 24-month hearing outcome. No significant association was found between baseline viral load and 6, 12 or 24-month hearing

		samples were available for 114 infants		outcome in patients treated for 6 weeks. No significant association was found between baseline viral load and neurodevelopmental outcome at 12 or 24 months. The authors concluded that “In infants with symptomatic congenital cytomegalovirus disease, higher whole blood viral load before initiation of antiviral therapy has no clinically meaningful predictive value for long-term outcomes”. This conclusion appears appropriate and is based on a relatively large cohort of infants. The study methods and length of follow-up were acceptable.
Meyer et al., 2017¹⁵⁶ Retrospective cohort study USA	Viral load in DBS	Children with SNHL N=15 with cCMV (of 57 children with unexplained SNHL)	SNHL Follow-up: 3 months to 10 years of age	When viral load was evaluated in the dried blood spots (DBS) of infants with retrospectively diagnosed cCMV, it was found to be $8.3 \pm 4.1 \times 10^4$ copies/μg DNA. No statistically significant correlation was found between viral load and SNHL severity at follow up. The authors concluded that “A retrospective DBS analysis demonstrated that 26% of patients presenting with unexplained SNHL in childhood had cCMV. DBS testing is useful in the retrospective diagnosis of cCMV, and may provide definitive diagnostic information about the etiology of SNHL”. This was a small retrospective study investigating causes of unexplained SNHL in children; only 15 children had cCMV and few results were reported relating to viral load and long-term sequelae.
Kakei et al., 2024¹⁵⁷ Prospective cohort study Japan	Viral load in blood and urine	Infants with cCMV N=24 (all symptomatic)	Response to treatment (hearing) Follow-up: at 6 months of treatment	All infants received antiviral treatment for 6 months. 4/24 infants had normal hearing on assessment of the ear with best hearing. Of those without normal hearing at baseline, hearing improved in 13/20 infants (responders) and remained the same in 7/20 infants (non-responders) after 6 weeks of treatment. After 6 months of treatment, 14/20 patients were responders and 6/20 were non-responders. The plasma CMV loads at baseline were significantly higher in non-responders than in responders at 6 weeks ($p=0.028$); the difference was no longer significant at 6 months ($p=0.124$). There was a trend of higher blood and urine CMV loads at baseline in non-responders than in responders. The authors concluded that “Infants with a low plasma viral load at screening tend to have an improvement in hearing impairment. Clinicians should be aware of neutropenia during valganciclovir treatment particularly in patients with a low neutrophil count during screening”.

				This was a small study that compared baseline characteristics between responders (improved hearing) and non-responders to antiviral treatment. The authors' conclusions that infants with a low plasma viral load tend to have an improvement in hearing impairment appears appropriate, but the study was very small with short-term follow-up.
Portet Sulla et al., 2024⁵⁶ Prospective cohort study France	Viral load in saliva	Newborns with cCMV detected by screening N=70 with cCMV (7 symptomatic)	Neurologic/hearing assessments Follow-up: 3-6 years	3/7 (42.8%) symptomatic infants later developed sequelae. 5/63 (7.9%) asymptomatic infants later developed sequelae; mainly SNHL and neurodevelopmental impairment. Overall, the occurrence of sequelae was associated with viral load in saliva, whether expressed as copies/mL ($p=0.039$) or copies/million cells ($p=0.006$). Amongst asymptomatic infants, the difference was only statistically significant when expressed as copies/million cells ($p=0.0044$; when expressed as copies/mL $p=0.0765$). The authors summarised that "Our results confirm that saliva swab is a suitable sample for universal neonatal screening. However, identifying newborns that will develop sequelae remains an issue in the management of cCMV infection". This study assessed the diagnostic accuracy of saliva swabs for cCMV screening and the association between viral load in saliva and symptomatology/outcome of cCMV. Whilst an association was found between viral load in saliva and the occurrence of sequelae, only 3 symptomatic infants and 5 asymptomatic infants developed sequelae, so the results are based on a very small number of children.
Fornier et al., 2015⁷⁷ Prospective cohort study Italy	Viral load in blood and urine (determined by real-time PCR)	Asymptomatic newborns with cCMV N=33 (none symptomatic)	Audiological, neurodevelopment and visual assessment Follow-up: 6 years of age	Ten of 33 infants developed late-onset sequelae between the age of 3 months and 5 years; SNHL (8 children) and/or psychomotor retardation (2 children), right-side hemiparesis (1 child). The mean viral load at birth was 17,045 copies/mL in babies who developed late-onset sequelae vs 1,770 copies/mL in those who remained symptom-free ($p=0.0002$). Multivariate logistic regression revealed that the risk of clinical disease crossed the level of 50% with a DNAemia at birth of $\geq 12,000$ copies/mL ($p=0.0002$) and the risk of hearing deficit crossed the level of 50% with a DNAemia at birth of $\geq 17,000$ copies/mL ($p=0.0001$). The authors concluded that "Asymptomatic newborns with a CMV DNAemia at birth of $\geq 12,000$ copies/mL were more likely to experience CMV-related

				sequelae. The risk of hearing deficit increased with a viral load in blood of $\geq 17,000$ copies/mL". The methods and length of follow-up appear appropriate. However, this was a small study including only 10 infants that developed long-term sequelae.
Salome et al., 2020¹⁵⁸ Prospective cohort study Italy	Viral load in blood and urine and clearance time	Newborns with asymptomatic cCMV N=125; 102 of which had follow-up of at least 12 months so included in study (none symptomatic)	SNHL Follow-up: median 2.8 years	Time to viral clearance was 104 days (range 11-659) in blood and 1,437 days (range 230-2,756) in urine. No infant developed a stable delayed SNHL and 88/102 children (86%) presented normal hearing function at all assessments. Only 14/102 (14%) presented a fluctuating hearing impairment, which was bilateral in 7 babies. The transient SNHL was mild in 12 children and moderate in 2. Children with fluctuating SNHL had a significantly higher urine viral load at baseline ($p=0.002$) and more often positive viremia ($p=0.015$) than children with stable normal hearing. None of the infants developed later sequelae such as neurologic disorders, visual loss, thrombocytopenia or liver disease during the follow-up period. The authors concluded that "CMV infected, asymptomatic neonates have a low risk of transient SNHL later in infancy. Positive viremia and high urine viral load at onset are significant risk factors for delayed fluctuating SNHL. These data are relevant for an appropriate follow up plan of these patients". The conclusions appear appropriate, however, only 14 children presented a fluctuating hearing impairment, so the comparison of viral load between those with stable normal hearing and fluctuating SNHL were based on a small number of children. This study is included in the Smyrli 2024 systematic review of impact of symptoms.
Wang et al., 2021¹⁵⁹ Prospective cohort study China	Viral load in saliva	"Likely asymptomatic" (based on the absence of symptoms, without neuroimaging or ophthalmic assessment) newborns with cCMV	Late-onset hearing loss Follow-up: Not reported (beyond 4 years for some children)	4/141 (2.8%) children had late-onset SNHL (2 bilateral and 2 unilateral, ranging from mild to moderate), diagnosed at 1-4 years of age. Children with late-onset SNHL had higher median CMV salivary viral loads at birth than children without hearing loss (median 5.3 vs 3.5 $\log_{10}$ copies/mL; $p=0.03$). Other maternal or childhood factors did not significantly differ between children with or without late-onset SNHL. The authors summarised that "After following 141 children with likely asymptomatic congenital cytomegalovirus infection in a highly immune population in China, 4 children (2.8%) were found to have late-onset hearing loss. No maternal or childhood factors, except higher saliva cytomegalovirus

		N=141 (none symptomatic)		viral load at birth ($p=0.03$), were associated with increased risk of developing a hearing loss". This study compared baseline characteristics between children with and without late-onset hearing loss. The conclusions appear appropriate, however, only 4 children had late-onset SNHL, so the comparisons were based on a very small number of children.
Genotype (n=6 studies)				
Chen et al., 2019¹⁶³ Prospective cohort study China (non-English language publication)	CMVgH1 genotype vs CMVgH2 genotype	Newborns with cCMV in NICU N=21 (all newborns were hospitalised in NICU, therefore, probably all symptomatic)	Clinical manifestations, laboratory examinations, hearing loss and neurological prognosis Follow-up: 3-5 years	14/21 (67%) infants were gH1 type and 7 (33%) were gH2 type. 20 infants (40 ears) had severe hearing loss; it was higher in gH1 group than gH2 group (4/22 vs 0/18 ears; $p=0.023$). The imaging Alarcon score was lower in the gH1 group than the gH2 group (0.4 vs 1.3; $p=0.024$). No significant difference was found in the probability of motor or language development lag in gH1 and gH2 groups. The authors concluded that "Compared with gH2 infection, gH1 infection in children has a younger gestational age. The major type of hearing loss in neonatal period is gH1 infection. Children with gH2 congenital infections are more likely to suffer from nervous systems damage". All infants were hospitalised in the neonatal intensive care unit, so represent a subgroup of cCMV infants. This information has been extracted from the abstract of this non-English language publication. Therefore, it is not possible to assess the validity of the results presented, although the number of infants in the different genotype groups was very small.
Dong et al., 2023¹⁶⁴ Prospective cohort study China	gB, gH and gN genotypes	Newborns with cCMV N=42 (all symptomatic)	Hearing loss and developmental delay Follow-up: Up to 3 years of age	The UL55 (gB), UL75 (gH) and UL73 (gN) genes were successfully sequenced for 32/42 (76%), 37/42 (88%) and 28/42 (67%) newborns, respectively. No significant association was found between CMVgB, gH or gN genotypes and hearing loss or developmental delay. No significant association was found between CMV genotypes and urine viral loads. The authors concluded that "Our findings demonstrate the overall distribution of gB, gH and gN genotypes in infants with symptomatic cCMV infection in Shanghai for the first time. The findings in our study may suggest a possible association between gH1 genotype and early infancy hearing loss. gB3 genotype was associated with a 6.5-fold increased risk of petechiae while gN4a strongly correlated with chorioretinitis due to cCMV infection. No

				significant correlation was found between urine viral loads and CMV genotypes or hearing impairment in cCMV infected infants”. Infants underwent audiological assessment within the first month of life and at 3, 6, 12 and 18 months of age. However, it is unclear whether the hearing loss results relate to newborn hearing loss or longer-term hearing loss (the outcome of interest to this review). Children underwent developmental assessment at least every 6 months; whilst the timing of diagnosis of developmental delay was not presented, it can be assumed that it was beyond the newborn period. The number of patients with different genotypes was very small (0 to 2), limiting the reliability of the study results.
Kasztelewicz et al., 2017¹⁶⁵ Prospective cohort study Poland	11 single nucleotide polymorphisms	Infants at a children's hospital N=72 with cCMV (61 symptomatic)	SNHL Follow-up: 6 months	A panel of 11 candidate single nucleotide polymorphisms (SNPs) was genotyped: TNF rs1799964, TNF rs1800629, TNFRSF1A rs4149570, IL1B rs16944, IL1B rs1143634, IL10 rs1800896, IL10RA rs4252279, IL12B rs3212227, CCL2 rs1024611, CCL2 rs13900, and CCR5 rs333. 22/72 newborns had early-onset SNHL. No significant association was found between severity of cCMV disease (i.e., illness score) and SNPs. However, when analysis was performed according to hearing outcome at birth as well as at the age of 6 months, a significant association between polymorphisms within CCL2 gene (rs13900) and SNHL was observed. In particular, carriers of CT or TT genotype of CCL2 rs13900 had increased risk of hearing loss (OR=3.38; p=0.033 and OR=4.0; p=0.04, for hearing status at birth and at age 6 months, respectively). The authors concluded that “The results of the present study suggest for the first time that polymorphisms in cytokine genes may influence the pathogenesis of cCMV infection. It has been shown that genetic polymorphisms in IL1B and TNF may modulate the susceptibility to intrauterine CMV infection, whereas CCL2 polymorphisms may influence the hearing outcome in children with cCMV infection. Further, larger studies and functional assessment of the SNPs are required to confirm their pathogenic role”. Whilst this was a reasonable sized study, the length of follow-up was relatively short and 16/72 infants who underwent hearing screening at birth did not have auditory brainstem response (ABR) data available for confirmation of hearing loss. In addition, 14 babies (7 with early-onset SNHL and 7 with normal hearing) were lost by the 6-month follow-up assessment.

Medoro et al., 2024 ¹⁶⁶ Prospective cohort study USA	T-cell cytokine responses	Infants with cCMV (and uninfected controls) N=30 with cCMV (22 symptomatic)	Neurodevelopmental outcomes Follow-up: at least 12 months	21/30 (70%) infants received antiviral therapy. 14/30 (47%) had any neurodevelopmental impairment; 9/30 (30%) had developmental delay, 9/30 (30%) had SNHL and 1 (3%) had 'unknown' neurodevelopmental impairment. Infants with subsequent developmental delay lacked detectable CMV-specific T cell responses, with patterns resembling those of uninfected infants, whereas those with normal developmental outcomes had high frequencies of the differentiated memory CD8+ T cells. Two children with progressive SNHL had high frequencies of PD-1+CD8+ T cells over the first year compared with children with nonprogressive SNHL and normal hearing. The authors concluded that "... a lack of in utero T cell differentiation was associated with developmental delay, and high frequencies of PD-1+CD8+ T cells persisted only in children with progressive SNHL. Further work is needed to define the specificity of these T cells and the mechanistic connection to these outcomes". A number of infants who had at least 12 months clinical follow-up had incomplete data on T-cell responses; therefore, the results are based on a small number of infants with relatively short-term follow-up.
Pati et al., 2013 ¹⁶⁷ Prospective cohort study USA	Genotype (UL55 (gB), UL73 (gN) and UL75 (gH)) and mixed vs non-mixed genotype	Children with cCMV V from CHIMES study and another cohort N=131 (37 symptomatic)	Hearing Follow-up: mean 29.6 months	The UL55, UL75 and UL73 genes were successfully amplified and genotype assigned for 118/131 (90%), 118/131 (90%) and 109/131 (83%) infants, respectively. No particular genotype of UL55, UL73, UL75 or US28 was associated with symptomatic disease or SNHL at birth. In the cohort of 52 infants with long-term hearing outcome data, 5 had hearing loss (3 at birth and 2 developed late-onset hearing loss at 12.4 months and 16.4 months of age). 3/5 (60%) children with hearing loss at follow up had infection with multiple strains, compared with 16/48 (33%) children with normal hearing at follow up (p=0.23). The authors concluded that "Mixed infection is common in infants with cCMV but is neither associated with symptomatic infection nor with SNHL". This conclusion appears appropriate and is based on a reasonable sized cohort of infants. However, only 52 infants had long-term hearing outcome data, of which only 5 had hearing loss. The study methods and length of follow-up were acceptable.
Rovito et al., 2018 ¹⁶⁸	Gene expression differences and viral load	Children retrospectively diagnosed with cC	Long-term impairments (hearing, visual, neurological, motor,	Differential expression of individual genes was compared between children with cCMV who developed long-term impairment and those that did not; there were no statistically significant differences between the groups. The

Retrospective cohort study (same cohort as Korndewal et al., 2017[ref]) Netherlands		MV from neonatal DBS N=12 with cCMV (5 symptomatic)	cognitive, speech-language) Follow-up: 6 years of age	differences in gene expression were assessed in relation to the logarithm of CMV viral load a continuous variable; there were no statistically significant differences observed. Pathway analysis suggested the involvement of innate immune response with higher CMV viral loads, and of anti-inflammatory markers in cCMV infected children who did not develop long-term impairments. The T cell exhaustion observed in infected newborns, in particular with higher viral load, did not correlate with long-term impairment, therefore other mechanisms are likely to be involved in the long-term immune dysfunction. The authors summarised that “Despite these data demonstrate limitation in determining prognostic markers for LTI by means of transcriptome analysis, this exploratory study represents a first step in unravelling the pathogenesis of cCMV, and the aforementioned pathways certainly merit further evaluation”. This was a very small study which included 12 children retrospectively diagnosed with cCMV using neonatal DBS (from a previous retrospective cohort study by Korndewal et al., 2017[ref]); 6 children with long-term impairments were compared against 6 children without long-term. Therefore, the results should be interpreted with caution.
Cerebrospinal fluid (CNS involvement) (n=2 studies)				
Czech-Kowalska et al., 2021 ¹⁶⁹ Prospective cohort study Poland	Detection of CMV DNA in cerebrospinal fluid (CSF) by PCR (CSF-CMV-PCR) as a marker of CNS involvement	Newborns with cCMV N=168 (162 symptomatic)	Clinical and laboratory assessment Follow-up: after 4-8 weeks of treatment. Audiologic assessment lasted at least 5 months	Antiviral treatment was used in 149 infants. Twenty-three (13.7%) infants had positive CSF-CMV-PCR results and 145 (86.3%) infants had negative CSF-CMV-PCR results. After 4-8 weeks of treatment, infants with positive CSF-CMV-PCR presented with a higher prevalence of the following clinical abnormalities: splenomegaly, microcephaly, seizures and opisthotonos. There were no differences between groups according to laboratory results apart from thrombocytopenia. The CSF-CMV-PCR results were negative in 15 infants, still positive in 5 infants and not available for 3 infants. The authors concluded that “The detection of CMV DNA in CSF is associated with a higher rate of CNS damage including abnormal MRI neuroimaging and severe hearing loss. Therefore, detection of CMV DNA in CSF may be considered as a marker of severe CNS injury in cCMV infection. However, the very low prevalence of the positive CSF-CMV-PCR results, even in infants with proven CNS involvement, may imply its limited role in clinical practice. This study reported very few results relating to longer term outcomes; clinical abnormalities after 4-8 weeks of treatment were presented only for infants

				with positive CSF-CMV-PCR results. The authors' cautious conclusion appears appropriate.
Goycochea-Valdivia et al., 2017 ¹⁷⁰ Retrospective cohort study Spain	Detection of human CMV DNA in CSF by PCR as a marker of CNS involvement (lumbar puncture performed at a median age of 5 days)	Newborns with cCMV N=136 (77 symptomatic)	Audiological and neurological outcomes Follow-up: 6 months	Antiviral treatment was used in 105 infants (all symptomatic infants and 28 asymptomatic infants). Twenty-one (15.4%) infants had positive CSF hCMV-PCR results and 115 (84.6%) had negative results. Significantly more infants in the positive group were symptomatic (17/21 [81%] vs 60/115 [52.2%] in the negative group; $p=0.01$). There were no differences between groups regarding the rate of microcephaly, neuroimaging abnormalities, neurological sequelae at 6 months of age or plasma viral load. SNHL at birth was associated with a positive CSF hCMV-PCR result ($OR=3.49$; $p=0.04$), although no association was found at 6 months of age ($OR=2.89$; $p=0.07$). The authors concluded that "A positive hCMV-PCR result in CSF is associated with symptomatic cCMV and SNHL at birth. However, no differences in neuroimaging studies, plasma viral load, or outcomes at 6 months were found. These results suggest that hCMV-PCR in CSF may not be a useful prognostic marker in cCMV. The authors' conclusion appears appropriate and is based on a relatively large cohort of infants, although only 21 infants had positive CSF hCMV-PCR results.
Multiple tests/characteristics assessed (n=18 studies)				
Poletti de Chaurand et al., 2024 ¹⁰¹ Prospective cohort study Italy	Antenatal characteristics and clinical characteristics at birth	Infants screened for cCMV (targeted screening) N=11 with cCMV (7 symptomatic)	Hearing, cognitive, motor and speech Follow-up: 6-36 months	An adverse clinical outcome was observed in 2/11 infants (1 infant died on day 11 and the other had cognitive delay and motor impairment). These 2 infants had the lowest gestational age (just under 34 weeks) of the cohort and both had abnormalities detected on neuroimaging (fetal or neonatal MRI). Two additional infants had mild abnormalities. All 4 infants with abnormalities were symptomatic at birth. The authors formed no conclusions in relation to associations between baseline factors and sequelae, as the study was not designed to investigate this. This was a very small study, without any formal analysis, and so findings related to associations between baseline factors and clinical sequelae should be interpreted with caution.

Puhakka et al., 2022¹¹⁸ Prospective cohort study Finland	Primary vs non-primary maternal infection and trimester of infection and symptoms at birth	Children with cCMV from a newborn screening study N=32 (3 symptomatic)	Hearing Follow-up: median age 3.1 years	Five children had unilateral hearing loss and 27 children had normal hearing. Unilateral hearing loss at 3-4 years of age was not associated with the type of maternal infection: 3/15 (20%) children born to mothers with non-primary infection had hearing loss at 3-4 years, compared to 2/10 (20%) children with primary maternal infection. In both infants with hearing loss following primary maternal infection, the infection was during the second to third trimester (2/4 infants infected in the second/third trimester had hearing loss vs 0/6 infants infected in the first trimester). Finally, unilateral hearing loss at 3-4 years of age appeared to have some association with symptoms at birth: 1/3 (33%) children who were symptomatic at birth had hearing loss at 3-4 years, compared to 4/29 (14%) children who were asymptomatic at birth. No attempt was made to examine interactions between the baseline variables and sequelae (for example, assessing if a particular combination of characteristics has an effect on outcome, that is over and above what could be inferred from the univariate results). The authors' conclusion was, "Adverse hearing outcome occurred both in children born to mothers with primary infection after the first trimester, and non-primary infection during pregnancy." This was a small study, in which only 5 infants had long-term hearing loss. Findings should therefore be treated with caution.
Noorbakhsh et al., 2022¹²⁰ Prospective cohort study Iran	Symptoms at birth and primary vs non-primary maternal infection	Newborns with cCMV admitted to NICU and from 2 other prospective screening studies in Iran N=22 (8 symptomatic)	SNHL Follow-up: 24 months of age	Hearing loss at follow up occurred in 2/8 (25%) children that were symptomatic at birth and in 1/14 (7.1%) children that were asymptomatic at birth. However a statistical analysis was not performed. Meanwhile, there was no significant association between the type of maternal infection and later hearing loss, with hearing loss in 1/6 (16.5%) children whose mothers had primary infection and in 1/11 (9.1%) children whose mothers had non-primary infection (p=0.4). There were also no significant associations observed between later hearing loss and each of maternal age (p=0.9), sex (p=0.3), preterm birth (p=0.7), jaundice at birth (p=0.9) or intrauterine growth restriction (p=0.2). No attempt was made to examine interactions between the different baseline variables and sequelae. Antivirals were provided to 5/8 symptomatic infants. The authors' conclusion was, "neonates with symptomatic CMV infection are more prone to develop HL."

				The authors' conclusion is difficult to unequivocally support. This was a very small study, and a lack of statistical analysis for the primary outcome, along with the failure to adjust for confounding, were major limitations.
Townsend et al., 2013 ¹²² Prospective cohort study UK and Sweden	Symptoms at birth and primary vs non-primary maternal infection	Newborns with cCMV (and controls) N=176 with cCMV (19 symptomatic)	Neurological sequelae Follow-up: at least age 5	8/19 (42%) children symptomatic at birth had sequelae (mild, moderate or severe), compared to 19/135 (14%) of those asymptomatic at birth ($p = 0.006$). 5/73 (6.8%) newborns whose mothers had presumed or confirmed primary infection had moderate or severe sequelae, compared to 9/39 (23.1%) newborns whose mothers had presumed or confirmed non-primary infection. This was not subjected to statistical analysis, and no interactions between variables were examined. No infants received antivirals. The authors concluded that, "Nonprimary infections contributed substantially to the burden of childhood congenital CMV disease". This relatively large study was weakened by the limited statistical analysis and the lack of adjustment, making the authors' conclusion somewhat speculative. In addition, no outcomes were reported for 22/176 (12.5%) children and the type of maternal infection was unknown for 42/154 (27.3%) of the children with outcomes reported. This study is included in the Maltezou 2020 systematic review of type of maternal infection.
Demmler-Harrison et al., 2020 ¹¹⁷ Prospective cohort study USA	Primary vs non-primary infection; symptomatic vs asymptomatic cCMV; time of maternal infection [first half vs second half]. The study also compared with 51 uninfected controls but these results are not included here.	Newborns with cCMV (and uninfected controls) N=186 with cCMV (77 symptomatic)	SNHL Follow-up: mean 15.4 years	57/77 (74.0%) children who were symptomatic at birth had hearing loss, compared to only 22/109 (20.2%) who were asymptomatic. In addition, 23/63 (36.5%) children whose mothers had primary infection had later hearing loss, compared to 2/22 (9.1%) whose mothers had non-primary infection. When considering the interaction between primary/non-primary and symptomatic/non-symptomatic, newborns were more likely to develop hearing loss if symptomatic and the mother had a primary infection (9/12) than symptomatic and the mother had a non-primary infection (0/4); and more likely to develop hearing loss if asymptomatic and the mother had a primary infection (14/51) than asymptomatic and the mother had a non-primary infection (2/18). Finally, 8/19 (42.1%) children whose mothers were infected in the first half of pregnancy had later hearing loss, compared to 14/38 (36.8%) whose mothers were infected in the second half of pregnancy. Statistical analyses were not conducted for these specific analyses.

