Cervical spinal cord injury: tailoring clinical trial endpoints to reflect meaningful functional improvements

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Abstract
Cervical spinal cord injury (SCI) results in partial to full paralysis of the upper and lower extremities. Traditional primary endpoints for acute SCI clinical trials are too broad to assess functional recovery in cervical subjects, raising the possibility of false positive outcomes in trials for cervical SCI. Endpoints focused on the recovery of hand and arm control (e.g., upper extremity motor score, motor level change) show the most potential for use as primary outcomes in upcoming trials of cervical SCI. As the field moves forward, the most reliable way to ensure meaningful clinical testing in cervical subjects may be the development of a composite primary endpoint that measures both neurological recovery and functional improvement.

Key Words: spinal cord injury; SCI; cervical; clinical trial; endpoint; Cethrin; UEMS

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Introduction
Acute spinal cord injury (SCI) is a serious unmet medical need. A traumatic lesion to the spinal cord results in sensory and motor impairment below the level of the injury and spontaneous recovery is very limited in neurologically complete injuries (Fawcett et al., 2007). Cervical SCI, the most devastating and most common type of SCI (Grossman et al., 2012), leaves individuals with impaired or absent function of the upper extremities (arms and hands) in addition to the lower body paralysis seen after thoracic/lumbar/sacral SCI. There are no approved drugs to foster repair after SCI, despite numerous preclinical studies showing the promise of regenerative medicine (Hollis and Tuszynski, 2011; Liu et al., 2011; Filli and Schwab, 2012; Dickenesher et al., 2013; Watzwaiickev et al., 2014).

Unlike other neurotrauma indications, there have been few clinical trials in SCI, and only very recent trials have focused exclusively on cervical SCI (NCT01828203, NCT01502631, NCT01597518). Clinical trials have traditionally used the International Standards for the Neurological Classification of Spinal Cord Injury (ISNCSCI) to broadly track motor/sensory changes in subjects after injury (Ditunno et al., 2005). The most common primary endpoints derived from this neurological assessment are American Spinal Injury Association (ASIA) Impairment Scale conversion and total motor score. These broad endpoints have limited value for the assessment of recovery after cervical injury, as conversion between grades or change in total motor score may occur without significant change in arm and hand control, an area of critical importance to individuals living with quadriplegia after cervical SCI (Anderson, 2004).

As the number of compounds entering late-stage clinical testing for SCI grows, the selection of meaningful primary endpoints for the evaluation of recovery after cervical injury becomes increasingly important. At present, two endpoints used in retrospective analyses of previous studies show the most promise: (1) Upper extremity motor score (UEMS) and (2) motor level change. These endpoints better reflect meaningful recovery in the arms and hands, but can still fail to distinguish functional benefit. For example, very small changes in multiple muscle groups may produce the same overall increase in UEMS score as a more beneficial, full recovery in fewer muscle groups. As the field moves forward, the development of a composite endpoint directly sensitive to both neurological recovery and functional improvement may permit a more meaningful evaluation of drugs for acute cervical SCI.

SCI
Traumatic SCI is a global problem. A SCI can instantly transform an otherwise healthy individual into a person facing a lifetime of disability, and more than 175,000 spinal cord injuries occur globally every year (Lee et al., 2014). In the United States, approximately 12,000 individuals suffer SCIs each year, most commonly from motor vehicle accidents or falls (NSCISC, 2013). Approximately 57–75% of U.S. SCIs are cervical (Grossman et al., 2012; Selvarajah et al., 2014), and complete cervical SCI leads to lifelong quadriplegia (Fawcett et al., 2007).
et al., 2007). Individuals with SCI also suffer comorbidities, including autonomic dysreflexia, bladder dysfunction, muscle spasticity, and chronic pain (Krassioukov et al., 2003). The combination of these medical complications and corresponding challenges for personal autonomy and community involvement lead individuals with SCI to consistently report a lower quality of life than the nondisabled community (Dijkers, 1997). There are currently no approved treatments to reduce paralysis and improve daily function after SCI.

Limitations of traditional endpoints for evaluation of cervical SCI

The most common primary endpoints in SCI clinical trials are total motor score and ASIA Impairment Scale (AIS) conversion, derivatives of the ISNCSCI assessment (Ditunno et al., 2005; Steeves et al., 2007). Total motor score and AIS conversion were originally designed to assess impairment after SCI (Maynard et al., 1997), but have been subsequently used in clinical trials as measures of drug-based neurological recovery (Steeves et al., 2007). Both endpoints can increase in a manner that may not signify improved function in cervical subjects.

