Congenital cytomegalovirus — time to diagnosis, management and clinical sequelae in Australia: opportunities for earlier identification

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ABSTRACT

Objectives: To report on the burden of disease in Australian infants with congenital cytomegalovirus (cCMV) infection in the era of neonatal hearing screening and improved diagnostic techniques.

Design, setting and participants: National data were collected from across Australia via the Australian Paediatric Surveillance Unit (APSU) with monthly reporting by > 1000 clinicians between January 1999 and February 2009. For each reported case, data on investigations and epidemiological and clinical features were analysed. Detailed clinical reviews were performed on 42 infants in two Sydney tertiary paediatric infectious diseases clinics.

Results: There were 195 infants with cCMV identified, including 126 definite and 69 probable cases. Of these, 175 (90%) were symptomatic and only 15 were treated with antiviral agents. Identification was delayed beyond 60 days of age in 30 cases (15%). During the period of study, neonatal hearing screening was introduced for most Australian infants. Detection of hearing loss increased from 19% of cCMV cases in 1999–2003 to 31% in 2004–2009. Of 42 infants whose cases were reviewed in detail, 33 (79%) had symptomatic disease. DNA detection of CMV, using polymerase chain reaction testing of newborn screening cards, was useful in retrospective identification, and was strongly correlated with the presence of clinical sequelae (15/18; 83%).

Conclusions: Congenital CMV is underdiagnosed, infrequently treated, and often manifests as isolated hearing loss. Delayed diagnoses both before and after the introduction of neonatal hearing screening represent missed treatment and management opportunities and are likely to lead to poorer, life-long outcomes for these children. Retrospective analysis of newborn screening cards for CMV should be undertaken for infants with sensorineural hearing loss, to identify unrecognised cCMV.

METHODS

APSU case series

Notifications of possible cases of cCMV were received through the APSU between January 1999 and February 2009. The APSU is an ongoing, active surveillance system that gathers national data on conditions affecting children, with over 1300 participating doctors, mainly paediatricians, now contributing.

Participating doctors are sent a monthly email or report card to record any new cases of the conditions listed on the card. These are returned to the APSU, and we are notified of reported cases and the contact details of the reporting doctor. We then ask doctors to provide data about the mothers and infants concerned, using de-identified questionnaires. Before the start of the national surveillance, participating doctors were sent an outline of the study and a protocol for reporting cases. Cases of cCMV were defined as follows.

• A definite case of cCMV (as previously described6) was defined as a child whom...
CMV was identified in the first 3 weeks of life, from urine, blood, saliva or any tissue taken by biopsy:

- A suspected case was defined as any child up to 12 months of age, from whom CMV was isolated, or who was positive for CMV IgM and in whom clinical features were suggestive of intrauterine CMV infection.6

### Clinical case review

We examined additional data from a subset of 42 of the infants: 22 who were referred to the Congenital Infections Clinic at Sydney Children's Hospital, and 20 who were referred to the Congenital and Perinatal Infection Clinic at the Children's Hospital at Westmead. Cases were referred to these clinics either because of maternal CMV infection during pregnancy or neonatal symptoms with subsequent confirmation of cCMV infection, using the above case definition. The investigators reviewed the infants' symptoms at identification and at follow-up, from available information at clinical review and from the APSU study enrolment form.

### Virological and serological methods

Cytomegalovirus was detected using cell culture, shell vial assays (direct immunofluorescence identification of immediate early CMV antigen) or by histopathological examination of tissue specimens.8 Serum IgM antibody to CMV was determined using an IgM capture enzyme immunoassay (Dia-Sorin, Saluggia, Italy). Additional infants with cCMV were identified as a result of another study on the seroprevalence of CMV among pregnant women and women identified with documented antenatally acquired CMV infection.8

The extraction of CMV DNA from newborn screening cards was performed as previously reported11 with few modifications.8 Three disks of 3 mm diameter were prepared from newborn screening cards using a Wal- lac DBS Puncher (product number 1296-071) (PerkinElmer, Turku, Finland). Blood was eluted from the newborn screening cards by incubating in 45 mL of minimum essential media at 55°C for 60 minutes, followed by heating at 100°C for 7 minutes. Samples were rapidly cooled, centrifuged at 10 000 g for 3 minutes and frozen at −80°C for at least 1 hour before further testing.

For extraction of CMV DNA from urine samples, a MagNA Pure semi-automated nucleic acid extraction machine (Roche Applied Science, Mannheim, Germany) was used. The samples were stored at −20°C before polymerase chain reaction (PCR) testing for the presence of the CMV glycoprotein gene gp58, and for the major immediate gene.8,12

For statistical analysis, comparisons between lines of diagnosis were performed using two-by-two table $\chi^2$ analysis. Comparisons between time periods were done using Yate's $\chi^2$ analysis.