				The authors' conclusions were, "Sensorineural hearing loss occurred in children born to mothers with both primary and non-primary CMV infections, and in both asymptomatic and symptomatic congenital CMV infection, but was more common after maternal primary infection". The authors' conclusions appear plausible in this relatively large study, despite the lack of statistical analysis or adjustment for confounding.
Turriziani Colonna et al., 2020¹¹⁴ Retrospective cohort study Italy	Symptomatic vs asymptomatic cCMV and trimester of infection	Newborns with cCMV treated with VGC N=36 (12 symptomatic)	Long-term clinical, audiological, visual, neurocognitive and behavioural outcome Follow-up: average age 4.23 years	There were some associations between symptoms at birth and a greater risk of later SNHL and language disorders: 5/12 (41.7%) symptomatic newborns went on to develop SNHL, compared to 1/24 asymptomatic newborns (4.2%) [p=0.012], whilst 3/12 (25%) symptomatic newborns went on to develop language disorders, compared to 3/24 asymptomatic newborns (12.5%). However, there were also associations favouring symptomatic newborns: 2/12 (16.7%) symptomatic newborns went on to present abnormal neuropsychological tests, compared to 9/12 (75%) asymptomatic newborns. Likewise, 2/12 (16.7%) symptomatic newborns went on to present abnormal behavioural test results, compared to 6/12 (50%) asymptomatic newborns. Cognitive impairment was equivalent between groups, being developed by 1/12 (8.4%) symptomatic newborns and 2/24 (8.4%) asymptomatic newborns. There were also associations between the trimester of maternal infection and abnormal performance on the neurocognitive and behaviour tests: 9/15 (60%) newborns whose mothers were infected in the 1st trimester had abnormal responses, compared 4/9 (44.4%) newborns whose mothers were infected in the 3rd trimester. The difference between 1st and 3rd trimester was non-significant [p>0.05]. No interactions between variables were considered. All infants received antivirals (VGC). The authors' conclusions were, "Our study shows that both symptomatic and asymptomatic newborns with cCMV infection develop long-term sequelae, particularly in the behavioural and communicative areas, independently from the trimester of maternal infection." The authors' conclusion appears to be partially backed up by the evidence from this small study for the specific association between symptoms at birth and later hearing outcomes. However, the analyses of most other sequelae were not subjected to statistical analysis and adjustment for confounding was not carried out in any analyses.

Zavattoni et al., 2014¹⁰⁵ Retrospective cohort study Italy	Timing of maternal infection, symptomatic vs asymptomatic cCMV and other fetal and neonatal parameters	Newborns with cCMV N=150 with cCMV (28 symptomatic) of which 115 had long-term data	Physical and clinical progress, hearing, neurologic and cognitive assessment and fundus oculi Follow-up: until school age	9/26 (34.6%) children who were symptomatic at birth developed sequelae, compared to 12/89 (13.5%) who were asymptomatic at birth [$p=0.021$]. 15/58 (25.9%) children whose mothers were infected in the 1st trimester developed sequelae, compared to 6/57 (10.5%) children whose mothers were infected later [$p=0.05$]. No significant effects on the risk of sequelae were seen for other fetal baseline variables (HCMV DNA in amniotic fluid and antigenaemia, viraemia, HCMV DNAemia, and IgM in fetal blood) and other neonatal variables (antigenaemia, HCMV DNAemia, IgM). No interactions were examined. The authors' conclusions were, "single clinical, virological, or instrumental features are poor predictors of clinical symptoms at birth and long-term sequelae, but the combined evaluation of maternal, fetal, and neonatal findings may increase prognostic significance." The authors' conclusions are a cautious but accurate summary of the findings. However, this large and well-conducted study appears to provide good evidence of the association of sequelae with symptoms at birth and timing of maternal infection.
Bilavsky et al., 2015⁷⁶ Retrospective cohort study Israel	Neuroimaging (brain ultrasound for detecting lentificulated vasculopathy [LSV]), hearing at birth and treatment	Infants with cCMV N=141 (89 symptomatic) This study contained 4 cCMV groups: group 1 had solitary LSV (without antiviral treatment), group 2 had solitary LSV (with antiviral treatment), group 3 had solitary LSV with abnormal hearing at birth (with antiviral treatment) and group 4 were asymptomatic.	Hearing deterioration/loss Follow-up: more than 1 year (up to 4 years)	For the 3 groups where there had been no hearing loss at birth, later hearing loss was observed in 11/13 (84.6%) in group 1, 0/51 in group 2, and 5/52 (9.6%) in group 4. Hearing deterioration was significantly higher in group 1 than in group 2 ($p<0.001$) and group 4 ($p<0.001$) and also in group 4 than in group 2 ($p=0.008$). The authors concluded that, "Lenticulostriated vasculopathy is a relatively common finding in infants with symptomatic cCMV, a sign of central nervous system involvement and further hearing deterioration." This conclusion is not fully supported by the data. Because the groups differed from each other by more than 1 independent variable (e.g., groups 2 and 3 received antiviral treatment) it is difficult to form conclusions about the effects of any single independent variable, apart from the likelihood that symptomatic infection leads to more long-term hearing loss than asymptomatic infection.

Mardegan et al., 2024⁹⁷ Retrospective cohort study Switzerland	Neuroimaging (ultrasound and MRI) and symptoms at birth	Infants with cCMV N=66 (56 symptomatic)	SNHL and developmental delay Follow-up: median age 34 months	Symptoms at birth (axial hypotonia [p=0.04] or petechia [p=0.004]) were associated with SNHL at follow up. There were also trends for an association between abnormal baseline US [p=0.06] or abnormal baseline MRI [p=0.1] with later SNHL. Vacuolization of anterior temporal lobe and ventricular septation on baseline MRI were associated with developmental delay [both p=0.05]. No US findings showed an association with this outcome. Microcephaly [p=0.05] and abnormal MRI [p=0.03] at baseline were associated with the composite later outcome of SNHL, developmental delay and/or abnormal vestibular function. Interactions between baseline variables were not evaluated. 20 infants received antivirals until 6 months of age (19 symptomatic and 1 asymptomatic) The authors' conclusions are, "this study highlights factors that are directly linked to a poor long-term outcome, including microcephaly and MRI abnormalities." The authors' conclusions are plausible, but adjustment for confounding was not performed in this moderately-sized study.
Oosterom et al., 2015⁹⁸ Retrospective cohort study Netherlands	Timing of maternal infection and MR /ultrasound	Infants diagnosed with cCMV admitted to NICU or maternity unit N=36 (26 symptomatic)	Developmental outcome and hearing loss Follow-up: until 5 years of age for hearing	Trimester of infection was known in 17/36 (47%) infants. Any adverse sequelae were observed in 8/10 (80%) children whose mothers had been infected in the first trimester, compared to 1/7 (14.3%) children whose mothers had been infected in the 2nd or 3rd trimester. Neurodevelopmental (ND) sequelae were observed in 15/15 (100%) children with severe US abnormalities at baseline, compared to 1/21 (4.8%) children with mild or no abnormalities. SNHL at follow up was observed in 7/15 (46.7%) children with severe US abnormalities at baseline, compared to 2/21 (9.5%) children with mild or no abnormalities. 6 children died; all had severe US abnormalities. ND sequelae were observed in 13/13 (100%) children with severe MRI abnormalities at baseline, compared to 2/7 (28.6%) children with mild abnormalities. SNHL at follow up was observed in 7/13 (53.8%) children with severe MRI abnormalities at baseline, compared to 2/7 (28.6%) children with mild abnormalities.

				No interactions between baseline variables were assessed. 5 children received antivirals, but no information was provided on which risk factor groups these belonged to. The authors concluded as follows: "Infants with a first trimester cCMV infection are most at risk of severe cerebral abnormalities and neurological sequelae. MRI and not cUS, enables an early diagnosis of migrational disorders, which can improve prediction of outcome" All infants were admitted to a neonatal intensive care unit or maternity unit, so represent a subgroup of cCMV infants. This was a small study, with no adjustments for confounding, and so the conclusions of this study should be interpreted with caution.
Kabani et al., 2023¹⁶¹ Prospective cohort study USA	Viral load in saliva and urine and symptoms at birth	Newborns with cCMV N=384 (39 symptomatic)	SNHL Follow-up: up to 48 months of age	Newborn saliva viral load was associated with the development of SNHL (viral load in those developing SNHL = 7.4×10^6 IU/mL; viral load in those with normal hearing at birth = 1.74×10^6 IU/mL; $p=0.004$). A significant effect was not observed in separate symptomatic or asymptomatic sub-groups [SYMPTOMATIC: viral load in symptomatic children with SNHL = 4.4×10^7 IU/mL; viral load in symptomatic children with normal hearing at birth = 4.1×10^6 IU/mL; $p=0.09$]; ASYMPTOMATIC: viral load in asymptomatic children with SNHL = 4.2×10^6 IU/mL; viral load in asymptomatic children with normal hearing at birth = 1.74×10^6 IU/mL; $p=0.19$). These data do not permit an evaluation of interactions between symptoms at birth and viral load. The authors concluded that, "VL is not a useful marker for hearing loss in cCMV, and should not be used for counseling patients and families regarding outcomes of cCMV." Although a relatively large study, the analysis of data did not adjust for confounding. The most significant limitation of the study was the ambiguity concerning the timing of SNHL measurement; whilst the paper implies that the SNHL value used in the analysis is measured at follow up this is not explicitly stated. This ambiguity is further increased by the comparator group being made up of 'people with normal hearing <i>at birth</i>.' A more useful comparator in terms of predicting <i>later</i> hearing loss would be 'people with normal hearing at <i>follow up</i>.'
Rovito et al., 2017a¹⁴³	Symptomatic at birth vs asymptomatic at birth and genotype	Children retrospectively diagnosed with cCMV	Long-term impairments (hearing, visual, neurological, motor,	Long-term impairments (LTI) in the child were more likely when mothers were HLA-G del/del than otherwise (OR 3.542, 95% CI 1.397–8.977, $p = 0.008$). HLA-C and HLE-E genotypes in the mother were not associated with LTI in

Retrospective cohort study (same cohort as previous Rovito 2017) Netherlands		MV from neonatal DBS N=96 (19 symptomatic)	cognitive, speech-language) Follow-up: 5 years of age	the child. None of the HLA-G, HLA-C or HLA-E genotypes in the child were related to LTI in the child. 11/19 (57.9%) symptomatic newborns went on to get 1 or more LTIs, compared to 16/77 (20.8%) asymptomatic newborns. Interactions between baseline variables were not evaluated. The authors' conclusion is as follows: "the potential role of maternal-child HLA in controlling CMV infection and cCMV-related disease, and the clinical value as predictor for long-term outcome certainly deserve further evaluation". The authors' tentative conclusion seems appropriate for this moderately sized and unadjusted study.
Rovito et al., 2017b¹⁴⁴ Retrospective cohort study (same cohort as previous Rovito 2017) Netherlands	Symptomatic at birth vs asymptomatic at birth and T and B cell markers	Children retrospectively diagnosed with cCMV from neonatal DBS N=99 with cCMV (16 symptomatic)	Long-term impairments (hearing, visual, neurological, motor, cognitive, speech-language) Follow-up: 5 years of age	Children who had lower numbers of B cells at birth (as measured by KREC copies per ml) were more likely to develop LTI than those with higher numbers of B cells ($p=0.005$). 8/16 (50%) symptomatic newborns went on to get 1 or more LTIs, compared to 14/83 (16.9%) asymptomatic newborns. Interactions between baseline variables were not evaluated. The authors' conclusion is as follows: "the potential protective role of B cells in controlling cCMV-related disease and the clinical value of this marker as a predictor of long-term outcome merit further evaluation". The authors' tentative conclusion seems appropriate for this moderately sized and unadjusted study.
Kawada et al., 2015¹⁶⁰ Prospective cohort study Japan	Viral load in blood and urine at diagnosis	Infants referred for CMV testing because of failed newborn hearing screening or suspected to have cCMV because of other clinical abnormalities	SNHL Follow-up: 1 year of age	Mean CMV viral load at diagnosis in whole blood was 6.6×10^2 copies/mL in newborns with SNHL at 1 year, and 4.2×10^2 copies/mL in those without SNHL at 1 year ($p>0.05$). Mean CMV viral load at diagnosis in urine was 8.8×10^5 copies/mL in newborns with SNHL at one year and 2.6×10^4 copies/mL in those without SNHL at 1 year ($p>0.05$). No baseline interactions were investigated. All 9 patients were treated with antivirals for 6 weeks.

		N=9 with cCMV (all symptomatic)		The authors' conclusion was, "Our results suggest that CMV load at diagnosis cannot predict the hearing outcome". This was an extremely small study, and so there is a possibility that type II errors may explain the lack of significant findings.
Capretti et al., 2017¹²¹ Prospective cohort study Italy	Clinical, laboratory and instrumental evaluations, including neuroimaging; symptomatic vs asymptomatic	Infants with cCMV N=48 (18 symptomatic)	Ophthalmological abnormalities and/or visual impairment Follow-up: mean 35 months	Visual impairment at last follow-up was suspected or detected in 4/18 (22%) symptomatic children and in 0/30 (0%) asymptomatic children ($p = 0.01$). Visual impairment was also associated with signs of central nervous system (CNS) involvement ($p < 0.001$). No association was found with the type of maternal infection. 12/18 (67%) symptomatic infants had SNHL; in 8 infants it was diagnosed at birth, while it was late-onset in 4 cases. No cases of SNHL were reported among asymptomatic infants. No interactions were described. 8/8 infants in the symptomatic group with ocular findings at birth were provided with antivirals (it is unclear whether infants with other symptoms at birth received antivirals). The authors concluded that, "ocular involvement seems to be common, even if often not severe, among symptomatic infants, while is rare among asymptomatic infants. Ophthalmological involvement is more common in infants with CNS involvement at birth, and both primary and non-primary maternal infections may be associated with symptomatic infections and ocular disease." This unadjusted study suggests that symptomatic newborns are at a slightly greater risk of visual impairment. The sample size was relatively small, but, despite this, findings were statistically significant.
De Cuyper et al., 2024¹¹⁵ Prospective cohort study (Flemish CMV registry) Belgium	Antenatal characteristics, clinical findings, imaging, viral load, hearing status at birth	Newborns with untreated cCMV N=387 (113 symptomatic)	Hearing evolution (long-term neurodevelopmental and audiological follow-up) Follow-up: at least 4 years	Children born to mothers infected in their first trimester had a greater odds of late-onset hearing loss than those born to mothers in their second or third trimester [OR (95% CI): 10.10 (2.90 to 34.48)]. None of the children born to mothers infected in their third trimester developed late-onset hearing loss. Late-onset hearing loss occurred in 12/135 ears (8.9%) of children with symptomatic infection and 31/548 ears (5.7%) of children with an asymptomatic infection. Children with unilateral hearing loss at birth had greater odds of developing late-onset hearing loss than those with normal hearing at birth [OR (95% CI): 2.68 (0.98-7.29)]. Children who went on to develop late-onset hearing loss had a higher viral load at baseline than those who did not develop such sequelae, though this was not statistically significant [Hedges g, 0.22 (-0.60 to 1.04)].

				Interactions between baseline variables were not assessed. None of the newborns received antiviral treatment. The authors concluded that, "Findings of this cohort study support that ongoing audiological follow-up for untreated children with congenital hearing loss is important, as the majority of patients had hearing deterioration. The timing of seroconversion was associated with the risk of developing late-onset hearing loss." The conclusion that hearing loss at birth is a risk factor for late-onset hearing loss is not fully supported by the evidence, which was uncertain. However the conclusion that the timing of maternal infection is related to late-onset hearing loss is supported by the data. However, the lack of adjustment for confounding prevents any assumption of causality. This study is included in the Vande Walle 2024 systematic review of MRI
Keymeulen et al., 2023¹¹⁶ Prospective cohort study (Flemish CMV registry) Belgium	Timing of seroconversion, fetal ultrasound and MRI, clinical features at birth, US, MRI, antiviral therapy	Children with cCMV N=753 (261 symptomatic)	Neurodevelopmental outcome Follow-up: most followed up beyond 1 year of age	Any long-term impairment was more likely in those who were severely symptomatic at birth (104/190) compared to those who were not severely symptomatic (99/533) ($p<0.001$). Any long-term impairment was also more likely in those children whose mothers were infected in the first trimester (95/233) compared to those who were not (44/262) ($p<0.001$). No interactions between baseline variables were investigated. Antivirals were received by 180/261 symptomatic children. The authors concluded that, "both symptomatic and asymptomatic children can develop sequelae, independent of the timing of seroconversion." This large study did not adjust for confounding. However, the authors' conservative conclusions are supported by their findings.
Soriano-Ramos et al., 2024⁹⁹ Prospective cohort study Spain	Type of maternal infection, newborn symptoms, US and MRI, T-cell immune responses	Newborns with cCMV N=64 (42 symptomatic)	Hearing loss and neurological impairment Follow-up: median age 25.3 months	Long-term sequelae were associated with the following baseline variables on a bivariate analysis: symptomatic cCMV ($p=0.012$), presence of abnormal physical examination at birth ($p=0.006$), abnormal findings in first cUS ($p=0.016$), presence of white matter abnormalities in cUS ($p=0.008$), and a failed hearing screening test ($p<0.001$). Sequelae were also associated with a lower total lymphocyte count ($p=0.013$), a lower CD4+ T-cell count ($p=0.020$), and a lower CD4+ / CD8+ ratio ($p=0.03$). No relationship was observed with CMV-specific IFN-γ-CD8+ or CD4+ lymphocyte levels. No interactions between baseline variables were investigated. 38 children received antivirals (17/18 with sequelae and 21/46 without sequelae).

				Multivariable analysis showed that sequelae were independently associated with lower total lymphocyte count (adjusted OR 0.549, 95% CI 0.323-0.833, $p=0.012$). There was also a trend toward an association with higher absolute count of CD8+ T cells ($p=0.090$). The authors concluded that, "CMV-specific IFN-γ-CD4+ and CD8+ T-lymphocyte responses in neonates with cCMV were not predictive of long-term sequelae." This study was enhanced by its rigorous analysis, though it is unclear if the moderate sample size was sufficient for such a large multivariable analysis. The authors' cautious conclusion is therefore justified, and further work in this area is warranted.
--	--	--	--	--

Appraisal of study quality and risk of bias

Table 29 presents the results of the ROBIS assessment of included systematic reviews (for reviews relevant to Criterion 7 and the Zheng review, relevant to Criterion 4). Table 30 presents the results of the QUADAS-2 evaluation of all included prognostic accuracy studies relevant to Criterion 7.

Table 29: ROBIS assessment of all included systematic reviews

STUDY	DOMAIN APPRAISAL
Vande Walle, 2024 ⁷⁹	DOMAIN 1: STUDY ELIGIBILITY CRITERIA
	Describe the study eligibility criteria, any restrictions on eligibility and whether there was evidence that objectives and eligibility criteria were pre-specified:
	Did the review adhere to pre-defined objectives and eligibility criteria? PN
	Were the eligibility criteria appropriate for the review question? PY
	Were eligibility criteria unambiguous? N
	Were any restrictions in eligibility criteria based on study characteristics appropriate (e.g. date, sample size, study quality, outcomes measured)? PN
	Were any restrictions in eligibility criteria based on sources of information appropriate (e.g. publication status or format, language, availability of data)? PN
	Concerns regarding specification of study eligibility criteria HIGH CONCERN
	DOMAIN 2: IDENTIFICATION AND SELECTION OF STUDIES
	Describe methods of study identification and selection (e.g. number of reviewers involved):
	Did the search include an appropriate range of databases/electronic sources for published and unpublished reports? Y
	Were methods additional to database searching used to identify relevant reports? N
	Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible? PN
	Were restrictions based on date, publication format, or language appropriate? PN
	Were efforts made to minimise error in selection of studies? PN
	Concerns regarding methods used to identify and/or select studies HIGH CONCERN
	DOMAIN 3: DATA COLLECTION AND STUDY APPRAISAL
	Describe methods of data collection, what data were extracted from studies or collected through other means, how risk of bias was assessed (e.g. number of reviewers involved) and the tool used to assess risk of bias:
	Were efforts made to minimise error in data collection? PY
	Were sufficient study characteristics available for both review authors and readers to be able to interpret the results? N
	Were all relevant study results collected for use in the synthesis? PY
	Was risk of bias (or methodological quality) formally assessed using appropriate criteria? Y
	Were efforts made to minimise error in risk of bias assessment? PN
	Concerns regarding methods used to collect data and appraise studies HIGH CONCERN

STUDY	DOMAIN APPRAISAL
	DOMAIN 4: SYNTHESIS AND FINDINGS
	Describe synthesis methods:
	Did the synthesis include all studies that it should? PN
	Were all pre-defined analyses reported or departures explained? NI
	Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies? Y
	Was between-study variation (heterogeneity) minimal or addressed in the synthesis? NA
	Were the findings robust, e.g. as demonstrated through funnel plot or sensitivity analyses? NA
	Were biases in primary studies minimal or addressed in the synthesis? NA
	Concerns regarding the synthesis and findings UNCLEAR
	OVERALL RISK OF BIAS IN THE REVIEW HIGH RISK OF BIAS
Chatzakis, 2020 ¹⁰²	DOMAIN 1: STUDY ELIGIBILITY CRITERIA
	Describe the study eligibility criteria, any restrictions on eligibility and whether there was evidence that objectives and eligibility criteria were pre-specified:
	Did the review adhere to pre-defined objectives and eligibility criteria? PY
	Were the eligibility criteria appropriate for the review question? PY
	Were eligibility criteria unambiguous? PN
	Were any restrictions in eligibility criteria based on study characteristics appropriate (e.g. date, sample size, study quality, outcomes measured)? Y
	Were any restrictions in eligibility criteria based on sources of information appropriate (e.g. publication status or format, language, availability of data)? Y
	Concerns regarding specification of study eligibility criteria LOW CONCERN
	DOMAIN 2: IDENTIFICATION AND SELECTION OF STUDIES
	Describe methods of study identification and selection (e.g. number of reviewers involved):
	Did the search include an appropriate range of databases/electronic sources for published and unpublished reports? PY
	Were methods additional to database searching used to identify relevant reports? N
	Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible? PN
	Were restrictions based on date, publication format, or language appropriate? PY
	Were efforts made to minimise error in selection of studies? PY
	Concerns regarding methods used to identify and/or select studies LOW CONCERN
	DOMAIN 3: DATA COLLECTION AND STUDY APPRAISAL
	Describe methods of data collection, what data were extracted from studies or collected through other means, how risk of bias was assessed (e.g. number of reviewers involved) and the tool used to assess risk of bias:
	Were efforts made to minimise error in data collection? Y
	Were sufficient study characteristics available for both review authors and readers to be able to interpret the results? Y
	Were all relevant study results collected for use in the synthesis? PY
	Was risk of bias (or methodological quality) formally assessed using appropriate criteria? Y

STUDY	DOMAIN APPRAISAL	
	Were efforts made to minimise error in risk of bias assessment?	Y
	Concerns regarding methods used to collect data and appraise studies	LOW CONCERN
	DOMAIN 4: SYNTHESIS AND FINDINGS	
	Describe synthesis methods:	
	Did the synthesis include all studies that it should?	PY
	Were all pre-defined analyses reported or departures explained?	PY
	Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies?	PN
	Was between-study variation (heterogeneity) minimal or addressed in the synthesis?	Y
	Were the findings robust, e.g. as demonstrated through funnel plot or sensitivity analyses?	N
	Were biases in primary studies minimal or addressed in the synthesis?	PY
	Concerns regarding the synthesis and findings	LOW CONCERN
Maltezou, 2020 ¹¹⁹	OVERALL RISK OF BIAS IN THE REVIEW	LOW RISK OF BIAS
	DOMAIN 1: STUDY ELIGIBILITY CRITERIA	
	Describe the study eligibility criteria, any restrictions on eligibility and whether there was evidence that objectives and eligibility criteria were pre-specified:	
	Did the review adhere to pre-defined objectives and eligibility criteria?	Y
	Were the eligibility criteria appropriate for the review question?	Y
	Were eligibility criteria unambiguous?	Y
	Were any restrictions in eligibility criteria based on study characteristics appropriate (e.g. date, sample size, study quality, outcomes measured)?	Y
	Were any restrictions in eligibility criteria based on sources of information appropriate (e.g. publication status or format, language, availability of data)?	PN
	Concerns regarding specification of study eligibility criteria	LOW CONCERN
	DOMAIN 2: IDENTIFICATION AND SELECTION OF STUDIES	
	Describe methods of study identification and selection (e.g. number of reviewers involved):	
	Did the search include an appropriate range of databases/electronic sources for published and unpublished reports?	PN
	Were methods additional to database searching used to identify relevant reports?	Y
	Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible?	PN
	Were restrictions based on date, publication format, or language appropriate?	PN
	Were efforts made to minimise error in selection of studies?	PY
	Concerns regarding methods used to identify and/or select studies	LOW CONCERN
	DOMAIN 3: DATA COLLECTION AND STUDY APPRAISAL	
	Describe methods of data collection, what data were extracted from studies or collected through other means, how risk of bias was assessed (e.g. number of reviewers involved) and the tool used to assess risk of bias:	
	Were efforts made to minimise error in data collection?	PY
	Were sufficient study characteristics available for both review authors and readers to be able to interpret the results?	PY

STUDY	DOMAIN APPRAISAL	
	Were all relevant study results collected for use in the synthesis?	Y
	Was risk of bias (or methodological quality) formally assessed using appropriate criteria?	Y
	Were efforts made to minimise error in risk of bias assessment?	NI
	Concerns regarding methods used to collect data and appraise studies	LOW CONCERN
	DOMAIN 4: SYNTHESIS AND FINDINGS	
	Describe synthesis methods:	
	Did the synthesis include all studies that it should?	Y
	Were all pre-defined analyses reported or departures explained?	Y
	Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies?	Y
	Was between-study variation (heterogeneity) minimal or addressed in the synthesis?	Y
	Were the findings robust, e.g. as demonstrated through funnel plot or sensitivity analyses?	PY
	Were biases in primary studies minimal or addressed in the synthesis?	N
	Concerns regarding the synthesis and findings	LOW CONCERN
	OVERALL RISK OF BIAS IN THE REVIEW	LOW RISK OF BIAS
Smyrli, 2024 ¹²⁵	DOMAIN 1: STUDY ELIGIBILITY CRITERIA	
	Describe the study eligibility criteria, any restrictions on eligibility and whether there was evidence that objectives and eligibility criteria were pre-specified:	
	Did the review adhere to pre-defined objectives and eligibility criteria?	Y
	Were the eligibility criteria appropriate for the review question?	Y
	Were eligibility criteria unambiguous?	Y
	Were any restrictions in eligibility criteria based on study characteristics appropriate (e.g. date, sample size, study quality, outcomes measured)?	Y
	Were any restrictions in eligibility criteria based on sources of information appropriate (e.g. publication status or format, language, availability of data)?	Y
	Concerns regarding specification of study eligibility criteria	LOW CONCERN
	DOMAIN 2: IDENTIFICATION AND SELECTION OF STUDIES	
	Describe methods of study identification and selection (e.g. number of reviewers involved):	
	Did the search include an appropriate range of databases/electronic sources for published and unpublished reports?	PN
	Were methods additional to database searching used to identify relevant reports?	N
	Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible?	Y
	Were restrictions based on date, publication format, or language appropriate?	Y
	Were efforts made to minimise error in selection of studies?	Y
	Concerns regarding methods used to identify and/or select studies	LOW CONCERN
	DOMAIN 3: DATA COLLECTION AND STUDY APPRAISAL	
	Describe methods of data collection, what data were extracted from studies or collected through other means, how risk of bias was assessed (e.g. number of reviewers involved) and the tool used to assess risk of bias:	

