Physiology-Based MR Imaging Assessment of CSF Flow at the Foramen Magnum with a Valsalva Maneuver

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ABSTRACT

BACKGROUND AND PURPOSE: MR imaging is currently not used to evaluate CSF flow changes due to short-lasting physiological maneuvers. The purpose of this study was to evaluate the ability of MR imaging to assess the CSF flow response to a Valsalva maneuver in healthy participants.

MATERIALS AND METHODS: A cardiac-gated fast cine-PC sequence with ≤15-second acquisition time was used to assess CSF flow in 8 healthy participants at the foramen magnum at rest, during, and immediately after a controlled Valsalva maneuver. CSF mean displacement volume \( V_{CSF} \) during the cardiac cycle and CSF flow waveform \( A_{pp} \) were determined. A work-in-progress real-time pencil-beam imaging method with temporal resolution ≤56 ms was used to scan 2 participants for 90 seconds during which resting, Valsalva, and post-Valsalva CSF flow, respiration, and HR were continuously recorded. Results were qualitatively compared with invasive craniospinal differential pressure measurements from the literature.

RESULTS: Both methods showed 1) a decrease from baseline in \( V_{CSF} \) and \( A_{pp} \) during Valsalva and 2) an increase in \( V_{CSF} \) and \( A_{pp} \) immediately after Valsalva compared with values measured both at rest and during Valsalva. Whereas fast cine-PC produced a single CSF flow waveform that is an average over many cardiac cycles, pencil-beam imaging depicted waveforms for each heartbeat and was able to capture many dynamic features of CSF flow, including transients synchronized with the Valsalva maneuver.

CONCLUSIONS: Both fast cine-PC and pencil-beam imaging demonstrated expected changes in CSF flow with Valsalva maneuver in healthy participants. The real-time capability of pencil-beam imaging may be necessary to detect Valsalva-related transient CSF flow obstruction in patients with pathologic conditions such as Chiari I malformation.

ABBREVIATIONS: \( A_{pp} \) = CSF flow waveform peak-to-peak amplitude; cine-PC = cine phase-contrast; HR = heart rate; PBI = pencil-beam imaging; \( V_{CSF} \) = CSF mean displacement volume

Invasive pressure studies have shown that physiologic challenges such as the Valsalva maneuver or coughing produce transient changes in intracranial and intraspinal pressures. Although such transients are rarely of significance in healthy participants, for patients with foramen magnum obstruction such as Chiari I malformation, these physiologic alterations may be responsible for producing headache and syringomyelia. In the past, attempts were made to use CSF pressure measurements with physiologic challenges as a clinical test. However, these methods have not gained widespread acceptance because they are invasive. Although noninvasive assessment of CSF flow pulsations by MR imaging that measure bidirectional CSF flow from pressure changes between the head and spine during a cardiac cycle can be used as a surrogate, all but one of the prior implementations of cine-PC MR imaging for CSF flow assessment required approximately 100 heartbeats, making it too long to use with a transient physiologic challenge. As a result, most cine-PC studies of CSF flow have only been used during quiet breathing, both in healthy participants and in patients with suspected CSF circulation abnormalities. The 1 exception is a 2006 abstract by Wentland et al reporting CSF flow during rest and Valsalva by using an undersampled projection reconstruction method with data acquisi-
MATERIALS AND METHODS

Nine healthy participants (mean age, 34.0 ± 11.1 years; 4 women) underwent MR imaging on a 3T scanner (Achieva; Philips Healthcare, Best, the Netherlands). Institutional review board approval was obtained for this Health Insurance Portability and Accountability Act–compliant prospective study, and each participant signed an informed consent. None of the participants had a history of cardiovascular or neurologic disorders.

CSF Flow-Imaging Sequences

CSF flow pulsations were assessed with 2 different techniques. The first flow-imaging technique was a fast cine-PC sequence that was achieved by use of a combination of parallel imaging (sensitivity encoding factor = 2.1), half Fourier acquisition (half-Fourier factor = 0.6), and multiple-phase encodes per cardiac phase (turbo field-echo factor = 3). Other imaging parameters were TR, 11 ms; TE, 7.3 ms; field of view, 15 × 15 cm; and voxel size, 1 × 1 × 5 mm. The velocity encoding was performed at 5 cm/s in all participants. This was repeated at 7 cm/s in a single patient because of aliasing. The sequence was retrospectively gated to the heartbeat via electrocardiographic leads.

