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A core outcome set for aphasia treatment research: the ROMA consensus statement

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Abstract

Background: A core outcome set (COS; an agreed, minimum set of outcomes) was needed to address the heterogeneous measurement of outcomes in aphasia treatment research and to facilitate the production of transparent, meaningful and efficient outcome data.Objective: The Research Outcome Measurement in Aphasia (ROMA) consensus statement provides evidence-based recommendations for the measurement of outcomes for adults with post-stroke aphasia within phase I-IV aphasia treatment studies.

Methods: This statement was informed by a four-year program of research which comprised investigation of stakeholder-important outcomes using consensus processes, a scoping review of aphasia outcome measurement instruments, and an international consensus meeting. This paper provides an overview of this process and presents the results and recommendations arising from the international consensus meeting. Results: Five essential outcome constructs were identified: Language, communication, patient-reported satisfaction with treatment and impact of treatment, emotional wellbeing, and quality of life. Consensus was reached for the following measurement instruments: Language: The Western Aphasia Battery Revised (WAB-R) (74% consensus); emotional well-being: General Health Questionnaire (GHQ)-12 (83% consensus); quality of life: Stroke and Aphasia Quality of Life Scale (SAQOL-39) (96% consensus). Consensus was unable to be reached for measures of communication (where multiple measures exist) or patient-reported satisfaction with treatment or impact of treatment (where no measures exist). Discussion: Harmonisation of the ROMA COS with other core outcome initiatives in stroke rehabilitation is discussed. Ongoing research and consensus processes are outlined. **Conclusion:** The WAB-R, GHQ, and SAQOL-39 are recommended to be routinely included within phase I-IV aphasia treatment studies. This consensus statement has been endorsed by the Collaboration of Aphasia Trialists, the British Aphasiology Society, the German Society for Aphasia Research and Therapy, and the Royal College of Speech Language Therapists.

A core outcome set for aphasia treatment research: the ROMA consensus statement

The Research Outcome Measurement in Aphasia (ROMA) consensus statement provides recommendations for a core outcome set (COS) for use in aphasia treatment studies. A COS is a minimum set of outcomes that should be measured and reported in research trials of a specific health condition or population (1). The use of a COS does not preclude the measurement of additional outcomes, but rather represents the minimum outcomes that should be collected and reported (2). A COS for aphasia was developed in response to a trend of heterogeneous outcome measurement in research and the merits of this initiative were debated in a published forum in 2014 (3-7). The ROMA consensus statement was informed by a four-year program of research in three phases: (1) investigation of stakeholder-important outcomes using consensus processes (8-11); (2) a scoping review to identify aphasia outcome measurement instruments (OMIs) and their psychometric properties (12); and (3) an international consensus meeting (results reported herein). The ROMA COS is intended to complement other existing and ongoing initiatives to standardise the measurement of stroke recovery (13-15).

Objective

The ROMA consensus statement provides evidence-based recommendations for the measurement of outcomes for adults with post-stroke aphasia within phase I-IV aphasia treatment studies.

Target users

The primary users of this consensus statement will be researchers involved in the design and conduct of aphasia treatment studies.

Methods

The research methods are based on the recommendations of the Core Outcome Measures in Effectiveness Trials (COMET) Initiative (2, 16) and are reported in alignment with the COS-STAR (Core Outcome Set-STAndards for Reporting) statement (17). The World Health Organization International Classification of Functioning, Disability and Health (ICF) (18) has been used as a conceptual framework and classification tool. This project is registered with the COMET Initiative (<u>http://www.comet-initiative.org/studies/details/287</u>).

Stage 1: Identification of Core Outcome Constructs

Outcome constructs were derived from three separate stakeholder consensus studies conducted with: people with aphasia and their families (9); aphasia clinicians and managers (8); and aphasia researchers (10). Outcomes prioritised by stakeholder groups were integrated using the framework of the ICF (19). Essential constructs were identified as: Language, communication, patient-reported satisfaction with treatment and impact of treatment, emotional wellbeing, and quality of life (11).