STUDY	DOMAIN APPRAISAL	
	Questions	
	Were efforts made to minimise error in data collection?	Y
	Were sufficient study characteristics available for both review authors and readers to be able to interpret the results?	Y
	Were all relevant study results collected for use in the synthesis?	Y
	Was risk of bias (or methodological quality) formally assessed using appropriate criteria?	Y
	Were efforts made to minimise error in risk of bias assessment?	NI
	Concerns regarding methods used to collect data and appraise studies	LOW CONCERN
	DOMAIN 4: SYNTHESIS AND FINDINGS	
	Describe synthesis methods:	
	Did the synthesis include all studies that it should?	PY
	Were all pre-defined analyses reported or departures explained?	Y
	Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies?	PN
	Was between-study variation (heterogeneity) minimal or addressed in the synthesis?	NA
	Were the findings robust, e.g. as demonstrated through funnel plot or sensitivity analyses?	NA
	Were biases in primary studies minimal or addressed in the synthesis?	NA
	Concerns regarding the synthesis and findings	LOW CONCERN
	OVERALL RISK OF BIAS IN THE REVIEW	LOW RISK OF BIAS
Buca,2021 ¹⁰³	DOMAIN 1: STUDY ELIGIBILITY CRITERIA	
	Describe the study eligibility criteria, any restrictions on eligibility and whether there was evidence that objectives and eligibility criteria were pre-specified:	
	Did the review adhere to pre-defined objectives and eligibility criteria?	Y
	Were the eligibility criteria appropriate for the review question?	Y
	Were eligibility criteria unambiguous?	Y
	Were any restrictions in eligibility criteria based on study characteristics appropriate (e.g. date, sample size, study quality, outcomes measured)?	Y
	Were any restrictions in eligibility criteria based on sources of information appropriate (e.g. publication status or format, language, availability of data)?	Y
	Concerns regarding specification of study eligibility criteria	LOW CONCERN
	DOMAIN 2: IDENTIFICATION AND SELECTION OF STUDIES	
	Describe methods of study identification and selection (e.g. number of reviewers involved):	
	Did the search include an appropriate range of databases/electronic sources for published and unpublished reports?	Y
	Were methods additional to database searching used to identify relevant reports?	Y
	Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible?	PY
	Were restrictions based on date, publication format, or language appropriate?	PY
	Were efforts made to minimise error in selection of studies?	Y
	Concerns regarding methods used to identify and/or select studies	LOW CONCERN

STUDY	DOMAIN APPRAISAL
	DOMAIN 3: DATA COLLECTION AND STUDY APPRAISAL Describe methods of data collection, what data were extracted from studies or collected through other means, how risk of bias was assessed (e.g. number of reviewers involved) and the tool used to assess risk of bias:
	Were efforts made to minimise error in data collection? Y
	Were sufficient study characteristics available for both review authors and readers to be able to interpret the results? Y
	Were all relevant study results collected for use in the synthesis? PY
	Was risk of bias (or methodological quality) formally assessed using appropriate criteria? Y
	Were efforts made to minimise error in risk of bias assessment? Y
	Concerns regarding methods used to collect data and appraise studies LOW CONCERN
	DOMAIN 4: SYNTHESIS AND FINDINGS Describe synthesis methods:
	Did the synthesis include all studies that it should? NI
	Were all pre-defined analyses reported or departures explained? NI
	Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies? NI
	Was between-study variation (heterogeneity) minimal or addressed in the synthesis? Y
	Were the findings robust, e.g. as demonstrated through funnel plot or sensitivity analyses? Y
	Were biases in primary studies minimal or addressed in the synthesis? PY
	Concerns regarding the synthesis and findings LOW CONCERN
	OVERALL RISK OF BIAS IN THE REVIEW LOW RISK OF BIAS
Vos, 2021 ¹²⁶	DOMAIN 1: STUDY ELIGIBILITY CRITERIA Describe the study eligibility criteria, any restrictions on eligibility and whether there was evidence that objectives and eligibility criteria were pre-specified:
	Did the review adhere to pre-defined objectives and eligibility criteria? Y
	Were the eligibility criteria appropriate for the review question? Y
	Were eligibility criteria unambiguous? Y
	Were any restrictions in eligibility criteria based on study characteristics appropriate (e.g. date, sample size, study quality, outcomes measured)? Y
	Were any restrictions in eligibility criteria based on sources of information appropriate (e.g. publication status or format, language, availability of data)? Y
	Concerns regarding specification of study eligibility criteria LOW CONCERN
	DOMAIN 2: IDENTIFICATION AND SELECTION OF STUDIES Describe methods of study identification and selection (e.g. number of reviewers involved):
	Did the search include an appropriate range of databases/electronic sources for published and unpublished reports? Y
	Were methods additional to database searching used to identify relevant reports? NI
	Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible? PY
	Were restrictions based on date, publication format, or language appropriate? Y

STUDY	DOMAIN APPRAISAL	
	Were efforts made to minimise error in selection of studies?	Y
	Concerns regarding methods used to identify and/or select studies	LOW CONCERN
	DOMAIN 3: DATA COLLECTION AND STUDY APPRAISAL	
	Describe methods of data collection, what data were extracted from studies or collected through other means, how risk of bias was assessed (e.g. number of reviewers involved) and the tool used to assess risk of bias:	
	Were efforts made to minimise error in data collection?	Y
	Were sufficient study characteristics available for both review authors and readers to be able to interpret the results?	N
	Were all relevant study results collected for use in the synthesis?	PY
	Was risk of bias (or methodological quality) formally assessed using appropriate criteria?	Y
	Were efforts made to minimise error in risk of bias assessment?	Y
	Concerns regarding methods used to collect data and appraise studies	LOW CONCERN
	DOMAIN 4: SYNTHESIS AND FINDINGS	
	Describe synthesis methods:	
	Did the synthesis include all studies that it should?	PY
	Were all pre-defined analyses reported or departures explained?	PY
	Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies?	Y
	Was between-study variation (heterogeneity) minimal or addressed in the synthesis?	NA
	Were the findings robust, e.g. as demonstrated through funnel plot or sensitivity analyses?	NA
	Were biases in primary studies minimal or addressed in the synthesis?	PY
	Concerns regarding the synthesis and findings	LOW CONCERN
	OVERALL RISK OF BIAS IN THE REVIEW	LOW RISK OF BIAS
Riga, 2018 ¹²⁷	DOMAIN 1: STUDY ELIGIBILITY CRITERIA	
	Describe the study eligibility criteria, any restrictions on eligibility and whether there was evidence that objectives and eligibility criteria were pre-specified:	
	Did the review adhere to pre-defined objectives and eligibility criteria?	PY
	Were the eligibility criteria appropriate for the review question?	Y
	Were eligibility criteria unambiguous?	PN
	Were any restrictions in eligibility criteria based on study characteristics appropriate (e.g. date, sample size, study quality, outcomes measured)?	Y
	Were any restrictions in eligibility criteria based on sources of information appropriate (e.g. publication status or format, language, availability of data)?	PY
	Concerns regarding specification of study eligibility criteria	LOW CONCERN
	DOMAIN 2: IDENTIFICATION AND SELECTION OF STUDIES	
	Describe methods of study identification and selection (e.g. number of reviewers involved):	
	Did the search include an appropriate range of databases/electronic sources for published and unpublished reports?	PY
	Were methods additional to database searching used to identify relevant reports?	Y

STUDY	DOMAIN APPRAISAL	
	Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible?	PN
	Were restrictions based on date, publication format, or language appropriate?	PY
	Were efforts made to minimise error in selection of studies?	PY
	Concerns regarding methods used to identify and/or select studies	HIGH CONCERN
	DOMAIN 3: DATA COLLECTION AND STUDY APPRAISAL	
	Describe methods of data collection, what data were extracted from studies or collected through other means, how risk of bias was assessed (e.g. number of reviewers involved) and the tool used to assess risk of bias:	
	Were efforts made to minimise error in data collection?	Y
	Were sufficient study characteristics available for both review authors and readers to be able to interpret the results?	PY
	Were all relevant study results collected for use in the synthesis?	Y
	Was risk of bias (or methodological quality) formally assessed using appropriate criteria?	Y
	Were efforts made to minimise error in risk of bias assessment?	NI
	Concerns regarding methods used to collect data and appraise studies	LOW CONCERN
	DOMAIN 4: SYNTHESIS AND FINDINGS	
	Describe synthesis methods:	
	Did the synthesis include all studies that it should?	PY
	Were all pre-defined analyses reported or departures explained?	PY
	Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies?	N
	Was between-study variation (heterogeneity) minimal or addressed in the synthesis?	N
	Were the findings robust, e.g. as demonstrated through funnel plot or sensitivity analyses?	NI
	Were biases in primary studies minimal or addressed in the synthesis?	PY
	Concerns regarding the synthesis and findings	HIGH CONCERN
	OVERALL RISK OF BIAS IN THE REVIEW	HIGH RISK OF BIAS
Zheng (2022) ³²	DOMAIN 1: STUDY ELIGIBILITY CRITERIA	
	Describe the study eligibility criteria, any restrictions on eligibility and whether there was evidence that objectives and eligibility criteria were pre-specified:	
	Did the review adhere to pre-defined objectives and eligibility criteria?	Y
	Were the eligibility criteria appropriate for the review question?	Y
	Were eligibility criteria unambiguous?	PN
	Were any restrictions in eligibility criteria based on study characteristics appropriate (e.g. date, sample size, study quality, outcomes measured)?	Y
	Were any restrictions in eligibility criteria based on sources of information appropriate (e.g. publication status or format, language, availability of data)?	PY
	Concerns regarding specification of study eligibility criteria	LOW CONCERN
	DOMAIN 2: IDENTIFICATION AND SELECTION OF STUDIES	
	Describe methods of study identification and selection (e.g. number of reviewers involved):	

STUDY	DOMAIN APPRAISAL	
	Did the search include an appropriate range of databases/electronic sources for published and unpublished reports?	Y
	Were methods additional to database searching used to identify relevant reports?	N
	Were the terms and structure of the search strategy likely to retrieve as many eligible studies as possible?	PY
	Were restrictions based on date, publication format, or language appropriate?	PY
	Were efforts made to minimise error in selection of studies?	NI
	Concerns regarding methods used to identify and/or select studies	LOW CONCERN
	DOMAIN 3: DATA COLLECTION AND STUDY APPRAISAL	
	Describe methods of data collection, what data were extracted from studies or collected through other means, how risk of bias was assessed (e.g. number of reviewers involved) and the tool used to assess risk of bias:	
	Were efforts made to minimise error in data collection?	Y
	Were sufficient study characteristics available for both review authors and readers to be able to interpret the results?	PY
	Were all relevant study results collected for use in the synthesis?	PY
	Was risk of bias (or methodological quality) formally assessed using appropriate criteria?	Y
	Were efforts made to minimise error in risk of bias assessment?	Y
	Concerns regarding methods used to collect data and appraise studies	LOW CONCERN
	DOMAIN 4: SYNTHESIS AND FINDINGS	
	Describe synthesis methods:	
	Did the synthesis include all studies that it should?	PY
	Were all pre-defined analyses reported or departures explained?	PY
	Was the synthesis appropriate given the nature and similarity in the research questions, study designs and outcomes across included studies?	Y
	Was between-study variation (heterogeneity) minimal or addressed in the synthesis?	PY
	Were the findings robust, e.g. as demonstrated through funnel plot or sensitivity analyses?	Y
	Were biases in primary studies minimal or addressed in the synthesis?	PY
	Concerns regarding the synthesis and findings	LOW CONCERN
	OVERALL RISK OF BIAS IN THE REVIEW	LOW RISK OF BIAS
Y = yes; PY = probably yes; PN = probably no; N = no; NI= no information		

Table 30: QUADAS-2 evaluation of all included prognostic accuracy studies

Paper	Domain 1: Patient selection		Domain 2: Index tests		Domain 3: Reference Standard		Domain 4: Flow and timing
		Applicability		Applicability		Applicability	
		Concerns about matching of patients to review question? (L/H/U)		Concerns about matching of index tests to review question? (L/H/U)		Concerns about matching of reference test to review question? (L/H/U)	
First author	ROB from selection (L/H/U)		ROB relating to index test (L/H/U)		ROB relating to reference test (L/H/U)		ROB from patient flow? (L/H/U)
Alarcon et al. 2013 ⁷⁵	L	H	H	L	L	L	U
Alarcon et al. 2016 ⁸⁰	L	H	U	L	L	L	U
Alarcon et al. 2024 ⁸¹	L	L	U	L	U	L	U
Blazquez-Gamero et al. 2019 ⁸²	L	H	U	L	U	L	U
Capretti et al. 2014 ⁸³	L	L	U	L	U	L	L
Castellanos et al. 2022 ⁸⁴	L	L	U	L	U	L	L
Chebib et al. 2022 ⁸⁵	L	L	H	L	U	L	L
Craeghs et al. 2021 ⁸⁶	L	L	U	L	U	L	L
De Juan et al. 2020 ⁸⁷	L	L	H	L	U	L	L
Dhondt et al. 2023 ¹⁵¹	L	L	U	L	U	L	H
Foulonnet al. 2019 ⁸⁸	L	L	U	L	U	L	U
Fourgeaud et al. 2024 ⁸⁹	L	L	U	L	U	L	U
Giannattasio et al. 2018 ⁹⁰	L	L	U	L	U	L	L
Kwak et al. 2018 ⁹¹	L	H	U	L	U	L	L

Paper	Domain 1: Patient selection		Domain 2: Index tests		Domain 3: Reference Standard		Domain 4: Flow and timing
		Applicability		Applicability		Applicability	
First author	ROB from selection (L/H/U)	Concerns about matching of patients to review question? (L/H/U)	ROB relating to index test (L/H/U)	Concerns about matching of index tests to review question? (L/H/U)	ROB relating to reference test (L/H/U)	Concerns about matching of reference test to review question? (L/H/U)	ROB from patient flow? (L/H/U)
Lucignani et al. 2021 ⁹²	L	L	H	L	L	L	L
Nishida et al. 2020 ⁹³	L	L	U	L	U	L	U
Ouellette et al. 2020 ¹⁶²	L	L	H	L	U	L	L
Vande Walle et al. 2024 ⁹⁴	L	L	H	L	U	L	U
Yamaguchi et al. 2022 ¹⁵⁴	L	L	H	L	U	L	U
L = low; H =High; U = unclear; ROB = risk of bias							

Appendix 4 – Relevant ongoing studies and unpublished studies

There were 8 ‘ongoing’ studies identified from clinical trial registers and PROSPERO (the international prospective register of systematic reviews), summarised in Table 31. Whilst most of these studies appear to have been completed (the study completion date has passed), no publications were identified relating to them.

Table 31: Summary of ongoing studies

Study details	Participant information	Intervention/ characteristic	Outcome (primary)
Ongoing systematic reviews			
Abrahao et al. 2022 ⁴³⁰ The congenital cytomegalovirus infection screening by urine: is it worthy? A systematic review PROSPERO: CRD42022326424 Anticipated completion date: 31/12/23	Newborns (without a known risk factor) within the first 2 weeks of life	Urine screening for cCMV	Hearing function (Brainstem Evoked Response Audiometry)
Wu et al. 2022 ⁴³¹ Meta-analysis of HCMV viral load level and the occurrence and outcome of hearing impairment PROSPERO: CRD42022301633 Anticipated completion date: 01/03/22	Children under the age of 18 who were diagnosed with cytomegalovirus infection	Viral load in different types of specimens such as saliva, blood, urine, and dried blood spots	Sensorineural hearing loss
Lopes et al. 2023 ⁴³² Impact of congenital cytomegalovirus infection on motor development in children: systematic review PROSPERO: CRD42023404868 Anticipated completion date: 01/11/24	Children with cCMV infection up to 5 years	Assessment of motor development (aspects of cCMV infection – diagnostic method, symptomatic or asymptomatic at birth and which symptoms, will be extracted)	Motor development: presence or absence of delay in motor development

Study details	Participant information	Intervention/ characteristic	Outcome (primary)
Shetty et al. 2024⁴³³ Evaluation of salivary sample in diagnosis, screening and management of cytomegalovirus (CMV) infection in newborns- systematic review and metanalysis PROSPERO: CRD42024603785 Anticipated completion date: 31/12/25	Neonates with cytomegalovirus	Salivary markers and screening test (including using pooled salivary samples) compared with gold standard tests on urine	Diagnostic accuracy (sensitivity, specificity, positive and negative predictive values)
Wang et al. 2023⁴³⁴ Accuracy and implementability of dried blood spot polymerase chain reaction assays for screening congenital cytomegalovirus infection: Systematic review and meta-analysis PROSPERO: CRD42023397836 Anticipated completion date: 30/06/24	Infants	Dried blood spot PCR assays collected among the first week of life compared with standard methods	Diagnostic accuracy (sensitivity, specificity, positive and negative likelihood ratios, positive and negative predictive values)
Ongoing primary studies			
Prognostic Value of Neonatal Markers for the Development of Neurosensorial Sequelae in Children Infected by Cytomegalovirus in Utero (CYMEPEDIA)⁴³⁵ Clinical trial identifier: NCT01923636 Study completion date: 16/05/22	Neonates less than 1 month with cCMV infection at birth objectified by the detection of CMV in a urine sample, in saliva or blood obtained in the first 10 days of life (n=254)	Prognostic score for the development of neurosensorineural sequelae, based on clinical, radiological and laboratory (virological and immunological) standardized reports	Prognostic value of neonatal markers (clinical, imaging and biological) for the development of neurosensorial sequelae at 1 year and 2 years of age
Markers for neurocognitive impairment in congenitally CMV infected infants: MARVELYS study⁴³⁶ Clinical trial identifier: NL-OMON43495 Study completion date: not reported (start date 17/05/16)	Neonates younger than 10 days of life with positive CMV PCR in urine or blood and/or positive CMV PCR in dried blood spot in children older than 10 days of age (n=100)	CSF and blood parameters	Neurocognitive performance, deviations in neuroimaging parameters, audiological and ophthalmologic measurements at the age of 1, 2, 5 and 8 years

Study details	Participant information	Intervention/ characteristic	Outcome (primary)
Targeted Screening for Congenital Cytomegalovirus-related Hearing loss in infants in Western Australia ⁴³⁷ Clinical trial identifier: ACTRN12621000484842 Study completion date: not reported (date of last data collection 07/12/22)	Infants born in Western Australia undergoing newborn hearing screening which results in a 'refer' (fail) result in 1 or both ears	Targeted testing of infants for congenital cytomegalovirus (cCMV). Infants are targeted using the newborn hearing screen result.	To assess whether targeted congenital cytomegalovirus testing is able to be completed by day 21 of life in Infants at high-risk of the virus across Western Australia.

Abbreviations: cCMV = congenital cytomegalovirus; CMV = cytomegalovirus; CSF = cerebrospinal fluid; PCR = polymerase chain reaction.

The electronic searches and supplementary searching identified 38 studies that have only been published as conference abstracts, summarised in Table 32. These studies have been cross-checked against the identified published studies, but do not appear to have been published in full.

Table 32: Summary of unpublished studies

Study details	Title
Criterion 4	
Brophy et al. 2024 ⁴³⁸	Congenital CMV AND HEARING in Ontario: Optimizing screening to improve child health outcomes (CAN HEAR ONTARIO)
Churyukina et al. 2023 ⁴³⁹	Predictive value of cytokines for preservation neurologic semiology in neonatal cytomegalovirus infection
Del Valle Penella et al. 2022 ⁴⁴⁰	Performance of dried blood spot (DBS) for the diagnosis of congenital cytomegalovirus (cCMV) in a single center: a real-world experience
Kakkar et al. 2024 ⁴⁴¹	Feasibility of dried urine on filter paper for universal congenital CMV screening
Kim et al. 2023 ⁴⁴²	CMV PCR agreement between blood and dried blood spots in infants with congenital cytomegalovirus infection
Kruc et al. 2021 ⁴⁴³	Does cytomegalovirus (CMV) viral load correlate with disease severity in the setting of congenital CMV (CCMV) infection? results from a universal CCMV screening study
Lazzarotto et al. 2013 ⁴⁴⁴	Pre-and post-natal diagnosis of congenital cytomegalovirus infection
Rohde et al. 2017 ⁴⁴⁵	Analytical and clinical performance of a real-time screening PCR assay identifying congenital CMV infection
Criterion 7	
Alarcon et al. 2022 ⁴⁴⁶	Isolated white matter abnormalities at birth as prognostic markers in congenital cytomegalovirus infection
Boscolo et al. 2024 ⁴⁴⁷	Correlation between neurodevelopmental outcomes and brain MRI severity score in patients with congenital cytomegalovirus infection: a monocentric retrospective study
Buonsenso et al. 2024 ⁴⁴⁸	Prognostic factors of late-onset hearing loss in infants with congenital cytomegalovirus and normal audiological assessment at birth
Castells Vilella et al. 2016 ⁴⁴⁹	Risk factors associated with hearing loss and neurologic impairment in the Spanish network of infants with congenital cytomegalovirus infection (REDICCMV)
Chung et al. 2024 ⁴⁵⁰	Predicting outcome in clinically inapparent congenital cytomegalovirus infection with hearing loss
Churyukina et al. 2021 ⁴⁵¹	Early predictors of preservation neurologic semiology in neonatal cytomegalovirus infection
Czech-Kowalska et al. 2022 ⁴⁵²	Predictors of sensorineural hearing loss in newborns with congenital CMV infection – a single-center experience
Eleni Loizou et al. 2024 ⁴⁵³	Vestibular dysfunction in asymptomatic children with congenital cytomegalovirus infection
Faure-Bardon et al. 2018 ⁴⁵⁴	Neonatal outcome and long-term sequelae in CMV infected neonates according to the timing of primary maternal infection
Fernandez-Rueda et al. 2024 ⁴⁵⁵	Risk factors associated with hearing loss in children with congenital cytomegalovirus infection
Fourgeaud et al. 2024 ⁴⁵⁶	Sequelae of congenital cytomegalovirus following maternal primary infections are limited to those acquired before 12 weeks

Study details	Title
Karagiannidou et al. 2024 ⁴⁵⁷	Delayed onset hearing loss (DOHL) in asymptomatic newborns with congenital cytomegalovirus infection (cCMV)
Kravchenko et al. 2014 ⁴⁵⁸	The outcome of cerebral pathology by the end of the first year of life in newborn babies with cytomegalovirus infection
Kravchenko et al. 2016 ⁴⁵⁹	Prognosis for the preservation of neurological symptoms by the end of the first year of life in Newborn babies, who had cytomegalovirus infection
Medoro et al. 2022 ⁴⁶⁰	Neonatal immune phenotypes and neurodevelopmental outcomes in infants with congenital cytomegalovirus infection (cCMV)
Morioka et al. 2017 ⁴⁶¹	Neurological outcomes at 18 months of age and neonatal risk factors for severe sequelae in congenitally cytomegalovirus-infected infants treated by antiviral treatment early in life
Nosan et al. 2022 ⁴⁶²	Genetic variations in toll-like receptor 2 gene in Slovenian newborns with congenital human cytomegalovirus infection
Salome et al. 2024 ⁴⁶³	Can viral load predict a symptomatic congenital CMV infection? A systematic review and meta-analysis
Sandoval Carmona et al. 2022 ⁴⁶⁴	Viral load kinetics as a prognostic indicator of outcomes among children with congenital cytomegalovirus infection in the Montreal cohort
Smiljkovic et al. 2016 ⁴⁶⁵	Use of quantitative polymerase chain reaction in blood and cerebrospinal fluid as a predictor of severity in congenital cytomegalovirus infection
Smithers-Sheedy et al. 2014 ⁴⁶⁶	Cerebral palsy attributed to congenital cytomegalovirus infection is associated with spastic quadriplegia and female sex
Vanwinkel et al. 2016 ⁴⁶⁷	Congenitally infected CMV fetuses following first trimester maternal infection: Neonatal and short-term outcome
Villaverde et al. 2024 ⁴⁶⁸	Association between congenital cytomegalovirus infection and autism spectrum disorder
Vossen et al. 2016 ⁴⁶⁹	Long-term impairment attributable to congenital cytomegalovirus infection
Criteria 12 and 13	
Carrillo-Marquez et al. 2017 ⁴⁷⁰	Programmatic congenital CMV universal screening programme
Landers et al. 2024 ⁴⁷¹	Analysis of dried blood spot and saliva PCR screening for detection of congenital cytomegalovirus infection informs implementation of a novel universal newborn screening program
Lund et al. 2024 ⁴⁷²	Targeted screening for congenital CMV in the capital region of Denmark
Penton et al. 2019 ⁴⁷³	Adherence to follow-up and CMV testing in infants who failed newborn hearing screens: Evaluation of new protocol to ensure follow-up and testing
Syridou et al. 2022 ⁴⁷⁴	Diagnostic ability of targeted and universal neonatal screening for cCMV-infection, to provide timely diagnosis of cCMV-infection associated neurodevelopmental delay
Webb et al. 2024 ⁴⁷⁵	Exploring parents' experiences of completing targeted congenital cytomegalovirus screening during their infant's newborn hearing screening

Appendix 5 – UK NSC reporting checklist for evidence summaries

All items on the UK NSC Reporting Checklist for Evidence Summaries have been addressed in this report. A summary of the checklist, along with the page or pages where each item can be found in this report, is presented in Table 33.

