Perinatal cortical infarction within middle cerebral artery trunks

Paul Govaert, Erika Matthys, Alexandra Zecic, Filip Roelens, Ann Oostra, Bart Vanzielegem

Abstract

Aim—To define neonatal pial middle cerebral artery infarction.

Methods—A retrospective study was made of neonates in whom focal arterial infarction had been detected ultrasonographically. A detailed study was made of cortical middle cerebral artery infarction subtypes.

Results—Forty infarctions, with the exception of those in a posterior cerebral artery, were detected ultrasonographically over a period of 10 years. Most were confirmed by computed tomography or magnetic resonance imaging. Factor V Leiden heterozygosity was documented in three. The onset was probably antepartum in three, and associated with fetal distress before labour in one. There were 19 cases of cortical middle cerebral artery stroke. The truncal type (n=13) was more common than complete (n = 5) middle cerebral artery infarction. Of six infarcts in the anterior trunk, four were in term infants and five affected the right hemisphere. Clinical seizures were part of the anterior truncal presentation in three. One of these infants, with involvement of the primary motor area, developed a severe motor hemisyndrome. The Bayley Mental Developmental Index was above 80 in all of three infants tested with anterior truncal infarction. Of seven patients with posterior truncal infarction, six were at or near term. Six of these lesions were left sided. Clinical seizures were observed in three. A mild motor hemisyndrome developed in at least three of these infants due to involvement of parieto-temporal non-primary cortex.

Conclusions—Inability to differentiate between truncal and complete middle cerebral artery stroke is one of the explanations for the reported different outcomes. Severe motor hemisyndrome can be predicted from neonatal ultrasonography on the basis of primary motor cortex involvement. Clinical seizures were recognised in less than half of the patients with truncal infarction; left sided presentation was present in the posterior, but not the anterior truncal type of infarction. Asphyxia is a rare cause of focal arterial infarction.

Keywords: cortical infarction; middle cerebral artery; severe motor hemisyndrome.

Focal arterial infarction is an important cause of permanent brain injury in the perinatal period. Estimates from brain imaging suggest an incidence of 0.2 per 1000 neonates. Precise definition of the vessel involved is often impossible, partly because of the difficulty of angiographic confirmation and limited correlations with post mortem findings.

The left middle cerebral artery tends to be affected predominantly. For infarction in the entire field of the middle cerebral artery the previously optimistic prognosis has been replaced with guarded realism. For partial cortical (pial) and lacunar infarction, the outcome is virtually unknown. Prognostic variability can be explained by unpredictable brain plasticity, and the variable accuracy with which lesions are anatomically defined. An additional reason may be overdiagnosis of infarction based on white matter attenuation with computed tomography or focal hyperattenuation within ischaemic but not necessarily necrotic tissue.

We report the retrospective analysis of 13 cases of infarction in one of the two major cortical trunks of the middle cerebral artery.

Methods

We retrospectively studied the ultrasound brain scans of infants born between 1988 and 1997, to identify those scans with the appearance of arterial infarction. Brain scans were routinely carried out on admission to our unit with few exceptions. All ventilated infants were scanned within 24 hours, and most within hours of admission. Subsequent scans were done up to three times in the first week on clinical indication, and thereafter weekly in the neonatal period. The scans were carried out by one of four neonatologists responsible for clinical surveillance, and all the images were reviewed by one of us (PG). The number of planes in which images were printed was not fixed, but adapted to the lesion seen in real time. The scanning machine used was an ATL Ultramark 4 with a mechanical sector scanhead providing 5, 7.5, and 10 MHz frequencies. Almost all the images were scanned using the 7.5 MHz transducer. The results of the scans were confirmed by computed tomography or magnetic resonance imaging in nearly all cases.