Stage 2: Identification of Outcome Measurement Instruments

A scoping review was conducted to identify OMIs which have been validated with people with aphasia. Primary searches were run using PUBMED, EMBASE, and CINAHL databases on 10 November 2015. The search strategy incorporated filters developed for the identification of studies reporting the measurement properties of health OMIs (see 20 and supplementary file). Inclusion criteria required that studies focused on

the psychometric properties of measurement instrument and included participants with aphasia or stroke patients where participants with aphasia were not specifically excluded. Studies reporting measurement instruments which primarily measure neurological function associated with, but not central to aphasia: e.g., consciousness; health; motor speech; cognition; memory; were excluded. Secondary searches were conducted for each OMI identified in the first search. In total, 184 references for 79 measurement instruments were identified (12). No measures of patient-reported treatment impact or patient-reported satisfaction were identified through this search.

Stage 3. Formation of Consensus Panel

Researchers who participated in the first phase of this project (n=80) (10) were invited to participate in the final consensus meeting. These researchers were purposively sampled from researchers whose trials were included with the Cochrane Collaboration review of "Speech and language therapy for aphasia following stroke"(21) and the 100 most highly published aphasia treatment researchers in the Web of Science database. In total, 23 researchers participated in a consensus meeting in London, UK (December, 2016). Panel members were experienced researchers with expertise in: the design and conduct of aphasia trials; measurement instrument development and testing; and clinical guidelines development (see table 1 and supplementary table 1). Authors Wallace, Worrall, Le Dorze and T. Rose facilitated the COS development process and did not participate in COS voting.

Table 1

Characteristics of researchers who participated in the international consensus panel (n=23)

Panel Characteristics

n (%)

Country

United Kingdom	9 (39)
United States of America	6 (26)
Australia	3 (13)
Canada	2 (9)
Germany	1 (4)
Sweden	1 (4)
Ireland	1 (4)
ICF component to which their own research relates (panel	
members could nominate more than one component)	
Body functions	16
Activity/Participation	21
Environmental factors	10
Personal factors	15
Quality of life*	12
Number of treatment studies published by participants	
1	2
2-5	8
6-10	4
more than 10	7
not specified	2

*nb. Quality of life is not defined as a component of the ICF

Stage 4. International Consensus Meeting

Ethical approval for the consensus meeting was gained from the Behavioural and Social Sciences Ethical Review Committee at The University of Queensland, Australia. The following process was used:

Prior to meeting

- (1) Panel members generated consensus-based criteria to enable an initial reduction of OMIs (see table 2).
- (2) The consensus-based criteria were applied to the list of OMIs identified in the stage 2 scoping review (n=79) to produce a short-list (n=50) (see supplementary table 2).
- (3) Panel members generated consensus-based feasibility criteria (see table 3).
- (4) The short-listed OMIs (see supplementary table 2) were assigned to panel members, who reviewed OMI feasibility and measurement properties prior to the consensus meeting.

During the meeting

(1) Panel members engaged in a whole-group discussion using an iterative process to apply feasibility criteria and eliminate OMIs.

- (2) Panel members divided to smaller groups to review the measurement properties for each OMI in the target population (people with aphasia). Properties considered included: acceptability/feasibility of use with people with aphasia, reliability (test-retest, inter- and intra- as applicable), construct validity, and sensitivity to change.
- (3) Each small group recommended two OMIs for voting. Panel members voted YES/NO for each OMI in a closed voting process with consensus defined a priori as agreement on each OMI for each outcome construct by \geq 70% of meeting participants, as suggested by the COMET initiative and GRADE working group (2). Potential conflicts of interest were managed through agreement that authors of OMIs under consideration could not participate in voting for that construct area.

Table 2

Criteria for initial reduction of outcome measurement instruments

Measures were excluded if:

- 1. The purpose of the measurement instrument was to screen for the presence of aphasia, rather than to measure outcomes.
- 2. The measurement instrument was published more than thirty years ago (i.e., prior to 1986) without subsequent revision and/or was not in current use.
- 3. The measurement instrument targeted only one severity level of aphasia.
- 4. For measures of language: the measurement instrument did not assess all modalities of language (e.g. reading only, writing only, comprehension only, verbal output only).

Table 3

Feasibility criteria

- 1. Availability in different languages or ease of translation/adaptation.
- 2. Cost.
- 3. Burden to respondents or researchers (ease of administration, length of outcome measurement instrument, completion time).
- 4. Ease of score calculation and provision of an aggregate score.

