ORIGINAL ARTICLE

Treatment requirements of infants with rhesus isoimmunisation within a geographically defined area

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Objective: To provide population based data on the treatment requirements of infants with rhesus isoimmunisation.

Setting: Twenty nine hospitals in South Thames in which 81 119 deliveries occurred between February 1999 and January 2000.

Design: Every month, a clinician identified in each of the hospitals sent back a postcard indicating whether or not an infant with RhD had required treatment in their institution. Antenatal and postnatal information was then requested from all those who gave positive responses.

Main outcome measures: Requirement for postnatal treatment for rhesus isoimmunisation.

Results: During the one year study period, only 26 infants required treatment for rhesus isoimmunisation. The median duration of phototherapy of the 26 infants was five days (range 1–12). Seven infants required at least one exchange transfusion (two required two exchange transfusions), and seven infants received one “top up” transfusion. None received erythropoietin and no infant died.

Conclusion: The results suggest that few infants require treatment for rhesus isoimmunisation.

The introduction of anti-D prophylaxis to prevent sensitisation of women lacking rhesus antigens resulted in an appreciable reduction in the incidence of rhesus isoimmunisation, with a current estimate of 10.6 cases per 10 000 live births. The maternal death rate was also reduced, from 15 per 10 000 births reported in 1963 to 0.54 per 10 000 births reported in 1983. The latter figure, however, is an underestimate of the true situation, as it does not include termination of pregnancy for severe Rh disease and spontaneous fetal losses before 28 weeks of gestation. Postnatal and sensitising event prophylaxis with anti-D is now undertaken. Almost 90% of the sensitisation that follows delivery can be prevented by administration of anti-D at delivery, but a residual level of D sensitisation occurs. Up to 2% of RhD negative women show evidence of antibody production despite prophylaxis. Antenatal administration of anti-D (500 IU of anti-D) at 28 and 34 weeks has been recommended in addition to current practices, as this would prevent isoimmunisation for virtually all non-sensitised RhD negative women. This, however, would require a fourfold increase in hyperimmune anti-D production with associated costs. Whether such an approach would be cost effective in the United Kingdom requires accurate information on the current outcome of infants with Rh disease. The aim therefore of this study was to undertake a prospective, population based survey of the treatment requirements of infants with rhesus isoimmunisation.

METHODS

Data were collected on all infants with rhesus D isoimmunisation (that is direct Coomb’s test positive infants born to Rh negative mothers) who were born within a geographically defined area during a 12 month period (February 1999 to January 2000 inclusive) and required postnatal treatment. Clinicians were identified at 29 hospitals in which deliveries occurred in South Thames (see Acknowledgements). The total live births at the hospitals during the study period was 81 119; about one sixth were to Rh negative women. Additional antenatal sensitisation with anti-D in the last trimester of pregnancy was not given during the study period in South Thames. Every month, each clinician was requested to send to the data collecting centre (the neonatal intensive care unit at King’s College Hospital) a postcard on which they indicated whether or not an infant with RhD disease had been delivered in their institution. Antenatal and postnatal outcome data were then requested from all those who gave positive results. From one hospital only were these data collected retrospectively. The data requested included: mother’s address, previous pregnancy outcomes, whether anti-D had been administered, and the number of intrauterine transfusions during the current pregnancy. In addition, information was requested on the gestational age at delivery, the duration of phototherapy, the requirement for exchange transfusion(s) and/or top up transfusions, whether the infant had received erythropoietin, and if the infant had died. Within the study period, late responders were chased, and, at the end of the year, each hospital was sent a summary of their results and requested to confirm its accuracy. Only the outcome of infants who required treatment for rhesus isoimmunisation and whose mothers were South Thames residents are reported.

RESULTS

During the 12 month period, 26 infants with Rh disease had at least received phototherapy. Only one of the 26 mothers was a primigravida. The data on anti-D administration in previous pregnancies were available in 14 cases; anti-D had not been given after three miscarriages and two terminations of pregnancy. Five of the 26 infants had undergone between two and 10 in utero transfusions. The 26 infants required a median duration of phototherapy of five days (range 1–12). Seven infants required at least one exchange transfusion, and two required two exchange transfusions. No infant received erythropoietin. Seven infants required one top up transfusion at a median age of 3 weeks (range 2 days to 12 weeks). No infant required a second top up transfusion. None of the 26 infants died. Ten infants were born prematurely; their median gestational age at delivery was 36 weeks (range 33–36). One infant was born at 33 weeks gestation, one at 34 weeks, and two at 35 weeks. The infant born at 33 weeks gestation required phototherapy for only one day, no exchange transfusion, and one top up transfusion at 14 days after birth. The
infant born at 34 weeks gestation required one exchange and one top up transfusion and 12 days of phototherapy.

**DISCUSSION**

In this population based survey over a 12 month period in which there were 81 119 live births, only 26 infants required treatment for Rh disease and no infant died. Deaths attributable to Rh haemolytic disease were reported in 1994 to be 1.3 per 100 000 live births. It has been suggested that the true death rate is closer to six deaths per 100 000, as certification data exclude abortions and five times as many deaths were uncertified as certified by the General Register Office in Scotland. Our survey only included live births, and thus we cannot comment on the total fetal and infant death rate attributable to rhesus isoimmunisation. The data, however, have the strength of being prospectively collected and population based, thus giving an accurate reflection of infant mortality. In a survey performed in Northern Ireland over a 30 month period and including about 65 000 deliveries, there was only one neonatal death. These and our data suggest that the current neonatal mortality from Rh disease is low.

It has been highlighted that a large proportion of the cost related to rhesus isoimmunisation is the cost of neonatal intensive care. We therefore prospectively documented treatment requirements of infants with Rh disease. We did not collect information on infants who did not require treatment as such infants would have no extra financial implications. In this population based survey, we found that 26 infants required treatment—that is, 0.03% of the live births surveyed. It is possible that infants who received phototherapy exclusively on the postnatal ward may have been missed, but we think that this is unlikely because in 28 of the 29 hospitals we recorded the data prospectively on a monthly basis. In addition, at the end of each year all the clinicians were sent a list of treated infants in their hospital and were asked to cross check their records for the whole year. In a previously published survey of the outcome of RhD antibody affected pregnancies in Northern Ireland, there were 59 neonatal unit admissions for rhesus isoimmunisation; 29 of the infants required exchange transfusion and 55 required phototherapy. That survey was carried out over a 30 month period during which time there were about 65 000 deliveries. These data therefore also suggest that, compared with the number of live births, only a small proportion of infants require neonatal treatment for rhesus isoimmunisation. Nevertheless, the treatment requirements of the infants who are rhesus isoimmunised may be substantial. In comparison with our data, a ment requirements of the infants who are rhesus isoimmunised may be substantial. In comparison with our data, a

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