Focal brain infarction may present in the early neonatal period as a triangular defect or lacuna filled with cerebrospinal fluid. Pathologists describe ventricular dilatation, shrunken gyri (ulegria), and cavities with glial membranes traversing the space left by absent tissue. If such a defect is found in the first two weeks of life prepartum onset is likely.
After peripartum or early neonatal insults the lesion presents in the neonatal period as a hyperechoic area confined to arterial anatomical boundaries. To be classified as peripartum or neonatal arterial infarction, the hyperechoic lesion had to be confined to the artery; in the case of a known recent event, the focus could not show signs of cystic change within the first 14 days of life; the lesion had to be unilateral, so as not to be confused with watershed infarction types like parasagittal cerebral injury and periventricular leucomalacia; haemorrhage had to be excluded, so parenchymal hyperechogenicity in association with ipsilateral germinal matrix haemorrhage was excluded; larger lesions had to have at least one linear margin. Analogous with adult stroke we did include smaller hyperechoic foci of ellipsoid shape in deep grey matter.

We defined the lesion as a cortical infarct when a pial area of the vessel involved was part of the infarct, even if deep structures were also affected. In contrast, we classified those involving deep grey nuclei or deep white matter without cortical involvement as deep infarcts. To be defined as complete anterior truncal middle cerebral artery infarction, the hyperechoic focus had to be a triangle, with its point at the insula and its base spread across the parasagittal sectors from frontal bone to around the central sulcus. The anterior insular gyri were also involved.

To be classified as complete posterior truncal middle cerebral artery infarction, the hyperechoic focus had to point towards the posterior insula on parasagittal section, and have a triangular extension with fuzzy base against the occiput. Viewed through the coronal plane, a blurred hyperechoic area was seen behind and underneath the circular groove of the insula. Parts of the long insular gyri were injured in some.

Clinical and ultrasonographic findings excluded primary haemorrhage, selective neuronal necrosis, venous infarction, tumour, abscess formation and phakomatosis. In some cases distinction from infarction within the area drained by a single small deep vein could not be made. These lesions were not excluded. Hyperpulsatility in the penumbra and high flow velocity in the (recanalised) artery pointing to the infarct were not considered mandatory. Early fetal arterial infarcts leading to schizencephaly, hydraencephaly, or Moebius’s syndrome were excluded.

OUTCOMES
The contralateral motor hemisyndrome was considered severe if the affected lower limb had no useful function (hemiplegia); moderate if some function was preserved; and mild if there was clumsiness with preservation of normal function. Manual function was assessed as follows: severe hemisyndrome: non-functioning hand; moderate: preservation of residual hand function; mild: independent finger movement. Follow up information was not available for all the children. The results of the Mental Developmental Index (MDI) determined by Bayley Scales of Infant Development were available for some children.

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Association between distribution of infarction and clinical factors</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deep (n=16)</td>
<td>Cortical (n=24)</td>
</tr>
<tr>
<td>&lt;37 weeks gestation</td>
<td>13 (81.3)</td>
</tr>
<tr>
<td>Neonatal seizures</td>
<td>2 (12.5)</td>
</tr>
<tr>
<td>Neonatal death</td>
<td>4 (25.0)</td>
</tr>
</tbody>
</table>

Results
From 1988 to 1997 40 cases of arterial brain infarction were identified (19 boys, 21 girls) involving one of the three major cerebral arteries. In 28 infants the site and characteristics of the lesion were confirmed by computed tomography, or magnetic resonance imaging. The distribution of lesions was as follows: anterior cerebral (n=4); middle cerebral (n=26); posterior cerebral (n=10). The prevalence of brain infarction between 1993 and 1997 was 12 per 1000 neonates admitted to the neonatal unit. Given that our referral population is about 15000 births a year, the incidence is probably about 0.35 per 1000 total births. Eighteen of the 40 infants (45%) had been born at <37 weeks’ gestation. The relation between the distribution of lesions, gestational age, and the occurrence of seizures is shown in table 1. Infants with cortical lesions were more likely to be term, to present with neonatal seizures, and to survive.

Putative causes of arterial infarction are summarised in table 2. There was no association between the vessel type and putative causal factors. The most common association (28% of cases) was pulmonary hypertension or shock, necessitating assisted ventilation. In 25% of cases there was no obvious “cause” for the infarction.