Ethics approval was granted by the human research ethics committees of the Northern Hospital Network, South Eastern Sydney Illawarra Area Health Service, and the Royal Alexandra Hospital for Children.

### RESULTS

#### APSU case series

There were 363 notifications of cCMV to the APSU from January 1999 to February 2009, and 237 (65%) follow-up questionnaires were returned by doctors. Of these, 195 were classified as definite or probable cCMV, using the case definition. The notifications were from across Australia, with a majority from New South Wales, which has about one-third of all Australian births13 (Box 2). There were 126 definite cases (65%) and 69 probable cases (35%); 104 (53%) were male, 88 (45%) were female, and in three cases (2%), sex was not stated on the report. There were two neonatal deaths (at ages 1 day and 7 days) and one stillbirth. Reported gestational age ranged from 27 to 40 weeks.

In total, 158 infants (81%) had suspected cCMV before age 60 days: 27 cases (14%) followed referral for antenatally diagnosed maternal CMV infection, 61 cases (31%) were suspected at birth and 70 cases (36%) were suspected between birth and 60 days of age. In 16 infants (8%), cCMV was identified between age 60 days and 1 year, and in 12 (6%), the identification was made after 1 year of age. For nine cases (5%), the age when cCMV was first suspected was not stated in the report. Of the total case series, 20 infants (10%) were asymptomatic, and the remaining 175 (90%) had symptoms (Box 3). Of a total of 175 symptomatic cases, 60 (34%) were male, 95 (54%) were female, and in three cases (2%), sex was not stated on the report. There were two neonatal deaths (at ages 1 day and 7 days) and one stillbirth. Reported gestational age ranged from 27 to 40 weeks.

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The clinical and laboratory features of cases (Box 3) were similar in definite and probable cases (data not shown). Nineteen infants had cataracts or microphthalmia reported as features of symptomatic cCMV. Most features of cCMV were evident within the first 60 days of life. Sensorineural hearing loss (SNHL) was present in 51 infants (26%) and was the sole presenting feature in 34 infants, two-thirds of those with SNHL overall. In 15 cases, SNHL was diagnosed after age 60 days. Twelve of these 15 (80%) were identified retrospectively by detection of CMV PCR on their newborn screening cards,9 or from stored cord blood samples. In total, 38 infants from this group had positive
CMV PCR results via testing of stored newborn screening cards or from stored blood samples. Box 1 shows cases of cCMV by year of birth, excluding four confirmed cases born before 1999, identified using CMV PCR on stored newborn screening cards. The overall proportion of symptomatic to total cases was not significantly different between the two study periods: 71/80 (89%) in 1999–2003 (when screening was partially introduced) and 105/115 (91%) in 2004–2009 (when screening was substantially introduced). SNHL was diagnosed in 36 of 115 cases (31%) in 2004–2009, compared with 15 of 80 cases (19%) in 1999–2003. This approached but did not reach significance (difference, 12%; 95% CI, −0.1% to 24.1%). Analysis by state (not shown) revealed more cases identified before and after age 60 days after the introduction of neonatal hearing screening programs. There were 94 cases identified in the first half of the study (1999–2003) compared with 67 cases in the second half of the study (2004–2009) (of 161 cases where location and age of identification were available), but this also failed to attain significance. Of the 28 total cases identified after 60 days of age, 16 (57%) were identified in states after introduction of neonatal hearing screening programs, seven (25%) before introduction of such programs and five (18%) were from unknown locations. Intracranial calcification is also included in Box 1 as a recognisable radiographic predictor of poor outcome, which did not change significantly during the study.

Clinical follow-up of a subset of infants
Forty-two infants were referred to two perinatal infection clinics and followed to a median age of 24 months (range, 1–108 months). In total, 18/42 infants (43%) had hearing loss and 17/42 (40%) had neurodevelopmental sequelae. Twelve cases (29%) had abnormal central nervous system imaging, and 9/12 (75%) had developmental delay. Of the 3/12 (25%) without developmental problems identified, two had hearing loss. Of 31 infants identified in their first 60 days of life, nine were referred because of an antenatal diagnosis of maternal CMV and 22 infants were referred because of symptoms. Details of clinical features are shown in Box 4. All 11 infants and children identified at over 60 days of age were referred because of symptomatic CMV disease. SNHL was present in 7/11 (64%) and was the sole feature of CMV disease for three of the 42 infants (7%). SNHL was identified after the newborn period in four of 11 infants (three born before the introduction of hearing screening).