Results

After compilation of votes, panel members reached consensus for measures of language, emotional wellbeing, and quality of life (refer to table

4). A consensus of \geq 70% was not reached for a measure of communication. Inability to gain consensus on a measure of communication may

relate to the multi-factorial nature of this construct, as well a lack of understanding and consensus around how 'effective communication' is best operationalised in treatment research.

Table 4

Construct	Measure*	Votes for
		inclusion
Language	The Western Aphasia Battery Revised (WAB-R)	74% (n=17)
	The Comprehensive Aphasia Test (CAT)	22% (n=5)
	Neither	4% (n=1)
Communication	The Scenario Test	57% (n=13)
	The Communication Effectiveness Index (CETI)	39% (n=9)
	Abstained	4% (n=1)
Emotional well-	General Health Questionnaire (GHQ)-12	83% (n=19)
being		
	Stroke Aphasic Depression Questionnaire (SADQ)	17% (n=4)
Quality of life	Stroke and Aphasia Quality of Life Scale (SAQOL-	96% (n=22)
	39)	
	Burden of Stroke Scale (BOSS)	0% (n=0)
	Abstained	4% (n=1)

Results of final voting to decide core outcome measurement instruments

Bolded figures indicate consensus criteria (\geq 70%) reached and OMI included in COS *Refer to supplementary tables 3 & 4 for OMI characteristics, properties and references.

Recommendations

It is recommended that the WAB-R, GHQ-12 and SAQOL-39 be included as core outcome measurement instruments in phase I-IV aphasia treatment studies for adults with post-stroke aphasia. These outcome measurement instruments and their psychometric properties are described in supplementary tables 3 & 4.

Discussion

The importance of implementing standardised approaches to outcome measurement in research trials is increasing acknowledged. In the field of stroke rehabilitation, the Stroke Recovery and Rehabilitation Roundtable (SRRR) (13) have provided consensus-based core recommendations for the measurement of sensorimotor recovery after stroke. Other initiatives have addressed the measurement of stroke outcomes in clinical practice (15) and there are ongoing works to standardise measures in arm rehabilitation trials after stroke (14). The ROMA COS has sought to provide recommendations specifically for the measurement of aphasia recovery post-stroke. Accordingly, some frequently used measures of global disability and health-related quality of life (e.g., EQ-5D) which do not contain communication-specific items or which have not been validated with stroke survivors with aphasia were not considered within this process. The ROMA COS seeks to harmonise with other existing stroke rehabilitation initiatives in addressing the need for standardised approaches to research trial outcomes measurement and its supplementary use may therefore be considered in any stroke study where people with aphasia are included.

Future Directions

The ROMA COS will be reviewed biennially. The next consensus meeting will focus on measures of communication and consider the development of measures of patient-reported satisfaction with treatment / impact of treatment. Factors relating to international COS implementation will be considered. New publications, initiatives and user feedback will also be considered in each review to: align this COS with other COSs; consider new OMIs; and to review the choice of OMIs based on user feedback.

Limitations

Participants in the international consensus meeting were predominately from English speaking countries. This may have impacted the consensus process and findings. Future meetings will seek to increase the diversity of participants with respect to cultural and linguistic background.

Funding

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Conflicts of Interest

Authors Babbit, Breitenstein, Cherney, Cruice, Enderby, and Hilari authored or adapted OMIs considered in this consensus process. These authors declared their conflict of interest during the meeting and did not participate in voting which related to their authored OMIs. Authors Wallace, Worrall, Le Dorze and T. Rose did not participate in voting on OMIs.

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Supplemental File Search Strategy

Search strategy (incorporates filters developed by Terwee and associates for the identification of studies reporting the measurement properties of health outcome measures; see Terwee CB, Jansma EP, Riphagen I, Vet HW. Development of a methodological PubMed search filter for finding studies on measurement properties of measurement instruments. Quality of Life Research 2009;18(8):1115-23.)