MIDDLE CEREBRAL ARTERY CORTICAL LESIONS
Cortical lesions predominated in infarctions involving the middle cerebral artery. Among the 19 patients with cortical infarction involving the middle cerebral artery 13 patients had lesions confined to the anterior (n=6) or posterior trunk (n=7). The other six patients had complete lesions in the distribution of the main artery (n=5) or very peripheral lesions involving only a small subpart of a trunk (n=1). Details of the 13 patients are shown in table 3. Six out of the seven patients with infarction in the posterior trunk had left sided lesions, in contrast to only one of the six patients with anterior trunk lesions. The gestational ages and incidence of seizures were similar in anterior and posterior truncal lesions. Typical images of the posterior truncal lesion are shown in fig 1, and of the anterior truncal variant in fig 2.

<table>
<thead>
<tr>
<th>Table 2</th>
<th>Putative causes of infarction in territories of major cerebral arteries</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pulmonary hypertension or shock requiring assisted ventilation</td>
<td>11 (28)</td>
</tr>
<tr>
<td>Unknown cause</td>
<td>10 (25)</td>
</tr>
<tr>
<td>Septicaemia</td>
<td>5 (13)</td>
</tr>
<tr>
<td>Fetal anaemia</td>
<td>4 (10)</td>
</tr>
<tr>
<td>IUGR and fetal distress</td>
<td>4 (10)</td>
</tr>
<tr>
<td>Factor V Leiden deficiency</td>
<td>3 (8)</td>
</tr>
<tr>
<td>Birth trauma</td>
<td>2 (5)</td>
</tr>
<tr>
<td>Severe intraventricular asphyxia</td>
<td>1 (3)</td>
</tr>
</tbody>
</table>
The outcome was known in 11 of the 13 infants with truncal middle cerebral artery lesions involving the cortex. Only one patient was hemiplegic and this was associated with an anterior truncal lesion involving the rolandic cortex. Three patients with posterior truncal lesions had a mild hemisyndrome. Most of the patients therefore did not have hemiplegia, a phenomenon previously reported as a sequel to infarction in the complete middle cerebral artery.

### Discussion

The estimated incidence of all types of focal cerebral infarction in our study (0.35 per 1000 total births) was 3–4 times higher than the results of studies in term infants who were diagnosed in later childhood. This was partly due to the inclusion in our study of infants with isolated deep infarctions. Aside from perinatal asphyxia, stroke is the most common cause of neonatal seizures at term. As anticipated, we observed that seizures were more frequently related to cortical than to deep infarcts. Like many others before us, we could not find a cause for neonatal stroke in a high proportion of cases. In our experience perinatal asphyxia is a rare cause of stroke. However, we confirmed an association between stroke and pulmonary hypertension.

Hemiplegia may follow infarction in areas of the brain supplied by the entire middle cerebral artery, in the complete set of cortical branches of the middle cerebral artery, in the anterior or posterior trunk of the middle cerebral artery, in

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**Table 3** Truncal middle cerebral artery infarction