Table 33 UK NSC reporting checklist

Section	Item	Page no.
Title and summaries		
Title Sheet	Identify the review as a UK NSC Evidence summary	Title page
Plain English summary	Plain English description of the executive summary.	7
Executive summary	Structured overview of the whole report. To include: the purpose/aim of the review; background; previous recommendations; findings and gaps in the evidence; recommendations on the screening that can or cannot be made on the basis of the review	8
Introduction and Approach		
Background and objectives	Background – Current policy context and rationale for the current review – for example, reference to details of previous reviews, basis for current recommendation, recommendations made, gaps identified, drivers for new reviews	14
	Objectives – What are the questions the current evidence summary intends to answer? – statement of the key questions for the current evidence summary, criteria they address, and number of studies included per question, description of the overall results of the literature search.	16
	Method – briefly outline the rapid review methods used.	18
		15
		17

Section	Item	Page no.
Eligibility for inclusion in the review	State all criteria for inclusion and exclusion of studies to the review clearly(PICO, dates, language, study type, publication type, publication status etc.) To be decided a priori	18
Appraisal for quality/ risk of bias tool	Details of tool/ checklist used to assess quality, e.g. QUADAS 2, CASP, SIGN, AMSTAR.	22
Search strategy and study selection		
Databases/ sources searched	Give details of all databases searched (including platform/ interface and coverage dates) and date of final search.	17
Search strategy and results	Present the full search strategy for at least one database (usually a version of Medline), including limits and search filters if used.	Appendix 1
	Provide details of the total number of (results from each database searched), number of duplicates removed, and the final number of unique records to consider for inclusion.	Appendix 2, Figure 4
Study selection	State the process for selecting studies – inclusion and exclusion criteria, number of studies screened by title/abstract and full text, number of reviewers, any cross checking carried out.	18, 24 and Appendix 2
Study level reporting of results (for each key question)		
Study level reporting, results and risk of bias assessment	For each study, produce a table that includes the full citation and a summary of the data relevant to the question (for example, study size, PICO, follow-up period, outcomes reported, statistical analyses etc.).	30-46 (Criterion 4): Tables 3-9
	Provide a simple summary of key measures, effect estimates and confidence intervals for each study where available. For each study, present the results of any assessment of quality/risk of bias.	Criterion 7: Tables 10-16 & Appendix 3 Criteria 11, 12, 13): Tables 17-21
Additional analyses	Describe additional analyses (for example, sensitivity, specificity, PPV, etc.) carried out by the reviewer. [Remove if not performed]	23

Question level synthesis

Section	Item	Page no.
Description of the evidence	For each question, give numbers of studies screened, assessed for eligibility, and inclusion in the review, with summary reasons for exclusion	Criterion 4: 25 onwards Criterion 7: 46 onwards Criteria 11, 12, 13: 75 onwards See also Appendix 2
Combining and presenting the findings	Provide a balanced discussion of the body of evidence which avoids over reliance on one study or set of studies. Consideration of four compartments should inform the reviewer's judgement on whether the criterion is "met", "not met" or "uncertain": quantity; quality; applicability and consistency.	onwards25 to 94
Summary of findings	Provide a description of the evidence reviewed and included for each question, with reference to their eligibility for inclusion. Summarise the main findings including the quality/ risk of bias issues for each question. Have the criteria addressed been "met", "not met" or "uncertain"?	44 (Criterion 4) 71 (Criterion 7) 91 (Criteria 11 & 13) 94 (Criterion 12)
Review Summary		
Conclusions and implications for policy	Do findings indicate whether screening should be recommended? IS further work warranted? Are there gaps in the evidence highlighted by the review?	10911 & 95
Limitations	Discuss limitations of the available evidence and of the review methodology if relevant.	11012 & 96

References

1. Advisory Committee on the Safety of Blood Tissues and Organs. *SaBTO report of the Cytomegalovirus Steering Group*. Department of Health and Social Care; 2012. Available from: <https://www.gov.uk/government/publications/sabto-report-of-the-cytomegalovirus-steering-group> [accessed 26 September 2024].
2. Revello MG, Gerna G. Diagnosis and management of human cytomegalovirus infection in the mother, fetus, and newborn infant. *Clin Microbiol Rev* 2002;**15**:680–715.
3. Ludwig A, Hengel H. Epidemiological impact and disease burden of congenital cytomegalovirus infection in Europe. *Eurosurveillance* 2009;**14**:26–32.
4. Office for National Statistics. *Births in England and Wales: 2023*. ONS; 2024. Available from: <https://www.ons.gov.uk/peoplepopulationandcommunity/birthsdeathsandmarriages/livebirths/bulletins/birthsummarytablesenglandandwales/2023> [accessed 21 May 2025].
5. Payne H, Aaltoranta M, Veikkolainen V, Kent N, Gkouleli T, Lennon A, et al. A high-sensitivity, high-throughput newborn screening assay for congenital cytomegalovirus-is it time for universal screening in the United Kingdom? *Front Pediatr* 2025;**13**:1543132.
6. Bartlett AW, McMullan B, Rawlinson WD, Palasanthiran P. Hearing and neurodevelopmental outcomes for children with asymptomatic congenital cytomegalovirus infection: a systematic review. *Rev Med Virol* 2017;**27**:e1938.
7. Costello Medical. *Newborn screening for cytomegalovirus. An evidence map to outline the volume and type of evidence related to screening for cytomegalovirus for the UK National Screening Committee*. London: UK National Screening Committee; 2022.
8. Leruez-Ville M, Chatzakis C, Lilleri D, Blazquez-Gamero D, Alarcon A, Bourgon N, et al. Consensus recommendation for prenatal, neonatal and postnatal management of congenital cytomegalovirus infection from the European congenital infection initiative (ECCI). *Lancet Reg Health Eur* 2024;**40**:100892.
9. Jones CE, Bailey H, Bamford A, Calvert A, Dorey RB, Drysdale SB, et al. Managing challenges in congenital CMV: current thinking. *Arch Dis Child* 2023;**108**:601–7.
10. Rodríguez-Muñoz MF, Martín-Martín C, Kovacheva K, Olivares ME, Izquierdo N, Pérez-Romero P, et al. Hygiene-based measures for the prevention of cytomegalovirus infection in pregnant women: a systematic review. *BMC Pregnancy Childbirth* 2024;**24**:172.
11. Revello MG, Tibaldi C, Masuelli G, Frisina V, Sacchi A, Furione M, et al. Prevention of primary cytomegalovirus infection in pregnancy. *EBioMedicine* 2015;**2**:1205–10.
12. Pesch MH, Kuboushek K, McKee MM, Thorne MC, Weinberg JB. Congenital cytomegalovirus infection. *BMJ* 2021;**373**:n1212.
13. Luck SE, Wieringa JW, Blazquez-Gamero D, Henneke P, Schuster K, Butler K, et al. Congenital cytomegalovirus: a European expert consensus statement on diagnosis and management. *Pediatr Infect Dis J* 2017;**36**:1205–13.
14. Schleiss MR. Congenital cytomegalovirus screening moves ahead. *JAMA Pediatr* 2025;**179**:241–3.
15. Canadian Agency for Drugs and Technologies in Health. *Newborn screening for congenital cytomegalovirus in Canada: environmental scan*. Canadian Agency for Drugs and Technologies in Health; 2024. Available from: <https://www.ncbi.nlm.nih.gov/books/NBK604835/> [accessed 21 May 2025].
16. Tanaka M, Saito H, Sakai K, Konomura K, Suzuki M, Tajima G, et al. EE125 Cost-effectiveness of targeted newborn screening for congenital cytomegalovirus infection in Japan. *Value Health* 2024;**27**:S78.
17. UK National Screening Committee. *Screening for cytomegalovirus*. London: UK National Screening Committee; 2022.
18. Bazian Ltd. *Newborn screening for cytomegalovirus. External review against programme appraisal criteria for the UK National Screening Committee (UK NSC)*. London: UK National Screening Committee; 2017.

19. Centre for Reviews and Dissemination. *Systematic reviews. CRD's guidance for undertaking reviews in health care*. York: CRD, University of York; 2009.
20. Page MJ, McKenzie JE, Bossuyt PM, Boutron I, Hoffmann TC, Mulrow CD, et al. The PRISMA 2020 statement: an updated guideline for reporting systematic reviews. *BMJ* 2021;**372**:n71.
21. McGowan J, Sampson M, Salzwedel DM, Cogo E, Foerster V, Lefebvre C. PRESS Peer Review of Electronic Search Strategies: 2015 Guideline Statement. *J Clin Epidemiol* 2016;**75**:40–6.
22. Thomas J, Graziosi S, Brunton J, Ghouze Z, O'Driscoll P, Bond M, et al. *EPPI-Reviewer: advanced software for systematic reviews, maps and evidence synthesis*. London: EPPI Centre, UCL Social Research Institute, University College London; 2022.
23. Whiting PF, Rutjes AW, Westwood ME, Mallett S, Deeks JJ, Reitsma JB, et al. QUADAS-2: a revised tool for the quality assessment of diagnostic accuracy studies. *Ann Intern Med* 2011;**155**:529–36.
24. Reitsma JB, Glas AS, Rutjes AW, Scholten RJ, Bossuyt PM, Zwinderman AH. Bivariate analysis of sensitivity and specificity produces informative summary measures in diagnostic reviews. *J Clin Epidemiol* 2005;**58**:982–90.
25. DerSimonian R, Laird N. Meta-analysis in clinical trials. *Control Clin Trials* 1986;**7**:177–88.
26. Higgins JP, Thompson SG, Deeks JJ, Altman DG. Measuring inconsistency in meta-analyses. *BMJ* 2003;**327**:557–60.
27. Wang L, Xu X, Zhang H, Qian J, Zhu J. Dried blood spots PCR assays to screen congenital cytomegalovirus infection: a meta-analysis. *Viral J* 2015;**12**:60.
28. Boppana SB, Ross SA, Shimamura M, Palmer AL, Ahmed A, Michaels MG, et al. Saliva polymerase-chain-reaction assay for cytomegalovirus screening in newborns. *N Engl J Med* 2011;**364**:2111–8.
29. Waters A, Jennings K, Fitzpatrick E, Coughlan S, Molloy EJ, De Gascun CF, et al. Incidence of congenital cytomegalovirus infection in Ireland: implications for screening and diagnosis. *J Clin Virol* 2014;**59**:156–60.
30. Freeman SC, Kerby CR, Patel A, Cooper NJ, Quinn T, Sutton AJ. Development of an interactive web-based tool to conduct and interrogate meta-analysis of diagnostic test accuracy studies: MetaDTA. *BMC Med Res Methodol* 2019;**19**:81.
31. Patel A, Cooper N, Freeman S, Sutton A. Graphical enhancements to summary receiver operating characteristic plots to facilitate the analysis and reporting of meta-analysis of diagnostic test accuracy data. *Res Synth Methods* 2021;**12**:34–44.
32. Zheng B, Wu FF, Li XX, Shen R, Zheng Z, Liu HY. Diagnostic test accuracy of PCR by saliva specimen for cytomegalovirus infection in newborn: a protocol for systematic review and meta-analysis. *Medicine (Baltimore)* 2022;**101**:e31776.
33. Alkhawaja S, Ismaeel A, Botta G, Senok AC. The prevalence of congenital and perinatal cytomegalovirus infections among newborns of seropositive mothers. *J Infect Dev Ctries* 2012;**6**:410–5.
34. Yamamoto AY, Mussi-Pinhata MM, Marin LJ, Brito RM, Oliveira PF, Coelho TB. Is saliva as reliable as urine for detection of cytomegalovirus DNA for neonatal screening of congenital CMV infection? *J Clin Virol* 2006;**36**:228–30.
35. Eventov-Friedman S, Manor H, Bar-Oz B, Averbuch D, Caplan O, Lifshitz A, et al. Saliva real-time polymerase chain reaction for targeted screening of congenital cytomegalovirus infection. *J Infect Dis* 2019;**220**:1790–6.
36. Barkai G, Ari-Even Roth D, Barzilai A, Tepperberg-Oikawa M, Mendelson E, Hildesheimer M, et al. Universal neonatal cytomegalovirus screening using saliva - report of clinical experience. *J Clin Virol* 2014;**60**:361–6.
37. Gunkel J, Wolfs TF, Nijman J, Schuurman R, Verboon-Macielek MA, de Vries LS, et al. Urine is superior to saliva when screening for postnatal CMV infections in preterm infants. *J Clin Virol* 2014;**61**:61–4.
38. Cardoso ES, Jesus BL, Gomes LG, Sousa SM, Gadelha SR, Marin LJ. The use of saliva as a practical and feasible alternative to urine in large-scale screening for congenital cytomegalovirus infection increases inclusion and detection rates. *Rev Soc Bras Med Trop* 2015;**48**:206–7.

39. Dollard SC, Dreon M, Hernandez-Alvarado N, Amin MM, Wong P, Lanzieri TM, et al. Sensitivity of dried blood spot testing for detection of congenital cytomegalovirus infection. *JAMA Pediatr* 2021;**175**:e205441.
40. Atwood EN, Esper FP, Bailey K, Doud MK, Benton AM, Friesen J, et al. False positive congenital cytomegalovirus saliva screening results with an FDA-approved assay at two institutions. *J Clin Virol* 2023;**166**:105527.
41. Dunn JJ, Selvarangan R, Maggert K, Young S, Leber AL. Multicenter evaluation of the DiaSorin Molecular Simplexa Congenital CMV Direct PCR Test on neonatal saliva and urine specimens. *J Clin Microbiol* 2023;**61**:e0028323.
42. Leruez-Ville M, Magny JF, Couderc S, Pichon C, Parodi M, Bussieres L, et al. Risk factors for congenital cytomegalovirus infection following primary and nonprimary maternal infection: a prospective neonatal screening study using polymerase chain reaction in saliva. *Clin Infect Dis* 2017;**65**:398–404.
43. Pinninti SG, Ross SA, Shimamura M, Novak Z, Palmer AL, Ahmed A, et al. Comparison of saliva PCR assay versus rapid culture for detection of congenital cytomegalovirus infection. *Pediatr Infect Dis J* 2015;**34**:536–7.
44. Ross SA, Michaels MG, Ahmed A, Palmer AL, Sanchez PJ, Bernstein DI, et al. Contribution of breastfeeding to false-positive saliva polymerase chain reaction for newborn congenital cytomegalovirus screening. *J Infect Dis* 2018;**217**:1612–5.
45. Tan NK, Pope CF, Carrington D. Performance evaluation of fully automated cobas 6800 CMV PCR for the detection and quantification of cytomegalovirus DNA in neonatal urine and saliva, and adult urine, saliva, and vaginal secretion. *J Med Virol* 2023;**95**:e29223.
46. Wang S, Wang T, Zhang W, Liu X, Wang X, Wang H, et al. Cohort study on maternal cytomegalovirus seroprevalence and prevalence and clinical manifestations of congenital infection in China. *Medicine (Baltimore)* 2017;**96**:e6007.
47. Wunderlich W, Sidebottom AC, Schulte AK, Taghon J, Dollard S, Hernandez-Alvarado N. The use of saliva samples to test for congenital cytomegalovirus infection in newborns: examination of false-positive samples associated with donor milk use. *Int J Neonatal Screen* 2023;**9**:46.
48. Gunlemez A, Kolayli F, Yazici Ozcelik E, Duranoglu A, Durgut M, Arisoy ES, et al. Congenital cytomegalovirus infection screening in newborns from saliva samples by real-time polymerase chain reaction analysis. *Turk Arch Pediatr* 2023;**58**:371–5.
49. Shlonsky Y, Smair NS, Mubarik R, Bamberger E, Hemo M, Cohen S, et al. Pooled saliva CMV DNA detection: a viable laboratory technique for universal CMV screening of healthy newborns. *J Clin Virol* 2021;**138**:104798.
50. Exler S, Daiminger A, Grothe M, Schalasta G, Enders G, Enders M. Primary cytomegalovirus (CMV) infection in pregnancy: diagnostic value of CMV PCR in saliva compared to urine at birth. *J Clin Virol* 2019;**117**:33–6.
51. Gantt S, Goldfarb DM, Park A, Rawlinson W, Boppana SB, Lazzarotto T, et al. Performance of the Alethia CMV assay for detection of cytomegalovirus by use of neonatal saliva swabs. *J Clin Microbiol* 2020;**58**:e01951–19.
52. Huang Y, Wang H, Li T, Li C, Tang J, Yu H, et al. Comparison of detection strategies for screening and confirming congenital cytomegalovirus infection in newborns in a highly seroprevalent population: a mother-child cohort study. *Lancet Reg Health West Pac* 2021;**12**:100182.
53. Izquierdo G, Guerra C, Reyes R, Araya L, Sepulveda B, Cabrera C, et al. Universal and expanded screening strategy for congenital cytomegalovirus infection: is pool testing by a rapid molecular test in saliva a new choice in developing countries? *Viruses* 2024;**16**:772.
54. Letamendia-Richard E, Perillaud-Dubois C, de La Guillonniere L, Thouard I, Cordier AG, Roque-Afonso AM, et al. Universal newborn screening for congenital cytomegalovirus infection: feasibility and relevance in a French type-III maternity cohort. *BJOG* 2022;**129**:291–9.
55. Pasternak Y, Oikawa MT, Mendelson E, Osovsky M, Klinger G, Bilavsky E. Diagnosing congenital cytomegalovirus by saliva on Guthrie paper. *J Clin Virol* 2020;**126**:104337.
56. Portet Sulla V, Kabore B, Rafek R, Chaghouri A, Decombe G, Lubrano Di Ciccone J, et al. Feasibility, performances and predictive value of congenital CMV neonatal screening on saliva swabs. *J Clin Virol* 2024;**174**:105713.

57. Ross SA, Ahmed A, Palmer AL, Michaels MG, Sanchez PJ, Bernstein DI, et al. Detection of congenital cytomegalovirus infection by real-time polymerase chain reaction analysis of saliva or urine specimens. *J Infect Dis* 2014;**210**:1415–8.
58. Stark A, Greenberg RG, Pittman R, Weimer KED. Concordance of cytomegalovirus saliva and urine testing in infants for the detection of congenital infection. *Pediatr Infect Dis J* 2024;**43**:1191–93.
59. Ross S, Long SS, Kimberlin DW. Closer to universal newborn screening for congenital cytomegalovirus infection but far away from antiviral therapy in all infected infants. *J Pediatr* 2018;**199**:7–9.
60. Atkinson C, Emery VC, Griffiths PD. Development of a novel single tube nested PCR for enhanced detection of cytomegalovirus DNA from dried blood spots. *J Virol Methods* 2014;**196**:40–4.
61. Del Valle Penella A, Miller J, Rochat R, Demmler-Harrison G. Utility of dried blood spots for the diagnosis of congenital cytomegaloviruses within the first 21 days of life in a single center. *Int J Neonatal Screen* 2023;**9**:44.
62. Fourgeaud J, Magny JF, Couderc S, Garcia P, Maillotte AM, Benard M, et al. Clinical value of serial quantitative analysis of cytomegalovirus DNA in blood and saliva over the first 24 months of life in congenital infection: the French Cymepedia Cohort. *J Pediatr* 2023;**253**:197–204.
63. Moteiki H, Isaka Y, Inaba Y, Motobayashi M, Nishio SY, Ohira S, et al. A rational approach to identifying newborns with hearing loss caused by congenital cytomegalovirus infection by dried blood spot screening. *Acta Otolaryngol* 2018;**138**:708–12.
64. Noorbakhsh S, Farhadi M, Rezaei F, Jomeh HE, Farahmand M, Haghighi F, et al. Prevalence of congenital cytomegalovirus infection among hospitalized neonates in Tehran, Iran. *Iran Red Crescent Med J* 2020;**22**:e66.
65. Ross SA, Ahmed A, Palmer AL, Michaels MG, Sanchez PJ, Stewart A, et al. Newborn dried blood spot PCR to identify infants with congenital cytomegalovirus-associated sensorineural hearing loss. *J Pediatr* 2017;**184**:57–61.
66. Vercauteren KOA, Keymeulen A, Mahieu L, Cossey V, Casaer A, Van Mol C, et al. Prospective multicenter comparison of urine culture with PCR on dried blood spots using 2 different extraction and PCR methods in neonates suspected for congenital cytomegalovirus infection. *Diagn Microbiol Infect Dis* 2020;**97**:115051.
67. Vives-Onos I, Codina-Grau MG, Noguera-Julian A, Blazquez-Gamero D, Fortuny C, Baquero-Artigao F, et al. Is polymerase chain reaction in neonatal dried blood spots reliable for the diagnosis of congenital cytomegalovirus infection? *Pediatr Infect Dis J* 2019;**38**:520–4.
68. Albanna EA, El-Latif RS, Sharaf HA, Gohar MK, Ibrahim BM. Diagnosis of congenital cytomegalovirus infection in high risk neonates. *Mediterr J Hematol Infect Dis* 2013;**5**:e2013049.
69. Coltella L, Ranno S, Pizzichemi G, Piccioni L, Chiavelli S, Onetti Muda A, et al. Real-time PCR versus shell vial culture on urine of patients with suspected congenital cytomegalovirus infection. *Future Virol* 2019;**14**:585–91.
70. De Monte A, Cannavo I, Caramella A, Ollier L, Giordanengo V. [Analytical performances of real-time PCR by Abbott RealTime CMV with m2000 for the detection of cytomegalovirus in urine]. *Ann Biol Clin (Paris)* 2016;**74**:244–50.
71. Ligozzi M, Poggi M, Saletti M, Gibellini D. Development of a real-time quantitative polymerase chain reaction assay for the detection of congenital human cytomegalovirus infection in urine samples. *Mol Cell Probes* 2016;**30**:50–2.
72. Wang XT, Dong ZB, Luo LM, Deng M. [Application of throat swab nested PCR in the diagnosis of congenital human cytomegalovirus infection in neonates]. *Zhongguo Dang Dai Er Ke Za Zhi* 2013;**15**:1086–8.
73. Barkai G, Barzilai A, Mendelson E, Tepperberg-Oikawa M, Roth DA, Kuint J. Newborn screening for congenital cytomegalovirus using real-time polymerase chain reaction in umbilical cord blood. *Isr Med Assoc J* 2013;**15**:347–51.
74. Reyes A, Taravillo I, Moral N, Moraleda C, Blazquez-Gamero D, Folgueira L. Feasible alternatives to DBS in the retrospective diagnosis of congenital cytomegalovirus infection. *J Clin Virol* 2020;**129**:104504.

75. Alarcon A, Martinez-Biarge M, Cabanas F, Hernanz A, Quero J, Garcia-Alix A. Clinical, biochemical, and neuroimaging findings predict long-term neurodevelopmental outcome in symptomatic congenital cytomegalovirus infection. *J Pediatr* 2013;**163**:828–34.
76. Bilavsky E, Schwarz M, Pardo J, Attias J, Levy I, Haimi-Cohen Y, et al. Lenticulostriated vasculopathy is a high-risk marker for hearing loss in congenital cytomegalovirus infections. *Acta Paediatr* 2015;**104**:e388–94.
77. Forner G, Abate D, Mengoli C, Palu G, Gussetti N. High cytomegalovirus (CMV) DNAemia predicts CMV sequelae in asymptomatic congenitally infected newborns born to women with primary infection during pregnancy. *J Infect Dis* 2015;**212**:67–71.
78. Goderis J, Keymeulen A, Smets K, Van Hoecke H, De Leenheer E, Boudewyns A, et al. Hearing in children with congenital cytomegalovirus infection: results of a longitudinal study. *J Pediatr* 2016;**172**:110–5.
79. Vande Walle C, Maris F, Schiettecatte E, Herregods N. The value of magnetic resonance imaging in congenital cytomegalovirus infection: a systematic review. *Pediatr Radiol* 2024;**54**:2157–74.
80. Alarcon A, Martinez-Biarge M, Cabanas F, Quero J, Garcia-Alix A. A prognostic neonatal neuroimaging scale for symptomatic congenital cytomegalovirus infection. *Neonatology* 2016;**110**:277–85.
81. Alarcon A, de Vries LS, Parodi A, Arnaez J, Cabanas F, Steggerda SJ, et al. Neuroimaging in infants with congenital cytomegalovirus infection and its correlation with outcome: emphasis on white matter abnormalities. *Arch Dis Child Fetal Neonatal Ed* 2024;**109**:F151–8.
82. Blazquez-Gamero D, Soriano-Ramos M, Martinez de Aragon A, Baquero-Artigao F, Frick MA, Noguera-Julian A, et al. Role of magnetic resonance imaging and cranial ultrasonography in congenital cytomegalovirus infection. *Pediatr Infect Dis J* 2019;**38**:1131–7.
83. Capretti MG, Lanari M, Tani G, Ancora G, Sciutti R, Marsico C, et al. Role of cerebral ultrasound and magnetic resonance imaging in newborns with congenital cytomegalovirus infection. *Brain Dev* 2014;**36**:203–11.
84. Castellanos ME, de la Mata Navazo S, Bermejo MC, Morín MG, Martín YR, Lozano JS, et al. Association between neuroimaging findings and neurological sequelae in patients with congenital cytomegalovirus infection. *Neurologia* 2022;**37**:122–9.
85. Chebib E, Maudoux A, Benoit C, Bernard S, Belarbi N, Parodi M, et al. Predictors of cochleovestibular dysfunction in children with congenital cytomegalovirus infection. *Eur J Pediatr* 2022;**181**:2909–18.
86. Craeghs L, Goderis J, Acke F, Keymeulen A, Smets K, Van Hoecke H, et al. Congenital CMV-associated hearing loss: can brain imaging predict hearing outcome? *Ear Hear* 2021;**42**:373–80.
87. de Juan Gallach A, Alemany Albert M, Marco Hernandez AV, Boronat Gonzalez N, Cernada Badia M, Tomas Vila M. Neurological sequelae in patients with congenital cytomegalovirus. *An Pediatr (Engl Ed)* 2020;**93**:111–7.
88. Foulon I, De Brucker Y, Buyl R, Lichtert E, Verbruggen K, Pierard D, et al. Hearing loss with congenital cytomegalovirus infection. *Pediatrics* 2019;**144**:e20183095.
89. Fourgeaud J, Magny JF, Couderc S, Garcia P, Maillotte AM, Benard M, et al. Predictors of the outcome at 2 years in neonates with congenital cytomegalovirus infection. *Pediatrics* 2024;**153**:e2023063531.
90. Giannattasio A, Bruzzese D, Di Costanzo P, Capone E, Romano A, D'Amico A, et al. Neuroimaging profiles and neurodevelopmental outcome in infants with congenital cytomegalovirus infection. *Pediatr Infect Dis J* 2018;**37**:1028–33.
91. Kwak M, Yum MS, Yeh HR, Kim HJ, Ko TS. Brain magnetic resonance imaging findings of congenital cytomegalovirus infection as a prognostic factor for neurological outcome. *Pediatr Neurol* 2018;**83**:14–8.
92. Lucignani G, Rossi Espagnet MC, Napolitano A, Figa Talamanca L, Calo Carducci FI, Auriti C, et al. A new MRI severity score to predict long-term adverse neurologic outcomes in children with congenital cytomegalovirus infection. *J Matern Fetal Neonatal Med* 2021;**34**:859–66.