Total motor score (TMS) measures contraction strength in five upper body muscle groups and five lower body muscle groups on either side of the body from 0 (total paralysis) to 5 (normal movement), for a total possible score of 100. The score does not weigh muscle groups by their functional potential or assess the functional value of a score increase. Clinical trial participants could therefore achieve a significant, 20 point improvement in total motor score over placebo in the year following injury due to an isolated recovery from paralysis (0) to normal function (5) in both ankle dorsiflexors and in both knee extensors. A subject could also increase 20 points by improving from full paralysis (0) to active movement with gravity eliminated (2) in all ten leg/foot muscle groups, a change offering similarly limited to nonexistent functional benefit for a quadriplegic individual. The use of a TMS endpoint could thus result in a false positive outcome in a clinical trial for cervical SCI, based on isolated large increases or widely dispersed small increases in lower body muscle groups.

The ASIA Impairment Scale ranks impairment according to body-wide ISNCSCI motor/sensory results, from AIS A (complete paralysis, no motor or sensory function below the level of the injury) to AIS B (complete motor paralysis, sensory function below injury level), AIS C (incomplete motor paralysis, more than half of key muscles below injury level score < 3), AIS D (incomplete motor paralysis, more than half of key muscles below injury level score ≥ 3), or AIS E (normal motor and sensory function). Transitions to higher AIS grades may similarly lack association with functional independence. For example, a two grade transition from AIS A to AIS C, generally regarded as a meaningful clinical outcome, could be due simply to an increase in sensation and the regain in normal control of one large toe (e.g., extensor hallucis longus: 0 to 5).

Both TMS and AIS grade can thus improve based on isolated or widely dispersed increases in lower body control. Though subtle changes in lower body control can hold meaning for individuals with an incomplete SCI or those with paraplegia from a thoracic injury, these changes may have little functional value for an individual with quadriplegia from a cervical SCI. In addition, such changes may simply reflect an incorrect classification of complete paralysis rather than incomplete paralysis during initial baseline measurements. The reliability of baseline measurements is a particular concern in trials of neuroprotective drugs that must be given within 12 hours of injury (Tadie et al., 2003; Casha et al., 2012; Grossman et al., 2014), as baseline assessments for these trials must be conducted while the spinal cord is still in a state of spinal shock (Ditunno et al., 2005). Overall, TMS and AIS conversion are unreliable outcome measures for assessing functional recovery in cervical subjects.

The limitations of these endpoints must be considered with particular care when designing or evaluating trials for drug repositioning, as primary endpoint analysis and scientific reporting in these trials may result in recommendations without the same robust regulatory oversight ensured during the approval process for a new drug. The clinical trial of methylprednisolone as a repositioned treatment for SCI reported total motor score recovery and remains controversial today (Bracken, 2001; Tator, 2006). More recent SCI clinical trials for drug repositioning have also reported primary analyses of total motor score and/or AIS conversion (e.g., the minocycline trial (Casha et al., 2012) and the riluzole trial (Grossman et al., 2014)). The need for standardized reporting of functionally meaningful endpoints in such trials is of critical importance for the evaluation of repositioned therapeutics for cervical SCI.

Meaningful endpoints for the evaluation of neurological recovery after cervical SCI

The selection of a meaningful primary endpoint for assessing recovery after cervical SCI requires an understanding of which drug-based neurological improvements would offer subjects the most functional benefit. Since individuals living with quadriplegia overwhelmingly agree that their quality of life would be most improved by a return of arm/hand function (Anderson, 2004), two ISNCSCI neurological assessment derivatives emerge as valuable endpoints for cervical subjects: (1) Upper extremity motor score (UEMS) and (2) Motor level change. Both endpoints have been used extensively in retrospective analyses of cervical subjects in previous SCI clinical trials and databases (Steeves et al., 2011, 2012; Kramer et al., 2012; McKerracher and Anderson, 2013). Though these endpoints are not impervious to the limitations seen in TMS/AIS conversion (e.g., increases in UEMS are also not weighted by comparative functional benefit), both offer a more focused measure of functionally meaningful increases in hand and arm control that could better serve as a primary endpoint to support drug approval for cervical SCI.

