Effect of maternal anticonvulsant treatment on neonatal blood coagulation

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Abstract
Aims—To investigate the impact of maternal anticonvulsant use on the ability of cord blood to coagulate.

Methods—Cord blood prothrombin times were measured, over 15 years in a consecutive series of 137 term babies born to women taking phenobarbitone, phenytoin, and/or carbamazepine while pregnant. The response to parenteral vitamin K was measured in 83 neonates.

Results—Only 14 of the 105 babies born to the mothers who had therapeutic anticonvulsant blood concentrations at birth had a prolonged prothrombin time (outside the 95% reference range). None had an overt bleeding tendency. The abnormality was corrected within 2 hours by 1 mg of parenteral vitamin K, but rapid intravenous prophylaxis produced complications in three infants.

Conclusions—A policy of giving vitamin K throughout the last third of pregnancy to all women being treated with anticonvulsants, as recently recommended, is not justified by the available evidence. The belief that there is a distinct, early form of neonatal vitamin K deficiency that is different from, and more dangerous than, the classic form of the disease, is not supported by a review of the published evidence.

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Forty years have elapsed since the first published report suggested that maternal anticonvulsant treatment might increase the vulnerability of a neonate to a form of coagulopathy—haemorrhagic disease of the newborn—correctable with vitamin K. However, the problem was generally considered rare until a study in 1970 which reported a high incidence of prothrombin deficiency among babies born to mothers taking anticonvulsant medication. This prompted a 15 year study of the coagulation status of all such babies in Newcastle using blood samples collected from the mother, and from the umbilical cord, at delivery. Because of a fear that intramuscular administration of vitamin K might itself cause bleeding, these babies were also admitted briefly to special care for the first 10 years of the study so they could be given vitamin K (1 mg of Konakion) intravenously, and further blood specimens were collected without trauma through an umbilical vein catheter 2–4 hours later. A review of this policy in 1979 revealed that significant prolongation of the prothrombin time was relatively rare, and umbilical catheterisation not without hazard, so a policy of immediate intramuscular prophylaxis was substituted.

In the light of the renewed debate about the indications for oral and intramuscular prophylaxis, the experience assembled during this previously unpublished study is now reported.

Methods
Blood was collected with a needle and syringe from the umbilical cord of 137 term babies whose mothers had been taking anticonvulsant drugs throughout pregnancy (care being taken to prevent contamination with Wharton’s jelly) and 2.5 ml of this blood added to 0.25 ml of 3.8% sodium citrate. The one stage prothrombin time was then measured on the spun sample using a rabbit brain thromboplastin (Dade, American Hospital Supply Corporation) from 1970 to 1980, and a human brain thromboplastin (Manchester reagent, Thrombosis Reference Centre, Manchester) thereafter. Citrated control plasma was obtained by pooling the plasma from 10 adult volunteers, and the prothrombin time expressed as a ratio of the time it took a simultaneous control plasma to clot. Blood was also collected from all but six of the mothers at delivery to determine anticonvulsant concentrations. Eighty three of the babies had a further specimen collected from an umbilical vein catheter 2 hours after receiving 1 mg of intravenous vitamin K.

Cord blood was collected during the first year of the study from 50 term babies whose mothers had not received any antenatal treatment, and from a further 50 control babies during the final year. The distribution of these ratios was normal after log transformation, and the two means and standard deviations were almost identical despite the change in thromboplastin two thirds of the way through the study.

Results
All the mothers had been taking phenobarbitone, phenytoin, or carbamazepine, or some combination of the three, but nearly a quarter (32/137) had anticonvulsant blood concentrations less than half the lower limit of the normally quoted therapeutic range at delivery, and subsequent enquiry revealed that many mothers had been taking treatment since childhood without medical review even though they had been free from seizures for five or even 10 years.
The mean prothrombin ratio for the 105 babies born to mothers receiving effective anticonvulsant treatment was higher than in the control babies, but only 14 had a prothrombin ratio above the 95% reference range and the amount of maternal medication taken, and no change in the mean ratio over time. The prothrombin ratio changed little in most of the babies after vitamin K administration, but it fell at least 25% in all 10 of the 68 with an initial cord blood ratio in excess of 1.7 who had been given intravenous vitamin K at birth. No baby had any overt evidence of a bleeding tendency. The suggestion that anticonvulsants could depress the fetal vitamin K dependent factor concentrations seriously enough to cause death from bleeding before birth deserves more study, but remains to be substantiated.