93. Nishida K, Fujioka K, Sugioka Y, Abe S, Ashina M, Fukushima S, et al. Prediction of neurodevelopmental impairment in congenital cytomegalovirus infection by early postnatal magnetic resonance imaging. *Neonatology* 2020;**117**:460–6.
94. Vande Walle C, Keymeulen A, Oostra A, Schiettecatte E, Dhooge I, Smets K, et al. Apparent diffusion coefficient values of the white matter in magnetic resonance imaging of the neonatal brain may help predict outcome in congenital cytomegalovirus infection. *Pediatr Radiol* 2024;**54**:337–46.
95. Giannattasio A, Di Costanzo P, Milite P, De Martino D, Capone E, Romano A, et al. Is lenticulostriated vasculopathy an unfavorable prognostic finding in infants with congenital cytomegalovirus infection? *J Clin Virol* 2017;**91**:31–5.
96. Vila-Bedmar S, de Aragon Calvo AM, Liebana-Rojas C, Pedrero-Tomas R, Camacho-Salas A, Nuñez-Enamorado N, et al. Brain abnormalities, neurodevelopmental milestones, and long-term follow-up in newborns with congenital cytomegalovirus identified through a neonatal screening program. *Pediatr Infect Dis J* 2024;**43**:e434–40.
97. Mardegan C, Van HC, Hanquinet S, Tolsa CB, Pittet LF, Posfay-Barbe KM. Outcome of children with congenital cytomegalovirus infection: a retrospective observational study. *J Pediatr Infect Dis* 2024;**19**:317–26.
98. Oosterom N, Nijman J, Gunkel J, Wolfs TF, Groenendaal F, Verboon-Macielek MA, et al. Neuro-imaging findings in infants with congenital cytomegalovirus infection: relation to trimester of infection. *Neonatology* 2015;**107**:289–96.
99. Soriano-Ramos M, Pedrero-Tome R, Gimenez-Quiles E, Albert E, Baquero-Artigao F, Rodriguez-Molino P, et al. T-cell immune responses in newborns and long-term sequelae in congenital cytomegalovirus infection (CYTRIC study). *J Pediatr* 2024;**272**:114084.
100. Vande Walle C, Keymeulen A, Oostra A, Schiettecatte E, Dhooge IJ, Smets K, et al. Implications of isolated white matter abnormalities on neonatal MRI in congenital CMV infection: a prospective single-centre study. *BMJ Paediatr Open* 2023;**7**:e002097.
101. Poletti de Chaurand V, Scandella G, Zicoia M, Arienti F, Fernicola F, Lanteri L, et al. Useful clinical criteria for identifying neonates with congenital cytomegalovirus infection at birth in the context of an expanded targeted screening program. *Viruses* 2024;**16**:1138.
102. Chatzakis C, Ville Y, Makrydimas G, Dinas K, Zavlanos A, Sotiriadis A. Timing of primary maternal cytomegalovirus infection and rates of vertical transmission and fetal consequences. *Am J Obstet Gynecol* 2020;**223**:870–83.
103. Buca D, Di Mascio D, Rizzo G, Giancotti A, D'Amico A, Leombroni M, et al. Outcome of fetuses with congenital cytomegalovirus infection and normal ultrasound at diagnosis: systematic review and meta-analysis. *Ultrasound Obstet Gynecol* 2021;**57**:551–9.
104. Amir J, Atias J, Linder N, Pardo J. Follow-up of infants with congenital cytomegalovirus and normal fetal imaging. *Arch Dis Child Fetal Neonatal Ed* 2016;**101**:F428–32.
105. Zavattoni M, Lombardi G, Rognoni V, Furione M, Klersy C, Stronati M, et al. Maternal, fetal, and neonatal parameters for prognosis and counseling of HCMV congenital infection. *J Med Virol* 2014;**86**:2163–70.
106. Elkan Miller T, Weisz B, Yinon Y, Weissbach T, De Castro H, Avnet H, et al. Congenital cytomegalovirus infection following second and third trimester maternal infection is associated with mild childhood adverse outcome not predicted by prenatal imaging. *J Pediatric Infect Dis Soc* 2021;**10**:562–8.
107. Faure-Bardon V, Magny JF, Parodi M, Couderc S, Garcia P, Maillotte AM, et al. Sequelae of congenital cytomegalovirus following maternal primary infections are limited to those acquired in the first trimester of pregnancy. *Clin Infect Dis* 2019;**69**:1526–32.
108. Lipitz S, Yinon Y, Malinger G, Yagel S, Levit L, Hoffman C, et al. Risk of cytomegalovirus-associated sequelae in relation to time of infection and findings on prenatal imaging. *Ultrasound Obstet Gynecol* 2013;**41**:508–14.
109. Massoud M, Chollet M, Cabet S, Butin M, Mekki Y, Lina-Granade G, et al. Predicting outcome of congenital cytomegalovirus infection by differentiating and revisiting severe versus mild prenatal imaging features. *Fetal Diagn Ther* 2023;**50**:143–57.
110. Nigro G, Adler SP. High-dose cytomegalovirus (CMV) hyperimmune globulin and maternal CMV DNAemia independently predict infant outcome in pregnant women with a primary CMV infection. *Clin Infect Dis* 2020;**71**:1491–8.

111. Puhakka L, Renko M, Helminen M, Peltola V, Heiskanen-Kosma T, Lappalainen M, et al. Primary versus non-primary maternal cytomegalovirus infection as a cause of symptomatic congenital infection - register-based study from Finland. *Infect Dis (Lond)* 2017;**49**:445–53.
112. Coscia A, Leone A, Rubino C, Galitska G, Biolatti M, Bertino E, et al. Risk of symptomatic infection after non-primary congenital cytomegalovirus infection. *Microorganisms* 2020;**8**:786.
113. Scaramuzzino F, Di Pastena M, Chiurciu S, Romani L, De Luca M, Lucignani G, et al. Secondary cytomegalovirus infections: how much do we still not know? Comparison of children with symptomatic congenital cytomegalovirus born to mothers with primary and secondary infection. *Front Pediatr* 2022;**10**:885926.
114. Turriziani Colonna A, Buonsenso D, Pata D, Salerno G, Chieffo DPR, Romeo DM, et al. Long-term clinical, audiological, visual, neurocognitive and behavioral outcome in children with symptomatic and asymptomatic congenital cytomegalovirus infection treated with valganciclovir. *Front Med (Lausanne)* 2020;**7**:268.
115. De Cuyper E, Acke F, Keymeulen A, De Leenheer E, Van Hoecke H, Padalko E, et al. Risk factors for natural hearing evolution in newborns with congenital cytomegalovirus infection. *JAMA Otolaryngol Head Neck Surg* 2024;**150**:30–8.
116. Keymeulen A, De Leenheer E, Casaer A, Cossey V, Laroche S, Mahieu L, et al. Neurodevelopmental outcome in children with congenital cytomegalovirus infection: a prospective multicenter cohort study. *Early Hum Dev* 2023;**182**:105777.
117. Demmler-Harrison GJ, Miller JA. Maternal cytomegalovirus immune status and hearing loss outcomes in congenital cytomegalovirus-infected offspring. *PLoS One* 2020;**15**:e0240172.
118. Puhakka L, Lappalainen M, Lonnqvist T, Nieminen T, Boppana S, Saxen H, et al. Hearing outcome in congenitally CMV infected children in Finland - results from follow-up after three years age. *Int J Pediatr Otorhinolaryngol* 2022;**156**:111099.
119. Maltezou PG, Kourlaba G, Kourkouni E, Luck S, Blazquez-Gamero D, Ville Y, et al. Maternal type of CMV infection and sequelae in infants with congenital CMV: systematic review and meta-analysis. *J Clin Virol* 2020;**129**:104518.
120. Noorbakhsh S, Joghataei MT, Farhadi M, Haghighi F, Emamjomeh H, Haghighi Hasanabad M. Assessment of hearing loss in two-year follow-up study of neonates with congenital cytomegalovirus infection. *Iran J Child Neurol* 2022;**16**:17–26.
121. Capretti MG, Marsico C, Guidelli Guidi S, Ciardella A, Simonazzi G, Galletti S, et al. Neonatal and long-term ophthalmological findings in infants with symptomatic and asymptomatic congenital cytomegalovirus infection. *J Clin Virol* 2017;**97**:59–63.
122. Townsend CL, Forsgren M, Ahlfors K, Ivarsson SA, Tookey PA, Peckham CS. Long-term outcomes of congenital cytomegalovirus infection in Sweden and the United Kingdom. *Clin Infect Dis* 2013;**56**:1232–9.
123. Giannattasio A, Di Costanzo P, De Matteis A, Milite P, De Martino D, Bucci L, et al. Outcomes of congenital cytomegalovirus disease following maternal primary and non-primary infection. *J Clin Virol* 2017;**96**:32–6.
124. Rovito R, Korndewal MJ, Schielen PCJ, Kroes ACM, Vossen ACTM. Neonatal screening parameters in infants with congenital cytomegalovirus infection. *Clin Chim Acta* 2017;**473**:191–7.
125. Smyrli A, Raveendran V, Walter S, Pagarkar W, Field N, Kadambari S, et al. What are the neurodevelopmental outcomes of children with asymptomatic congenital cytomegalovirus infection at birth? A systematic literature review. *Rev Med Virol* 2024;**34**:e2555.
126. Vos B, Noll D, Whittingham J, Pigeon M, Bagatto M, Fitzpatrick EM. Cytomegalovirus - a risk factor for childhood hearing loss: a systematic review. *Ear Hear* 2021;**42**:1447–61.
127. Riga M, Korres G, Chouridis P, Naxakis S, Danielides V. Congenital cytomegalovirus infection inducing non-congenital sensorineural hearing loss during childhood; a systematic review. *Int J Pediatr Otorhinolaryngol* 2018;**115**:156–64.
128. Airlangga TJ, Bashiruddin J, Mangunatmadja I, Pandelaki J, Bardosono S, Ibrahim F, et al. Hearing instability and abnormal auditory pathways in infants with congenital cytomegalovirus infection: an audiological and radiological single-centre prospective cohort analysis. *Med J Malaysia* 2024;**79**:414–20.

129. Auriti C, Bucci S, De Rose DU, Coltella L, Santisi A, Martini L, et al. Maternal-fetal infections (cytomegalovirus, toxoplasma, syphilis): short-term and long-term neurodevelopmental outcomes in children infected and uninfected at birth. *Pathogens* 2022;**11**:1278.
130. Capretti MG, Marsico C, Chiereghin A, Gabrielli L, Aceti A, Lazzarotto T. Immune monitoring using quantiFERON R-CMV assay in congenital cytomegalovirus infection: correlation with clinical presentation and CMV DNA load. *Clin Infect Dis* 2021;**73**:367–73.
131. Dhondt C, Maes L, Rombaut L, Martens S, Vanaudenaerde S, Van Hoecke H, et al. Vestibular function in children with a congenital cytomegalovirus infection: 3 years of follow-up. *Ear Hear* 2021;**42**:76–86.
132. Hu BF, Liu XX, Zhan CY, Yuan TM, Chen LH, Liang JF, et al. [A multicenter study on the effects of congenital cytomegalovirus infection on hearing loss]. *Zhonghua Er Ke Za Zhi* 2024;**62**:721–6.
133. Jin HD, Demmler-Harrison GJ, Miller J, Edmond JC, Coats DK, Paysse EA, et al. Cortical visual impairment in congenital cytomegalovirus infection. *J Pediatr Ophthalmol Strabismus* 2019;**56**:194–202.
134. Lanzieri TM, Chung W, Leung J, Caviness AC, Baumgardner JL, Blum P, et al. Hearing trajectory in children with congenital cytomegalovirus infection. *Otolaryngol Head Neck Surg* 2018;**158**:736–44.
135. Lo TH, Lin PH, Hsu WC, Tsao PN, Liu TC, Yang TH, et al. Prognostic determinants of hearing outcomes in children with congenital cytomegalovirus infection. *Sci Rep* 2022;**12**:5219.
136. Royackers L, Rector E, Verhaert N, Desloovere C. Long-term audiological follow-up of children with congenital cytomegalovirus. *B-ENT* 2013;**9**:57–64.
137. Salome S, Ciampa N, Giordano M, Raimondi R, Capone E, Grieco C, et al. Ophthalmological impairment in patients with congenital cytomegalovirus infection. *Front Pediatr* 2023;**11**:1251893.
138. Seneviratne M, Fernando ME, Kandasamy Y, White A, Sabesan V, Norton R. Cytomegalovirus infection in a single-centre Australian neonatal cohort. *J Paediatr Child Health* 2022;**58**:1136–44.
139. Zavattoni M, Rustico M, Tassis B, Lombardi G, Furione M, Piralla A, et al. Risk of congenital disease in 46 infected fetuses according to gestational age of primary human cytomegalovirus infection in the mother. *J Med Virol* 2016;**88**:120–6.
140. Pinninti SG, Rodgers MD, Novak Z, Britt WJ, Fowler KB, Boppana SB, et al. Clinical predictors of sensorineural hearing loss and cognitive outcome in infants with symptomatic congenital cytomegalovirus infection. *Pediatr Infect Dis J* 2016;**35**:924–6.
141. Simonazzi G, Curti A, Murano P, Cervi F, Contoli M, Lazzarotto T, et al. Congenital cytomegalovirus infection and small for gestational age infants. *Prenat Diagn* 2014;**34**:765–9.
142. Tapasak B, Cronkite DA, Hustedt-Mai AR, Morlet TM, Parkes WJ, Maul TM, et al. Hearing outcomes in children with congenital cytomegalovirus: a multi-center, single-enterprise experience. *Int J Pediatr Otorhinolaryngol* 2022;**163**:111376.
143. Rovito R, Claas FHJ, Haasnoot GW, Roelen DL, Kroes ACM, Eikmans M, et al. Congenital cytomegalovirus infection: maternal-child HLA-C, HLA-E, and HLA-G affect clinical outcome. *Front Immunol* 2017;**8**:1904.
144. Rovito R, Korndewal MJ, van Zelm MC, Ziagkos D, Wessels E, van der Burg M, et al. T and B cell markers in dried blood spots of neonates with congenital cytomegalovirus infection: B cell numbers at birth are associated with long-term outcomes. *J Immunol* 2017;**198**:102–9.
145. Minsart AF, Rypens F, Smiljkovic M, Kakkar F, Renaud C, Lamarre V, et al. Prenatal findings, neonatal symptoms and neurodevelopmental outcome of congenital cytomegalovirus infection in a university hospital in Montreal, Quebec. *J Perinat Med* 2020;**48**:234–41.
146. Korndewal MJ, Oudesluys-Murphy AM, Kroes ACM, van der Sande MAB, de Melker HE, Vossen A. Long-term impairment attributable to congenital cytomegalovirus infection: a retrospective cohort study. *Dev Med Child Neurol* 2017;**59**:1261–8.

147. Bartlett AW, Hall BM, Palasanthiran P, McMullan B, Shand AW, Rawlinson WD. Recognition, treatment, and sequelae of congenital cytomegalovirus in Australia: an observational study. *J Clin Virol* 2018;**108**:121–5.
148. Koyano S, Morioka I, Oka A, Moriuchi H, Asano K, Ito Y, et al. Congenital cytomegalovirus in Japan: more than 2 year follow up of infected newborns. *Pediatr Int* 2018;**60**:57–62.
149. Topham JD, Miller JA, Wright GW, Turcich M, Vinson SS, Iovino I, et al. Inattention and hyperactivity in children with symptomatic and asymptomatic congenital cytomegalovirus. *J Dev Behav Pediatr* 2019;**40**:743–50.
150. Maes L, De Kegel A, Van Waelvelde H, De Leenheer E, Van Hoecke H, Goderis J, et al. Comparison of the motor performance and vestibular function in infants with a congenital cytomegalovirus infection or a connexin 26 mutation: a preliminary study. *Ear Hear* 2017;**38**:e49–56.
151. Dhondt C, Maes L, Martens S, Vanaudenaerde S, Rombaut L, Sucaet M, et al. Predicting early vestibular and motor function in congenital cytomegalovirus infection. *Laryngoscope* 2023;**133**:1757–65.
152. Forli F, Capobianco S, Berrettini S, Bruschini L, Lorenzoni F, Fiori S, et al. Long-term outcomes of congenital cytomegalovirus infection in children early identified by extended hearing-targeted screening. *Int J Pediatr Otorhinolaryngol* 2024;**184**:112070.
153. Rohren L, Shanley R, Smith M, Yue M, Huang TC, Nelson P, et al. Congenital cytomegalovirus-associated sensorineural hearing loss in children: identification following universal newborn hearing screening, effect of antiviral treatment, and long-term hearing outcomes. *Ear Hear* 2024;**45**:198–206.
154. Yamaguchi M, Kawada JI, Torii Y, Haruta K, Suzuki T, Horiba K, et al. Quantitative assessment of viral load in the blood and urine of patients with congenital cytomegalovirus infection using droplet digital PCR. *J Med Virol* 2022;**94**:4559–64.
155. Marsico C, Aban I, Kuo H, James SH, Sanchez PJ, Ahmed A, et al. Blood viral load in symptomatic congenital cytomegalovirus infection. *J Infect Dis* 2019;**219**:1398–406.
156. Meyer L, Sharon B, Huang TC, Meyer AC, Gravel KE, Schimmenti LA, et al. Analysis of archived newborn dried blood spots (DBS) identifies congenital cytomegalovirus as a major cause of unexplained pediatric sensorineural hearing loss. *Am J Otolaryngol* 2017;**38**:565–70.
157. Kakei Y, Morioka I, Imai T, Itohara K, Yano I, Takahashi N, et al. Assessment of patients' characteristics associated with the efficacy and safety of oral valganciclovir treatment for infants with symptomatic congenital cytomegalovirus disease. *J Infect Chemother* 2024;**30**:971–7.
158. Salome S, Giannattasio A, Malesci R, Marciano E, Dolce P, Portella G, et al. The natural history of hearing disorders in asymptomatic congenital cytomegalovirus infection. *Front Pediatr* 2020;**8**:217.
159. Wang C, Liu X, Wang S, Zhang W, Wang H, Ma W, et al. Late-onset hearing loss from congenital cytomegalovirus infection after newborn period in a highly immune population in China. *Pediatr Infect Dis J* 2021;**40**:70–3.
160. Kawada J, Torii Y, Kawano Y, Suzuki M, Kamiya Y, Kotani T, et al. Viral load in children with congenital cytomegalovirus infection identified on newborn hearing screening. *J Clin Virol* 2015;**65**:41–5.
161. Kabani N, Pinninti S, Boppana S, Fowler K, Ross S. Urine and saliva viral load in children with congenital cytomegalovirus infection. *J Pediatric Infect Dis Soc* 2023;**12**:230–3.
162. Ouellette CP, Sanchez PJ, Xu Z, Blankenship D, Zeray F, Ronchi A, et al. Blood genome expression profiles in infants with congenital cytomegalovirus infection. *Nat Commun* 2020;**11**:3548.
163. Chen LY, Li W, Xu JL, Tao R, Li HM, Liu LF, et al. [Relationship between gH genotyping and clinical characteristics of children with congenital cytomegalovirus infection]. *Zhonghua Er Ke Za Zhi* 2019;**57**:597–602.
164. Dong N, Cao L, Zheng D, Su L, Lu L, Dong Z, et al. Distribution of CMV envelope glycoprotein B, H and N genotypes in infants with congenital cytomegalovirus symptomatic infection. *Front Pediatr* 2023;**11**:112645.

165. Kasztelewicz B, Czech-Kowalska J, Lipka B, Milewska-Bobula B, Borszewska-Kornacka MK, Romanska J, et al. Cytokine gene polymorphism associations with congenital cytomegalovirus infection and sensorineural hearing loss. *Eur J Clin Microbiol Infect Dis* 2017;**36**:1811–8.
166. Medoro AK, Dhital R, Sanchez PJ, Flint K, Graber B, Pifer T, et al. T cell responses and clinical symptoms among infants with congenital cytomegalovirus infection. *JCI Insight* 2024;**9**:e171029.
167. Pati SK, Pinninti S, Novak Z, Chowdhury N, Patro RK, Fowler K, et al. Genotypic diversity and mixed infection in newborn disease and hearing loss in congenital cytomegalovirus infection. *Pediatr Infect Dis J* 2013;**32**:1050–4.
168. Rovito R, Warnatz HJ, Kielbasa SM, Mei H, Amstislavskiy V, Arens R, et al. Impact of congenital cytomegalovirus infection on transcriptomes from archived dried blood spots in relation to long-term clinical outcome. *PLoS ONE* 2018;**13**:e0200652.
169. Czech-Kowalska J, Jedlinska-Pijanowska D, Kasztelewicz B, Klodzinska M, Pietrzyk A, Sarkaria E, et al. The limitations of cytomegalovirus DNA detection in cerebrospinal fluid of newborn infants with congenital CMV infection: a tertiary care neonatal center experience. *Pediatr Infect Dis J* 2021;**40**:838–45.
170. Goycochea-Valdivia WA, Baquero-Artigao F, del Rosal T, Frick MA, Rojo P, Echeverria MJ, et al. Cytomegalovirus DNA detection by polymerase chain reaction in cerebrospinal fluid of infants with congenital infection: associations with clinical evaluation at birth and implications for follow-up. *Clin Infect Dis* 2017;**64**:1335–42.
171. Yamada H, Tanimura K, Fukushima S, Fujioka K, Deguchi M, Sasagawa Y, et al. A cohort study of the universal neonatal urine screening for congenital cytomegalovirus infection. *J Infect Chemother* 2020;**26**:790–4.
172. Dreher AM, Arora N, Fowler KB, Novak Z, Britt WJ, Boppana SB, et al. Spectrum of disease and outcome in children with symptomatic congenital cytomegalovirus infection. *J Pediatr* 2014;**164**:855–9.
173. Hilditch C, Liersch B, Spurrier N, Callander EJ, Cooper C, Keir AK. Does screening for congenital cytomegalovirus at birth improve longer term hearing outcomes? *Arch Dis Child* 2018;**103**:988–92.
174. Pollick SA, Mansour Y, Pesch MH. Newborn congenital cytomegalovirus screening and hearing outcomes: a systematic review of current literature. *Curr Opin Otolaryngol Head Neck Surg* 2024;**32**:329–38.
175. Puhakka L, Lappalainen M, Lonnqvist T, Niemensivu R, Lindahl P, Nieminen T, et al. The burden of congenital cytomegalovirus infection: a prospective cohort study of 20,000 infants in Finland. *J Pediatric Infect Dis Soc* 2019;**8**:205–12.
176. Chierighin A, Pavia C, Turello G, Borgatti EC, Baiesi Pillastrini F, Gabrielli L, et al. Universal newborn screening for congenital cytomegalovirus infection - from infant to maternal infection: a prospective multicenter study. *Front Pediatr* 2022;**10**:909646.
177. Dunn JKE, Chakraborty P, Reuvers E, Gallagher L, Kernohan KD, Lacaria M, et al. Outcomes of a population-based congenital cytomegalovirus screening program. *JAMA Pediatr* 2025;**179**:332–9.
178. Yamamoto AY, Anastasio ART, Massuda ET, Isaac ML, Manfredi AKS, Cavalcante JMS, et al. Contribution of congenital cytomegalovirus infection to permanent hearing loss in a highly seropositive population: the Brazilian cytomegalovirus hearing and maternal secondary infection study. *Clin Infect Dis* 2020;**70**:1379–84.
179. Kaye T, Dufort EM, Rosendahl SD, Hullerman Umar J, Pavan A, Tricas K, et al. Notes from the field: universal newborn screening and surveillance for congenital cytomegalovirus - Minnesota, 2023-2024. *MMWR Morb Mortal Wkly Rep* 2024;**73**:703–5.
180. Kruc RM, Osterholm EA, Holm T, Nestrail I, Lanzieri TM, Schleiss MR. Cranial ultrasound findings in infants with congenital cytomegalovirus infection in a universal newborn screening study in Minnesota. *J Pediatric Infect Dis Soc* 2024;**13**:413–20.
181. Yamaguchi A, Oh-Ishi T, Arai T, Sakata H, Adachi N, Asanuma S, et al. Screening for seemingly healthy newborns with congenital cytomegalovirus infection by quantitative real-time polymerase chain reaction using newborn urine: an observational study. *BMJ Open* 2017;**7**:e013810.

