Spina bifida is a congenital malformation in which the neural tube fails to develop and close properly, resulting in a defect in early neurulation. Open spina bifida, or myelomeningocele (MMC), is the most common and severe form of spina bifida. Due to altered neurodevelopment, individuals with MMC often have multiple central nervous system (CNS) abnormalities that include the spinal cord and multiple brain regions. We aim to describe these CNS abnormalities and discuss the range of possible neurobehavioral outcomes that can occur in individuals with MMC. These patterns should be recognized by the treatment team, including the neurosurgeon, so as to positively impact patient outcome and identify patients who may be experiencing a decline related to reversible neurosurgical disease.

Anatomical Basis for Neurobehavioral Changes

MMC affects multiple anatomical levels, causing abnormalities in the spinal cord, cerebellum, brainstem, corpus callosum, and cerebral ventricles (Fig. 1).

Spinal Cord Lesion

Individuals with MMC typically exhibit sensory and motor deficits below the level of the spinal lesion. The lesion can occur at different levels, with greater impacts on functional outcome associated with lesions located at higher spinal levels. Patients experience a range of deficits, including lower-limb weakness or paralysis, sensory loss, incontinence, and musculoskeletal anomalies. Thoracic-level lesions are associated with more severe brain abnormalities, which are in turn associated with worse neurobehavioral outcomes. Motor impairments may alter the developmental trajectory from birth because movement restrictions limit environmental exploration and learning experiences.

Chiari II Malformation

Chiari II malformation (CM-II) consists of several structural changes, most significantly deformity of the brainstem and cerebellum. Approximately 96% and 90% of children with spinal lesions at T12 or higher and below T12 meet criteria for CM-II, respectively. The CM-II unified theory posits that a cascade of interrelated, time-
dependent defects unfolds during embryonic and fetal development, resulting in pathophysiology. Leakage of CSF through the spinal defect prevents normal distention of the embryonic ventricular system, limiting growth of the posterior fossa. The cerebellum develops in a smaller posterior fossa, resulting in displacement of the cerebellar tonsils, vermis, and brainstem through the foramen magnum into the spinal canal. Consequently, abnormalities may occur in other cerebral areas as well, including corpus callosum (CC) hypoplasia, tectal beaking, polymicrogyria, and cortical heterotopias. Thus, CM-II can lead to a constellation of brain abnormalities that impact neurodevelopment and compromise aspects of neurobehavioral functioning.

Cerebellum

The cerebellum has long been considered important for the timing and coordination of movement. Damage to the cerebellum generally leads to difficulties with motor control, including truncal and axial movements. More recently, the cerebellum has emerged as critical in cognition as well, particularly executive functioning, due to corticocerebellar pathways. Individuals with MMC typically have a reduced cerebellar volume, with further reductions seen with higher-level lesions. Cerebellar reorganization also results in a larger anterior portion and smaller posterior-inferior region. While the anterior lobe of the cerebellum is generally associated with motor function, individuals with MMC often show deficits in the timing and precision of truncal, upper-limb, and eye movements.

Brainstem and Midbrain

Brainstem and midbrain structures can develop abnormally due to overcrowding in the small posterior fossa and herniation of hindbrain structures through the foramen magnum. Approximately 65% of individuals with MMC have distortion of the midbrain, often marked by tectal beaking in which the colliculi fuse into a single beak. The medulla is elongated and kinked at the spinomedullary junction in approximately 70% of patients. The cerebellum and midbrain structures are both involved in control of eye movements, which tend to be impaired in individuals with MMC. Abnormalities in the midbrain and tectum are often associated with impairment in attention orienting as well.

Corpus Callosum

The CC is the largest subcortical white matter tract involved in interhemispheric transfer and integration of information from one or both hemispheres. Abnormalities of the CC are seen in approximately 70%–90% of individuals with CM-II and usually consist of hypoplasia or hypogenesis. The CC has four subdivisions: rostrum, genu, body, and splenium. In MMC, CC abnormalities primarily occur in the dorsal part (i.e., splenium), whereas the anterior parts tend to be intact. However, significant variability has been observed in the spectrum of CC abnormalities in MMC, which include a combination of congenital and acquired (from complications like hydrocephalus) effects on cerebral white matter. Individuals with MMC demonstrate deficits on cognitive tasks requiring recruitment of posterior networks and interhemispheric transfer of information, and severity of impairment is partially dependent on the integrity of the CC.