Upper extremity motor score (UEMS) measures contraction strength in five key arm and hand muscle groups on either side of the body from 0 (total paralysis) to 5 (active movement), for a maximal possible score of 50. UEMS correlates more closely than total motor score or lower extremity motor score with improvements in self-care and mobility measured by the Functional Independence Measure (FIM) (Marino and Graves, 2004). Increases in UEMS also correspond
directly to improvements in functional activities of daily living and self-care measured with the Spinal Cord Independence Measure (SCIM) (Kramer et al., 2012). Furthermore, individuals with a higher UEMS have an increased capacity for self-feeding, as measured by the Quadriplegia Index of Function (QIF) (Marino et al., 1995).

The Motor Level of a cervical SCI corresponds to the lowest spinal segment from which extending neurons permit muscle movement against gravity, and above which motor function is normal. In particular, the key muscle group innervated by the 'motor level' must receive an ISNCSCI motor score of at least 3 (contraction against gravity alone), while all key muscle groups innervated by spinal segments above this level receive a 5 (normal function). Since the cervical levels of the spinal cord innervate the arms and hands, the motor level of a cervical subject is a direct measure of arm/hand regions under voluntary control. The recovery of two or more motor levels in the year following a cervical SCI is associated with a significant improvement in functional independence, as measured by the SCIM self-care subcategory (personal grooming, feeding, bathing, and dressing) (Kramer et al., 2012).

As more compounds move into later-stage clinical trials for SCI, the importance of evaluating cervical subjects based on such functionally meaningful primary endpoints must be emphasized. To date, only one trial including acute cervical SCI subjects has selected a primary endpoint with the capacity to measure functional arm and hand recovery (NCT01502631, ≥ 2 motor level recovery). Standardized reporting of a primary UEMS endpoint in future SCI clinical trials will allow assessment of meaningful recovery in cervical subjects and facilitate comparative analyses of drug efficacy between trials. Such reporting will be particularly important for repositioned drug trials, where presented analyses of UEMS recovery have previously been secondary or absent (Bracken et al., 1997; Casha et al., 2012; Grossman et al., 2014).

**Meaningful endpoint selection in the phase II/III trial of Cethrin for cervical SCI**

The biologic drug Cethrin is a Rho antagonist designed to promote neuroregeneration and neuroprotection when delivered as a topical adjunct to decompression surgery after SCI (McKerracher and Guertin, 2013). An open-label, phase I/IIa trial demonstrated that Cethrin was well-tolerated, and offered a preliminary assessment of efficacy on the traditional SCI endpoints of AIS grade and total motor score (Fehlings et al., 2011). Though improvements by Cethrin-treated cervical subjects on these endpoints were promising (McKerracher and Anderson, 2013), it is clear that an increase in AIS grade conversion or total motor score may not signify improved daily function.

To permit a more meaningful assessment of phase I/IIa Cethrin trial results, the original ISNCSCI assessment data was recently re-analyzed on the endpoint of upper extremity motor score (McKerracher and Anderson, 2013). During the first year after injury, the sixteen Cethrin-treated cervical subjects in all five tested dose groups (0.3, 1, 3, 6, 9 mg) improved an average of 12.2 ± 2.6 points in UEMS from a baseline of complete paralysis (Figure 1). The nine cervical subjects in the three highest dose groups (3, 6, and 9 mg) demonstrated an average UEMS recovery of 14 ± 2.4 points. These improvements would confer functional benefit over the expected spontaneous recovery of 8.8 ± 0.5 points (Model Systems (Marino et al., 2011)) to 9.6 ± 0.4 points (Sygen database (Steeves et al., 2011)) seen in historical cervical individuals with complete paralysis. Even a two point improvement in UEMS recovery can result in increases in hand and arm strength with a tremendous functional impact for subjects with cervical SCI (Steeves et al., 2012).

The upcoming phase II/III Cethrin trial will examine the ability of the drug to reduce paralysis and promote functional recovery after acute cervical SCI (NCT02053883). Upper extremity motor score has been selected as the primary endpoint for this trial. The use of this meaningful primary endpoint for a pivotal trial in cervical SCI will help to shift the field toward the choice of functionally relevant endpoints for future cervical trials.