182. Demortier J, Fourgeaud J, Abasse S, Lambrecht L, Gromand M, Boumahni B, et al. A prospective study evaluating congenital CMV infection in Mayotte and La Reunion Islands (France). *J Clin Virol* 2021;**138**:104793.
183. Merav L, Ofek Shlomai N, Oiknine-Djian E, Caplan O, Livneh A, Sido T, et al. Implementation of pooled saliva tests for universal screening of cCMV infection. *Nat Med* 2024;**30**:1111–7.
184. Papaevangelou V, Christoni Z, Vliora C, Kottaridi C, Fotiou A, Malamitsi-Puchner A, et al. Neonatal screening for congenital CMV infection stresses the importance of maternal nonprimary infection even in an area where prenatal serology testing is common. *J Matern Fetal Neonatal Med* 2019;**32**:1901–4.
185. Zeytinoglu A, Terek D, Arslan A, Erensoy S, Altun Koroglu O, Bozdemir T, et al. [Investigation of congenital CMV infection with the presence of CMV DNA in saliva samples of new born babies]. *Mikrobiyol Bul* 2019;**53**:53–60.
186. Ari-Even Roth D, Lubin D, Kuint J, Teperberg-Oikawa M, Mendelson E, Strauss T, et al. Contribution of targeted saliva screening for congenital CMV-related hearing loss in newborns who fail hearing screening. *Arch Dis Child Fetal Neonatal Ed* 2017;**102**:F519–24.
187. Chung PK, Schornagel F, Oudesluys-Murphy AM, de Vries LS, Soede W, van Zwet E, et al. Targeted screening for congenital cytomegalovirus infection: clinical, audiological and neuroimaging findings. *Arch Dis Child Fetal Neonatal Ed* 2023;**108**:F302–8.
188. Ronner EA, Glovsky CK, Herrmann BS, Woythaler MA, Pasternack MS, Cohen MS. Congenital cytomegalovirus targeted screening implementation and outcomes: a retrospective chart review. *Otolaryngol Head Neck Surg* 2022;**167**:178–82.
189. Medoro AK, Malhotra PS, Shimamura M, Findlen U, Gerth H, Hounam G, et al. Timing of newborn hearing screening in the neonatal intensive care unit: implications for targeted screening for congenital cytomegalovirus infection. *J Perinatol* 2021;**41**:310–4.
190. Vancor E, Shapiro ED, Loyal J. Results of a targeted screening program for congenital cytomegalovirus infection in infants who fail newborn hearing screening. *J Pediatric Infect Dis Soc* 2019;**8**:55–9.
191. Beswick R, McHugh L, Clark JE. Integrating congenital cytomegalovirus screening within a newborn hearing screening program: is it worthwhile? *Int J Pediatr Otorhinolaryngol* 2021;**142**:110594.
192. Fowler KB, McCollister FP, Sabo DL, Shoup AG, Owen KE, Woodruff JL, et al. A targeted approach for congenital cytomegalovirus screening within newborn hearing screening. *Pediatrics* 2017;**139**:e20162128.
193. Wei D, Sardesai SR, Barton L. The C in TORCH: a cost-effective alternative to screening small-for-gestational-age infants. *Neonatology* 2014;**106**:24–9.
194. Kadambari S, Luck S, Davis A, Walter S, Agrup C, Atkinson C, et al. Evaluating the feasibility of integrating salivary testing for congenital CMV into the Newborn Hearing Screening Programme in the UK. *Eur J Pediatr* 2015;**174**:1117–21.
195. Diener ML, Zick CD, McVicar SB, Boettger J, Park AH. Outcomes from a hearing-targeted cytomegalovirus screening program. *Pediatrics* 2017;**139**:e20160789.
196. Fourgeaud J, Boithias C, Walter-Nicolet E, Kermorvant E, Couderc S, Parat S, et al. Performance of targeted congenital cytomegalovirus screening in newborns failing universal hearing screening: a multicenter study. *Pediatr Infect Dis J* 2022;**41**:478–81.
197. Rawlinson WD, Palasanthiran P, Hall B, Al Yazidi L, Cannon MJ, Cottier C, et al. Neonates with congenital cytomegalovirus and hearing loss identified via the universal newborn hearing screening program. *J Clin Virol* 2018;**102**:110–5.
198. Williams EJ, Kadambari S, Berrington JE, Luck S, Atkinson C, Walter S, et al. Feasibility and acceptability of targeted screening for congenital CMV-related hearing loss. *Arch Dis Child Fetal Neonatal Ed* 2014;**99**:F230–6.
199. Masarweh K, Felszer-Fisch C, Shinwell E, Hasanein J, Peniakov M, Weiner SA, et al. The yield of targeted examination for the detection of symptomatic congenital cytomegalovirus infection. *Isr Med Assoc J* 2021;**23**:318–22.
200. McCrary H, Shi K, Newberry IC, Hamilton C, Ostrander B, O'Brien E, et al. Outcomes from an expanded targeted early cytomegalovirus testing program. *J Pediatr Infect Dis* 2020;**15**:189–94.

201. Palma S, Botti C, Roversi MF, Bettini M, Pietrosevoli P, Berardi A, et al. What happens when the newborn hearing screening program is integrated with congenital cytomegalovirus infection screening? Preliminary results in a tertiary hospital. *Hearing Balance Commun* 2021;**19**:175–9.
202. Fujii T, Oka A, Morioka I, Moriuchi H, Koyano S, Yamada H, et al. Newborn congenital cytomegalovirus screening based on clinical manifestations and evaluation of DNA-based assays for in vitro diagnostics. *Pediatr Infect Dis J* 2017;**36**:942–6.
203. Zhang Y, Egashira T, Egashira M, Ogiwara S, Tomino H, Shichijo A, et al. Expanded targeted screening for congenital cytomegalovirus infection. *Congenit Anom (Kyoto)* 2023;**63**:79–82.
204. Lu CY, Tsao PN, Ke YY, Lin YH, Lin YH, Hung CC, et al. Concurrent hearing, genetic, and cytomegalovirus screening in newborns, Taiwan. *J Pediatr* 2018;**199**:144–50.
205. Tan YY, Chan CY, Sothirasan K, Tan PL, Maiwald M, Thoon KC, et al. Outcomes of a targeted congenital cytomegalovirus infection screening approach among infants born ≥ 35 weeks gestation. *Ann Acad Med Singap* 2023;**52**:643–4.
206. Webb E, Hodgson J, Gillespie AN, Jones CA, Poulakis Z, Wong J, et al. Hearing screening for congenital cytomegalovirus-exploring parents' experiences of completing targeted congenital cytomegalovirus screening at the time of their infants' newborn hearing screening. *J Clin Med* 2024;**13**:4367.
207. Forli F, Lazzerini F, Canelli R, Lorenzoni F, Franciosi B, Berrettini S, et al. Extended-hearing targeted screening for congenital cytomegalovirus infection. *Minerva Pediatr (Torino)* 2024;**76**:590–8.
208. Akiko U, Hitomi I, Yutoku S, Yuki S, Tokuro S, Kenji T, et al. Clinical factors associated with congenital cytomegalovirus infection in pregnant women: a cohort study at a perinatal center. *J Obstet Gynaecol Res* 2021;**47**:2926.
209. Akiva MH, Hyde De Souza H, Lamarre V, Boucoiran I, Gantt S, Renaud C, et al. Identifying clinical criteria for an expanded targeted approach to screening for congenital cytomegalovirus infection - a retrospective study. *Int J Neonatal Screen* 2023;**9**:24.
210. Albano MS, Ciubotariu R, Dobrila L, Tarnawski M, Watanabe C, Madry MM, et al. Development and validation of a real-time polymerase chain reaction (RT-PCR) to detect human cytomegalovirus (CMV) viremia in cord blood (CB)-derived cell therapy products. *Transfusion* 2014;**54**:22A.
211. Albano MS, Ciubotariu R, Watanabe C, Krishnan S, Tarnawski M, Dobrila L, et al. Cytomegalovirus (CMV) viral load in cord blood (CB) donors and impact of congenital viremia on markers of hematopoietic progenitor cell (HPC) potency. *Blood* 2016;**128**:2174.
212. Albano MS, Ciubotariu R, Dobrila L, Tarnawski M, DeLeon M, Watanabe C, et al. Cytomegalovirus viral load in cord blood and impact of congenital infection on markers of hematopoietic progenitor cell potency. *Transfusion* 2017;**57**:2768–74.
213. Alifleraki S, Payne H, Hathaway C, Tan RWY, Lyall H. Delays in diagnosis and treatment initiation for congenital cytomegalovirus infection - Why we need universal screening. *Front Pediatr* 2022;**10**:988039.
214. Almeida S, Gouveia P, Jorge A, Fortuna A, Binda S, Barbi M, et al. Diagnosing congenital cytomegalovirus infections using archived dried blood spots: a 15-year observational study, Portugal. *J Clin Virol* 2023;**165**:105516.
215. Alwan SN, Shamran HA, Ghaib AH, Kadhim HS, Al-Mayah QS, Al-Saffar AJ, et al. Genotyping of cytomegalovirus from symptomatic infected neonates in Iraq. *Am J Trop Med Hyg* 2019;**100**:957–63.
216. Amin MM, Wong P, McCann M, Dollard SC. Detection of cytomegalovirus in urine dried on filter paper. *J Pediatric Infect Dis Soc* 2021;**10**:958–61.
217. Carter A, Rawlinson W, Ward M, Palasanthiran P, Shand A. Retrospective audit of cytomegalovirus (CMV) testing in neonates and rationale for testing. *J Paediatr Child Health* 2023;**59**:65–6.
218. Anonymous. Erratum: Hearing in children with congenital cytomegalovirus infection: results of a longitudinal study. *J Pediatr* 2016;**177**:335.
219. Arancibia J, Coqueiro A, Sgardoli I, Lamas A, Mesquita P, Benicio R, et al. Incorporating the screening for congenital cytomegalovirus into a newborn genetic testing

- performed on saliva swab: an intersection between human genetics and infectious diseases. *Clin Chem* 2023;**69**:i114.
220. Arellano-Galindo J, Villanueva-Garcia D, Cruz-Ramirez JL, Yalaupari-Mejia JP, Uribe-Gutierrez G, Velazquez-Guadarrama N, et al. Detection and gB genotyping of CMV in Mexican preterm infants in the context of maternal seropositivity. *J Infect Dev Ctries* 2014;**8**:758–67.
221. Banjac L, Banjac G. Cranial ultrasound in congenital cytomegalovirus infection. *Current Pediatric Research* 2022;**26**:1473–5.
222. Barriga EM, Delpino MV, Berberian G, Sagarzazú EL, Ruhle M, Rosanova MT, et al. Congenital cytomegalovirus infection: diagnosis assessment and challenges in early detection. *Acta Bioquímica Clínica Latinoamericana* 2024;**58**:169–78.
223. Ben Shoham A, Schlesinger Y, Miskin I, Kalderon Z, Michaelson-Cohen R, Wiener-Well Y. Cytomegalovirus (CMV) seroprevalence among women at childbearing age, maternal and congenital CMV infection: policy implications of a descriptive, retrospective, community-based study. *Isr J Health Policy Res* 2023;**12**:16.
224. Berg C, Friis MB, Rosenkilde MM, Benfield T, Nielsen L, Luttichau HR, et al. Development of highly efficient protocols for extraction and amplification of cytomegalovirus DNA from dried blood spots for detection and genotyping of polymorphic immunomodulatory genes. *PLoS ONE* 2019;**14**:e0222053.
225. Bergevin A, Zick CD, McVicar SB, Park AH. Cost-benefit analysis of targeted hearing directed early testing for congenital cytomegalovirus infection. *Int J Pediatr Otorhinolaryngol* 2015;**79**:2090–3.
226. Beswick R, David M, Higashi H, Thomas D, Nourse C, Koh G, et al. Integration of congenital cytomegalovirus screening within a newborn hearing screening programme. *J Paediatr Child Health* 2019;**55**:1381–8.
227. Bilavsky E, Pardo J, Attias J, Levy I, Magny JF, Ville Y, et al. Clinical implications for children born with congenital cytomegalovirus infection following a negative amniocentesis. *Clin Infect Dis* 2016;**63**:33–8.
228. Bilavsky E, Watad S, Levy I, Linder N, Pardo J, Ben-Zvi H, et al. Positive IgM in congenital CMV infection. *Clin Pediatr (Phila)* 2017;**56**:371–5.
229. Blazquez-Gamero D, Soriano-Ramos M, Vicente M, Pallas-Alonso CR, Perez-Rivilla A, Garcia-Alvarez M, et al. Prevalence and clinical manifestations of congenital cytomegalovirus infection in a screening program in Madrid (PICCSA Study). *Pediatr Infect Dis J* 2020;**39**:1050–6.
230. Blazquez-Gamero D, Sanchez B, Folgueira MD. Dried blood spot testing for detection of congenital cytomegalovirus. *JAMA Pediatrics* 2021;**175**:865–6.
231. Boppana S, Ross S, Pati S, Britt WJ, Fowler K. Breastfeeding and newborn congenital cytomegalovirus screening using saliva polymerase chain reaction. *Open Forum Infect Dis* 2016;**3**:S1.
232. Boscarino G, Romano R, Tegoni F, Iotti C, Perrone S, Esposito S, et al. Congenital cytomegalovirus severity definitions and treatment decisions around the world: a systematic scoping review of the literature. *J Clin Med* 2024;**13**:5997.
233. Branas P, Blazquez-Gamero D, Galindo A, Prieto C, Olabarrieta I, Cuadrado I, et al. Cytomegalovirus genotype distribution among congenitally and postnatally infected patients: association of particular glycoprotein (g)B and gN types with symptomatic disease. *Open Forum Infect Dis* 2015;**2**:ofv151.
234. Brunet J. *Pertinence de l'ajout du dépistage universel de l'infection congénitale au cytomégalo virus (CMV) au programme québécois de dépistage néonatal*. Quebec, Canada: Institut national d'excellence en sante et en services sociaux (INESSS); 2024.
235. Campione A, Lanzieri TM, Ricotta E, Grosse SD, Kadri SS, Nussenblatt V, et al. Missing diagnoses of congenital cytomegalovirus infection in electronic health records for infants with laboratory-confirmed infection. *Curr Med Res Opin* 2022;**38**:273–5.
236. Canadian Agency for Drugs and Technologies in Health. *Newborn screening for congenital cytomegalovirus: rapid review*. Ottawa: CADTH; 2024.
237. Central Hospital Nancy France. *Diagnosis of congenital cytomegalovirus infection in newborn with particular risk*. ClinicalTrials.gov. Bethesda (MD): National Library of Medicine

(US); 2023. Available from: <https://clinicaltrials.gov/study/NCT05754879> [accessed 20 November 2024].

238. Chasqueira MJ, Fernandez C, Marques A, Rodrigues L, Marcal M, Tuna M, et al. Pooling saliva sample as an effective strategy for the systematic CMV screening of newborns - a multicentric prospective study. *Pediatr Infect Dis J* 2023;**42**:1117–20.
239. Chauhan A, Poomkudy JT, Stellato N, Sood SK. A large health system wide approach to early identification of congenital CMV. *Pediatrics* 2021;**147**:114–5.
240. Chierighin A, Pavia C, Gabrielli L, Piccirilli G, Squarzone D, Turello G, et al. Clinical evaluation of the new Roche platform of serological and molecular cytomegalovirus-specific assays in the diagnosis and prognosis of congenital cytomegalovirus infection. *J Virol Methods* 2017;**248**:250–4.
241. Chisholm KM, Folkins AK, Guo F, McDowell M, Aziz N, Pinsky BA. Diagnosis of congenital CMV using PCR performed on formalin-fixed, paraffin-embedded placental tissue. *Pediatr Dev Pathol* 2013;**16**:52.
242. Chisholm KM, Aziz N, McDowell M, Guo FP, Srinivas N, Benitz WE, et al. Evaluation of serial urine viral cultures for the diagnosis of cytomegalovirus infection in neonates and infants. *Pediatr Dev Pathol* 2014;**17**:176–80.
243. Choudhary A, Pati SK, Patro RK, Deorari AK, Dar L. Comparison of conventional, immunological and molecular techniques for the diagnosis of symptomatic congenital human cytomegalovirus infection in neonates and infants. *Indian J Med Microbiol* 2015;**33**:15–9.
244. Correa C, Kouri V, Perez L, Soto Y, Limia C. Diagnosis, gB genotype distribution and viral load of symptomatic congenitally infected CMV patients in Cuba. *J Perinatol* 2016;**36**:837–42.
245. Cushing SL, Purcell PL, Papaiaonnou V, Neghandi J, Daien M, Blaser SI, et al. Hearing instability in children with congenital cytomegalovirus: evidence and neural consequences. *Laryngoscope* 2022;**132**:S1–24.
246. Czech-Kowalska J, Jedlinska-Pijanowska D, Pleskaczynska AK, Niezgoda A, Gradowska K, Pietrzyk A, et al. Single nucleotide polymorphisms of interleukins and toll-like receptors and neuroimaging results in newborns with congenital HCMV infection. *Viruses* 2021;**13**:1783.
247. Dallaire S, Filippov G, Veling M, Dougherty E, Miller M, Xie P, et al. Congenital cytomegalovirus detection in newborn screening: an Eonis™ assay. *Mol Genet Metab* 2021;**132**:S187–8.
248. Dallaire S, McCunn K, Ravenscroft J, Kennedy J, Hegde M. [eP498] Detection of congenital cytomegalovirus infection on high-risk newborn population. *Genet Med* 2022;**24**:S316.
249. Dar L, Namdeo D, Kumar P, Thakar A, Kant S, Rai S, et al. Congenital cytomegalovirus infection and permanent hearing loss in rural North Indian children. *Pediatr Infect Dis J* 2017;**36**:670–3.
250. De Cuyper E, Acke F, Keymeulen A, De Leenheer EMR, Van Hoecke H, Padalko E, et al. Risk Factors for hearing loss at birth in newborns with congenital cytomegalovirus infection. *JAMA Otolaryngol Head Neck Surg* 2023;**149**:122–30.
251. de Vries JJC, van Zwet EW, Dekker FW, Kroes ACM, Verkerk PH, Vossen A. The apparent paradox of maternal seropositivity as a risk factor for congenital cytomegalovirus infection: a population-based prediction model. *Rev Med Virol* 2013;**23**:241–9.
252. Del Valle Penella A, Rochat RH, Demmler-Harrison GJ. Clinical characteristics of children with congenital cytomegalovirus diagnosed retrospectively by dried blood spot. *Open Forum Infect Dis* 2023;**10**:S750–1.
253. Demmler-Harrison GJ. Newborn dried blood spot testing for congenital cytomegalovirus screening: the little engine that could. *JAMA Pediatr* 2021;**175**:e205445.
254. Dharnidhar A, Kousalya V, Barani R, Mani M, Gopalakrishnan K, Padmasani LN, et al. Detection of cytomegalovirus by real time PCR among pediatric patients in different samples. *BMC Infect Dis* 2019;**20**:68.
255. Dimopoulou D, Kourlaba G, Antoniadou A, Mariolis L, Kavatha D, Stoungioti S, et al. Low birth weight and head circumference as potential biomarkers of sensorineural hearing loss in asymptomatic congenitally CMV-infected infants. *J Clin Virol* 2020;**129**:104471.

256. Dobbins GC, Patki A, Chen D, Tiwari HK, Hendrickson C, Britt WJ, et al. Association of CMV genomic mutations with symptomatic infection and hearing loss in congenital CMV infection. *BMC Infect Dis* 2019;**19**:1046.
257. Dougherty E, Veling M, McCunn K, Ravenscroft J, Dallaire S, Kennedy J, et al. [eP487] Universal newborn screening of congenital cytomegalovirus using dried blood spots and qPCR. *Genet Med* 2022;**24**:S310.
258. Dreher AM, Fowler KB, Boppana SB, Ross SA, Britt WJ. The importance of newborn screening in defining the natural history of congenital cytomegalovirus infection. *Biopolym Cell* 2013;**29**:31.
259. Ebrahimi-Rad M, Shakeri TS, Shirvani F, Shahrokhi K, Shahrokhi N. Prevalence of congenital cytomegalovirus infection in symptomatic newborns under 3 weeks in Tehran, Iran. *BMC Infect Dis* 2017;**17**:688.
260. Eres Saritas Z, Peker BO, Colak D, Saglik I, Sarinoglu RC, Turhan M, et al. Congenital cytomegalovirus infection cases and follow-up findings in Antalya, Turkey. *Marmara Medical Journal* 2023;**36**:290–6.
261. Eventov-Friedman S, Zaharan S, Geal-Dor M, Wolf D, Bar-Oz B. Predictive values of prenatal and neonatal testing indications for the diagnosis of congenital cytomegalovirus (CMV) infection. *Arch Dis Child* 2014;**99**:A424.
262. Farnan SK, Sintim-Aboagye E, Quach H, Punia S, Mejia-Plazas M, Verma N, et al. Characterization of the placental proteome during congenital CMV infection. *Open Forum Infect Dis* 2023;**10**:S170–1.
263. Faure-Bardon V, Millischer AE, Deloison B, Sonigo P, Grevent D, Salomon L, et al. Refining the prognosis of fetuses infected with cytomegalovirus in the first trimester of pregnancy by serial prenatal assessment: a single-centre retrospective study. *BJOG* 2020;**127**:355–62.
264. Fernandes C, Marques A, de Jesus Chasqueira M, Braz MC, Ferreira AR, Neto AS, et al. Saliva pools for screening of human cytomegalovirus using real-time PCR. *Eur J Pediatr* 2021;**180**:1067–72.
265. Folkins AK, Chisholm KM, Guo FP, McDowell M, Aziz N, Pinsky BA. Diagnosis of congenital CMV using PCR performed on formalin-fixed, paraffin-embedded placental tissue. *Am J Surg Pathol* 2013;**37**:1413–20.
266. Gandhoke I, Hussain SA, Pasha ST, Chauhan LS, Khare S. Glycoprotein B genotyping in congenital/perinatal cytomegalovirus infection in symptomatic infants. *Indian Pediatr* 2013;**50**:663–7.
267. Gantt S, Goldfarb DM, Dionne F, Bulman D, Doutre SM. In reference to should infants who fail their newborn hearing screen undergo cytomegalovirus testing? *Laryngoscope* 2018;**128**:E267.
268. Garnham J, Gaur P, Basheer N, Lyall H, Jan W, Kachramanoglou C. Evolution of the intracranial features of congenital cytomegalovirus on MRI. *Clin Radiol* 2023;**78**:e451–7.
269. Gillespie AN, Dalziel K, Webb E, Wong J, Jones CA, Sung V. Targeted screening for congenital cytomegalovirus: a micro-costing analysis. *J Paediatr Child Health* 2023;**59**:64–71.
270. Goderis J, De Leenheer E, Van Hoecke H, Smets K, Keymeulen A, Dhooge I. Audiological findings in congenital cytomegalovirus infection: preliminary results of a prospective cohort study. *B-ENT* 2013;**20**:45.
271. Goel A, Chaudhari S, Sutar J, Bhonde G, Bhatnagar S, Patel V, et al. Detection of cytomegalovirus in liver tissue by polymerase chain reaction in infants with neonatal cholestasis. *Pediatr Infect Dis J* 2018;**37**:632–6.
272. Goetzl L, Eldar-Yedidia Y, Hillel M, Darbinian N, Merabova N, Valeri A, et al. Non-invasive diagnosis of CMV infection with fetal central nervous system derived exosomes. *Reproductive Sciences* 2019;**26**:237A.
273. Gonzalez Lopez M, Mesa Fernandez A, Martin Cantero M, Gomez Robles C, Salguero Garcia E, Rodriguez Vives M. Cytomegalovirus infection in neonate with risk: Importance of an early diagnosis and monitoring. *Journal of Perinatal Medicine. Conference: 12th World Congress of Perinatal Medicine* 2015;**43**.