Hydrocephalus

Hydrocephalus is often conceptualized as a secondary CNS insult related to the CM-II in MMC. It may be caused by aqueductal occlusion, obstruction of the fourth ventricle, or eradication of the subarachnoid space by the overcrowded posterior fossa. Hydrocephalus occurs in approximately 85%–90% of individuals with MMC without fetal neurosurgical intervention. Expansion of the ventricles typically occurs in a posterior-to-anterior direction and can result in the stretching of parietal and occipital lobe axons around the dilated posterior horns of the lateral ventricles. Hydrocephalus is often treated with shunt...
placement or endoscopic third ventriculostomy. Unfortunately, shunts can become infected or blocked, and hydrocephalus can recur. Individuals with shunts may require multiple surgical revisions, placing them at risk for continued cerebral damage and infection. Children with MMC and either arrested hydrocephalus (no shunt) or shunted hydrocephalus tend to exhibit worse neurobehavioral outcomes than children with MMC who do not have hydrocephalus.23,34

Other Regions of the Cerebrum

Additional cerebral abnormalities can occur in MMC but are much less common. Gray matter heterotopias occur in about 19% of MMC cases.56 Seizures occur in approximately 15%–25% of MMC patients and are likely related to heterotopias and polymicrogyria.11,15 Seizures and additional brain abnormalities have added deleterious effects on neurobehavioral outcomes.

Neurobehavioral Profile of MMC

As the result of these neurological and medical sequelae, children with MMC are at increased risk of neurobehavioral difficulties. A pattern of specific neurobehavioral challenges constitutes a “characteristic” profile often seen in MMC; that said, cognitive difficulties can be significantly impacted by environmental, psychosocial, and demographic factors, including medical/developmental interventions, family socioeconomic status (SES), and race/ethnicity, which leads to individual variability in specific difficulties. Existing research about neurocognitive profiles in MMC has led to the development of a model of information processing and cognitive phenotypes to explain this variability.23 Briefly, this model describes two modes of processing: assembled processing and associative processing. Assembled processing refers to integration of information into larger unified concepts, while associative processing refers to the grouping or categorization of information.23 Assembled processing is typically thought to be negatively affected by the underlying neuropathophysiology and associated core deficits of MMC. In contrast, associative processing is typically less affected by the underlying neurophysiological changes of MMC but is more vulnerable to differences in the environment, like family SES and education. For a more detailed discussion of this model, see an overview written by Fletcher and Dennis.31 The following sections describe how differential strengths and weaknesses in assembled and associative processing tend to manifest in each neurobehavioral domain.

Intelligence

Individuals with MMC generally have slightly lower IQs than those of the general population, though the presence of complications affects outcomes. Individuals with MMC without hydrocephalus have an IQ about one-half a standard deviation below the mean but still remain within the average range.34,52 The presence of hydrocephalus is associated with a decrease of about 1 full standard deviation, leading to mean scores in the low-average range.34,52 Shunt-related complications can further negatively impact IQ, as well.34 Children with MMC also tend to show relatively stronger verbal reasoning and relatively weaker non-verbal reasoning.23,34

Attention

Attention is generally conceptualized as a 3-part system, with distinct networks guiding 3 key aspects: alerting, orienting, and executive control.59 This system is divided into anterior and posterior networks, which control alerting and sustained attention, and attention orientation, respectively. While these systems certainly overlap in day-to-day attention, children with MMC generally have more difficulty with orienting attention and shifting focus, likely due to structural and functional abnormalities in brain regions involved in the posterior attention system. They show slowed orientation to both internal and external cues and take longer to reorient their attention to an area they have already attended to (also known as inhibition of return) as compared with healthy controls.18,19 In contrast, sustained attention is usually fairly age appropriate in MMC,13 which is thought to reflect a more typical anterior attention system.51

