Disorders of CSF dynamics such as syringomyelia and obstructive hydrocephalus can be caused by thin mobile obstructive lesions not visible on traditional MRI sequences. New imaging techniques with balanced steady-state free precession (bSSFP) and dynamic imaging with bSSFP cine allow visualization of these pulsatile structures within the CSF space. The authors present 2 cases involving pediatric patients—one who developed presumed idiopathic syringomyelia and one with presumed communicating hydrocephalus in association with Pfeiffer syndrome—who harbored thin dynamic obstructive lesions seen on bSSFP cine studies using 1.5-T MRI.

In combination with traditional CSF cine studies and bSSFP, bSSFP cine sequence was able to detect dynamic membranous adhesions not seen on traditional MRI sequences. These previously undetectable lesions on traditional MRI sequences were the etiology of CSF obstruction, and tailored surgical approaches were performed to avoid shunting in both patients. These reports demonstrate the clinical utility for using these novel imaging tools for the detection of thin adhesions and dynamic lesions in the central nervous system. Balanced SSFP cine sequences can supplement conventional MR modalities to identify these otherwise poorly visualized lesions responsible for presumed communicating hydrocephalus or idiopathic syringomyelia.

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Disorders of CSF dynamics such as syringomyelia and hydrocephalus are relatively common disorders in children and frequently involve obstruction of CSF flow. CSF flow obstruction can often be caused by thin arachnoid adhesions or membranes. While effects of CSF obstruction can manifest as ventriculomegaly and syringomyelia that are easily identifiable on traditional T1- and T2-weighted MRI, the thin lesions themselves, which can be pulsatile, are nearly impossible to visualize with these conventional imaging techniques.

Recently, new MRI sequences have been developed which may offer a solution. Steady-state free precession (SSFP) is a type of gradient-echo steady-state coherent MRI that produces high spatial resolution images with high signal-to-noise and high contrast-to-noise ratios. This sequence utilizes a very short repetition time, employs radiofrequency pulses with a large flip angle, and balances gradients in all 3 directions. Commercial names for balanced SSFP (bSSFP) include: fast imaging employing steady-state acquisition (FIESTA) sequence for GE scanners, balanced fast field echo (bFFE) for Philips, and true fast imaging with steady-state precession (TrueFISP) for Siemens. These gradient echo sequences yield high signal of blood vessels and body fluid without using contrast material and can be used to highlight signals from the CSF space. MR cisternography using bSSFP sequence can effectively visualize the neurovascular structures within the cerebellopontine cistern without CSF pulsation artifacts in a short examination time. Balanced SSFP has been demonstrated to be extremely useful for detailed anatomy of the cisternal segments of the lower cranial nerves as well as cerebellopontine angle lesions. Balanced SSFP is also useful in detecting neurovascular compression of the trigeminal nerve.

However, bSSFP and other traditional MRI techniques are limited by their susceptibility to motion artifacts caused...
by heartbeat, breathing, or swallowing. While bSSFP is effective at detecting detailed structures within the CSF space, dynamic thin membranous adhesions are difficult to visualize using a non time-resolved approach since they move synchronously with CSF pulsations. A cardiac gated sequence called bSSFP cine has been reported to be an effective method for the functional diagnosis of cardiac disease. Similar to bSSFP, bSSFP cine uses shorter echo time relative to the repetition time. Balanced SSFP cine has been used in abdominal imaging as well as arteriography and venography to visualize structures that move with the cardiac cycle. A recent report showed that cardiac-gated bSSFP cine was able to detect pulsatile arachnoid spinal adhesions not seen on other MRI sequences in an adult patient with presumed idiopathic syringomyelia. This raises the possibility for its broader application in the pediatric population as part of the evaluation for dynamic CSF disorders.

Here, we present 2 cases—one involving a child with presumed idiopathic syringomyelia and the other involving a child with presumed communicating hydrocephalus—in which bSSFP and bSSFP cine studies were performed to search for dynamic obstructive lesions. Using these imaging techniques, we were able to localize the adhesions causing the CSF flow obstruction and tailor the surgical treatments to avoid shunting. These illustrative cases show that bSSFP may have great clinical value in the preoperative evaluation for conditions of altered CSF dynamics to localize obstructive lesions.