PUBMED

Aphasia OR dysphasia AND stroke

AND

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EMBASE

aphasia OR dysphasia AND stroke

AND

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aphasia OR dysphasia AND stroke

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ROMA consensus meeting facilitators

Sarah J. Wallace PhD BSpPath(Hons)	Linda Worrall PhD BSpThy FSPA	Guylaine Le Dorze Ph.D MSc (A)	Tanya Rose PhD BSpPath(Hons)
GradCert Gerontology CPSP	Speech Pathologist, Teaching and	Teaching and Research Academic,	GradCert Higher Ed CPSP
Certified Practising Speech Pathologist	Research Academic, School of Health	Speech-Language Pathologist, School of	Certified Practising Speech Pathologist
and Teaching and Research Academic,	and Rehabilitation Sciences, The	Speech-Language Pathology and	and Teaching and Research Academic,
School of Health and Rehabilitation	University of Queensland, Australia.	Audiology, Faculty of Medicine,	School of Health and Rehabilitation
Sciences, The University of Queensland.	Expertise: post-stroke aphasia	Université de Montréal.	Sciences, The University of Queensland.
Expertise: post-stroke aphasia	rehabilitation, ICF, aphasia trial design	Expertise: post-stroke aphasia	Expertise: Post-stroke aphasia
rehabilitation, core outcome set	and conduct, consumer perspective,	rehabilitation, participation, single-	rehabilitation, paediatric and adult
development, stakeholder perspectives,	aphasia rehabilitation guideline	subject designs, qualitative methods.	language, accessible health information,
consensus processes, ICF.	development.		mixed-methods research.

ROMA consensus panel

Edna Babbitt PhD CCC-SLP BC-	Arpita Bose PhD MSc (Speech and	Marian Brady PhD BSc	Caterina Breitenstein PhD academic
ANCDS	Hearing) BSc (Audiology and Speech	Speech and language therapist, Director	degrees in Clinical Psychology and
Research Speech-Language Pathologist	Rehabilitation). Speech and Language	Stroke Rehabilitation Research,	Cognitive Neuroscience.
Assistant Research Professor,	Therapist, Teaching and Research	NMAHP Research Unit, Glasgow	Teaching and Research Academic, Dept.
Department of Physical Medicine and	Academic, School of Psychology and	Caledonian University, Glasgow,	of Neurology, University of Muenster,
Rehabilitation, Feinberg School of	Clinical Language Sciences, University	Scotland.	Germany.
Medicine, Northwestern University,	of Reading, Reading, UK.	Expertise: Stroke rehabilitation, design,	Expertise: Development and national
Chicago, USA & Shirley Ryan	Expertise: Post-stroke aphasia	development and evaluation of complex	adaptations of communication outcome
AbilityLab, Chicago, USA.	assessment and rehabilitation,	multidisciplinary interventions, survey,	measures, clinical trials methodology,
Expertise: Post-stroke aphasia	bilingualism, single subject	mixed methods, systematic review, meta-	intervention studies in post-stroke
assessment and rehabilitation.	experimental designs, quality of life	analyses and the use of randomised	aphasia rehabilitation.
	issues in aphasia, SLT training in	controlled trial archives.	
	decision-making in aphasia.		
Leora R. Cherney PhD CCC-SLP	David Copland PhD BSpPath (Hons)	Madeline Cruice PhD BSpPath	Pam Enderby PhD MBE DSc (Hons)
BC-ANCDS. Research Scientist and	Speech Pathologist, Principal Research	(Hons)	MSc FRCSLT
Speech and Language Pathologist.	Fellow, School of Health &	Registered Speech and Language	Speech and Language Therapist,
Shirley Ryan AbilityLab (formerly the	Rehabilitation Sciences and Centre for	Therapist, Reader, Teaching and	Professor Emeritus of Community
Rehabilitation Institute of Chicago) and	Clinical Research, The University of	Research Academic, School of Health	Rehabilitation, University of Sheffield,
Northwestern University, Chicago, IL	Queensland, Brisbane, Australia.	Sciences, City University of London,	Sheffield. UK.
USA. Expertise: Post-stroke aphasia		London, UK.	