The second flow-imaging technique was PBI. This pulse sequence excites a narrow cylinder or “pencil” region and has been described in detail previously.16,17 Because this sequence is not available for routine clinical use, a research version was used. The sequence consisted of a nominal 20-mm diameter cylindrical excitation by using a 2D spatially localized radio-frequency pulse with 8-turn spiral gradients, followed by a bipolar velocity-encoding gradient and a readout gradient applied along the axis of the cylinder (pencil beam). PBI thus provides spatial resolution only along the axis of the cylinder. The pencil beam was 64 mm in length, with a velocity encoding of 5 cm/s along the superior to inferior direction. Other imaging parameters were TR, 25–28 ms; TE, 3.8 ms; and flip angle, 10°. The bipolar phase-encoding gradient was alternated on sequential TRs, making the effective temporal resolution 2 × TR (ie, 50–56 ms). No gating was used, but HR (from the electrocardiogram) and respiration (from the respiratory bellows) were continuously monitored by using the scanner’s physiologic recording system.

Confirmation of Bipolar Phase-Encoding Gradient Velocity Encoding

Using a MR imaging-compatible flow pump (Compuflow 1000; Shelly Medical Technologies, Ontario, Canada), we performed a phantom study. The pump was programmed to provide waveforms with various shapes, flow rates, and amplitudes (in the physiologic range of CSF flow rates and amplitudes). The MR-measured flow in the phantom experiments was in excellent agreement (± 5%) with the actual flow programmed in the pump.

Valsalva Maneuver

A nonmagnetic mechanical bellows device was designed and built to perform a controlled Valsalva maneuver in the MR imaging environment (Fig 1). It consists of flexible plastic bellows tubing closed at 1 end, and with the extension of the closed end calibrated for pressures ranging from 20–40 cm H2O. To perform a Valsalva maneuver, a participant was asked to blow into the tube until the pressure reached a target pressure, typically 40 cm H2O, indicated by the bellows extension to a mark on the cylinder. The participant was asked to

FIG1. Schematic of Valsalva device used for experiments. Participants were directed to exhale into the plastic tubing with sufficient force to hold the end of the bellows at the 40-cm H2O mark on the clear plastic cylinder. This provided a consistent and reproducible Valsalva-generated pressure.
The On-line figure shows PBI results in the 2 participants studied. Although PBI can display flow at various positions along the 6.4-cm-long excitation cylinder, we focused on measurements restricted to the level of the C1 vertebra for this preliminary study. Although both PBI and fast cine-PC can display flow at various positions along the 6.4-cm-long excitation cylinder, we focused on measurements restricted to the level of the C1 vertebra for this preliminary study. Although both PBI and fast cine-PC can display flow at various positions along the 6.4-cm-long excitation cylinder, we focused on measurements restricted to the level of the C1 vertebra for this preliminary study.
Table 1: CSF flow displacement volume per cardiac cycle, peak-to-peak amplitude, and heart rate at rest, during, and post-Valsalva maneuver, from fast cine-PC measurements

<table>
<thead>
<tr>
<th></th>
<th>R</th>
<th>VAL</th>
<th>P-VAL</th>
<th>R vs VAL</th>
<th>VAL vs P-VAL</th>
<th>R vs P-VAL</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean CSF $V_{CSF}$ (mL) per cardiac cycle</td>
<td>0.57 ± 0.24</td>
<td>0.37 ± 0.17</td>
<td>0.68 ± 0.19</td>
<td>&lt;.001</td>
<td>&lt;.001</td>
<td>&lt;.05</td>
</tr>
<tr>
<td>$A_{pp}$ (mL/s)</td>
<td>4.51 ± 1.61</td>
<td>3.52 ± 1.59</td>
<td>5.28 ± 1.49</td>
<td>&lt;.007</td>
<td>&lt;.001</td>
<td>&lt;.01</td>
</tr>
<tr>
<td>Heart rate (beats/min)</td>
<td>67.1 ± 10.4</td>
<td>78.5 ± 9.9</td>
<td>64.22 ± 6.6</td>
<td>&lt;.002</td>
<td>&lt;.001</td>
<td>NS</td>
</tr>
</tbody>
</table>

NS indicates not significant; P-VAL, post-Valsalva; R, resting; VAL, Valsalva.

Note: All values are expressed as mean ± SD.

**FIG 3.** Cardiac cycle–dependent CSF flow before, during, and after a Valsalva maneuver in 1 participant by using fast cine-PC imaging. Flow in milliliters per second is shown as a function of time, normalized to the cardiac cycle. During Valsalva, $V_{CSF}$ and $A_{pp}$ decreased compared with resting. After Valsalva, both $V_{CSF}$ and $A_{pp}$ increased compared with during the maneuver, rebounding to values larger than resting.

**Qualitative Comparison with Previous Invasive Pressure Measurements**

Both flow-imaging methods showed a CSF flow response to Valsalva consistent with invasive craniospinal differential pressure measurements by Williams,1 (Fig 4, trace 4). The advantage of measuring CSF flow in real-time with PBI compared with fast cine-PC can be most appreciated by comparing the On-line figure with Fig 4. Here, similarities between transient changes in CSF flow waveforms during the resting, Valsalva, and post-Valsalva periods and the craniospinal differential pressure tracing (trace 4) are most apparent.

