Microvessel ultrasound of neonatal brain parenchyma: feasibility, reproducibility, and normal imaging features by superb microvascular imaging (SMI)

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Abstract

Objectives To evaluate the feasibility and reproducibility of superb microvascular imaging (SMI) of the neonatal brain and to describe normal imaging features.

Methods We performed transcranial ultrasound with SMI in 19 healthy term-born neonates. SMI was done according to a structured examination protocol, using two linear 18 MHz and 14 MHz transducers. Superficial and deep scans were acquired in the coronal and sagittal planes, using the left and right superior frontal gyri as anatomical landmarks. All SMI views were imaged by monochrome and colour SMI and evaluated with respect to visibility of extrastriatal (i.e. cortical and medullary) and striatal microvessels.

Results We have described normal morphologic features of intraparenchymal brain microvasculature as “short parallel” cortical vessels, “smoothly curved” medullary vessels, and deep striatal vessels. In general, SMI performance was better on coronal views than on sagittal views. On superficial coronal scans, cortical microvessels were identifiable in 90–100%, medullary microvessels in 95–100%. On deep scans, cortical and medullary microvessels were visible in all cases, while striatal microvessels were identifiable in 71% of cases.

Conclusions Cerebral SMI ultrasound is feasible and well-reproducible and provides a novel non-invasive imaging tool for the assessment of intraparenchymal brain microvasculature (extrastriatal and striatal microvessels) in neonates without the use of contrast.

Key Points

• Superb microvascular imaging (SMI) of the neonatal brain is feasible and reproducible.
• SMI depicts extrastriatal and striatal microvessels.
• SMI detects two types of extrastriatal microvessels: cortical and medullary.

Keywords Ultrasound imaging · Microvasculature · Neonate · Brain

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Abbreviations

cSMI  Colour SMI
CT     Computed tomography
CV     Cortical microvessels
F1     Superior frontal gyrus (surgically referred to as F1) [1]
MRI    Magnetic resonance imaging
mSMI   Monochrome SMI
MV     Medullary microvessels
SMI    Superb microvascular imaging

Introduction

Intraparenchymal brain vasculature consists of a hierarchical network of small arteries, veins, and capillaries. Initial anatomical descriptions date back to the mid-sixteenth century [2]. Anatomical studies addressing the macro- and microanatomy of the neonatal human cerebral vasculature have been conducted on post-mortem specimens and sections [3–7]. Main arterial trunks supply the cerebrum by giving rise to basal perforating arteries proximally and pial cortical arteries peripherally [3]. Pial arteries arborise into intracortical, subcortical, and medullary arteries which supply the cortex, while the cerebral white matter is exclusively supplied by medullary arteries [8–10].

Apart from invasive procedures, such as digital subtraction angiography, in vivo visualisation of neonatal cerebral vasculature is possible using computed tomography (CT) and magnetic resonance imaging (MRI). CT is limited by the use of radiation and the need for intravenous contrast material and MRI by the demanding logistics and the use of sedation, when imaging neonates. With diameters of 100–200 μm, intraparenchymal brain vessels typically remain beneath the spatial resolution of clinical imaging methods including CT and MRI (field strengths up to 3 Tesla) [11].

In neonates with patent fontanels, transcranial ultrasound is used as the first-line imaging method due to wide availability with the option of bedside use, low cost, a patient-friendly safety profile, and high image resolution [12]. Targeting blood flow in vascular structures, Doppler investigations can further enhance the clinical value of transcranial ultrasound [13]. Recently, vascular ultrasound has been given a new twist by the establishment of an innovative method referred to as “Superb microvascular imaging (SMI)” [14]. SMI has evolved as a novel tool for non-invasive imaging of microvasculature without the use of intravenous contrast material [14]. The method uses advanced clutter suppression and processes low flow Doppler signals that are otherwise filtered and removed as “clutter”. Advantages are high resolution and frame rate, and visualisation of low-velocity flow. Two modes are available: monochrome SMI (mSMI) and colour SMI (cSMI). cSMI displays low flow components in colour overlaid on the grey-scale image with high temporal and spatial resolution simultaneously. mSMI reveals microvasculature with even higher sensitivity by subtracting the anatomical background.

Ishikawa et al used SMI to image tumour vessels and tumour margins during open brain surgery in adults [15]. Regarding paediatric applications, SMI has been shown helpful in the assessment of vesico-urethral reflux [16] as well as in undescended testes [17, 18].