assessment and rehabilitation, development and evaluation of novel aphasia treatments, single subject and RCT design, systematic reviews.	Expertise: Post-stroke aphasia assessment and rehabilitation, aphasia trial design and conduct, neuroimaging in aphasia.	<i>Expertise: Post-stroke aphasia</i> <i>rehabilitation, therapeutic process and</i> <i>evaluation, quality of life evaluation in</i> <i>research and clinical practice,</i> <i>behaviour change.</i>	Expertise: Aphasia management, Clinical Evaluation of Interventions, RCTs, Psychometric Properties of Outcome Measures.
Deborah Hersh PhD MSc BSc(Hons) GradCert Higher Ed FSPA. Speech Pathologist, Teaching and Research Academic, School of Medical and Health Sciences, Edith Cowan University, Perth, Australia. Expertise: Post-stroke aphasia rehabilitation, consumer perspective, aphasia rehabilitation guideline development.	Katerina Hilari PhD MRCSLT MHPC Psychologist, Registered Speech and Language Therapist, Teaching and Research Academic, School of Health Sciences, City, University of London, UK. Expertise: Outcome measurement development, validation and cultural adaptation, post-stroke aphasia rehabilitation, feasibility RCTs, clinical guideline development.	Tami Howe PhD MHSc BEd SLP(C) Speech Pathologist and Teaching and Research Academic, School of Audiology and Speech Sciences, University of British Columbia, Vancouver, Canada. <i>Expertise: Aphasia rehabilitation,</i> <i>ICF, accessibility, goal setting, social</i> <i>participation, impact of aphasia on</i> <i>family members.</i>	Helen Kelly PhD MRCSLT Registered Speech and Language Therapist, Teaching and Research Academic, Department of Speech and Hearing Sciences, University College Cork, Cork, Ireland. Expertise: Post-stroke aphasia assessment and management, single subject and RCT feasibility aphasia trial design and conduct, consumer perspective.
Swathi Kiran PhD CCC-SLP Speech Language pathologist, Teaching and Research Academic. Professor, Associate Dean for Research Sargent College of Health and Rehabilitation Sciences, Boston University, Boston, MA, USA. <i>Expertise: Aphasia rehabilitation,</i> <i>neuroimaging, bilingualism, single</i> <i>subject experimental design.</i>	Ann-Charlotte Laska MD A/Professor Department of Clinical Science Karolinska Institutet Danderyd Hospital, Sweden Expertise: Post-stroke aphasia, study design and conduct, RCT.	Jane Marshall PhD Post Grad Diploma in Clinical Communication Studies BA FRCSLT Registered Speech and Language Therapist, Teaching and Research Academic, School of Health Sciences, City, University of London, UK. Expertise: Post-stroke aphasia rehabilitation, the development and evaluation of novel treatments.	Marjorie Nicholas PhD CCC-SLP Professor and Interim Chair Dept. of Communication Sciences and Disorders, MGH Institute of Health Professions, Boston, MA, USA. Expertise: Aphasia rehabilitation, nonverbal cognition in aphasia, Life Participation Approach to Aphasia and community aphasia program design, ICAP design.
Janet Patterson PhD CCC-SLP ASHA Fellow Chief, Audiology & Speech-Language Pathology Service, VA Northern California Health Care System Practicing Speech-Language Pathologist, Teaching and Research Academic. <i>Expertise: Post-stroke aphasia</i>	Gill Pearl MPhil Dip Hum Commun. Certified practicing speech and language therapist in role as Chief Executive Officer of Speakeasy - specialist aphasia centre, UK. Expertise: Development and evaluation of novel approaches to providing long term aphasia support and therapy, facilitator of consumer involvement in	Elizabeth Rochon PhD MSc (A) Reg CASLPO SLP(c) Speech Pathologist, Teaching and Research Academic, Department of Speech-Language Pathology and Rehabilitation Sciences Institute, University of Toronto, Canada. <i>Expertise: Post-stroke aphasia</i> <i>assessment and rehabilitation,</i> <i>development of aphasia treatment</i>	Miranda Rose PhD BSpPath FSPA Speech pathologist, Teaching and Research Academic, School of Allied Health, La Trobe University, Victoria, Australia. Expertise: Post-stroke aphasia rehabilitation, aphasia trial design and conduct, single subject designs, consumer perspective, aphasia rehabilitation guideline development.