**DISCUSSION**

The results show that both fast cine-PC and PBI can be used to assess changes in CSF flow in response to a physiologic challenge, and lead to objective analysis of differences in response. CSF flow between the head and the spine decreases during Valsalva and increases past the resting flow rate in the post-Valsalva period. These findings are consistent with previous experiments performed by Williams1 by using simultaneous invasive measure-ments of CSF pressure in the head and spine. Table 2 shows the expected respiratory and cardiovascular changes as well as expected changes in intracranial and spinal pressure and CSF flow in 4 phases of the Valsalva maneuver.1,18 During performance of the Valsalva maneuver (phases I–II), spinal CSF pressure increases because of transmission of high intrathoracic pressure and consequent distension of the epidural veins.1 The intracranial pressure also increases, but this increase is much less because of the associated decrease in cardiac output and reduced systolic arterial inflow to the head.1,18 The higher spinal pressure relative to intracranial pressure reduces CSF flow across the foramen magnum. Immediately after normal breathing resumes (phases III–IV), spinal pressure decreases because of reduction in intrathoracic pressure and collapse of the epidural veins. However, increased cardiac output and cranial arterial inflow momentarily increase intracranial pressure above resting value, resulting in the post-Valsalva rebound.1 The higher intracranial pressure relative to spinal pressure increases CSF flow across the foramen magnum.

Although fast cine-PC was able to show changes in CSF flow averaged during its 15-second acquisition time, PBI captured many dynamic features of CSF flow, including transients synchronized with the beginning and end of the Valsalva. In addition, our PBI results show that it is possible to noninvasively demonstrate heartbeat-to-heartbeat changes in CSF flow induced by differences in craniospinal pressure during Valsalva.1 This finding is evident by comparing CSF flow changes seen in the On-line Figure to invasive pressure measurements of Williams1 shown in Fig 4 (trace 4). Specifically, the gradual decrease in pressure during the Valsalva followed by the sharp rebound in pressure fluctuation amplitude on its termination is mirrored in the flow behavior revealed by the PBI approach. This is consistent with flow, especially in its transient behavior, being driven by corresponding fluctuations in pressure in the spinal column coincident with the heartbeat. It is important to note that these transient flow changes associated with the onset and termination of the Valsalva maneuver are seen with PBI and not with cine-PC. Finally, unlike fast cine-PC, PBI can simultaneously assess changes both above and below the level of the epidural veins.1 The intracranial pressure also increases, but this increase is much less because of the associated decrease in cardiac output and reduced systolic arterial inflow to the head.1,18 The higher spinal pressure relative to intracranial pressure reduces CSF flow across the foramen magnum. Immediately after normal breathing resumes (phases III–IV), spinal pressure decreases because of reduction in intrathoracic pressure and collapse of the epidural veins. However, increased cardiac output and cranial arterial inflow momentarily increase intracranial pressure above resting value, resulting in the post-Valsalva rebound.1 The higher intracranial pressure relative to spinal pressure increases CSF flow across the foramen magnum.

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of the foramen magnum, along the length of the excitation cylinder, and it does not require cardiac gating. This eliminates the possibility of gating interruption because of sudden changes in HR during the Valsalva maneuver (as described in the Results section). We believe that the real-time capability of PBI may be necessary to detect pathologically altered Valsalva-induced transient CSF flow changes in patients with pathologic conditions such as Chiari I malformation. PBI may also prove useful in evaluating a differential CSF flow response between pathology and health for other physiologic challenges such as cough, the Mueller maneuver, or neck flexion. Also, future work can include a Fourier analysis of the CSF flow waveform to determine changes synchronous with respiration during free breathing.

Although the physiology-based challenge approach we describe here to study alterations in CSF flow by MR imaging is not a novel idea, its potential has not been fully explored. The justification for this approach comes from numerous previous experimental studies on CSF pressure in humans that used a physiologic challenge to assess the CSF pressure dynamics. Because CSF flow estimation by MR imaging is currently done in resting conditions, limited clinical usefulness has been found for quantitative CSF flow imaging in patient populations. This is likely because of a high variability in CSF flow pulsation amplitudes among patients because of differences in cerebral arterial and venous flow amplitudes, intracranial compliance, and the anatomy of CSF pathways. The major benefit of our approach is that by comparing cardiac cycle–related CSF flow for a participant at rest with that during and after a Valsalva maneuver, compensatory CSF flow changes can be assessed on an individualized basis, eliminating the need to compare CSF flow across participants.