Executive Functions

Executive functions are a broad group of cognitive skills that help an individual work toward complex goals, including initiation, inhibition, sequencing, fluency, planning, integration, and organization.26 While frontal lobe networks have long been implicated in executive functions, cerebellum and corticocerebellar pathways also play a role.50 In MMC, altered cerebellar structure affects several aspects of executive functions, including timing, sequencing of information, and organization,23 with executive function deficits persisting in individuals with MMC even when IQ is controlled for.34,64 Some aspects of executive function difficulty, such as speed and integration of context to understand metaphors, are thought to be associated with CC dysgenesis.43

Language

Language is often hailed as a strength for children with MMC. In the aspects of language involving more concrete knowledge, such as grammar and lexicon, individuals with MMC generally show age-appropriate performance.28 However, they often struggle more with higher-level tasks, such as understanding others’ meaning, using language in context, and speaking in a complex manner.7 These difficulties tend to appear in both spoken and written language.7

Memory

Explicit memory refers to intentional recollection of prior experiences or previously learned information, while implicit memory is the use of learned information without conscious awareness. Prospective memory is the concept of remembering to do something in the future (such as doing a task at a specific time). Sometimes categorized as part of executive function, working memory is a system that holds information in mind for immediate processing. These specific learning and memory functions appear to be differentially impacted in MMC, such that individu-
als with MMC often have strengths in rote memorization and show fairly typical implicit memory but have more difficulty with explicit and prospective memory, which is thought to be due to the disrupted development of subcortical structures and associated networks. Individuals with MMC also struggle to use previously learned information flexibly or in novel ways because of their executive function difficulties. Furthermore, these patients show difficulties with reconstructive memory as opposed to recognition or repetition. Working memory appears to generally be adequate with simple information but decreases with complex demands.

Visuospatial Functions
Children with MMC often show weaknesses in visuospatial functions, including perception, construction, and analysis. Many of the structural and functional changes that occur in neurodevelopment with MMC affect the posterior portion of the brain, where many networks involved in visual function are based. Hydrocephalus also tends to exert greater impact on posterior brain regions. Furthermore, the cerebellum is important for aspects of visuospatial functions, including timing and sequencing of information. Weaker visuospatial skills in MMC may also be related to the transfer and integration of information between visual areas of both hemispheres via the CC.

Motor Functions
Children with MMC have impaired movement and motor functions. These difficulties are thought to arise from physiological changes to the cerebellum and midbrain, as well as spinal lesion level. Although individuals with MMC typically retain the ability to learn motoric tasks, they have much more difficulty with precise timing and the smoothness of motor functions, resulting in imprecise and arrhythmic movements often manifesting as dysmetria, ataxia, or dysarthria. The combined motor and visuospatial deficits in MMC can lead to difficulties with real-world tasks such as handwriting and drawing.

Academic Functions
Children with MMC have an increased risk of learning and academic difficulties, which can in turn yield negative psychological and occupational outcomes. They generally decode adequately and read fluently, but they struggle to understand the meaning of passages and stories. Similarly, they can memorize math facts appropriately but struggle to apply numeric knowledge conceptually and use the information flexibly. In both instances, children can perform in an age-appropriate manner on basic assessments (spelling/math worksheets) but may struggle with real-world tasks or schoolwork involving concepts and flexibility.

Psychosocial and Adaptive Concerns
Individuals with MMC show increased risk of psychological difficulties, including both internalizing (e.g., depression, anxiety) and externalizing (e.g., impulsive behavior) symptoms. They also exhibit social difficulties, including decreased acceptability by peers and social competence, and lower-quality friendships. The social difficulties are thought to arise both from underlying neurocognitive difficulties and awareness of overt differences from typically developing peers. Furthermore, adaptive functioning outcomes are important metrics of independence and quality of life. Both the individual’s specific pathophysiology and aspects of their environment affect adaptive outcomes. Lesion level and severity of motor deficits, as well as higher number of treatments needed in cases with hydrocephalus, significantly impact adaptive and occupational functioning. Additionally, parents of children with MMC have been found to show higher levels of intrusiveness, psychological control, and authoritarian-style parenting which are associated with lower decision-making autonomy in the child and more behavioral concerns. The cognitive difficulties common in MMC can also impact adaptive functioning. In particular, poorer executive functions, like planning and cognitive flexibility, have been linked to decreased adaptive functioning.

