Application of transcranial transmission ultrasound in the detection of vasospasm in patients with aneurysmal subarachnoid hemorrhage: illustrative cases

Maximilian Schwendner, MD,1,2 Leonie Kram, MSc,1,2 Haosu Zhang, MD, PhD,1,2 Ann-Kathrin Joerger, MD,1 Raimunde Liang, MD,1 Chiara Negwer, MD,1 Bernhard Meyer, MD,1 and Maria Wostrack, MD1

1Department of Neurosurgery, Technical University of Munich, School of Medicine, Klinikum rechts der Isar, Germany; and 2Department of Neurosurgery, Heidelberg University Hospital, Ruprecht-Karls-University Heidelberg, Heidelberg, Germany

BACKGROUND Effective management of patients with aneurysmal subarachnoid hemorrhage (aSAH) demands vigilant monitoring and treatment, given the risks of complications such as cerebral vasospasm and delayed ischemic neurological deficits (DINDs). Transcranial transmission ultrasound (TTUS) is a well-established technique for assessing brain pulsatility. This pilot study aims to explore the utility of TTUS in detecting impaired intracerebral blood flow associated with DINDs.

OBSERVATIONS The authors examined 2 male patients, ages 45 and 52 years, with aSAH Hunt and Hess grades 4 and 2, respectively, who developed DINDs during their clinical course. Simultaneous recordings of arterial blood pressure, heart rate, and TTUS measurements were obtained in the intensive care unit. TTUS analysis revealed abnormal arrhythmic wave patterns during DIND episodes, whereas baseline measurements on DIND-free days showed no abnormalities. Following endovascular spasmolysis, TTUS demonstrated a normalization of abnormal waves, returning to baseline levels, alongside the resolution of neurological symptoms.

LESSONS TTUS, a noninvasive method for assessing brain pulsatility, shows promise as a novel tool for monitoring aSAH patients, potentially aiding in prompt diagnostics and additional therapeutic interventions. Its capacity to provide further insights for individuals at risk of delayed cerebral ischemia warrants further investigation in clinical studies.

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KEYWORDS aneurysmal subarachnoid hemorrhage; vasospasm; noninvasive; ultrasound

Aneurysmal subarachnoid hemorrhage (aSAH) is a devastating disease because of its high mortality rate of over 30% and its high incidence of severe disability or cognitive impairment in surviving patients.1,2 Five percent of all strokes result from ruptured intracranial aneurysms and the subsequent subarachnoid hemorrhage.1

Cerebral vasospasm (CVS) is the most frequent complication after aSAH and occurs between days 3 and 14 after intracranial aneurysm rupture, with maximal expression between days 6 and 8. Angiographically apparent CVS occurs in more than half of all patients, but the relationship between angiographic CVS and clinical symptoms due to impaired cerebral perfusion is not fully understood.3,5 The feared clinical sequelae of CVS is delayed ischemic neurological deficit (DIND) and delayed cerebral ischemia (DCI), occurring in roughly 30% of aSAH patients with CVS.6,7

Although CVS after aSAH has been the subject of several studies, the exact mechanisms of its development remain largely unclear to date. The pathogenesis is obviously multifactorial and highly complex.6 Recently, in addition to a radiographically evident macrovasospasm, microvasospasm and cortical spreading depression, as well as inflammatory processes, have been discussed.6 Concurrently with vasospasm, electrolyte imbalances, notably hyponatremia, natriuresis, and hypovolemia, frequently emerge, contributing to the development of DINDs.10,11

Reducing the incidence of DINDs and DCI resulting from CVS can be accomplished in certain cases through measures such as administering oral nimodipine, elevating arterial blood pressure, and, as salvage therapy, conducting endovascular vasospasmolysis.12 Still, a prerequisite of therapeutic success remains the early detection of...
symptomatic CVS. Transcranial Doppler (TCD) ultrasonography, digital subtraction angiography (DSA), computed tomography (CT) angiography, CT perfusion, and perfusion magnetic resonance imaging (MRI) can document changes in vessel caliber either directly or indirectly. However, each technique has its own disadvantages. Imaging techniques like DSA, CT, and MRI pose logistical challenges and provide only a snapshot of vascular status. Additionally, CT and DSA involve significant exposure to ionizing radiation, making them unsuitable for daily monitoring diagnostics.

