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Nicholas J. Cioe Assumption College, nj.cioe@assumption.edu

Gary S. Seale *Centre for Neuro Skills* 

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## TITLE

## **Brain Injury Rehabilitation Outcome Measurement: Challenges and Future Directions**

### **AUTHORS**

Nicholas J. Cioe, Ph.D., CRC, CBIST Assumption College

Gary S. Seale, Ph.D., LPA, LCDC Centre for Neuro Skills

1	The Brain Injury Association of America (BIAA) published a position paper
2	(http://www.biausa.org/biaa-position-papers.htm) that describes the state of outcomes research in
3	the field of brain injury and identifies the factors to consider when evaluating existing studies
4	and embarking on future outcomes research. This commentary summarizes that paper by
5	highlighting three major challenges for outcome measurement in brain injury rehabilitation. It
6	also presents opportunities for improving outcomes research through improved research design,
7	standardization of measurement tools across the continuum with utilization of national databases,
8	and an evidence-based approach to providing care to help move brain injury rehabilitation
9	outcome measurement in a positive direction.
10	Outcome measurement in brain injury rehabilitation has progressed. The Interagency
11	Traumatic Brain Injury (TBI) Outcomes workgroup recommended the use of multiple measures
12	with sound psychometric properties that demonstrate utility with the brain injury population in
13	TBI outcomes research.[1] At a minimum, measures selected should assess global level of
14	function, neuropsychological impairment, psychological status, activity limitations and
15	participation restrictions, and perceived health-related quality of life. To accomplish this, most
16	rehabilitation programs employ multiple measures to assess function in several domains as part
17	of a comprehensive outcomes or program evaluation system.[2]
18	Despite this progress, there remain several major challenges for outcome measurement in
19	brain injury rehabilitation. First, there are no universally accepted outcome measurement tools.
20	Second, outcome measurement research has not translated to better predictability of outcome.
21	Third, most persons with brain injury do not have adequate funding to support access to the full
22	continuum of care necessary for value-based service delivery models.
23	Absence of universally accepted outcome measurement tools used throughout the industry

23 Absence of universally accepted outcome measurement tools used throughout the industry

The World Health Organization's (WHO) International Classification of Functioning,
Disability, and Health (ICF) provides a standard language and framework for the description of
health and health-related states. It classifies outcome measure in three domains: *impairment* (e.g.,
memory, attention, balance, swallowing, executive functioning, etc.), *activity limitations* (e.g.,
ambulation, eating, dressing, grooming, talking, etc.), and *participation restrictions* (e.g.,
working, volunteer, home maker, parent, etc.).

Consistent with the language of the WHO ICF, a number of global outcome measures 30 have been developed for the brain injury population and have demonstrated sound psychometric 31 32 properties, for example, the Mayo-Portland Adaptability Inventory and the Community Integration Questionnaire. Likewise, a number of domain-specific measures have been 33 developed, such as the Disability Rating Scale and the Supervision Rating Scale. Unfortunately, 34 given the range of potential outcomes important to brain injury rehabilitation, particularly at the 35 post-acute level of care (i.e., ADL performance, community integration, employment, life 36 satisfaction and quality of life), there is no agreed-upon single measure or set of measures to 37 evaluate outcome following brain injury rehabilitation. In a survey conducted by the Brain 38 Injury Interdisciplinary Special Interest Group (BI-ISIG) of the American Congress of 39 40 Rehabilitation Medicine (ACRM) involving 49 community-based brain injury rehabilitation programs across 23 states, little consistency was found in terms of outcomes measurement tools 41 used. Fifty-nine percent of programs surveyed employed the Mayo-Portland Adaptability 42 Inventory as an outcome management tool, but over 31% of programs reported using a "home 43 grown" measure. The authors concluded that the "absence of universally accepted outcome 44 45 measures limits the availability of a common language and the ability to benchmark against like 46 programs or define best practices for community-based treatment." [3]

The Uniform Data System for Medical Rehabilitation (UDSMR), established in 1987, has 47 positively influenced outcome measurement at the acute care level, including the required use of 48 the Functional Independence Measure (FIM) as the primary tool for outcome measurement. 49 More than 600 articles have been published using UDSMR instruments and data, and the 50 51 research suggested that functional assessment could be developed into a science to provide the knowledge for understanding the "biology of disability," a term that implies possible dominant 52 (and therefore expected) response patterns to disability based on the biological, social, and 53 environmental factors that influence outcome. The "biology of disability" concept could help 54 55 clinicians distinguish patterns of expected functional limitations from unexpected functional limitations.[9] 56