**Case Reports**

**Case 1**

This 10-year-old boy with Pfeiffer syndrome presented with hydrocephalus. He had been born with severe facial dysmorphism, and although a head CT performed shortly after his birth showed open sutures, by 16 months he had developed cranial pansynostosis. He underwent calvarial vault expansion and frontoorbital remodeling. Imaging performed in the perioperative period showed no hydrocephalus. However, at 6 years of age he developed headaches, and a CT scan revealed ventriculomegaly, which was presumed to be due to communicating hydrocephalus. Standard MRI sequences revealed a mild Chiari malformation and a stable posterior fossa arachnoid cyst (Fig. 2A), and a right occipital ventriculoperitoneal shunt was placed. The patient subsequently underwent Le Fort III midface osteotomy and midface advancement.

He was doing well until he presented with severe headache and fever and was found to have community-acquired Streptococcus pneumoniae meningitis from CSF cultures at the age of 10 years. A shunt tap confirmed the infection. The shunt was removed and an external ventricular drain was placed. He was treated with intravenous antibiotic therapy. The external ventricular drain continued to have high output (400–500 ml/day) despite weaning, and he underwent additional imaging for further evaluation. A CSF cine study showed flow obstruction from the sylvian aqueduct to the fourth ventricle with flow through the foramen magnum (Fig. 2B and Video 3).

**Case 2**

This 10-year-old boy with Pfeiffer syndrome presented with hydrocephalus. He had been born with severe facial dysmorphism, and although a head CT performed shortly after his birth showed open sutures, by 16 months he had developed cranial pansynostosis. He underwent calvarial vault expansion and frontoorbital remodeling. Imaging performed in the perioperative period showed no hydrocephalus. However, at 6 years of age he developed headaches, and a CT scan revealed ventriculomegaly, which was presumed to be due to communicating hydrocephalus. Standard MRI sequences revealed a mild Chiari malformation and a stable posterior fossa arachnoid cyst (Fig. 2A), and a right occipital ventriculoperitoneal shunt was placed. The patient subsequently underwent Le Fort III midface osteotomy and midface advancement.

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**VIDEO 2.** Case 1. Video clip from the cardiac-gated bSSFP cine sequence demonstrating mobile membrane at the dorsal cervical cord responsible for the syringomyelia. Copyright Doris D. Wang. Published with permission. Click here to view with Quicktime.

T1-weighted MRI, however, did not reveal aqueductal stenosis (Fig. 2A). Balanced SSFP MRI showed narrowing...
of the aqueduct (Fig. 2C) but did not definitively identify the obstructive lesion for performance of an endoscopic third ventriculostomy (ETV). On the other hand, bSSFP cine imaging revealed a membrane between the anterior tectum and the dorsal midbrain occluding the sylvian aqueduct (Fig. 2D and Video 4).

VIDEO 4. Case 2. Video clip from the cardiac-gated bSSFP cine sequence demonstrating adhesion between dorsal midbrain and tectum. Copyright Doris D. Wang. Published with permission. Click here to view with Media Player. Click here to view with Quicktime.

Given the definitive imaging evidence of aqueductal obstruction, an ETV was performed. This procedure successfully treated the patient's hydrocephalus, and as of this writing, he has had no clinical or imaging evidence of recurrent hydrocephalus for 20 months.

Discussion
Advances in neuroimaging techniques have been essential in the diagnosis and treatment of central nervous system disorders. With the advent of high-resolution, high-contrast MRI sequences, we are now able to discern small vessels, cranial nerves, and other structures that were once difficult to visualize. Here, we introduce the use of gradient echo sequences bSSFP and bSSFP cine study to evaluate dynamic lesions of the central nervous system that are extremely difficult to visualize on conventional MRI sequences.