TCD ultrasonography is a portable examination technique that is free of ionizing radiation, enabling assessment of the large vessels in the circle of Willis. However, several studies have demonstrated reliable identification of CVS sufficient for clinical application, primarily in the middle cerebral artery (MCA). Additionally, correct diagnosis is even more difficult in unconscious or sedated patients, in whom DCI is often detected only after irreversible ischemic changes have already occurred. Therefore, there is great clinical interest in a noninvasive, portable method that allows for the assessment of clinically relevant CVS.

Transcranial transmission ultrasound (TTUS) is a noninvasive technology that acquires changes in the density and volume of brain tissue arising from cerebral blood flow via ultrasound pulses propagating along the human brain. For selected frequencies, a dispersive character of the brain tissue has been demonstrated. Dispersion is an effect in which the medium's nonlinear, frequency-dependent compression modulus results in different propagation velocities for different ultrasound frequencies. In this study, we report on 2 male patients with aSAH who underwent TTUS measurements as an additional diagnostic tool during DINDs. To our knowledge, this is the first report on the application of TTUS to evaluate DINDs in aSAH.

Illustrative Cases

Data Acquisition

For the data analysis, only patients experiencing DINDs were included. Arterial blood pressure, heart rate, and TTUS measurements were simultaneously recorded in conditions with and without DIND. A DIND was defined as the occurrence of a new focal neurological deficit and/or a decrease of at least 2 points on the Glasgow Coma Scale (GCS) after excluding other potential causes such as postoperative hematoma or seizures. TTUS measurements (UltraEasy 3ACG, Sonovum GmbH) were performed daily within a short interval after the TCD measurements were acquired in the clinical routine. The setup to perform TTUS measurements consisted of a transmitting and a receiving ultrasound probe set up perpendicular to the head surface (Fig. 1). Ultrasound gel was applied to achieve sufficient coupling of the ultrasound. A series of 80 ultrasound waveform packages were accumulated to a single data point to increment the signal-to-noise ratio, leading to the accumulated time-of-flight signal. The accumulated time of flight was then registered. TCD measurements were recorded as the mean velocity (Vmean) of the flow in the anterior communicating artery, MCA, and posterior communicating artery.

Pathological TTUS measurements, TCD measurements showing an increased Vmean, and the occurrence of DIND events were recorded. The timeline illustration provides a combined overview of these parameters (Fig. 2). The preprocessing of TTUS measurements involved the identification and exclusion of episodes with high noise based on a frequency analysis of pattern > 10 Hz using fast Fourier transformation. Further data analyses were done using Python 3.19.14. After preprocessing, 2 data scientists manually evaluated the TTUS measurements in a blinded manner for characteristic patterns associated with the DIND events.

Findings

Two male patients, aged 52 and 45 years, were examined between January 2022 and April 2023. Initial aSAH grades according to the Hunt and Hess classification were 2 and 4, respectively. The aneurysms were in the pericallosal artery and anterior communicating artery, and both aneurysms were occluded via an endovascular approach. In the first patient, an external ventricular drain was placed for monitoring intracranial pressure (ICP) and drainage of cerebrospinal fluid, which did not indicate elevated ICP throughout all TTUS measurements performed. Nine TTUS measurements were performed in the first patient, and 5 were performed in the second patient. No adverse events were reported regarding TTUS measurements.
DINDs occurred twice in the first patient on days 10 and 13 after aSAH onset, respectively, and once in the second patient on day 8. Symptoms included a decrease in the GCS score and new motor deficits, respectively. Endovascular spasmolysis was performed in all cases of DIND according to the internal standard-of-care protocol. TTUS measurements were available for both patients before, during the onset of, and after the DIND events. During the days without neurological decline, typical rhythmic waves were recorded in these patients, as shown in Fig. 3A. Data analysis did not reveal any apparent changes in TTUS signals preceding the onset of a DIND. However, a distinct signal displaying arrhythmic waveforms was consistently detected whenever TTUS was performed concurrently with the DIND episodes (Fig. 3B). Notably, the TTUS waveforms promptly normalized following resolution of the DIND after effective spasmolysis interventions (Fig. 3C).