Currently, little is known about the in vivo morphology of the microvascular architecture of the human brain after birth. Our study aims to visualise microvessels of the neonatal brain using SMI ultrasound. Firstly, the feasibility and reproducibility of transfontanellar SMI is assessed in a cohort of term-born neonates. Secondly, normal SMI features in the healthy neonatal brain are described.

Methods

Study design and participants

We conducted a prospective single-centre imaging study in term-born neonates: for the main study, a total of 19 newborns
were included at the Department of Paediatrics and Adolescent Medicine of the Medical University of Vienna, Austria during a 1-year study period. Inclusion criteria were defined as follows: term-born neonates with good postnatal adaptation without neurological deficits or suspected cerebral pathologies. Medical history during pregnancy had to be unremarkable, prenatal ultrasound of the central nervous system was required to be normal. Therefore, newborns with chromosomal abnormalities, postnatal neurological deficits, or any cerebral lesion diagnosed by prenatal or postnatal ultrasound were excluded from the study participation. Informed consent was obtained in all patients. The study was approved by the Ethics Committee of the Medical University of Vienna (EK 1530/2015). In order to design and plan the main study, five newborns, which were not part of the present study, underwent a structured exploratory application of SMI, which was compared with normal colour Doppler.

**Standard transcranial ultrasound**

All participants underwent standard transcranial ultrasound using a Toshiba Aplio 400 scanner (Canon Medical Systems Corporation) during the first weeks of life in an awake state. A small curved array transducer (11 MHz) was placed on the patent anterior fontanel to acquire coronal and sagittal standard views according to the Austrian and German Society for Ultrasound in Medicine [19]. In addition, a mid-sagittal colour Doppler image and resistive indices were obtained from the internal carotid artery and the anterior cerebral artery. Images were stored and read using a radiologic picture archiving and communication system (PACS, AGFA HealthCare). Routine radiology reports were generated for all participants.

**SMI ultrasound—image acquisition and interpretation**

SMI ultrasound was performed and documented using a predefined examination protocol by two board-certified paediatric radiologists (JP and AH), with more than 5 years of experience in neonatal head ultrasound and anatomical knowledge of the angioarchitecture of brain parenchyma. The examination protocol included superficial and deep scans. Superficial scans were acquired using a linear 18 MHz transducer (Toshiba 18L7) and captured coronal and sagittal views.
of the left and right superior frontal gyri (neuro-surgically abbreviated and referred to as F1 [1]; Figs. 1 and 2). F1 was chosen because of proximity to the anterior fontanel and thus optimal sonographic accessibility. Maximum zoom depth of a superficial scan was 2.5 cm. SMI box depth of superficial scan was 1.7 cm. SMI setting were 7.2 MHz (Doppler frequency), 21 kHz (Pulse repetition frequency), and colour gain 45–50. Deep scans were acquired only in the coronal plane using a linear 14 MHz (Toshiba 14L5) transducer (Fig. 3). SMI box depth of deep scans was 5–6 cm. SMI setting were 7 MHz (Doppler frequency), 9 kHz (Pulse repetition frequency), and colour gain 45–50. For a typical ultrasound exam, safety parameters were as follows: mechanical index 0.8–1.5, bone thermal index 0.6–0.8, and soft tissue thermal index 0.6–0.8.

All predefined views were imaged by monochrome and colour SMI using setting recommendations provided by the manufacturer. Grey-scale video sequences were acquired and documented twice consecutively, while colour sequences were acquired and documented once per anatomical sub-region. The minimum duration of each video sequence was 5 s. SMI video sequences were stored and reviewed using AGFA PACS. Based on radiologic feature observations made during image acquisition, a structured semi-quantitative reading scheme was developed and used (Supplementary Material): following this scheme, vessels were classified as visible or invisible. The classification and nomenclature of the angioarchitecture was adopted from Nelson et al [20] and Okudera et al [7]. Specifically, we distinguished two main vascular territories: extrastriatal vessels (supplying cortex and white matter) and striatal vessels (supplying caudate and lentiform nucleus). These vessels were rated as visible, if they appeared as bright echogenic curvilinear structures on mSMI video sequences or as red curvilinear structures on cSMI sequences.

To test inter-rater and intra-rater reproducibility, images were read off-line twice by the main reader (JP) with a time interval of 6 months between readings and once by a second reader (KG).