<table>
<thead>
<tr>
<th>Case No</th>
<th>Gestational age</th>
<th>Clinical details</th>
<th>Site</th>
<th>Outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>Entire posterior trunk of the middle cerebral artery</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>36</td>
<td>PROM for 3 days, congenital pneumonia due to early onset group B streptococcal infection, shock</td>
<td>Left</td>
<td>Mild hemisyndrome</td>
</tr>
<tr>
<td>2</td>
<td>39</td>
<td>Vacuum extraction, cephalhaematoma, focal seizures on day 1, early onset enterobacter septicaemia, no shock</td>
<td>Left</td>
<td>No sequelae</td>
</tr>
<tr>
<td>3</td>
<td>38</td>
<td>Second twin, cord prolapse, difficult vacuum delivery, subgaleal bleeding, basd subdural haematoma, focal EEG changes</td>
<td>Left</td>
<td>Mild hemisyndrome</td>
</tr>
<tr>
<td>4</td>
<td>40</td>
<td>PROM for 3 days, vacuum delivery, Apgar scores 4/6, general seizures on day 1, day 1 Hb 10.2 g/dl, metabolic acidosis</td>
<td>Left</td>
<td>Mild hemisyndrome, speech difficulties</td>
</tr>
<tr>
<td>5 (fig 1)</td>
<td>37</td>
<td>Twin, left renal vein thrombosis and adrenal haemorrhage, apc resistance</td>
<td>Right</td>
<td>No sequelae</td>
</tr>
<tr>
<td>6</td>
<td>38</td>
<td>Breach delivery, focal seizures on day 2</td>
<td>Left</td>
<td>One epileptic insult at 3 months, no hemisyndrome</td>
</tr>
<tr>
<td>7</td>
<td>33</td>
<td>Caesarean section for fetal distress superimposed on growth retardation; no IRDS; APC resistance; early cystic stage of the haemorrhagic infarction on day 1</td>
<td>Left</td>
<td>No follow up</td>
</tr>
<tr>
<td>Entire anterior trunk of the middle cerebral artery</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8 (fig 2)</td>
<td>34</td>
<td>IUGR, lost heart to heart beat variation, brief postnatal acidosis (the area involved span the right frontal and part of the parietal lobe’s convexity, including rolandic cortex)</td>
<td>Right</td>
<td>Severe hemiplegia, MDI 42</td>
</tr>
<tr>
<td>9</td>
<td>40</td>
<td>Focal seizures on day 2, APC resistance in propositus and asymptomatic father (presentation was with a right frontal lobe haematoma; the evolution showed this to be infarction with late neonatal regression with multiple cysts)</td>
<td>Right</td>
<td>Normal at 12 months, right predominance</td>
</tr>
<tr>
<td>Within part of the anterior trunk of the middle cerebral artery</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>30</td>
<td>Death of co-twin at 26 weeks, IRDS (the affected vessel was probably a branch of the prefrontal artery)</td>
<td>Right</td>
<td>Unknown</td>
</tr>
<tr>
<td>11</td>
<td>38</td>
<td>Second day seizures in a healthy infant</td>
<td>Left</td>
<td>Unknown</td>
</tr>
<tr>
<td>12</td>
<td>38</td>
<td>Coarctation aorta, oesophageal atresia, systemic acidosis (haemorrhagic leukomalacia and right anterior middle cerebral artery infarction were associated in this child)</td>
<td>Right</td>
<td>Unknown</td>
</tr>
<tr>
<td>13</td>
<td>40</td>
<td>Oligohydramnios and stunted fetal growth for 2 weeks, chronic fetomaternal transfusion (Kleihauer-Betke 6%), focal seizures on day 2 (the area involved may have been within the prefrontal and precentral branches of the middle cerebral artery or may have been a watershed zone with the anterior cerebral artery)</td>
<td>Right</td>
<td>Normal at 12 months</td>
</tr>
</tbody>
</table>

Key: PROM = Premature rupture of membrane; IRDS = Idiopathic respiratory distress syndrome; APC = Activated protein C resistance (factor V Leiden).

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**Figure 1** Sonograms (7.5 MHz) through the coronal (top) and parasagittal planes of the right hemisphere and T1 weighted axial magnetic resonance image in the late neonatal period in case 5. The position of the central groove is shown in relation to the infarct in the posterior section of the middle cerebral artery. The infarct is pial as the striatum is not affected.

**Figure 2** Sonograms (7.5 MHz) on day 3 showing coronal (top) and parasagittal planes (bottom) of case 8. Infarction is evident within the anterior trunk of the middle cerebral artery. Parasagittal plane shows involvement of the central groove which explains the subsequent hemiplegia; necrosis within the entire anterior trunk involves not only the central area but also the insular triangle.
the lenticulo-striate branches of the middle cerebral artery or in the anterior choroidal artery. The main purpose of this study was to draw attention to the occurrence of cortical infarction in areas of the cortex supplied by the anterior and posterior trunks of the middle cerebral artery. The precise anatomy of the branches of the middle cerebral artery varies, but the usual arrangement is the common stem dividing at the insula into anterior and posterior trunks (fig 3). The anterior trunk splits into prefrontal, precentral, central and anterior parietal arteries. It is not uncommon for the central and precentral, central and anterior parietal posterior trunks (fig 3) to extend as far as the motor cortex, this is uncommon. The central groove can be localised with ultrasonography (figs 1 and 2).