rehabilitation, systematic reviews of literature, single subject designs.	research, feasibility studies, case series studies, RCT design and conduct.	studies, feasibility studies, single subject and RCT design, systematic reviews.	
Karen Sage PhD Dip DisHumComm BA (Hons) HCPC Registered Speech and Language Therapist, MRCSLT; Teaching and Research Academic, Department of Allied Health Professions, Sheffield Hallam University, Sheffield, UK. Expertise: Aphasia assessment and management, stroke rehabilitation, single case, case series, mixed methods.	Steven L. Small PhD MD Professor of Neurology, University of California, Irvine <i>Expertise: Neurobiology of Language,</i> <i>Cognitive Neurology.</i>	Janet Webster PhD MRCSLT Registered Speech and Language Therapist, Teaching and Research Academic, Newcastle University, UK Expertise: Post-stroke aphasia assessment and management, single subject design.	

OMIs (n=50) identified in scoping review and retained following application of the

consensus-based criteria

Construct	Outcome measurement instrument
Language	 The Comprehensive Aphasia Test (CAT) (1) The Western Aphasia Battery Revised (WAB-R) (AQ+LQ) (2) Therapy Outcome Measures (TOM) (3-5) The Aphasia Checklist (ACL) (6) Aachen Aphasia Test (AAT) (7) Aphasia Language Assessment Test (ALA) (8) The Thai Aphasia Language Performance Scales (ALPS) (9) Bilingual Aphasia Test (BAT) (10) The Boston Diagnostic Aphasia Examination (BDAE) (11) Ege Aphasia Test (I2) Kentucky Aphasia Test (KAT) (13) Montreal-Toulouse Language Assessment Battery (MTL) (14) The Norsk Grunntest for Afasi (NGTA) (15)
Emononal well-being	 The Norsk Grunntest for Afasi (NGTA) (15) Communication Confidence Rating Scale for Aphasia (CCRSA) (16) Hospital Anxiety and Depression Scale (HADS) (17) Montgomery-Asberg Depression Rating Scale (MADRS) (18) Geriatric Depression Scale (GDS) 15 item / 30 item (19, 20) Warwick and Edinburgh mental well-being scale (21) Geriatric anxiety scale (22) Stroke and Aphasia (SAD) Scale (23) Signs of Depression Scale (SODS) (24) Stroke Aphasic Depression Questionnaire (SADQ) (25) Visual Analogue Self-Esteem Scale (VASES) (26) Centre for Epidemiology Depression Scale –Revised (27)

-		Aphasia Communication Outcome Measure (ACOM) (35)
		• American Speech-Language and Hearing Association Functional Assessment
		of Communication Skills for Adults (ASHA-FACS) (36)
		 Amsterdam-Nijmegen Everyday Language Test (ANELT) (37)
		• The Communication Activity Log (CAL) (38)
		• The Communication Outcome After Stroke (COAST) (39)
	Ę	• The Communicative Activities Checklist (COMACT) (40)
	Itio	• The Social Activities Checklist (SOCACT) (40)
	Communication	• The Communication Disability Profile (CDP) (41)
	Inu	• The Communication Effectiveness Index (CETI) (42)
	III	• Community Integration Questionnaire (CIQ-R) (43)
	C	 Communication Activities of Daily Living (CADL) (44)
		• The Functional Outcome Questionnaire for Aphasia (FOQ-A) (45)
		• Measure of participation in conversation (MPC) (46)
		• The Scenario Test (47)
		• The Speech Questionnaire (48)
		• Therapy Outcome Measures (TOM) (29-31)
		• The Communication Participation Item Bank (49)
-	ife	Aachen Life Quality Inventory (ALQI) (50)
	f Li	• Burden of Stroke Scale (BOSS) (51)
	y o	• The Newcastle Stroke-Specific Quality of Life Measure (NEWSQOL) (52)
	Quality of Life	• Short Form 36 Health Survey (SF-36) (53)
	Qu	• Stroke and Aphasia Quality of Life Scale (SAQOL-39) (54, 55)