### Statistical analysis

Statistical analysis was performed using IBM SPSS Statistics for Windows 23.0 (IBM Corporation). Metric data are described using means ± SD if normally distributed or as median (IQR) for skewed data. Nominal data are shown as counts and percentages. Because of the exploratory character of the study, no detailed statistical tests were performed. Due to small sample size and binary data yield of SMI readings, agreement was

| Table 1 | Descriptive data |
|---------|------------------|
|         | Mean ± SD; median (IQR) |
| Gestational age (weeks) | 38.74 ± 1.54 |
| Weight (g) | 2954.00 ± 539.68 |
| Length (cm) | 49.58 ± 2.78 |
| Head circumference (cm) | 34.03 ± 1.45 |
| Percentile weight | 30.16 ± 26.65 |
| Percentile length | 49.79 ± 31.33 |
| Percentile head circumference | 49.37 ± 24.39 |
| Apgar minute 1 | 9 (9–9) |
| Apgar minute 5 | 10 (10–10) |
| Apgar minute 10 | 10 (10–10) |
| Umbilical artery pH | 7.33 ± 0.06 |
| Delivered by caesarean section | 12/19 (63%) |
| Delivered by vaginal delivery | 7/19 (37%) |
| Female | 11/19 (58%) |
| Male | 8/19 (42%) |
| Singleton pregnancies | 15/19 (79%) |
| Twins | 4/9 (21%) |
assessed as percentage of concordat ratings instead of using Cohen’s kappa or ICC.

**Results**

In order to design and plan the main study, five neonates underwent a structured exploratory application of SMI which was compared with normal colour Doppler ultrasound: while normal colour Doppler was able to depict some striatal and extrastriatal microvessels, there was striking superiority of SMI with regard to anatomical detail and density of visible microvessels in all five cases (Fig. 4).

For the main study, 19 term born neonates born with a mean gestational age of 38.7 weeks (± 1.5 weeks SD) were included. Newborns included in this study were mainly hospitalised due to maternal problems and not due to neonatal disease. Neonatal diagnoses included transient hypoglycaemia and mild hyperbilirubinaemia. Seventeen (17/19, 90%) were inborn, two (2/19, 11%) were outborn and were transferred to our unit for monitoring or further treatment (e.g. glucose infusion, phototherapy). SMI examination was performed at a median age of 3 days (IQR 2–4) postpartum. Further patient characteristics are provided in Table 1.

Superficial mSMI showed extrastriatal and striatal microvessels (Figs. 1 and 2). Extrastriatal vessels were subdivided into cortical and medullary. Cortical microvessels (CV) appeared as short hyperechoic, parallel streaks perpendicular to the brain surface on coronal and sagittal views. Medullary microvessels (MV) appeared as curvilinear hyperechogenicities within the white matter displaying a characteristic “fountain-like” morphology on coronal views [7, 21]. On sagittal views, MV displayed a straight course. Deep coronal SMI showed striatal

**Table 2  Visibility of extrastriatal vessels on superficial scans**

|                      | Cortical microvessels | Medullary microvessels |
|----------------------|-----------------------|------------------------|
|                      | Coronal              | Sagittal               | Coronal              | Sagittal               |
|                      | Right F1             | Left F1                | Right F1             | Left F1                |
| mSMI                 | 19/19 (100%)         | 18/19 (95%)            | 12/19 (63%)          | 14/19 (74%)            | 19/19 (100%)         | 19/19 (100%)         | 17/19 (90%)          | 16/19 (84%)           |
| cSMI                 | 17/19 (90%)          | 19/19 (100%)           | 5/17 (29%)           | 8/17 (47%)             | 18/19 (95%)          | 19/19 (100%)         | 15/17 (88%)          | 14/17 (82%)           |

F1 superior frontal gyrus, mSMI monochrome SMI, cSMI colour SMI
microvessels appearing as “lotus-flower” shaped curvilinear hyperechogenicities coursing through the thalamus and basal ganglia (Figs. 3 and 5).

On superficial scans, visibility of CV and MV was comparable between the left and right scan regions. In general, SMI performance was better on coronal views than on sagittal views, as both CV and MV could be identified more frequently. On sagittal views, MV were easier to visualise than CV (Table 2). On deep scans, CV and MV were visible in all cases (17/17, 100%) and striatal microvessels in 12 cases (12/17, 71%) (Tables 3 and 4). Regarding repeated monochrome acquisitions and comparability of mSMI and cSMI, visibility of MV and CV was excellent on coronal scans and lower in the sagittal plane (Table 5).

We found excellent inter- and intra-rater agreement: both values were best for MV on coronal superficial scans and worst for CV on sagittal superficial scans (Table 3). On deep scans, concordant ratings were high for all different types of microvessels (Table 4).