The phase II/III Cethrin trial will also include secondary endpoints to assess the impact of Cethrin on the serious comorbidities of SCI, such as respiratory impairment, bowel and bladder dysfunction, pain, muscle spasticity, pressure ulcers, and autonomic dysreflexia (Krassioukov et al., 2003; Stein et al., 2010). In addition to the standard International SCI Data Sets, the trial will use a newly developed questionnaire to directly assess personal perceptions of comorbidity severity in Cethrin-treated versus placebo subjects. As the field moves forward, the development and use of other new tools to evaluate SCI comorbidities may permit improved clinical assessments (e.g., an additional test of the ability to generate an effective cough might enhance standard respiratory assessments of forced expiratory volume). By complementing a meaningful primary endpoint (UEMS) with carefully designed secondary tests of SCI comorbidities, the phase II/III Cethrin trial will set the tone for a more patient-centric analysis of cervical SCI treatments.

**Developing a composite endpoint for cervical SCI**

With an overall goal of determining the functional benefit of a trial therapeutic for cervical SCI, the natural complement to selecting an appropriate neurological assessment is the additional use of a direct test to measure functional ability. Tests of functional independence, such as FIM and, more recently, SCIM, have been incorporated as secondary endpoints in SCI clinical trials (Bracken et al., 1997; Casha et al., 2012, NCT01828203). However, these tests are unsuited for independent use as primary endpoints for cervical SCI, as improvement in a specific category may reflect external factors/rehabilitation, rather than drug-based improvements in body control. For example, a cervical subject could advance in the dressing category of SCIM from a score of 1 (requiring partial assistance with clothing without buttons, zippers, or laces) to a score of 2 (independent with clothing without buttons, zippers, or laces; requires adaptive devices and/or specific settings) by acquiring a relevant adaptive device.

The ideal solution for future clinical testing in acute cervical SCI may be the development of a composite endpoint that incorporates both a relevant neurological assessment...
and a direct test of functional autonomy. Composite endpoints have become increasingly prevalent in clinical trials for other neurological indications (Cutter et al., 1999; Elm and Investigators, 2012; Kozauer and Katz, 2013). For example, the Multiple Sclerosis Functional Composite evaluates leg function/ambulation (via a timed, 25-foot walk), arm/hand function (via a test involving placing pegs in a pegboard), and cognitive function (via a test of the subject’s ability to add numbers heard at 2–3 second intervals) (Cutter et al., 1999). The use of a well-designed composite for cervical SCI could ensure that an approved therapeutic offers neurological improvements conferring direct functional benefit.

The upcoming phase II/III Cethrin trial will take the first step toward the development of such a composite endpoint. The trial has been designed to assess whether the following measures of arm and hand recovery can be combined to form a composite primary endpoint: (1) UEMS, (2) Graded Redefined Assessment of Strength Sensibility and Prehension (GRASSP) (Kalsi-Ryan et al., 2012), (3) Capabilities of Upper Extremity Test (CUE-T) (Marino et al., 2012), and (4) SCIM III Self-Care Subscore (Itzkovich et al., 2007). GRASSP and CUE-T were recently developed to directly measure the recovery of hand and arm capabilities (e.g. reaching forward, pushing down, pinching a key), and the ability to perform basic functional tasks (e.g. pouring water, lifting a container). The Self-Care subsection of SCIM measures a subject’s ability to perform complex tasks of daily living, such as feeding, dressing, bathing, and grooming. UEMS offers a functionally relevant assessment of drug-based neurological changes in arm and hand control. UEMS, GRASSP or CUE-T, and the SCIM Self-Care subscore could together form a composite endpoint sensitive to drug-based functional improvement that transforms the trajectory of future cervical SCI clinical trials.

**Conclusions**

Traditional primary endpoints for acute SCI clinical trials do not adequately assess functional recovery in cervical subjects, raising the possibility of false positive outcomes in trials for cervical SCI. Endpoints focused on the recovery of hand/arm control (UEMS, motor level change) show the most potential for use as primary outcomes for cervical SCI in the immediate future. Selection of the upper extremity motor score as the primary endpoint for the upcoming phase II/III trial of Cethrin will set the tone for the use of this endpoint in clinical development programs. The most reliable way to ensure meaningful clinical testing in acute cervical SCI may be the development of a composite primary endpoint that directly measures both neurological recovery and functional improvement.

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