The required use of a single tool or set of outcome measurement tools, consistent with the 57 WHO ICF, from the point of discharge in an acute setting through the post-acute continuum of 58 care, would support the understanding of the biology of disability following brain injury. 59 OutcomesInfo, a web-based database system developed through National Institutes of Health 60 (NIH) Small Business Technology Transfer (STTR), maintains a national database platform to 61 support the collection of brain injury outcome measurement data. At this point, use of outcome 62 63 measurement tools supported by OutcomesInfo's platform and submission of data to the national database is voluntary. 64

### 65 Outcome measurement research has not translated to better predictability of outcome

The expectation is better measurement of outcome will allow for analyses that support a better understanding of the factors that influence outcome. While true in theory, such analyses require the ability to control for variables that influence outcome. Unfortunately, brain injury rehabilitation outcomes research is characterized by vast differences in the demographics of

individuals who are injured and in their access to care, substantial heterogeneity in etiology,

severity, chronicity, and disease progression following injury, and important distinctions in

72 treatment settings, provider expertise, intervention types and intensities, and measurement

73 tools.[2]

Alone or in combination, each of the aforementioned variables can significantly impact the outcome of brain injury rehabilitation [10] making it extremely difficult to perform research studies that adequately control for these variables. The use of common outcome measurement tools across the continuum of care (see Figure 1), and the entry of those outcome data into a national database, would help provide a data set of adequate size and content to support research that can identify practices to improve outcome predictability.

80

# INSERT CONTINUUM OF CARE (Figure 1) HERE

# 81 Inadequate funding to support access to the full continuum of care necessary for value-

## 82 based service delivery models

Although there remains significant room for improvement in brain injury rehabilitation, 83 adequate knowledge and tested clinical interventions exist to treat brain injury effectively for 84 85 improved outcome. [4-8] However, few individuals who sustain a brain injury have access to the full array of services along the TBI continuum of care (as depicted in Figure 2). Given the 86 numerous variables that influence outcome and the often unorganized and inaccessible array of 87 88 services along the continuum, it is extremely difficult to compare one brain injury outcome to another. Consider the difference in outcome between former Congresswoman Gabby Giffords 89 90 and an 18-year-old gang member. While both experienced a bullet wound to the head during an 91 assault, the many pre-existing and injury related variables (e.g., age, pre-injury education level,

SES, length of loss of consciousness, treatment access) must be considered before treatment
related factors can be reliably analyzed.

An alternative approach to outcome research would allow each person who receives a brain injury to access a coordinated array of services along the continuum of care during which time a standardized set of data is collected. By standardizing – in so much as providing the proper treatment intensity for the appropriate duration at the right time in the right setting (see Figure 2) – the treatment experience, individual demographics and heterogeneity in etiology, severity, chronicity, and disease progression can be analyzed to provide better outcome predictability.

101

# INSERT OUTCOME = ILLUSTRATION (Figure 2) HERE

102 The current political landscape supports health care service delivery that maximizes 103 value, that is, improved health outcomes for the money spent. Brain injury services can be 104 extremely expensive making them a prime target for criticism and funding restrictions. Unlike 105 treatment for a torn knee ligament, there is not a widely accepted and adequately funded course 106 of treatment following brain injury. The absence of such inhibits a meaningful and accurate 107 value-based analysis of services.

The field of brain injury rehabilitation continues to develop. Increased appreciation for the moral and ethical responsibility to allocate adequate resources to support post-acute care should help improve access to the full continuum of care. Commensurate attention to improving interventions, improvements in imaging, and increased knowledge about genetic factors should help move best practice forward. There is a sense among leaders in the field that the future is bright.

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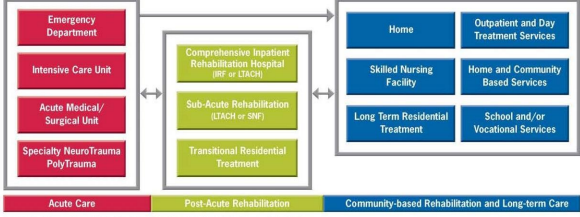
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### **ILLUSTRATIONS**

### Figure 1

### Traumatic Brain Injury Continuum of Care



Developed by H. Reyst for The Essential Brain Injury Guide 5th Edition (p. 13)

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## Figure 2



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