Variability in Presentation
While the strengths and weaknesses described above are typically what are seen in MMC, presentation varies significantly due to variability in physiological and environmental factors. Indeed, most neurobehavioral outcomes tend to be worse in individuals with MMC who develop hydrocephalus than in those who do not. These differences are thought to result from both the presence of hydrocephalus, which itself is known to significantly affect cognitive functioning, and the underlying physiological changes that lead to hydrocephalus, like posterior fossa crowding, CM-II, and hindbrain herniation. While hydrocephalus treatment varies by child, with some children needing little to no intervention and others requiring serial shunt placements and revisions, the overall presence of hydrocephalus generally signals more impairment in neurobehavioral functioning.

Additionally, when considering characteristics of the common neurobehavioral profile seen with MMC, it is important to take into account environmental factors that impact function and outcomes. The child and family’s race/ethnicity, SES, and family cohesion or conflict have been repeatedly found to influence outcomes. These environmental aspects are thought to be associated with outcomes because of the multidirectional nature of how a child and his/her environment influence one another, such that children with specific experiences (e.g., living in a household with limited resources or experiencing increased stress from family conflict) interact with their environments differently, in turn shaping their learning experiences.

Current Neurosurgical Treatments
The primary treatment for MMC is surgical closure of the spinal lesion. Additional treatments involve medical and surgical management of hydrocephalus, spinal cysts, and tethered cord. While closure of the spinal lesion is most commonly done in the immediate postnatal period, the emergence of new surgical techniques in the last 2 decades now allows for in utero closure in a subset of patients.
meeting specific criteria. Only a fraction of patients diagnosed with MMC during pregnancy qualify for prenatal surgery, and the majority of patients continue to undergo postnatal treatments. Prenatal closure is associated with improved motor functioning, reversal of hindbrain herniation, and decreased risk for a shunt by 30 months of age as compared with postnatal closure. Since the majority of research on neurobehavioral outcomes was done with children who underwent postnatal closure, updated studies examining outcomes in children with MMC who underwent prenatal closure are needed. One concern related to prenatal closure is the increased risk of preterm birth, which can have its own deleterious effects on neurocognitive development. Ongoing research is examining the efficacy of alternate approaches for in utero closure (i.e., modifications of surgical techniques, fetoscopic methods) in attempts to mitigate this increased risk for preterm birth.

Additionally, recent changes in the treatment of hydrocephalus are also likely to impact medical and neurodevelopmental outcomes. Historically, around 80% of children with postnatal MMC closure have required treatment of hydrocephalus with shunt placement. However, hydrocephalus treatments have evolved over time such that some neurosurgical centers now tend to avoid shunting as long as possible. Ongoing research and debate regarding the appropriate threshold for intervention continues with toleration of larger ventricles and use of intracranial pressure and neurobehavioral symptoms as new metrics for determining when to intervene. An additional change is the increasing use of endoscopic third ventriculostomy with choroid plexus coagulation, which is associated with fewer follow-up procedures when successful, in lieu of a shunt. Furthermore, since prenatal closure is associated with a decreased risk for hydrocephalus compared with postnatal closure, rates of hydrocephalus among all children with MMC will continue to change as more prenatal closures are conducted. Management protocols may continue to shift over time, as well. The implications of larger ventricles in patients treated with endoscopic third ventriculostomy or in patients who do not require shunting but still have larger ventricles than normal will need to be analyzed in future studies with prospective neurobehavioral evaluation.