**Patient Informed Consent**

The necessary patient informed consent was obtained in this study.

**Discussion**

**Observations**

Apart from urgent occlusion of the ruptured aneurysm, intensive care unit treatment and extensive neurological monitoring are crucial in the treatment of aSAH. Early detection of neurological deterioration is essential to prevent irreversible damage resulting in persisting disabilities or death. Symptomatic CVS requires immediate diagnostics and additional treatment. However, especially in patients with a compromised neurological status and impaired consciousness, performing detailed neurological monitoring is notably challenging. Further diagnostic tools to allow for early identification of patients at risk and tools to enable a more thorough examination of patients with an impaired neurological status are warranted.

TCD ultrasonography, as a standard cerebrovascular monitoring tool, often falls short of reliably identifying clinically relevant CVSs.\(^{15,16}\) Additionally, the routine assessment of smaller vessels is challenging, leading to the underdiagnosis of microvasospasms.
In the present study, we evaluated the diagnostic value of TTUS in assessing CVS in patients with ASAH. The application of TTUS was straightforward and easily applicable in an intensive care unit setting, as previously described. During multiple measurements, no adverse events or technical issues were observed.

Although no linear change in the TTUS measurements allowing for direct correlation or automated analysis was observed, specific arrhythmic pulse waveforms occurred in both patients during DINDs as compared to baseline wave patterns and resolved with improved DINDs (Fig. 3). In these cases, TTUS was able both to identify DIND and to determine treatment success by resolving neurological symptoms. This suggests that TTUS may serve as a tool for identifying patients experiencing neurological decline due to the perfusion deficit associated with CVS beyond merely measuring the apparatus-detectable CVS.

No clear early changes in TTUS were identified before the onset of a DIND in this study. However, the measurements were conducted intermittently, potentially introducing gaps in data acquisition. Larger-scale investigations are needed to explore the potential prognostic value of TTUS, where alterations in wave patterns may indicate the onset of a DIND at a later stage.

A significant question lingers regarding the physiological explanation of the phenomenon of how TTUS can detect the onset of neurological deficits associated with perfusion deficits. In our opinion, the most plausible theory is related to changes in brain compliance: CVSs induce blood-brain barrier opening, cytotoxic edema, ischemic components, and altered cerebrovascular autoregulation, resulting in changes in blood flow, a shift in volume, and by this, alterations in compliance. These changes probably result in wave pattern alterations as assessed by TTUS.

Lessons

To our knowledge, the present case study is the first to explore the potential of TTUS as a noninvasive method for assessing clinically relevant CVS. TTUS demonstrated promise in identifying patients with DINDs associated with CVS. The dynamic changes in TTUS waveforms correlated with the occurrence of DINDs and reflected treatment success following interventional spasmolysis. These findings suggest that TTUS may serve as an additional novel tool for monitoring neurological decline following CVS. Further investigations with standardized evaluation protocols are warranted to explore the potential prognostic value of TTUS.

References


Disclosures

Dr. Schwender reported that the transcranial transmission ultrasound device was provided by Sonovum GmbH. Dr. Meyer reported being a shareholder in Sonovum GmbH during the conduct of the study; grants from Brainlab and Zeiss; royalties from Spineart and Medacta; and personal fees from Companionspine outside the submitted work. Dr. Wostrack reported that the ACG device for ACG measurements was provided free of charge by Sonovum GmbH during the study period.
Author Contributions

Correspondence
Maximilian Schwendner: Heidelberg University Hospital, Ruprecht-Karls-University Heidelberg, Germany. maximilian.schwendner@med.uni-heidelberg.de.