Although physiology-based assessment of CSF flow may have a variety of clinical applications, we believe that the most immediate and obvious application is in the assessment of CSF flow in patients with Chiari I malformation. It is generally agreed that many of the symptoms and signs associated with Chiari I malformation are attributed to abnormal CSF circulation between the head and the spine secondary to obstruction of the foramen magnum. Williams and others have previously demonstrated this in patients with Chiari I malformation. On the basis of these results, we hypothesize that in patients with Chiari I malformation with a significant foramen magnum obstruction, the immediate post-Valsalva rebound in CSF flow shown here will be either absent or significantly reduced. We further expect that PBI will be able to observe this altered CSF flow transient and be of diagnostic value. We plan to address this hypothesis in future work.

Although we believe that PBI is superior to fast cine-PC in studying transient alterations in CSF flow in response to a physiologic challenge, it is still an emerging technology for quantitative estimation of CSF flow. As such, there are several limitations. First, the MR phase signal is not completely determined by flow within the spinal cord, as the nominal boundaries of the pencil beam may extend outside of the cord, allowing contributions from other flow sources such as the epidural veins. The shape and size of the pencil beam need to be selected so that non-CSF flow is minimally included in the PBI cross-section. Second, quantitative estimates of volume displacement similar to that obtained with cine-PC require determination of the size of the CSF pathway being assessed, which may also change during the Valsalva maneuver. This will also be the subject of future work. Finally, PBI at present is available only as a research sequence that limits its wide-

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**Table 2: Effects of Valsalva maneuver on intrathoracic pressure, cardiovascular system, intracranial and spinal pressures, and CSF flow**

<table>
<thead>
<tr>
<th>Mechanical Effect</th>
<th>Phase I</th>
<th>Phase II</th>
<th>Phase III</th>
<th>Phase IV</th>
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<tbody>
<tr>
<td>Intrathoracic pressure</td>
<td></td>
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<tr>
<td>Venous return</td>
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<tr>
<td>Left ventricular stroke volume</td>
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<tr>
<td>Blood pressure</td>
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<tr>
<td>Pulse rate</td>
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<tr>
<td>Intracranial pressure</td>
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<tr>
<td>Intraspinal pressure</td>
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<tr>
<td>CSF flow across foramen magnum</td>
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</table>

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**FIG 4.** Figure shows invasive pressure measurements vs time. Trace 4 shows heartbeat-by-heartbeat changes in lumbar minus ventricular pressure (craniospinal pressure differential) with Valsalva. Note the similarities between trace 4 and CSF flow waveforms (panel 2) in the On-line figure. Note further that PBI captured many dynamic CSF flow features including transients precisely synchronized with timing of Valsalva similar to that seen here. Reproduced from Williams B. Simultaneous cerebral and spinal fluid pressure recordings. I. Technique, physiology, and normal results. Acta Neurochir (Wein) 1981; 58:167–85 with kind permission from Springer Science and Business Media.
spread use. However, we believe that research may stimulate MR imaging vendors to make this sequence more readily available for routine clinical use.

CONCLUSIONS
Both fast cine-PC and PBI demonstrate a CSF flow response to Valsalva, consistent with previous invasive pressure measurements. Because cine-PC is a 2D technique, one can determine the CSF cross-sectional area allowing computation of absolute CSF flow. PBI must rely on a separate anatomic scan to translate measured average velocity over the pencil beam cross-section to flow. On the contrary, PBI captures transient changes in flow that cine-PC cannot. We believe that the real-time capability of PBI may be necessary to detect transient CSF flow changes in a pathologically altered Valsalva response in a patient population.

Disclosures: Rafeeque Bhadelia—RELATED: Grant: Conquer Chiari Foundation,* Comments: money to pay for MR imaging scans, equipment, and volunteers. Neel Madan—RELATED: Grant: Conquer Chiari Foundation,* Comments: The institution received money for the research MR imaging scans. No money otherwise to me or my institution; UNRELATED: Consultancy: Near Infrared Imaging. Comments: Company developing a medical imaging device using photoacoustic tomography. There is no relevance to this study, other than it is in the broad category of medical imaging. Received stocks/options; Expert Testimony: expert testimony on imaging, although no cases relevant to this study. Yansong Zhao—UNRELATED: Employment: I am an employee of Philips Healthcare, Cleveland, Ohio. Mark Waghal—UNRELATED: Payment for Lectures (including service on speaker bureau): DoD,* NIH,* James Butler—RELATED: Consulting Fee or Honorarium: Beth Israel Deaconess Hospital. Samuel Patz—RELATED: Consulting Fee or Honorarium: Beth Israel Deaconess Hospital. Comments: I was paid as a Consultant Investigator for assistance with the research reported in the submitted manuscript. This also covered my assistance with data analysis and preparation of this manuscript. The Consultant fee was budgeted in a grant to Dr. Bhadelia (first author) and awarded to Beth Israel Deaconess Hospital from the Conquer Chiari Foundation. *(Money paid to institution.)

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