Shifts in hydrocephalus management with fewer shunt procedures also impact psychosocial outcomes for patients and families. In addition to lower infection and complication risks, families have fewer medical appointments, as well as decreased stress and overall burden, all of which can affect neurodevelopment and child coping. Since changes in hydrocephalus management have occurred fairly recently, follow-up studies to assess for changes in medical, neurodevelopmental, and psychosocial outcomes in relation to hydrocephalus management are ongoing.

Current Recommendations for Neurodevelopmental Follow-Up

As part of the healthcare of individuals with MMC, neurodevelopmental and neurobehavioral follow-up is recommended to assess for the presence of motor and cognitive impairments and to help guide planning for interventions or accommodations. The Spina Bifida Association has produced standard-of-care guidelines identifying primary, secondary, and tertiary outcomes to be assessed at specific ages. These outcomes involve development and acquisition of cognition, language, academic abilities, participation in school, vocational settings, broader society, and independence. Based on these guidelines and the literature reviewed above, the following time points are recommended for neurobehavioral follow-up. Changes or concerns encountered during evaluations should be communicated to the treatment team, particularly the spina bifida and neurosurgery teams, so that any needed interventions can be performed in a timely fashion.

Age-Specific Recommendations

Ages 0 to 1 Year

Infants with MMC are typically monitored carefully for hydrocephalus during the 1st year. In conjunction with this, assessment of neurodevelopmental progress using standardized comprehensive infant development instruments (i.e., the Bayley Scales of Infant Development) can provide information about development and acquisition of abilities to inform medical/treatment planning and guide decision-making about parenting approaches and eligibility for early intervention. The child’s motor, language, and cognitive functions throughout the 1st year are important metrics of medical status and developmental progress, and multiple evaluations during the 1st year (e.g., at 6 and 12 months) help provide information about the child’s developmental trajectory.

Ages 1 to 3 Years

All children with MMC should be referred for early intervention (speech/language, occupational, and physical therapies) during this timeframe, particularly if there are changes in developmental status. Standardized assessments at ages 1–3 years are helpful in determining what supports children need to explore their environment (e.g., supportive seating) and participate in group activities, both of which are important for neurocognitive and psychosocial development. Use of standardized infant and toddler developmental instruments continues to be indicated as comprehensive measures are more sensitive to subtle weaknesses or changes in developmental progress, which may also signal a change in medical status.

Ages 3 to 5 Years

As children with MMC progress to preschool and kindergarten, an additional goal of neurodevelopmental/neurobehavioral follow-up is determining readiness for preacademic and academic skill acquisition. Conducting a comprehensive neuropsychological evaluation around the time the child begins kindergarten is useful for guiding educational planning, including implementation of special education programs like an individualized education program or Section 504 plan.

School Age

As the child moves from early childhood into school
age, evaluations should use a comprehensive battery of standardized neurobehavioral instruments assessing a broad range of neurocognitive/behavioral domains, as outlined above. Specific goals for school-age follow-up assessments change as expectations and demands change over the years. Scheduling neuropsychological evaluations around the time of major transitions in school is recommended. These time points all generally involve changes in demands for the child, such as the shift from learning to read to reading to learn around third grade; the shift toward more autonomy and self-regulation that occurs during middle school; the continued increase in autonomy, independence, and self-management that occurs during high school; and finally the transition from high school to a career or post-secondary education in 12th grade.  

Transitioning to Adulthood

Managing the transition to young adulthood is critical for individuals with MMC, who may be less likely to pursue higher education, work full time, live independently, or develop romantic relationships than healthy peers. Predictors of success with these milestones include pathology (no hydrocephalus, greater mobility), executive functioning abilities, family factors (SES, parental involvement), and logistical factors (financial stability, access to job training). When transitioning to adult healthcare, it is important for the healthcare team to provide direct referrals to adult providers, logistical support (vocational rehabilitation or transitioning health insurance), and active teaching to help develop the skills needed to transition.  

Neurocognitive monitoring during adulthood also remains important, particularly if changes in cognitive status occur (which may signal neurological changes).  

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Author Contributions
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Correspondence
Jane E. Schreiber: Children’s Hospital of Philadelphia, PA. schreiberj@email.chop.edu.