274. Gorzer I, Trajanoski S, Popow-Kraupp T, Puchhammer-Stockl E. Analysis of human cytomegalovirus strain populations in urine samples of newborns by ultra deep sequencing. *J Clin Virol* 2015;**73**:101–4.
275. Gunes O, Gulhan B, Uckardes F, Guney AY, Coskun ZN, Ozen S, et al. Assessment of blood viral load in asymptomatic and symptomatic cases of congenital cytomegalovirus infection. *J Med Virol* 2024;**96**:e29853.
276. Gupta A, Lawrence SM, Fraley SI. A broad-based probe-free qPCR assay for detection and discrimination of three human herpes viruses. *J Virol Methods* 2023;**322**:114824.
277. Gutierrez Posso JD, Anta Escuredo JA, Aguirre Unceta-Barrenechea A, Zabala Lopez de Maturana JA. Importance of congenital cytomegalovirus in the neonatal hearing screening program. *Acta Otorrinolaringol Esp* 2023;**74**:346–51.
278. Gwee A, Curtis N, Garland SM, Connell TG, Daley AJ. Question 2: which infants with congenital cytomegalovirus infection benefit from antiviral therapy? *Arch Dis Child* 2014;**99**:597–601.
279. Hadar E, Salzer L, Dorfman E, Amir J, Pardo J. Antenatal risk factors for symptomatic congenital CMV disease following primary maternal CMV infection. *J Perinat Med* 2016;**44**:339–44.
280. Hadar E, Dorfman E, Bardin R, Gabbay-Benziv R, Amir J, Pardo J. Symptomatic congenital cytomegalovirus disease following non-primary maternal infection: a retrospective cohort study. *BMC Infect Dis* 2017;**17**:31.
281. Hamid NA, Ismail N, Hashim AMB, Mohamad S, Salmuna ZN. The prevalence of cytomegalovirus (CMV) infection among infants and correlation between CMV PCR with clinical outcomes in a tertiary teaching hospital in Malaysia. *Sains Malays* 2022;**51**:239–47.
282. Hamilton ST, van Zuylen W, Shand A, Scott GM, Naing Z, Hall B, et al. Prevention of congenital cytomegalovirus complications by maternal and neonatal treatments: a systematic review. *Rev Med Virol* 2014;**24**:420–33.
283. Hasan AH, Hasany HJ, Al-Abbasi AM, Aziz SS, Samanje J. Impact of cytomegalovirus infection as a cause of sensorineural hearing loss (SNHL) in children of Basrah, Southern Iraq. *Current Pediatric Research* 2021;**25**:929–35.
284. Hernandez-Alvarado N, Bierle CJ, Schleiss MR. Droplet digital PCR (ddPCR) does not enhance the sensitivity of detection of cytomegalovirus (CMV) DNA in newborn dried blood spots evaluated in the context of newborn congenital CMV (cCMV) screening. *Int J Neonatal Screen* 2023;**10**:1.
285. Hirth D, Pfisterer L, Stover T, Kramer S, Schuetz L, Buxmann H. Hearing test-triggered screening of CMV-infection in newborns. *Laryngorhinootologie* 2018;**97**:S300.
286. Hirth D, Stover T, Kramer S. Late-onset hearing loss in children with congenital CMV infection. *Laryngorhinootologie* 2021;**100**:S252.
287. Hranilovich J, Park A, Bale J. Brain abnormalities in children with congenital cytomegalovirus infection identified through newborn hearing screening. *Ann Neurol* 2016;**80**:S350–1.
288. Hranilovich JA, Park AH, Knackstedt ED, Ostrander BE, Hedlund GL, Shi K, et al. Brain magnetic resonance imaging in congenital cytomegalovirus with failed newborn hearing screen. *Pediatr Neurol* 2020;**110**:55–8.
289. Hu HB, Wu JG, Sun JJ, Peng QY, Shang XP. Cytomegalovirus genotype and virulence in infants with congenital infection. *J Pediatr Infect Dis* 2021;**16**:171–7.
290. Huang Y, Tang J, Yu H, Song Q, Hao M, Wang H, et al. Reconsideration of maternal serological testing for predicting congenital CMV infection. *J Infect Dis* 2024;**229**:1817–22.
291. Ikuta K, Minematsu T, Inoue N, Kubo T, Asano K, Ishibashi K, et al. Cytomegalovirus (CMV) glycoprotein H-based serological analysis in Japanese healthy pregnant women, and in neonates with congenital CMV infection and their mothers. *J Clin Virol* 2013;**58**:474–8.
292. Imafuku H, Yamada H, Uchida A, Deguchi M, Shirakawa T, Sasagawa Y, et al. Clinical and ultrasound features associated with congenital cytomegalovirus infection as potential predictors for targeted newborn screening in high-risk pregnancies. *Sci Rep* 2020;**10**:19706.
293. Ishida Y, Takeshita M, Morishita N, Morichi S, Morishima Y, Oana S, et al. Clinical analysis of 7 children with congenital cytomegalovirus infection. *No To Hattatsu* 2015;**47**:S268.

294. Ismail A, Murphy AM, Qadri S, Sreenan C. Sensitivity of torch screen vs urine for CMV in diagnosing intrauterine infection in midwestern region. *Intensive Care Med* 2013;**39**:S83–4.
295. Izquierdo G, Farfan MJ, Villavicencio L, Montecinos L, Tarque F, Acevedo W, et al. Optimizing congenital cytomegalovirus detection by pool testing in saliva by a rapid molecular test. *Eur J Pediatr* 2023;**182**:5131–6.
296. Javid N, Cheraghali F, Moradi A, Kelishadi M, Tabarraei A. Newborn screening for congenital cytomegalovirus infection in Iran. *Pediatr Infect Dis J* 2016;**35**:1052.
297. Jedlinska-Pijanowska D, Kasztelewicz B, Czech-Kowalska J, Jaworski M, Charusta-Sienkiewicz K, Dobrzanska A. Association between single nucleotide polymorphisms (SNPs) of IL1, IL12, IL28 and TLR4 and symptoms of congenital cytomegalovirus infection. *PLoS One* 2020;**15**:e0233096.
298. Jedlinska-Pijanowska D, Kasztelewicz B, Dobrzanska A, Dzierzanowska-Fangrat K, Jaworski M, Czech-Kowalska J. Association between single nucleotide polymorphisms and viral load in congenital cytomegalovirus infection. *J Mother Child* 2021;**24**:9–17.
299. Kachramanoglou C, Jan W, Jones B, Papachatz E, Zombori L, Khan F, et al. Diagnostic analysis of baseline brain MRI features in infants with congenital cytomegalovirus infection: a simplified scoring system. *Clin Radiol* 2021;**76**:942.e7–e14.
300. Kakkar F, Renaud C, Valois S, Lapointe N, Lamarre V. Implementation of a CMV screening program for infants of HIV infected mothers by salivary PCR and/or culture: results from a single center. *Can J Infect Dis Med Microbiol* 2015;**26**:72B.
301. Karamchandani U, Ahmed U, Rufai SR, Tan N, Tan W, Petrushkin H, et al. Long-term ocular and visual outcomes following symptomatic and asymptomatic congenital CMV infection: a systematic review protocol. *BMJ Open* 2022;**12**:e059038.
302. Karimi A, Manaberi M, Azimi L, Taleghani NT, Shirvani F, Shariati MK, et al. Incidence and prognosis of congenital cytomegalovirus infection, a retrospective single-center experience, Tehran, Iran. *Arch Pediatr Infect Dis* 2023;**11**:e135767.
303. Kawano Y, Kawada J, Kamiya Y, Suzuki M, Torii Y, Kimura H, et al. Analysis of circulating human and viral microRNAs in patients with congenital cytomegalovirus infection. *J Perinatol* 2016;**36**:1101–5.
304. Keymeulen A, De Leenheer E, Casaer A, Cossey V, Herregods N, Laroche S, et al. Cranial ultrasound and MRI: complementary or not in the diagnostic assessment of children with congenital CMV infection? *Eur J Pediatr* 2022;**181**:911–20.
305. Khalil A, Scala C, D'Antonio F. *Prenatal risk factors for symptomatic congenital CMV infection*. PROSPERO; 2016. Available from: http://www.crd.york.ac.uk/PROSPERO/display_record.asp?ID=CRD42016045397 [accessed 19 November 2024].
306. Kim JH, Robles V, Weimer KED, Gehtland LM, Kucera KS. Improved dried blood spot PCR assay for universal congenital cytomegalovirus screening in newborns. *Microbiol Spectr* 2023:e0404122.
307. Kipfmüller F, Jungbluth K, Eis-Hubinger AM, Reber U, Holdenrieder S, Gembruch U, et al. Evaluation of cellulose pads as a method to detect cytomegalovirus DNA in neonatal urine. *Ann Clin Biochem* 2018;**55**:553–60.
308. Kitamura A, Toriyabe K, Hagimoto-Akasaka M, Ikejiri M, Minematsu T, Suga S, et al. Congenital cytomegalovirus infection and maternal primary cytomegalovirus infection in universal newborn hearing screening referral patients: a prospective cohort study. *Clin Exp Obstet Gynecol* 2022;**49**:259.
309. Kobayashi Y, Morioka I, Koda T, Nakamachi Y, Okazaki Y, Noguchi Y, et al. Low total IgM values and high cytomegalovirus loads in the blood of newborns with symptomatic congenital cytomegalovirus infection. *J Perinat Med* 2015;**43**:239–43.
310. Kohmer N, Nagel A, Berger A, Enders M, Hamprecht K, Korn K, et al. Laboratory diagnosis of congenital CMV infection in newborns: impact of pre-analytic factors. *J Clin Virol* 2019;**115**:32–6.
311. Koontz D, Baecher K, Amin M, Nikolova S, Gallagher M, Dollard S. Evaluation of DNA extraction methods for the detection of cytomegalovirus in dried blood spots. *J Clin Virol* 2015;**66**:95–9.

312. Korndewal MJ, Vossen AC, Cremer J, Van Binnendijk RS, Kroes AC, Van Der Sande MA, et al. Disease burden of congenital cytomegalovirus infection at school entry age: study design, participation rate and birth prevalence. *Epidemiol Infect* 2016;**144**:1520–7.
313. Kravchenko LL, Levkovich MA, Krukier II, Pyatikova MV, Zaurava LM, Afonin AA. Diagnosis of an infectious process in newborns born by mothers with chronic inflammatory gynaecological diseases. *Arch Dis Child* 2015;**100**:A186.
314. Kravchenko LV. [Prognosis of severe cytomegalovirus infection in newborns]. *Infektsiia Immun* 2021;**11**:745–51.
315. Krishna S, Nemerofsky SL, Iyare A, Ramdhanie MA, Nassar M, Nafday S. Early extended neonatal screening for congenital cytomegalovirus infection: a quality improvement initiative. *Jt Comm J Qual Patient Saf* 2020;**46**:516–23.
316. Kruc R. Does cranial ultrasound differentiate asymptomatic congenital CMV (cCMV) infection from cCMV with central nervous system disease in infants? Results from a universal screening study in Minnesota, USA. *J Pediatric Infect Dis Soc* 2022;**11**:S5.
317. Kyriakopoulou A, Papaevangelou V, Argyropoulou M, Papathanasiou M, Xydis V, Giorgi M, et al. Fetal brain imaging provides valuable information in cCMV infected infants. *J Matern Fetal Neonatal Med* 2023;**36**:2220564.
318. Lakhani A, Gie A, Rhode D, Mfingwana L, Parker N, Goussard P. Cytomegalovirus viral load as predictor of the clinical course of hypoxic pneumonia in children. *Int J Tuberc Lung Dis* 2023;**27**:49–54.
319. Lam T, Jones B, Basheer SN, Lyall H. Congenital cerebral cytomegalovirus infection: postnatal and paediatric MR imaging. *Dev Med Child Neurol* 2017;**59**:103–4.
320. Lee H, Kim M, Nam Y, Cho E, Yang J, Lee M, et al. Comparison of the three real-time CMV assay kits for the quantitation of cytomegalovirus DNA. *J Mol Diagn* 2013;**15**:888.
321. Leruez-Ville M, Ngini S, Guilleminot T, Kfutwah A, Moussa S, Tran T, et al. Detection of cytomegalovirus DNA on dried blood spots collected from infants infected with HIV: an in-house method adaptable in resource-limited settings. *J Virol Methods* 2013;**193**:503–7.
322. Leruez-Ville M, Guilleminot T, Stirnemann J, Salomon LJ, Spaggiari E, Faure-Bardon V, et al. Quantifying the burden of congenital cytomegalovirus infection with long-term sequelae in subsequent pregnancies of women seronegative at their first pregnancy. *Clin Infect Dis* 2020;**71**:1598–603.
323. Levit Y, Dym L, Yochpaz S, Manor Y, Adler A, Halutz O, et al. Assessment of risk indicators for targeted cytomegalovirus screening in neonates. *Neonatology* 2020;**117**:750–5.
324. Lewald ZS, Crisan R, Pifer T, De Alba Romero C, Shimamura M, Mejias A, et al. "False-Positive" cytomegalovirus (CMV) screening tests in neonates: a real dilemma! *Open Forum Infect Dis* 2023;**10**:S752–3.
325. Liao EN, Stephans J, Taketa E, Mohamad NI, Kaur Khalsa I, Naugle K, et al. Factors associated with congenital cytomegalovirus infection detected by dried blood spot testing in children with hearing loss. *Int J Pediatr Otorhinolaryngol* 2023;**165**:111446.
326. Lilleri D, Tassis B, Pugni L, Ronchi A, Pietrasanta C, Spinillo A, et al. Prevalence, outcome, and prevention of congenital cytomegalovirus infection in neonates born to women with preconception immunity (CHILd Study). *Clin Infect Dis* 2023;**76**:513–20.
327. Lim BG, Clark RH, Kelleher AS, Lin Z, Spitzer AR. Utility of genetic testing for the detection of late-onset hearing loss in neonates. *Am J Audiol* 2013;**22**:209–15.
328. Lipitz S, Elkan Miller T, Yinon Y, Weissbach T, De-Castro H, Hoffman C, et al. Revisiting short- and long-term outcome after fetal first-trimester primary cytomegalovirus infection in relation to prenatal imaging findings. *Ultrasound Obstet Gynecol* 2020;**56**:572–8.
329. Lopez AS, Lanzieri TM, Claussen AH, Vinson SS, Turcich MR, Iovino IR, et al. Intelligence and academic achievement with asymptomatic congenital cytomegalovirus infection. *Pediatrics* 2017;**140**.
330. Luck SE, Emery VC, Atkinson C, Sharland M, Griffiths PD. Compartmentalized dynamics of cytomegalovirus replication in treated congenital infection. *J Clin Virol* 2016;**82**:152–8.
331. Mack I, Burckhardt MA, Heininger U, Prufer F, Schulzke S, Wellmann S. Symptomatic congenital cytomegalovirus infection in children of seropositive women. *Front Pediatr* 2017;**5**:134.

332. Mahajan V, Patel J. Presentations of and testing practices for congenital CMV. *Pediatrics* 2018;**144**:674.
333. Maltezou PG, Kourlaba G, Kourkouni E, Luck S, Blázquez-Gamero D, Ville Y, et al. *Long-term sequelae in infants with congenital cytomegalovirus infection born to mothers with primary versus non-primary infection: a systematic review and meta-analysis*. PROSPERO; 2019. Available from: http://www.crd.york.ac.uk/PROSPERO/display_record.asp?ID=CRD42019125179 [accessed 19 November 2024].
334. Mappa I, De Vito M, Flacco ME, di Mascio D, D'Antonio F, Rizzo G. Prenatal predictors of adverse perinatal outcome in congenital cytomegalovirus infection: a retrospective multicenter study. *J Perinat Med* 2023;**51**:102–10.
335. Mareri A, Coclite E, Barone E, Gasparroni G, Iapadre G, Santini MS, et al. Subependymal and choroid plexus cysts as possible ultrasound markers of cerebropathy in congenital CMV infection. *J Perinat Med* 2013;**41**:515.
336. Martinez SAD, Elmubarak I, Tutaryebwa L, Princewill N, Predmore L, Russell B. Results of a protocol for targeted testing of congenital cytomegalovirus infection in babies who fail the newborn hearing screen. *Pediatrics* 2021;**149**:718.
337. Mastutik G, Kurniasari N, Rahniayu A, Rahaju AS, Ruslan SEN, Ilmiah K, et al. Detection of cytomegalovirus in urine specimen of cholestatic infants by polymerase chain reaction. *Res J Pharm Technol* 2022;**15**:2151–7.
338. Matsuo K, Morioka I, Oda M, Kobayashi Y, Nakamachi Y, Kawano S, et al. Quantitative evaluation of ventricular dilatation using computed tomography in infants with congenital cytomegalovirus infection. *Brain Dev* 2014;**36**:10–5.
339. McCrary HC, Newberry CI, Hamilton C, Nielson C, Calvo VD, Brown AC, et al. Implementation of a symptom-based cytomegalovirus screening protocol. *Otolaryngol Head Neck Surg* 2019;**161**:P285.
340. Medoro A, Malhotra P, Shimamura M, Hounam G, Findlen U, Wozniak P, et al. "Targeted" screening for cytomegalovirus (CMV)-related hearing loss: it's time for universal CMV screening in the NICU! *Open Forum Infect Dis* 2017;**4**:S691.
341. Medoro A, Shimamura M, Hanlon CT, Kaptan I, Malhotra MDS, Findlen UM, et al. Saliva screening for congenital cytomegalovirus infection in the neonatal intensive care unit: beware! *Open Forum Infect Dis* 2019;**6**:S801–2.
342. Medoro AK, Shimamura M, Malhotra MDS, Findlen UM, Hounam G, Hanlon CT, et al. Newborn dried blood spot for retrospective diagnosis of congenital cytomegalovirus (CMV) infection: it's time for universal screening! *Open Forum Infect Dis* 2019;**6**:S802.
343. Medoro AK, Hanlon CT, Pifer T, Escamilla MR, Shimamura M, Findlen UM, et al. Late-onset hearing loss and antiviral therapy for congenital cytomegalovirus infection. *Open Forum Infect Dis* 2020;**7**:S599.
344. Melamed R, Shemer-Avni Y, Shany E, Kurtzman L, Goral R, Landau D. Targeted and universal screen in term and preterm infants for congenital CMV infection. *Infect Dis (Lond)* 2020;**52**:730–5.
345. Meogrossi N, Calandra D, Di Sebastiano F, Pio Buca DI, Liberati M, D'Antonio F. Prenatal predictors of adverse perinatal outcome in congenital cytomegalovirus infection: a multicenter study. *Ital J Gynaecol Obstet* 2023;**35**:59.
346. Merino-Hernandez A, Sanchez-Barriopedro L, Villar-Castro S, Aguado-Del Hoyo A, Marsinyach-Ros I, Sanchez-Luna M. Cost-effectiveness of a cytomegalovirus screening strategy in neonates born after 34 weeks small for gestational age. *An Pediatr (Barc)* 2023;**98**:41–7.
347. Minami SB, Yamanobe Y, Nakano A, Sakamoto H, Masuda S, Takiguchi T, et al. A high risk of missing congenital cytomegalovirus-associated hearing loss through newborn hearing screening in Japan. *J Clin Med* 2021;**10**:29.
348. Mirsalehi N, Yavarian J, Ghavami N, Naseri M, Khodakhah F, Shatizadeh Malekshahi S, et al. Congenital cytomegalovirus infection in newborns suspected of congenital rubella syndrome in Iran: a cross-sectional study. *BMC Pediatrics* 2024;**24**:31.
349. Morimoto C, Nishikubo T, Nishimura T, Onishi T, Takeyama M, Uchida Y, et al. Late-onset and congenital hearing loss detected using AABR due to congenital cytomegalovirus infection that improved with valganciclovir. *Congenit Anom (Kyoto)* 2023;**63**:40–3.

350. Mu H, Qiao W, Zou J, Zhang H. Human cytomegalovirus glycoprotein B genotypic distributions and viral load in symptomatic infants. *J Infect Dev Ctries* 2023;**17**:1806–13.
351. Mujtaba G, Khurshid A, Sharif S, Alam MM, Aamir UB, Shaukat S, et al. Distribution of cytomegalovirus genotypes among neonates born to infected mothers in Islamabad, Pakistan. *PLoS ONE* 2016;**11**:e0156049.
352. Nagel A, Dimitrakopoulou E, Teig N, Kern P, Lucke T, Michna D, et al. Characterization of a universal screening approach for congenital CMV infection based on a highly-sensitive, quantitative, multiplex real-time PCR assay. *PLoS One* 2020;**15**:e0227143.
353. Namdeo D, Kumar P, Kumar A, Rai S, Thakar A, Deorari AK, et al. Cytomegalovirus infection and hearing loss in a highly seropositive rural population. *Indian J Virol* 2013;**24**:138.
354. Nishida K, Morioka I, Yamada H. Neurological outcomes in congenital cytomegalovirus-infected infants with small for gestational age. *J Reprod Immunol* 2016;**118**:119.
355. Nishida K, Morioka I, Nakamachi Y, Kobayashi Y, Imanishi T, Kawano S, et al. Neurological outcomes in symptomatic congenital cytomegalovirus-infected infants after introduction of newborn urine screening and antiviral treatment. *Brain Dev* 2016;**38**:209–16.
356. Niz Xavier PC, Goncalves Vieira P, de Souza Arantes T, Yano M, Martinelli Tavares LV, Duarte Miglioli AM, et al. Cytomegalovirus identification in blood and urine of newborns by nested polymerase chain reaction. *West Indian Med J* 2015;**65**:291–4.
357. Noorbakhsh S, Farhadi M, Haghighi F, Minaeian S, Hasanabad MH. Neonatal screening for congenital cytomegalovirus infection in Tehran, Iran, using Guthrie cards. *Iran J Microbiol* 2020;**12**:198–203.
358. Noorbakhsh S, Farhadi M, Minaeian S, Haghighi Hasanabad M. [Prevalence of congenital cytomegalovirus infection in newborns admitted to the intensive care units in Tehran, Iran]. *Tehran University Medical Journal* 2023;**81**:217–24.
359. Norton ME. Hearing loss with congenital cytomegalovirus infection. *Obstet Gynecol Surv* 2020;**75**:5–7.
360. Ohyama S, Fujioka K, Fukushima S, Abe S, Ashina M, Ikuta T, et al. Diagnostic value of cytomegalovirus IgM antibodies at birth in PCR-confirmed congenital cytomegalovirus infection. *Int J Mol Sci* 2019;**20**:3239.
361. Onoko M, Otieno NA, Dollard SC, Lanzieri TM. Neurodevelopmental outcomes of infants with congenital cytomegalovirus infection in Western Kenya. *J Clin Virol* 2020;**128**:104367.
362. Orb QT, Pesch M, Allen CM, Wilkes A, Ahmad I, Alfonso K, et al. Congenital cytomegalovirus testing outcomes from the ValEAR trial. *Otolaryngol Head Neck Surg* 2024;**170**:1430–41.
363. Palma S, Roversi MF, Bettini M, Mazzoni S, Pietrosemoli P, Lucaccioni L, et al. Hearing loss in children with congenital cytomegalovirus infection: an 11-year retrospective study based on laboratory database of a tertiary paediatric hospital. *Acta Otorhinolaryngol Ital* 2019;**39**:40–5.
364. Palma S, Forli F, Rossi C, Filice R, D'Adamo C, Roversi MF, et al. The audiological follow-up of children with symptomatic congenital cytomegalovirus infection: an experience in two Italian centers. *Children (Basel)* 2023;**10**:1136.
365. Paradowska E, Jablonska A, Studzinska M, Suski P, Kasztelewicz B, Zawilinska B, et al. Distribution of cytomegalovirus gN variants and associated clinical sequelae in infants. *J Clin Virol* 2013;**58**:271–5.
366. Paradowska E, Jablonska A, Plociennikowska A, Studzinska M, Suski P, Wisniewska-Ligier M, et al. Cytomegalovirus alpha-chemokine genotypes are associated with clinical manifestations in children with congenital or postnatal infections. *Virology* 2014;**462**–**463**:207–17.
367. Paradowska E, Jablonska A, Studzinska M, Kasztelewicz B, Wisniewska-Ligier M, Dzierzanowska-Fangrat K, et al. Distribution of the CMV glycoprotein gH/gL/gO and gH/gL/pUL128/pUL130/pUL131A complex variants and associated clinical manifestations in infants infected congenitally or postnatally. *Sci Rep* 2019;**9**:16352.
368. Parsons E, Teague K, Schaefer L, Hogue J, Kunz AN, Sainato RJ. Congenital CMV screening in infants with failed newborn hearing screens. *Pediatrics* 2021;**147**:1033–4.

369. Pathirana J, Groome M, Dorfman J, Kwatra G, Boppana S, Cutland C, et al. Prevalence of congenital cytomegalovirus infection and associated risk of in utero Human Immunodeficiency Virus (HIV) acquisition in a high-HIV prevalence setting, South Africa. *Clin Infect Dis* 2019;**69**:1789–96.
370. Pellegrinelli L, Galli C, Primache V, Alde M, Fagnani E, Di Berardino F, et al. Diagnosis of congenital CMV infection via DBS samples testing and neonatal hearing screening: an observational study in Italy. *BMC Infect Dis* 2019;**19**:652.
371. Peterson J, Nishimura C, Smith RJH. Genetic testing for congenital bilateral hearing loss in the context of targeted cytomegalovirus screening. *Laryngoscope* 2020;**130**:2714–8.
372. Pitlick MM, Orr K, Momany AM, McDonald EL, Murray JC, Ryckman KK. Determining the prevalence of cytomegalovirus infection in a cohort of preterm infants. *J Neonatal Perinatal Med* 2015;**8**:137–41.
373. Rahniayu A, Mastutik G, Rahaju AS, Ruslan SEN, Wiratama PA, Sulistiyani E, et al. Polymerase chain reaction of human cytomegalovirus from liver and urine compared with serological test in cholestasis infants. *J Infect Dev Ctries* 2022;**16**:1630–6.
374. Raynor E, Holmes C, Crowson M, Peskoe S, Planey A, Lantos PM. Loss to follow up of failed hearing screen and missed opportunities to detect congenital cytomegalovirus are better identified with the implementation of a new electronic health record system protocol. *Int J Pediatr Otorhinolaryngol* 2021;**148**:110818.
375. Reina J, Weber I, Riera E, Busquets M, Morales C. [Usefulness of a real-time quantitative polymerase-chain reaction (PCR) assay for the diagnosis of congenital and postnatal cytomegalovirus infection]. *An Pediatr (Barc)* 2014;**80**:299–303.
376. Renaud C, Smiljkovic M, Boucoiran I, Valois S, Tapiero B, Lamarre V, et al. Results of a targeted neonatal screening program for congenital cytomegalovirus infection in Montreal, Quebec. *Open Forum Infect Dis* 2018;**5**:S3–4.
377. Reynders M, Kerkhofs K, Heyndrickx A, Noerens K, Foulon I. Neurodevelopmental impact of congenital cytomegalovirus in children with cochlear implants. *Int J Pediatr Otorhinolaryngol* 2024;**180**:111939.
378. Ronchi A, Zeraf F, Lee LE, Owen KE, Shoup AG, Garcia F, et al. Evaluation of clinically asymptomatic high risk infants with congenital cytomegalovirus infection. *J Perinatol* 2020;**40**:89–96.
379. Roy D, Chatterjee A, Mishra L, Chakraborty N. Progression of retinal choroidal neovascularization by latent human cytomegalovirus infection and immunological signaling among neonatal patients admitted to tertiary care hospital. *J Med Virol* 2024;**96**:e29478.
380. Rymer-Haskel N, Barkai G, Duvdevani N, Weisz B, Lipitz S, Yinon Y. 651: Neurological outcome of congenital cytomegalovirus infection following primary maternal infection and a negative amniocentesis. *Am J Obstet Gynecol* 2019;**220**:S432.
381. Sahiner F, Cekmez F, Cetinkaya M, Kaya G, Kalayci T, Gunes O, et al. Congenital cytomegalovirus infections and glycoprotein B genotypes in live-born infants: a prevalence study in Turkey. *Infect Dis (Lond)* 2015;**47**:465–71.
382. Sakamoto A, Moriuchi H, Matsuzaki J, Motoyama K, Moriuchi M. Retrospective diagnosis of congenital cytomegalovirus infection in children with autism spectrum disorder but no other major neurologic deficit. *Brain Dev* 2015;**37**:200–5.
383. Sarkar A, Das D, Ansari S, Chatterjee RP, Mishra L, Basu B, et al. Genotypes of glycoprotein B gene among the Indian symptomatic neonates with congenital CMV infection. *BMC Pediatr* 2019;**19**:291.
384. Schleiss MR, Osterholm E, Hernandez-Alvarado N, Kruc R, Herd H, Sidebottom A, et al. Universal newborn testing for congenital cytomegalovirus (cCMV) infection comes of age: clinical sensitivity of screening tests and infant outcomes in a cCMV screening study in Minnesota. *Open Forum Infect Dis* 2023;**10**:S1264.
385. Schleiss MR, Panther L, Basnet S, Workneh M, Diaz-Decaro J. Comparison of overall sensitivity and specificity across different newborn screening algorithms for congenital cytomegalovirus. *Int J Neonatal Screen* 2023;**9**:33.
386. Shahar-Nissan K, Oikawa Tepperberg M, Mendelson E, Bilavsky E. Retrospective identification of congenital cytomegalovirus infection using dried blood samples - missed opportunities and lessons. *J Clin Virol* 2022;**152**:105186.