These imaging techniques have potential widespread clinical value in identifying dynamic CSF obstructive lesions in pediatric patients in order to avoid the need for CSF shunting. As shown by these 2 illustrative cases, identifying dynamic obstructive lesions not seen on standard MRI sequences can alter the surgical management. In the case of syringomyelia, the pathophysiology usually involves CSF flow alteration or spinal cord tethering. However, obstructive lesions caused by arachnoid membranes cannot be directly identified by means of standard MRI techniques due to their mobile nature. Without removal or fenestration of the lesion causing flow obstruction, placement of syringopleural or syringosubarachnoid shunts is the standard treatment for syringomyelia. However, insertion of these shunts may be associated with morbidity, and the shunts are often prone to failure. Us-
ing bSSFP and bSSFP cine makes it possible to identify mobile adhesions and visualize their pulsations in real time, allowing for direct surgical treatment of the CSF obstruction without shunting. In our Case 1, it is interesting to note that the obstructive lesion was rostral to the syrinx at the craniocervical junction, which has also been described in cases where syringomyelia has improved following craniocervical decompression in patients without hindbrain herniation (Chiari 0). In some Chiari 0 patients, a fourth ventricular veil and intradural adhesions were encountered intraoperatively. Surgical lysis of these lesions can lead to resolution of the syringomyelia. While these lesions themselves were not seen on standard MRI, they may now be visualized with bSSFP.

In the diagnosis of hydrocephalus, using bSSFP and bSSFP cine in combination may have clinical value in detecting lesions that cause CSF flow obstruction. The etiology of hydrocephalus in craniostenosis is thought to be variable and multifactorial. The patient in our Case 2 was assumed to have communicating hydrocephalus based on a standard MRI study. Although there was a mild Chiari malformation, in the absence of aqueductal stenosis, the patient was initially treated with a ventriculoperitoneal shunt. However, further imaging with bSSFP suggested that there was an aqueductal obstruction, and bSSFP cine was able to confirm the presence of obstructive aqueductal lesion or stenosis not seen on the previous MRI. With the diagnosis of obstructive hydrocephalus, the patient was able to undergo successful treatment with an ETV instead of ventriculoperitoneal shunt placement.

The combination of CSF cine, bSSFP, and dynamic bSSFP cine studies offers several advantages. While cine study can detect CSF flow or obstruction, the image resolution is extremely poor. Balanced SSFP can accurately detect thin membranous adhesions not identifiable on traditional T1- and T2-weighted MR images. However, in the case of dynamic lesions, mobile membranes will appear as streak artifacts on bSSFP sequences. Cardiac-gated bSSFP cine sequences, on the other hand, have lower resolution compared with bSSFP, but their ability to discern dynamic components is uniquely useful. In addition, bSSFP sequences are relatively quick to obtain on a 1.5-T MR scanner; a cardiac-gated bSSFP sequence has an acquisition time of 1.5 minutes and does not require intravenous contrast administration. Therefore, when used in combination, these imaging techniques can not only detect CSF flow obstructions, but also localize the lesion, even down to the thinnest arachnoid adhesions.

Application of bSSFP sequences can identify obstructive lesions and allow for effective direct surgical treatment of these lesions without the need for CSF shunting. Currently, at Oakland Children’s Hospital, our protocol utilizes the combined CSF cine, bSSFP, and dynamic bSSFP cine studies in patients with newly diagnosed hydrocephalus, those with shunt failure who are being considered for ETV, and those with idiopathic syringomyelia. Future prospective studies can be designed to elucidate the frequency of adhesion or lesion detection with these sequences in presumed communicating hydrocephalus and idiopathic syringomyelia.

Balanced SSFP and dynamic bSSFP cine are exciting imaging tools that allow clinicians and researchers to visualize “invisible” lesions of the central nervous system. These imaging techniques can be extremely useful in evaluating patients with disorders of CSF flow dynamics.

References

Author Contributions
Conception and design: Wang, Sun. Acquisition of data: Wang, Martin. Analysis and interpretation of data: Wang. Drafting the article: Wang. Critically revising the article: all authors. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Wang. Study supervision: Sun.

Supplemental Information
Videos


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