It is now widely held that maternal anticonvulsant treatment leaves babies vulnerable to a distinct early form of vitamin K deficiency bleeding that is different from, and more lethal than, the classic disease, but this is to misread the published case reports. It would be more correct to say that such babies are at greater risk of severe bleeding if they sustain serious trauma during delivery, and they are also more likely to develop classic symptoms later in the first week of life, unless given prophylactic vitamin K at birth. It was a demonstrable reduction in the number of deaths associated with lethal early intracranial and intraperitoneal bleeding, just as much as a reduction in the number showing evidence of a generalised but less serious bleeding tendency later in the first few days of life, that motivated the Scandinavian paediatricians who first recommended universal (oral) prophylaxis 50 years ago. Units that later adopted a policy of selective intramuscular prophylaxis (often for lack of any licensed oral alternative) were reflecting this double objective. The target was babies who had sustained a traumatic delivery, and babies who were unlikely to get much milk in the first few days of life. The slow transformation of this logical strategy into a policy that resulted in intramuscular vitamin K being given to every baby having a caesarean or forceps delivery, and every baby going to

**Key messages**
- Few babies born to women taking phenobarbitaline, phenytoin and/or carbamazepine have an overt bleeding tendency at birth, although 13% have a prothrombin time that is longer than normal (above the 95% reference range)
- A significant number of women seem to be taking subtherapeutic doses of these potentially teratogenic anticonvulsants in an unsupervised (and probably unnecessary) way
- Parenteral treatment at delivery suffices to correct the coagulation abnormality within 2 hours (the adequacy of oral treatment remains undetermined)

![Figure 1](http://fn.bmj.com/)

Figure 1 Cord blood prothrombin ratio in 105 babies whose mothers probably had therapeutic blood anticonvulsant concentrations at delivery. The reference range (within which 95% of the control ratios lay) is shown by the dotted lines. Although the mean ratio is higher in the babies of mothers receiving anticonvulsant treatment, it is the tail of high values that is of clinical significance.

Nevertheless, of the 38 published cases, less than a third can be counted as established cases by the criteria most would now judge appropriate in any epidemiological study of this condition. Many of the more adequately documented cases (with a measured prothrombin time and a response to vitamin K administration) only had mild symptoms, and half first presented when more than 48 hours old. Only two of these babies developed symptomatic bleeding despite the administration of prophylactic intramuscular vitamin K at birth: both had sustained trauma to the liver, and only one had evidence of a generalised bleeding tendency. The suggestion that anticonvulsants could depress the fetal vitamin K dependent factor concentrations seriously enough to cause death from bleeding before birth deserves more study, but remains to be substantiated.

Discussion

Overt bleeding due to vitamin K deficiency is a rare but well described phenomenon in babies born to mothers taking phenytoin or a barbiturate anticonvulsant for epilepsy.
special care, shows what can happen when staff no longer understand the reasons underpinning unit protocol.

Five of the 16 babies born to mothers receiving anticonvulsant treatment in the study reported by Mountain et al had a low cord blood prothrombin (factor II) concentration. Deblay et al encountered bleeding in 7% (8/115) of babies not given vitamin K at birth. They also found a significant reduction in mean prothrombin concentration in a prospective study of 74 babies,22 and argued that the best way to prevent this was to give the mother additional oral vitamin K. Vitamin K does not cross the placenta readily, so large doses have to be given. Treatment with 20 mg a day for two weeks before delivery was thought to be associated with a higher prothrombin concentration in the 14 babies born to mothers so treated in this report, but the study was too small for the difference to be statistically convincing.22

Treatment with 10 mg every day for a month abolished all evidence of subclinical vitamin K deficiency in a more recent study of 16 babies,23 but prothrombin times were not reported. Maternal prophylaxis does not reduce the cord blood prothrombin time of normal preterm babies.24 That some babies born to mothers receiving phenobarbitone or phenytoin do have marginally prolonged prothrombin times at birth is confirmed by this study, but symptomatic deficiency of vitamin K dependent factors was not seen, and seems to be rarer than early reports have suggested. As immediate prophylaxis with intramuscular vitamin K at birth might well have prevented all the published cases of documented symptomatic factor deficiency unassociated with trauma to the liver or spleen, many clinicians will prefer to give a physiological dose of vitamin K to the baby at delivery, rather than a pharmacological dose to the mother before delivery, as is frequently recommended,4 17 20 26 especially while there remains any continuing possibility that high doses could be carcinogenic.27 28 The problem should become less common as other drugs come to be used in the management of adult epilepsy—valproate, in particular, does not seem to cause neonatal hypoprothrombinaemia.3

This study was initiated 20 years ago by the late Dr Gerald Nelson, with advice and laboratory support from the late Professor William Walker, and undertaken at the Princess Mary Maternity Hospital in Newcastle, which has now closed. I am grateful to Dr M Reid and Dr M Cornilssen for helpful comment and advice.


15 Speidel BD, Meadow SR. Maternal epilepsy and abnormalities of the fetus and newborn. Lancet 1972;i:839–43.