Description of recommended outcome measurement instruments

Outcome	Development /	Aims/instrument	Number	Du	ration	Sc	oring system	Training	Cost*/	Language
instrument and	alternate versions	description	of items						availability	translations
abbreviation										
Western	Developed by	Primary: Assessment	>300	•	Bedside WAB-	•	Aphasia Quotient	Administration:	Testing	Cantonese (57)
Aphasia	Kertesz in 1979	of linguistic skills in			R: 15 min		(AQ): a weighted	"some training"	materials:	Korean (58)
Battery	based on the	aphasia:			(comprises half		average of the	required	+++	Bangla (59)
Revised	original format of	1. Spontaneous speech			of the items of		WAB spoken	according to		Tagalog (60)
(WAB-R) (2)	the Boston	2. Auditory verbal			WAB-R Part 1)		language subtest	developers.	Available	Brazilian
	Diagnostic	comprehension		•	Part 1: 30-45		scores.	actorperst	from:	Portuguese (61)
	Aphasia	3. Repetition			min	•	Cortical Quotient	Scoring	https://ww	Japanese (62)
	Examination (56).	4. Naming and word		•	Part 2: 45-60		(CQ): a weighted	procedures	w.pearsonc	Hungarian
		finding			min		average of both	require training.	linical.com	French
	Revisions	5. Reading					the language and		<u>Innear.com</u>	Turkish (63)
	published in 1982	6. Writing					non-language			Hebrew
	and 2006 (WAB-	7. Apraxia					subtest scores.			Spanish (64)
	R):	8. Constructional,				•	The Language			Spanish (01)
	Supplemental	visuospatial, and					Quotient (LQ):			
	tasks, revision of	calculation tasks					reflects auditory			
	15 items and	9. Supplemental					comprehension,			
	testing materials	writing and reading					oral expression,			
	(e.g. spiral-bound	tasks: reading and					reading, and			
	stimulus book	writing of irregular					writing			
	replacing loose	and non-words					performance.			
	stimulus cards), as	(WAB-R only)					1			
	well as revised	Secondary: Assessment								
	directions and	of non-linguistic skills								
	scoring guidelines	in aphasia:								
	for clarity.	drawing, block design,								
		calculation, and praxis								
	The WAB-R also	1. Additional aims:								
	includes a bedside	Classification of 8								
	screening tool	aphasia types:								
	(Bedside WAB-	Global, Broca's,								
	R).	Transcortical motor,								
		Wernicke's,								

	 Transcortical sensory, Mixed transcortical, Conduction, and Anomic Assessment of aphasia severity Used to determine the location of the lesion 						
Aphasia Quality of Life Scale (SAQOL-39; SAQOL-39g) (54, 55)39 is the form of SAQOL is itsel adapta the SS (Stroke specifi Quality scale).The SA 39 was origina tested people chroni aphasi measu 	AQOL- he shortInterview-administered self-report measure, SAQOL-39 comprises 39 questions, in four quality of life (QoL) domains:0, which ation of 6-QOL1. Physical (17 items) 2. Communication (7 items)2. Communication (7 items)3. Psychosocial (11 items)3. Psychosocial (11 items)3. Psychosocial (11 items)4. Energy (4 items)AQOL- s ally in icSAQOL 39g comprises the same 39 questions, in three quality of life (QoL) domains:a (the ic c ia (the cal, osocial unicatio1. Physical (16 items) 2. Communication (7 items)3. Psychosocial (16 items)3. Psychosocial (16 items)3. Psychosocial (16 items)	39	15-20 min (depending on severity of aphasia)	 Twenty-one of the items ask the respondents how much trouble they have had with activities (e.g., getting dressed, speaking). The response format for these questions is a 5- point scale that varies from 1='couldn't do it at all' to 5='no trouble at all'. The rest of the items (18) ask about feelings (e.g., 'did you feel irritable?') and other activities (e.g., 'did you see your friends less often than you would like?'). Their response format 	Administration: Guidance is provided in administration guidelines. Administrators need to have skills in communicating with people with aphasia Scoring procedures: no training required	Free. Available from: https://blog s.city.ac.uk /cityaccess /saqol- description /	Chilean (68) Chinese (69) Chinese mandarin (70) Dutch (71) Greek (72, 73) Hindi (74) Italian (75) (76) Japanese (77) Kannada (78) Korean (79) Malayalam (80) Persian (81) Portuguese (82) Spanish (83) Turkish (84)