Infarction within the posterior trunk affects the convex posterior temporal, parietal, and occipital lobe. Video recording of sonograms may permit retrospective examination of involvement of the rolandic area. The sonograms were strikingly similar in all patients with this type of infarction, and comparable cases have been reported, for it to be considered a separate entity. In spite of a reported normal outcome, we recorded contralateral motor and orientation disturbances in half our infants with infarction in the posterior trunk. Case 3 was compared with her healthy male co-twin at 3 years of age by her mother: the account is typical: “She still cannot turn off a tap, invariably turning it the wrong way; she was much older before she mastered her tricycle because she pedalled backward; she cannot recognise the face in a photo when it was cut into pieces; inside the toilet, once the door is closed, she found it difficult to differentiate the door from the walls, whereas her brother easily remembers where the door was.”

The substrate for linguistic specialisation is present at birth, so that left and not right hemispheric lesions may cause permanent symptoms. Residual epilepsy seems to be rare. Smaller neonatal infarctions in the posterior trunk may well go undetected. For example, one infant without a hemisyndrome (case 2) had a normal follow up computed tomogram, suggesting that the neonatal hyperechoic focus (and attenuated corresponding computed tomographic area) had been ischaemic rather than necrotic and had completely resolved.

Caution is needed when assessing the probable perinatal cause of hemiplegia from brain imaging in childhood. One confounder is the existence of trans-synaptic anterograde neuronal atrophy. Cortical infarction over an extensive frontoparietal area perfused by the middle cerebral artery will cause delayed thalamic atrophy and macroscopic ipsilateral atrophy of the corticospinal tract at brainstem level above the decussation. Precise delineation of the original focus of damage may not be straightforward when reviewed subsequently.

As fewer than half the survivors with infarction of truncal distribution developed severe hemiplegia it is clear that the incidence of focal infarction in the perinatal period will be underestimated by basing estimates on observations in children with hemiplegia. Infarction involving the complete middle cerebral artery leads to contralateral hemiplegia, with the arm being more affected than the leg, and with astereognosis. Cognitive dysfunction tends to coincide with residual epilepsy. Details are available of perceptual performance and communication problems and language.
difficulties. The risk of residual epilepsy is between 25 and 50%.

It would be impossible to deduce the number of lesions we missed by using ultrasonography as a screening tool. On the other hand, there is no other technique available for screening large numbers of sick infants. As clinical markers for arterial infarction are insensitive, except, perhaps, for seizures, limb pallor, and apnoea in term neonates, sono-graphic surveillance assumes greater importance. The chance of finding a cerebral infarct is 30% in term newborn infants with seizures, and a five minute Apgar score of at least seven is likely. The limits of ultrasonography are in the description of posterior cerebral artery infarction and infarction between the fontanelles. Similarly, it is unlikely that ultrasonography will be helpful in brainstem or cerebellar arterial infarction.

More widespread sonographic detection might facilitate collaborative studies of the treat-ment of neonatal stroke. So far, attempts at treatment have been erratic. Infarction is likely if systemic hypoperfusion is associated with variable anatomy of the cerebral arteries. The study of the normal arterial anatomy of the newborn brain with Doppler or spectral analysis might facilitate collaborative studies of the treat-ment of neonatal stroke. So far, attempts at treatment have been erratic. Another major advance would be illustration of the vessel itself without invasive angiography. Magnetic reso-nance angiography looks promising in this regard, at least for vessels of more than 1 mm in width. Additional early information may be gathered with colour Doppler sonography, and magnetic resonance spectroscopy.

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