### Table 3 Inter-rater and intra-rater agreement on superficial scans

|                 | A Cortical microvessels |               | Medullary microvessels |
|-----------------|-------------------------|---------------|------------------------|
|                 | Coronal | Sagittal | Coronal | Sagittal |
| Inter-rater     |         |         |         |         |
| 1st scan        | 95 (90–100) | 87 (84–90) | 100 (100–100) | 97 (95–100) |
| 2nd scan        | 97 (95–100) | 97 (95–100) | 100 (100–100) | 97 (95–100) |
| cSMI            | 97 (95–100) | 88 (82–94) | 100 (100–100) | 97 (94–100) |
| Intra-rater     |         |         |         |         |
| 1st scan        | 95 (95–95) | 87 (84–90) | 100 (100–100) | 92 (90–95) |
| 2nd scan        | 97 (95–100) | 79 (79–79) | 100 (100–100) | 90 (90–90) |
| cSMI            | 92 (90–95) | 82 (81–82) | 97 (95–100) | 100 (100–100) |

Data are shown as mean (min–max)
cSMI colour SMI

### Table 4 Inter-rater and intra-rater agreement on deep scans

|                 | B Cortical microvessels | Medullary microvessels | Striatal microvessels |
|-----------------|-------------------------|------------------------|-----------------------|
|                 | Coronal | Sagittal | Coronal | Sagittal |
| Inter-rater     |         |         |         |         |
| 100 (100–100)   | 100 (100–100) | 100 (100–100) |
| Intra-rater     |         |         |         |         |
| 94 (94–94)      | 100 (100–100) | 88 (88–88) |

Data are shown as mean (min–max)

### Table 5 Agreement of the visibility between two monochrome acquisitions (1st and 2nd scans) and between monochrome and colour SMI

|                 | Cortical microvessels | Medullary microvessels |
|-----------------|-----------------------|------------------------|
|                 | Coronal | Sagittal | Coronal | Sagittal |
| 1st and 2nd scans |         |         |         |         |
| Right F1        | 19/19 (100%) | 11/19 (58%) | 12/19 (63%) | 15/19 (79%) |
| Left F1         | 17/19 (90%) | 19/19 (100%) | 16/19 (84%) | 16/19 (84%) |
| mSMI and cSMI   | 16/19 (84%) | 18/19 (95%) | 5/17 (29%) | 15/17 (88%) |

F1 superior frontal gyrus, mSMI monochrome SMI, cSMI colour SMI

### Discussion

Transfontanellar SMI ultrasound in neonates was feasible and well reproducible. From our standpoint, SMI appears to be an extremely robust and reliable technique in the setting of neonatal ultrasound. Since initial reports in March 2016 [14], SMI has been mainly used in oncologic conditions (especially breast lesions) [22–26], infectious diseases [16, 27, 28], gastrointestinal disorders [29–32], obstetric ultrasound [33, 34], musculoskeletal [35], vascular [36], and endocrine disorders [14]. In paediatric imaging, SMI has been used to detect direction of urinary flow in patients with vesico-urethral reflux without the use of contrast [16] and to demonstrate differences in vascular flow grades between normal and undescended testes, which were invisible by Power Doppler [17, 18]. Our paper is not the first reporting the use of SMI in cerebral imaging, but it is the first with a dedicated neuropediatric imaging approach. So far, only one report described the use of SMI in the adult brain: Ishikawa et al performed intra-operative ultrasound with SMI in subjects undergoing open brain surgery. They were able to recognise tumour vessels and thus to differentiate tumour from surrounding healthy tissue [15].

Our observations are in line with anatomical literature describing the morphology and course of intraparenchymal small hemispheric arteries and veins [7, 37]: vessel patterns seen in the present study are highly reminiscent of those found in neonatal brain specimens. Others have used high-resolution brain ultrasound, specifically ultrafast Doppler for in vivo...
assessment of cerebral hemodynamics: Demené et al have shown similar microvascular anatomy but have focused on functional imaging based on neurovascular coupling. They were able to show subtle variation in local and global brain perfusion during different sleep stages and epileptic activity [38, 39]. As opposed to them, we are using SMI, a vendor-developed tool.

In order to be consistent with existing terminology, we have decided to use the neuro-anatomic nomenclature of Nelson et al [20] and Okudera et al [7], which only differentiates two main intraparenchymal microvascular territories: striatal and extrastriatal vessels. Based on strikingly distinct SMI morphology, we have further subclassified extrastriatal microvessels as cortical and medullary.