We were able to examine 23/42 infants using CMV PCR. Of 18 infants with a positive CMV PCR test result, 15/18 (83%) had signs of cCMV disease and 3/18 (17%) were asymptomatic. Ten of 18 (56%) had neurodevelopmental sequelae, and 7/18 (39%) had SNHL. Of those with negative PCR test results, 2/5 (40%) were symptomatic, one with isolated retinitis and one with isolated SNHL. This equated to a sensitivity of 88%, specificity of 50% and positive likelihood ratio of 1.76 for symptomatic disease with a positive CMV PCR test result. Overall, identification of cCMV was confirmed in five infants.
cases with retrospective PCR using stored newborn screening cards.

**DISCUSSION**

We have found that congenital cytomegalovirus infection is underreported in Australia. Based on a live birth rate of 285,000 per year in Australia, and a CMV birth prevalence of around 0.64% (1,2,10) the predicted incidence is about 1800 infected infants per year, which is substantially more than the national rates of 5–25 cases per year. (1,2,10)

Critically, most of these infected infants will be asymptomatic at birth and are unlikely to be identified without screening. About 13.5% (240 per year) will develop permanent sequelae. (3)

With an additional estimated 200–230 cases (11%–12.7%) (3,4) symptomatic at birth every year, over 440 children per year may be affected by this disease in Australia, with no routine newborn screening currently in place.

Only a minority of our cases (15/195; 8%) received antiviral therapy. Although ganciclovir has been demonstrated to positively influence the clinical course of SNHL when commenced before 30 days of age in symptomatic infants (5), it is acknowledged that a safer and more practical effective treatment is needed. Valganciclovir, an oral prodrug of ganciclovir (therefore not requiring central venous access), has demonstrated pharmacokinetic equivalence with intravenous ganciclovir in neonates with symptomatic disease. (6,7) A clinical trial is ongoing to determine whether valganciclovir improves hearing and neurological outcomes in these infants. (8) When the results of this study are available, however, early identification will remain important, as cCMV cannot be reliably diagnosed after 21 days of age, except by recourse to CMV PCR testing of newborn screening cards, where available. Twenty-eight (14%) of our cases were identified after 60 days of age, which precluded consideration of evidence-based antiviral therapy to prevent hearing loss or early intervention.

In our subgroup of 42 infants for whom we undertook clinical review, one-quarter were not identified with cCMV until after 60 days of age. Neurodevelopmental problems and SNHL were common in this group, indicating the importance of considering cCMV in children with SNHL, particularly if identified late. The utility of PCR testing of newborn screening cards in diagnosing cCMV infection has been demonstrated by others. (9)

In our case series, positive CMV PCR test results were strongly associated with clinical disease, although negativity did not exclude disease.

A limitation of our study is referral bias, as symptomatic children are more likely to be identified and referred for review. The data presented on these infants, however, demonstrate the burden of disease and sequelae in a contemporary population of symptomatic infants.

During the course of this study, neonatal hearing screening was introduced in most Australian states (Ms M Wilkinson, Head of Audiology, Sydney Children’s Hospital, Randwick, personal communication, 2009) (Box 2). We found a trend of increasingly reported SNHL, which may indicate increased case ascertainment. The introduction of these screening programs was staggered across the country and, in some states, program implementation was partial, so some infants may have lived in areas without access to such screening. Children identified with SNHL are not currently routinely screened for CMV in Australia and this is arguably an important gap within the program. Neonatal hearing screening is cost-effective and allows for early intervention with hearing aids and education, which can result in functional language improvements. (10)

These criteria meet the World Health Organization preconditions for a screening program. (11) The cost-effectiveness of incorporating cCMV screening into this algorithm urgently needs to be determined. Potential benefits include the opportunity for targeted antiviral therapy in infants with hearing loss and cCMV. This must be weighed against issues of cost and availability of timely CMV testing, as well as parental distress at identification of an infection which may or may not result in neurodevelopmental sequelae.

A routine referral program for cCMV screening in infants with hearing impairment is currently operating in Queensland, incorporated into the state’s universal neonatal hearing screening program. Internationally, there is ongoing debate about incorporation of neonatal testing for CMV in order to allow for targeted clinical follow-up of affected infants. (12–23) This view is further supported by estimates that cCMV contributes to 15%–20% of moderate-to-profound bilateral hearing loss (hearing loss of > 40 decibels) in children. (24) Hearing loss in cCMV may be progressive or fluctuating, however, and more than half of infants with cCMV and SNHL may be missed by neonatal hearing screening alone. (25)

In centres where newborn hearing screening programs are in place, early CMV testing is important. A positive result has implications of possible hearing loss and other neurodevelopmental sequelae, (26) and may allow for planned developmental follow-up as well as consideration of antiviral therapy. Furthermore, the identification of CMV is highly relevant when doctors are counseling parents about future pregnancies. Alternatively, CMV PCR testing or salivary culture could be included in routine newborn screening of all infants to allow for targeted follow-up of infants who test positive, as they are more likely to develop symptomatic disease, and SNHL in particular, beyond the neonatal age group.

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**COMPETING INTERESTS**

None identified.

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