387. Shahrokhi N, Shakeri TS, Shirvani F, Seifi K. Rapid detection of cytomegalovirus and listeria monocytogenes infection in symptomatic newborns by fluorescence end-point detection PCR (FEP PCR). *Journal of Comprehensive Pediatrics* 2020;**11**:e86340.
388. Shears A, Yan G, Mortimer H, Cross E, Sapuan S, Kadambari S, et al. Vestibular and balance dysfunction in children with congenital CMV: a systematic review. *Arch Dis Child Fetal Neonatal Ed* 2022;**107**:630–6.
389. Smiechura M, Struzycka M, Konopka W. Congenital and acquired cytomegalovirus infection and hearing evaluation in children. *Otolaryngol Pol* 2014;**68**:303–7.
390. Smiljkovic M, Renaud C, Tapiero B, Lamarre V, Kakkar F. Head ultrasound or MRI? The role of neuroimaging in the assessment of symptomatic and asymptomatic infants with congenital CMV infection. *Open Forum Infect Dis* 2017;**4**:S691.
391. Smiljkovic M, Renaud C, Tapiero B, Lamarre V, Kakkar F. Head ultrasound, CT or MRI? The choice of neuroimaging in the assessment of infants with congenital cytomegalovirus infection. *BMC Pediatr* 2019;**19**:180.
392. Smiljkovic M, Le Meur JB, Malette B, Boucoiran I, Minsart AF, Lamarre V, et al. Blood viral load in the diagnostic workup of congenital cytomegalovirus infection. *J Clin Virol* 2020;**122**:104231.
393. Smyrli A, Raveendran V, Walter S, Pagarkar W, Field N, Kadambari S, et al. *Neurodevelopmental outcomes of children with asymptomatic congenital cytomegalovirus infection at birth: a systematic literature review*. PROSPERO; 2022. Available from: http://www.crd.york.ac.uk/PROSPERO/display_record.asp?ID=CRD42022331535 [accessed 19 November 2024].
394. Southern Illinois University. *Screening for congenital cytomegalovirus infection in newborns (CMV)*. ClinicalTrials.gov. Bethesda (MD): National Library of Medicine (US); 2016. Available from: <https://clinicaltrials.gov/study/NCT02683096> [accessed 20 November 2024].
395. Spalletti-Cernia D, Barbato S, Sorrentino R, Vallefucio L, Rocco C, Di Costanzo P, et al. Evaluation of the automated QIA Symphony SP/AS workflow for cytomegalovirus DNA extraction and amplification from dried blood spots. *Intervirology* 2016;**59**:211–6.
396. Ssentongo P, Hehnly C, Birungi P, Roach MA, Spady J, Fronterre C, et al. Congenital cytomegalovirus infection burden and epidemiologic risk factors in countries with universal screening: a systematic review and meta-analysis. *JAMA Netw Open* 2021;**4**:e2120736.
397. Stoyell SM, Elison JT, Graupmann E, Miller NC, Emerick J, Ramey E, et al. Neurobehavioral outcomes of neonatal asymptomatic congenital cytomegalovirus infection at 12-months. *J Neurodev Disord* 2024;**16**:19.
398. Stoykova Z, Ivanova L, Yordanova D, Cvetkova S, Kalchev K. Relevance of CMV DNA detection in symptomatic children up to 3 months of age. *C R Acad Bulg Sci* 2022;**75**:1195–201.
399. Suarez D, Nielson C, McVicar SB, Sidesinger M, Ostrander B, O'Brien E, et al. Analysis of an expanded targeted early cytomegalovirus testing program. *Otolaryngol Head Neck Surg* 2023;**169**:679–86.
400. Thompson CR, Cohen HL, Meals E, Tomlinson R, Harrison L, Lane C, et al. Frequency and severity of cranial ultrasound abnormalities in a cohort of newborns molecularly screened for congenital cytomegalovirus infection. *J Investig Med* 2020;**68**:541.
401. Torii Y, Suzuki T, Fukuda Y, Haruta K, Yamaguchi M, Horiba K, et al. MicroRNA expression profiling of urine exosomes in children with congenital cytomegalovirus infection. *Sci Rep* 2024;**14**:5475.
402. Torre P, Zeldow B, Yao TJ, Hoffman HJ, Siberry GK, Purswani MU, et al. Newborn hearing screenings in human immunodeficiency virus-exposed uninfected infants. *J AIDS Immune Res* 2016;**1**:102.
403. Turner KM, Lee HC, Boppana SB, Carlo WA, Randolph DA. Incidence and impact of CMV infection in very low birth weight infants. *Pediatrics* 2014;**133**:e609–15.
404. Uematsu M, Haginoya K, Kikuchi A, Hino-Fukuyo N, Ishii K, Shiihara T, et al. Asymptomatic congenital cytomegalovirus infection with neurological sequelae: a retrospective study using umbilical cord. *Brain Dev* 2016;**38**:819–26.

405. Vande Walle C, Keymeulen A, Schiettecatte E, Acke F, Dhooge I, Smets K, et al. Brain MRI findings in newborns with congenital cytomegalovirus infection: results from a large cohort study. *Eur Radiol* 2021;**31**:8001–10.
406. Vargas J, Schiavo V, Medina G, Espinosa C, Randis T, Balakrishnan M. Targeted cytomegalovirus screening in a level IV neonatal intensive care unit. *J Investig Med* 2022;**70**:717.
407. Viswanathan R, Bafna S, Mergu R, Deshpande G, Gunjekar R, Gaikwad S, et al. Direct saliva real-time polymerase chain reaction assay shows low birth prevalence of congenital cytomegalovirus infection in urban Western India. *Pediatr Infect Dis J* 2019;**38**:e65–8.
408. Vives-Onos I, Soler-Palacin P, Codina-Grau MG, Martin-Nalda A, Lopez-Galera RM, Marin-Soria JL, et al. [Can we rule out a congenital cytomegalovirus infection when the result of polymerase chain reaction in dried blood spots is negative?]. *Enferm Infecc Microbiol Clin* 2014;**32**:570–3.
409. Wang G, Feng D. Dynamic relationship between infantile hepatitis syndrome and cytomegalovirus infection. *Exp Ther Med* 2017;**13**:3443–7.
410. Wang Y, Geng M, Zhang H, Ping K. [Effects of cytomegalovirus infection on infants' hearing and speech development]. *Lin Chuang Er Bi Yan Hou Tou Jing Wai Ke Za Zhi* 2022;**36**:163–6.
411. Webb E, Gillespie AN, Poulakis Z, Gartland T, Buttery J, Casalaz D, et al. Feasibility and acceptability of targeted salivary cytomegalovirus screening through universal newborn hearing screening. *J Paediatr Child Health* 2022;**58**:288–94.
412. Williams EJ, Gray J, Luck S, Atkinson C, Embleton ND, Kadambari S, et al. First estimates of the potential cost and cost saving of protecting childhood hearing from damage caused by congenital CMV infection. *Arch Dis Child Fetal Neonatal Ed* 2015;**100**:F501–6.
413. Wissel M, Demmler G, Gandaria C, Cravens S, Berg A, Widen A, et al. Detection of cytomegalovirus (CMV) in saliva of congenitally infected neonates. *Open Forum Infect Dis* 2017;**4**:S357.
414. Pooled saliva testing for newborns enhances early detection of congenital cytomegalovirus. *Infection Control Today* 2024;**28**:13–4.
415. Wu PH, Zhou Y, Wu KQ, Yin BB, Zhu B. [Correlation between serum IgM antibody and viral load with clinical symptoms in neonates infected with cytomegalovirus]. *Zhonghua Yu Fang Yi Xue Za Zhi* 2022;**56**:1642–7.
416. Wu PH, Lee CY, Huang JY, Yang SF, Shih CP. The correlation between neonatal parameters and late-onset inner ear disorders in congenital cytomegalovirus infection: a 10-year population-based cohort study. *Clin Otolaryngol* 2022;**47**:107–14.
417. Wujcicka W, Wilczynski J, Paradowska E, Studzinska M, Nowakowska D. The role of single nucleotide polymorphisms, contained in proinflammatory cytokine genes, in the development of congenital infection with human cytomegalovirus in fetuses and neonates. *Microb Pathog* 2017;**105**:106–16.
418. Yamaguchi M, Suzuki T, Kidokoro H, Iwada KI, Fukuda Y, Haruta K, et al. Identification of plasma biomarkers of neurological complications in patients with congenital cytomegalovirus infection by proteome analysis. *Open Forum Infect Dis* 2023;**10**:S752.
419. Yamaguchi M, Suzuki T, Kidokoro H, Iwata KI, Fukuda Y, Haruta K, et al. Proteomic analysis reveals novel plasma biomarkers for neurological complications in patients with congenital cytomegalovirus infection. *J Pediatric Infect Dis Soc* 2023;**12**:525–33.
420. Yang TH, Huang HM, Hsu WC, Tsao PN, Liu TC, Hsu CJ, et al. The prevalence and demographic features of congenital cytomegalovirus infection in an urban area of East Asia: a population-based study. *PLoS One* 2021;**16**:e0248801.
421. Yao HT. Sero-epidemiological survey of cytomegalovirus infection among pregnant women in Taiwan. *Int J Gynaecol Obstet* 2018;**143**:708.
422. Zavattoni M, Furione M, Arossa A, Iasci A, Spinillo A, Lombardi G, et al. Diagnosis and counseling of fetal and neonatal HCMV infection. *Early Hum Dev* 2014;**90**:S29–31.
423. Zavattoni M, Lombardi G, Garofoli F, Scalia G, Rizzo A, Angelini M, et al. Neonatal HCMV-related polymicrogyria in seroimmune women: what is the optimal pregnancy management? *J Clin Virol* 2018;**108**:141–6.

424. Zeng ZC, Chang Q, Sun ZW, Song MM, Jin XL, Jiang SY, et al. Detection of cytomegalovirus (CMV) infection in wheezing infants by urine DNA and serum IgG testing. *Med Sci Monit* 2017;**23**:1242–6.
425. Zhang SX, He XZ, Wang SW, Wang XF. [The comparison of two newborn cytomegalovirus IgG antibody screening ELISA kits]. *Zhonghua Shi Yan He Lin Chuang Bing Du Xue Za Zhi* 2013;**27**:392–4.
426. Zhang MJ, Yuan TM, Wang LZ. [Risk factors for hearing impairment induced by cytomegalovirus infection]. *Zhongguo Dang Dai Er Ke Za Zhi* 2016;**18**:224–8.
427. Zheng B, Liu HY, Shen R, Li XX. *Diagnostic test accuracy of PCR by saliva specimen for cytomegalovirus infection in newborn: systematic review and meta-analysis*. PROSPERO; 2022. Available from: http://www.crd.york.ac.uk/PROSPERO/display_record.asp?ID=CRD42022337717 [accessed 19 November 2024].
428. Stagno S, Pass RF, Cloud G, Britt WJ, Henderson RE, Walton PD, et al. Primary cytomegalovirus infection in pregnancy. Incidence, transmission to fetus, and clinical outcome. *JAMA* 1986;**256**:1904–8.
429. Lipitz S, Hoffmann C, Feldman B, Tepperberg-Dikawa M, Schiff E, Weisz B. Value of prenatal ultrasound and magnetic resonance imaging in assessment of congenital primary cytomegalovirus infection. *Ultrasound Obstet Gynecol* 2010;**36**:709–17.
430. Abrahao N, Rodrigues V, Castilho A. *The congenital cytomegalovirus infection screening by urine: it is worthy? A systematic review*. PROSPERO; 2022. Available from: <http://www.crd.york.ac.uk/PROSPERO/view/CRD42022326424> [accessed 19 November 2024].
431. Wu L, Hu J, Wang Z, Deng H, Chen Y, Wang B. *Meta-analysis of HCMV viral load level and the occurrence and outcome of hearing impairment*. PROSPERO; 2022. Available from: <https://www.crd.york.ac.uk/PROSPERO/view/CRD42022301633> [accessed 19 November 2024].
432. Aira da Silva Lopes I, de Santana Bispo Melo A, Saquetto MB, Neto MG, Moreira LMO. *Impact of congenital cytomegalovirus infection on motor development in children: systematic review*. PROSPERO; 2023. Available from: <https://www.crd.york.ac.uk/PROSPERO/view/CRD42023404868> [accessed 19 November 2024].
433. Shetty JK, Rady SM, Alameddine M, Abdelmoati MI. *Evaluation of salivary sample in diagnosis, screening and management of cytomegalovirus (CMV) infection in newborns-systematic review and metanalysis*. PROSPERO; 2024. Available from: <http://www.crd.york.ac.uk/PROSPERO/view/CRD42024603785> [accessed 19 November 2024].
434. Wang M, Wang X, Zhang W, Xiong Y, Chen M, Zhao J, et al. *Accuracy and implementability of dried blood spot polymerase chain reaction assays for screening congenital cytomegalovirus infection: systematic review and meta-analysis*. PROSPERO; 2024. Available from: http://www.crd.york.ac.uk/PROSPERO/display_record.asp?ID=CRD42023397836 [accessed 19 November 2024].
435. Assistance Publique - Hôpitaux de Paris. *Prognostic value of neonatal markers for the development of neurosensory sequelae in children infected by cytomegalovirus in utero (CYMEPEDIA)*. ClinicalTrials.gov. Bethesda (MD): National Library of Medicine (US); 2013. Available from: <https://clinicaltrials.gov/study/NCT01923636> [accessed 20 November 2024].
436. Academisch Medisch Centrum. *Markers for neurocognitive impairment in congenitally CMV infected infants: MARVELYS study*. Overview of Medical Research in the Netherlands (OMON). The Hague: Central Committee on Research Involving Human Subjects (CCMO); 2016. Available from: <https://onderzoekmetmensen.nl/en/trial/43495> [accessed 20 November 2024].
437. Kuthubutheen J. *Targeted screening for congenital cytomegalovirus-related hearing loss in infants in Western Australia*. Australian New Zealand Clinical Trials Registry; 2021. Available from: <https://anzctr.org.au/ACTRN12621000484842.aspx> [accessed 2 January 2025].

438. Brophy J, Moazin Z, Gallager L, Kernohan K, Lacaria M, O'Sullivan C, et al. *Congenital CMV AND HEARING in Ontario: Optimizing screening to improve child health outcomes (CAN HEAR ONTARIO)*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 54.
439. Churyukina E, Kravchenko L, Levkovich M, Afonin A, Krukier I, Berezhanskaya S, et al. Predictive value of cytokines for preservation neurologic semiology in neonatal cytomegalovirus infection. *Allergy* 2023;**78**:254.
440. Del Valle Penella AP, Rochat RH, Miller JA, Demmler-Harrison GJ. Performance of Dried Blood Spot (DBS) for the diagnosis of congenital cytomegalovirus (cCMV) in a single center: a real-world experience. *Open Forum Infect Dis* 2022;**9**:S860–1.
441. Kakkar F, Govender K, Renaud C, Gantt S, Lamarre V, Boucoiran I, et al. *Feasibility of dried urine on filter paper for universal congenital CMV screening*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 26.
442. Kim J, Wang H, Medoro AK, Pifer T, Crisan R, Leber A, et al. CMV PCR agreement between blood and dried blood spots in infants with congenital cytomegalovirus infection. *Open Forum Infect Dis* 2023;**10**:S43.
443. Kruc R, Osterholm EA, Hernandez-Alvarado N, Rosendahl S, McCann M, Sidebottom A, et al. Does cytomegalovirus (CMV) viral load correlate with disease severity in the setting of congenital CMV (cCMV) infection? Results from a universal cCMV screening study. *J Pediatric Infect Dis Soc* 2021;**21**:S13–4.
444. Lazzarotto T. Pre-and post-natal diagnosis of congenital cytomegalovirus infection. *Biochim Clin* 2013;**37**:S55.
445. Rohde B, Carillo-Marquez M, Tomlinson R, Blanch J, Kim-Hoehamer YI, Patel H, et al. Analytical and clinical performance of a real-time screening PCR assay identifying congenital CMV infection. *Open Forum Infect Dis* 2017;**4**:S356.
446. Alarcón A, de Vries LS, Arnáez J, Noguera-Julian A, Ríos-Barnés M, Fortuny C, et al. *Isolated white matter abnormalities at birth as prognostic markers in congenital cytomegalovirus infection*. In: European Congenital Cytomegalovirus Initiative (ECCI), 20-21 October 2022. Athens, Greece; 2022. p. 16–7.
447. Boscolo L, Spadavecchia A, Garnerio A, Leone A, Morana G, Coscia A. *Correlation between neurodevelopmental outcomes and brain MRI severity score in patients with congenital cytomegalovirus infection: a monocentric retrospective study*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 31.
448. Buonsenso D, Pedrero Tomé R, Raimondi F, Salomé S, Papaevangelou V, Syridou G, et al. *Prognostic factors of late onset hearing loss in infants with congenital cytomegalovirus and normal audiological assessment at birth*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 73.
449. Castells Vilella L, Diez R, Vilas J, Olabarrieta I, Medina A, Galan P, et al. Risk factors associated with hearing loss and neurologic impairment in the Spanish network of infants with congenital cytomegalovirus infection (REDICCMV). *J Matern Fetal Neonatal Med* 2016;**29**:33.
450. Chung PK, Schornagel F, Goeman J, Vossen A. *Predicting outcome in clinically inapparent congenital cytomegalovirus infection with hearing loss*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 61.
451. Churyukina EV, Kravchenko LV, Levkovich MA, Afonin AA, Berezhanskaya SB, Panova IV. Early predictors of preservation neurologic semiology in neonatal cytomegalovirus infection. *Allergy* 2021;**76**:381.
452. Czech-Kowalska J, Jedlińska-Pijanowska D, Kasztelewicz B, Pietrzyk A, Sokołowska M, Mesjasz M. *Predictors of sensorineural hearing loss in newborns with congenital CMV infection - a single-center experience*. In: European Congenital Cytomegalovirus Initiative (ECCI), 20-21 October 2022. Athens, Greece; 2022. p. 54.
453. Loizou CE, Karagiannidou S, Syridou G, Korres G, Papaevangelou V. *Vestibular dysfunction in asymptomatic children with congenital cytomegalovirus infection*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 71.

454. Faure-Bardon V, Stirneman J, Cornet N, Magny J, Nicloux M, Leruez-Ville M, et al. OC27.01 Neonatal outcome and long-term sequelae in CMV infected neonates according to the timing of primary maternal infection. *Ultrasound Obstet Gynecol* 2018;**52**:62.
455. Fernández-Rueda M, Vergas Gutierrez JJ, García Fernández A, Pedrero Tomé R, Blazquez Gamero D. *Risk factors associated with hearing loss in children with congenital cytomegalovirus infection*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 63.
456. Fourgeaud J, Vauloup-Fellous C, Bourgon N, Portet-Sulla V, Veyrenche N, Magny JF, et al. *Sequelae of congenital cytomegalovirus following maternal primary infections are limited to those acquired before 12 weeks*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 30.
457. Karagiannidou S, Syridou G, Loizou CE, Karapatoglou A, Nikolopoulos T, Papaevangelou V. *Delayed onset hearing loss (DOHL) in asymptomatic newborns with congenital cytomegalovirus infection (ccmv)*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 69.
458. Kravchenko L, Levkovich M, Zaurava L, Demidova M, Afonin A. The outcome of cerebral pathology by the end of the first year of life in newborn babies with cytomegalovirus infection. *J Matern Fetal Neonatal Med* 2014;**27**:224–5.
459. Kravchenko LV, Afonin AA. Prognosis for the preservation of neurological symptoms by the end of the first year of life in newborn babies, who had cytomegalovirus infection. *Arch Dis Child* 2016;**101**:A39.
460. Medoro AK, Dhital R, Flint K, Graber B, Pifer T, Honegger J, et al. Neonatal immune phenotypes and neurodevelopmental outcomes in infants with congenital cytomegalovirus infection (cCMV). *Open Forum Infect Dis* 2022;**9**:S859–S60.
461. Morioka I, Fukushima S, Ohyama S, Nishida K, Yamana K, Fujioka K, et al. Neurological outcomes at 18 months of age and neonatal risk factors for severe sequelae in congenitally cytomegalovirus-infected infants treated by antiviral treatment early in life. *J Reprod Immunol* 2017;**124**:80.
462. Nosan G, Hovnik T, Trebušak Podkrajšek K, Paro Panjan D. *Genetic variations in toll-like receptor 2 gene in Slovenian newborns with congenital human cytomegalovirus infection*. In: European Congenital Cytomegalovirus Initiative (ECCI), 20-21 October 2022. Athens, Greece; 2022. p. 60.
463. Salomè S, Gammella R, Coppla C, Dolce P, Blázquez-Gamero D, Capasso L, et al. *Can viral load predict a symptomatic congenital CMV infection? A systematic review and meta-analysis*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 79.
464. Carmona AS, Kakkar F, Gantt S. Perinatal cytomegalovirus infection. *Curr Treat Options Pediatr* 2022;**8**:395–411.
465. Smiljkovic M, Renaud C, Leduc D, Boucoiran I, Lamarre V, Kakkar F. Use of quantitative polymerase chain reaction in blood and cerebrospinal fluid as a predictor of severity in congenital cytomegalovirus infection. *Open Forum Infect Dis* 2016;**1**:137.
466. Smithers-Sheedy H, Raynes-Greenow C, Badawi N, Gibson C, Reid S, Meehan E, et al. Cerebral palsy attributed to congenital cytomegalovirus infection is associated with spastic quadriplegia and female sex. *Dev Med Child Neurol* 2014;**56**:29–30.
467. Vanwinkel S, De Keersmaecker B, Devlieger R, Naulaers G, De Catte L. Congenitally infected CMV fetuses following first trimester maternal infection: neonatal and shortterm outcome. *J Matern Fetal Neonatal Med* 2016;**29**:125.
468. Villaverde S, Villegas C, Pedrero Tomé R, Ríos M, Noguera Julian A, Fortuny C, et al. *Association between congenital cytomegalovirus infection and autism spectrum disorder*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 35–6.
469. Vossen ACTM, Korndewal MJ, Oudesluys-Murphy AM, Kroes ACM, van der Sande MAB, De Melker HE. Long-term impairment attributable to congenital cytomegalovirus infection. *J Clin Virol* 2016;**82**:S2–S3.
470. Carrillo-Marquez M, Rohde B, Tomlinson R, Blanch J, Bradford L, Kim-Hoehamer YI, et al. Programmatic congenital CMV universal screening program. *Open Forum Infect Dis* 2017;**4**:S23–4.

471. Landers D, Osterholm E, Herd H, Huang T, Hernandez-Alvarado N, Kruc R, et al. Analysis of dried blood spot and saliva PCR screening for detection of congenital cytomegalovirus infection informs implementation of a novel universal newborn screening program. *Am J Obstet Gynecol* 2024;**230**:S630–1.
472. Lund S, Bille M, Jordan K, Hansen B, Nygaard U. *Targeted screening for congenital CMV in the capital region of Denmark*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 58.
473. Penton M, Herzog A, Pan J, Manopla C, Otto C, Kohlhoff S. Adherence to follow-up and CMV testing in infants who failed newborn hearing screens: evaluation of new protocol to ensure follow-up and testing. *Open Forum Infect Dis* 2019;**6**:S802.
474. Syridou G, Mavridi A, Douros K, Giorgi M, Fryganas A, Siafakas N, et al. *Diagnostic ability of targeted and universal neonatal screening for cCMV-infection, to provide timely diagnosis of cCMV-infection associated neurodevelopmental delay*. In: European Congenital Cytomegalovirus Initiative (ECCI), 20-21 October 2022. Athens, Greece; 2022. p. 20–1.
475. Webb E, Hodgson J, Gillespie A, Jones CA, Poulakis Z, Wong J, et al. *Exploring parents' experiences of completing targeted congenital cytomegalovirus screening during their infant's newborn hearing screening*. In: European Congenital Cytomegalovirus Initiative (ECCI), 9-11 October 2024. Leiden, The Netherlands; 2024. p. 55.