Fig. 6 Pathologic example: 3-month old patient with mitochondrial disease (Leigh disease) imaged by MRI (a), routine ultrasound (b and e), and SMI (c, d, f, and g). While MRI showed metabolic infarcts and signal alterations of the basal ganglia and cortex (a; white arrow), coronal mSMI showed posthypoxic cortical hyperperfusion (c and d; yellow arrow). In addition, SMI showed elongated and tortuous medullary microvessels (f and g; yellow arrowhead), which were confirmed by histology (not shown).
Although SMI depicted intraparenchymal microvasculature at extraordinary detail, neither monochrome nor colour SMI could differentiate arterial from venous microvessels. Vessel characterisation by spectral Doppler had not been defined as a priori-aim of the current study. Nevertheless, we have been able to obtain characteristic arterial and venous signals in extrastriatal, specifically MV in some cases (not shown). Based on the clear superiority of SMI to normal colour Doppler ultrasound (Fig. 4), we opted against a comparative study design. We found coronal SMI views easier to be obtained and read than sagittal views. Moreover, microvessels were better depicted on coronal views, which might be explained by the intrinsic three-dimensional anatomy of cerebral microvasculature [5, 40]. Comparing the performance of monochrome and colour SMI, our data suggest that mSMI was slightly superior to cSMI with regard to depiction of extrastriatal microvessels and image/data reproducibility. Only at a single assessment region, cSMI improved overall SMI performance. Superior performance of mSMI is most likely explained by methodical reasons: cSMI is technically optimised for the suppression of motion artefacts (both from patient and/or probe) and displays flow components in low velocity ranges in colour overlaid on the grey-scale image with high temporal and spatial resolution simultaneously. mSMI is further optimised for the detection of low flow: within the region of interest, B-mode (i.e. anatomical background) is removed from the image, resulting in high sensitivity for the depiction of microvasculature. Although we found mSMI generally superior to cSMI, cSMI might still be a relevant add-on for visual display.

From a clinical standpoint, it should be stressed that there is great potential for the use of SMI in the neonatal brain: hypoxia-ischemic encephalopathy, cerebral malformations, infections, and preterm birth are conditions associated with microvascular abnormalities. After ischemia, SMI might allow the identification and bedside monitoring of posthypoxic hyper-perfusion (Fig. 6) [6, 41]. In preterm neonates with intraventricular haemorrhage, SMI might allow the early identification of venous congestion and periventricular infarction following an intraventricular haemorrhage. In combination with neuro-physiological methods, the right timepoint for pressure-reducing interventions could be identified in the presence of post-haemorrhagic hydrocephalus.

The comparison between SMI and MRI as clinical gold standard is warranted in future studies. Bearing in mind that the development of intraparenchymal brain vasculature is a continuous process starting at embryonal stages of human life and continuing throughout infancy [42], longitudinal SMI studies including preterm infants are needed.

Our study has several limitations: the sample size was small. From a practical perspective, it should be mentioned that the size of the anterior fontanel is an important limiting factor for the diagnostic quality of transcranial ultrasound. Nevertheless, in our study cohort, no patient had to be excluded due to small anterior fontanel or bad quality images. Spontaneous movements represent another limiting factor to the feasibility of cerebral SMI in newborns. We have solved this issue by instructing the parent/accompanying nurse to gently stabilise the head in a neutral position with both hands on either side, by the use of a pacifier and/or oral sucrose, and by having two operators perform the exam: one focused on image acquisition, while the other was responsible for parameter setting and documentation according to the study protocol. We are aware that in daily practise, a single operator is most likely to be performing the exam but opted for this strict approach to ensure high standards in initial reporting.

In summary, we have demonstrated practical feasibility and excellent reproducibility of transfontanellar cerebral SMI ultrasound in healthy term-born neonates. Using superficial and deep scanning modes, we have described normal morphologic features of intraparenchymal brain microvasculature as “short parallel” cortical vessels, “smoothly curved” medullary vessels, and deep striatal vessels. We conclude from our study that SMI ultrasound provides a novel non-invasive imaging tool for the assessment of intraparenchymal brain vasculature in neonates without the use of contrast material.

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Compliance with ethical standards

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Statistics and biometry No complex statistical methods were necessary for this paper.

Informed consent Written informed consent was obtained from all subjects (patients) in this study.

Ethical approval Institutional Review Board approval was obtained.

Methodology

- Prospective
- Cross-sectional study
- Performed at one institution
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