TEACHING CASE REPORT

Congenital rubella syndrome despite maternal antibodies

THE CASE: In January 2004 a female infant was delivered by cesarean section at 36 weeks’ gestation because of severe intrauterine growth restriction. The mother was a healthy 31-year-old gravida 4 para 2 woman who had been travelling in Sri Lanka from before conception until 6 weeks’ gestation. Blood drawn for antenatal tests at 6 weeks’ gestation upon her return to Canada was not initially tested for rubella titre. A rubella titre at 12 weeks’ gestation (425 IU/mL) was interpreted as indicating immunity. The pregnancy was uncomplicated, without a history of fever or rash.

The mother had immigrated to Canada from Sri Lanka in 1993, at which point her rubella titre had been 17 IU/mL. She did not recall a history of rubella infection or of rubella vaccination. She had delivered a healthy child in 1996, but rubella screening had not been requested during that pregnancy. In her other 2 pregnancies, which did not carry to term, her rubella titre had not been tested.

Findings on initial physical examination of the daughter born in 2004 were unremarkable. Her birth weight was 1.58 kg, head circumference 28.3 cm and length 41 cm (all below the third percentile). Hematologic studies revealed a normal leukocyte count and hemoglobin concentration but a low platelet count (49 × 10⁹/L). During the first few days, the infant had hypoglycemia, hypocalcemia and hyponatremia, with a single seizure being attributed to hyponatremia. Ultrasonography of the head yielded normal findings. A murmur was noted on day 3 of life, and echocardiography demonstrated a large patent ductus arteriosis, a patent foramen ovale and severe pulmonary valvular stenosis. The infant was subsequently given a diagnosis of bilateral congenital cataracts. On the first day of the infant’s life, her rubella IgG titre was 243 IU/mL (immunity is indicated at a titre of 10–15 IU/mL or greater) and the rubella IgM titre was 1.123 index units (a positive result is 0.3 or more index units [Behring Enzygnost, Behring Diagnostics, Marburg, Germany]). Rubella virus was cultured from the urine and from both extracted cataracts.

After delivery, the stored antenatal serum that had been collected at 6 weeks’ gestation was performed at 12 weeks’ gestation upon her return to Canada was not initially tested for rubella titre. A rubella-specific IgM test was negative. The pregnancy was uncomplicated, without a history of fever or rash.

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After delivery, the stored antenatal serum that had been collected at 6 weeks’ gestation was tested, and the rubella IgG and IgM titres were both positive (281 IU/mL and 0.619 index units respectively). All rubella titres were performed at the Ontario Central Public Health Laboratory.

Althought congenital rubella syndrome (Box 1) is rare in Canada, it does still occur. Many women immigrate to Canada from countries without a rubella vaccination program and may not have adequate immunity. In 2002, only 124 (58%) of 214 countries or territories around the world had implemented a rubella vaccination program.1 An Australian study found that women born in Asia, sub-Saharan Africa and South America were 5 times as likely as other women to be seronegative for rubella virus. In a review of all cases of infants with congenital rubella syndrome in the United States reported to the National Congenital Rubella Syndrome Registry from 1997 to 1999, 83% (20/24) were born to Hispanic mothers and 91% (21/23) were born to foreign-born mothers.

Sri Lanka does not have a rubella vaccination program; therefore, if our patient’s low positive titre from 1993 was a true-positive result, it was likely the result of natural immunity. Since serologic testing for rubella was documented only once preceding this pregnancy, we strongly suspect, but cannot confirm, that this case of congenital rubella syndrome resulted from rubella reinfection: the IgG titre had increased markedly, from 17 IU/mL in 1993 to 281 IU/mL 10 years later, with a positive IgM titre and no history of revaccination. A rubella-specific IgM test performed at 12 weeks’ gestation would have confirmed infection rather than immunity.

Numerous cases of congenital rubella syndrome due to rubella reinfection have been reported.2 All involved exposure to rubella in the first trimester of pregnancy after both natural and vaccine-induced immunity, usually with subclinical infection in the mother. The immune response may involve more than humoral immunity, but it is clear that rubella antibodies decline over time and may increase the risk of reinfection. In a study involving Korean children, 18.8% of those who had been vaccinated and 13.8% of those with natural immunity were found to be seronegative for rubella virus after 3 years. An

Key points

- Consider rubella vaccination of nonpregnant immigrant women at their first encounter with the health care system.
- Women with a low positive IgG titre (e.g., 10–15 IU/mL) before pregnancy may benefit from repeat rubella vaccination.
- Women should be screened for rubella susceptibility at each pregnancy, because immunity (from previous vaccination or natural immunity) can wane.
- If a pregnant woman with a low positive IgG titre has recently been exposed to rubella, measurement of her IgM titre can help distinguish between new infection and immunity.
- Seronegative pregnant women need to be given rubella vaccine after delivery.
Italian study showed that 9.8% of vaccinated girls were reinfected by wild-type rubella virus within 5 years.

Prenatal serologic testing for rubella usually includes only the IgG titre, and the result is reported as “immune” or “nonimmune” (Box 2). There is controversy over what level of titre confers immunity. Bullens and coauthors documented that 8 of 16 mothers whose infants had congenital rubella syndrome had rubella IgG titres of at least 15 IU/mL at the start of pregnancy, a level considered to be protective. Their finding questions whether the current cutoff of 10–15 IU/mL is too low to be considered protective and whether women with low positive rubella titres before pregnancy might benefit from repeat vaccination.

In the absence of quantitative IgG reporting, clinicians do not know which women have low positive titres. A further problem is that a single IgG titre without an IgM titre cannot differentiate recent infection from immunity, as demonstrated by the case we have reported. Processing all rubella serologic tests in a centralized laboratory with quantitative reporting, clinicians do not consider offering measles–mumps–rubella vaccination to nonpregnant immigrant women from developing countries at their first contact with the healthcare system.3 Serologic screening before vaccination is not recommended and may result in a missed opportunity to vaccinate if pregnancy occurs in the interim or if the woman is lost to follow-up. Consultations before travel may represent an opportunity to review rubella immunization status of women of childbearing age.

Because rubella antibodies wane over time, screening for rubella susceptibility is recommended at the first prenatal visit for all pregnant women in Canada. In a Canadian study, 2.3%–13.3% of pregnant women were seronegative for rubella virus. A Quebec study found that only half of the 8.4% women who were rubella seronegative received the vaccine after delivery. Standing orders for rubella vaccination after delivery may increase immunization rates.

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