	Testing the SAQOL-39 in generic stroke population (n=87) resulted in the SAQOL-39g, which has the same items as the SAQOL-39 but three domains (all energy items groups with the psychosocial domain). There are alternative forms for proxy administration (65, 66) and for postal and telephone administration (67)	Multi-modal presentation, i.e., patients can both read and listen to the questions. People with expressive aphasia can point to their responses instead of verbally responding.			 varies from 1='definitely yes' to 5='definitely no'. Calculation of: 1. total score: mean score of all 39 items 2. Domain scores: mean score of all items relating to the respective domain 			
General Health Questionnaire (GHQ) 12	Developed in 1972. Current version published in 2011) Alternate versions: • GHQ-60: 60- item questionnaire • GHQ-30: a short form without items relating to	 Primary: Screening device for identifying minor psychiatric disorders in the general population and within community or non- psychiatric clinical settings such as primary care or general medical out-patients. 12 questions relating to symptoms of various psychiatric conditions, assesses the respondent's 	12	2 min administration time (in non-language impaired samples)	 4-scale response options (exact wording depends on item): 1. 'better/healthier than normal' 2. 'same as usual' 3. 'worse/more than usual' 4. 'much worse/more than usual' 	Administration: no training required. Scoring procedures: no training required.	Testing materials: + Available from: https://ww w.gl- assessment .co.uk	Italian (85) Arabic (86) Turkish (87) Persian (88) Portuguese (89) Kannada (90) Hindi (91) Spanish (92) A number of other unvalidated translations are available. The MAPI Research

physical illnesscurrent state and asks if that differs from his or• GHQ-28: a 28 item scaledher usual state, and is therefore sensitive to short-term psychiatric disorders.version - assesses somatic symptoms, anxiety and insomnia, social dysfunction and severe depression (7 items for each of the four scales)	4 possible methods of scoring. GHQ scoring (0-0-1-1) is advocated by the test author. GHQ-12 yields only an overall total score (range: 0 to 12 points with standard scoring procedure).	Trust distributes translated versions on behalf of GL Assessment. Contact: <u>PROinformation</u> <u>@mapi-trust.org</u>
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* Free, + Up to US\$100, ++ Up to US\$200, +++ > US\$200

Properties of recommended outcome measurement instruments

	Western Aphasia Battery – Revised (WAB-R)	Stroke and Aphasia Quality of Life Scale (SAQOL- 39/39g)	General Health Questionnaire (GHQ-12)
Objectivity	 During assessment: Limited because no audio recordings of verbal stimulus material available During scoring: Limited for spontaneous speech and written output subtests 	 During assessment: Moderate (interaction between assessor and patient frequently required because of physical stroke symptoms (arm paresis) and lack of pictorial task instructions (written sentences only) During scoring: High 	 During assessment: High if assessor does not interact with patient During scoring: High
Internal consistency	High: Cronbach's alpha of total score= 0.91 (93).	High: Cronbach's alpha of total score= 0.93; Cronbach's alpha of subscale scores= 0.74–0.94 (54). SAQOL-39g: High: Cronbach's alpha of total score= 0.95; Cronbach's alpha for subscale scores=	High (in general population): Cronbach's alpha of total score= 0.79-0.91 (94-96). Cronbach's alpha of subscale scores= 0.80-0.92.
		0.92-0.95 (55)	
Test-retest reliability*	Excellent test-retest reliability: r >0.90 Acute stage post stroke:	Good to excellent test-retest reliability ICC=0.89- 0.98	Acceptable to excellent test-retest reliability
	 Korean version; (58); 5-day test–retest interval (n=20 people with aphasia; Aphasia Quotient: r=0.976; Language Quotient: r=0.977; Cortical Quotient: r=0.920; Spontaneous Speech: r=0.96; Auditory Comprehension: r=0.967; Repetition: r=0.952; Naming: r=0.934; Reading: r=0.986; Writing: r=0.988; Praxis, r=0.908; Construction: r=0.922). Chronic stage post stroke: 	 English version; 2 to 14 days; n=17 people with aphasia; ICC=0.98 overall, 0.94–0.98 subscales (54). English generic stroke version (SAQOL-39g); 7 ± 4 day test-retest interval; n=18 people with stroke/ stroke and aphasia; ICC= 0.96 overall; ICC= 0.92–0.98 subscales (55) Other translated versions: Chilean version; ICC=0.95 (67) Chinese ICC=0.97(69) 	 General population: ICC=0.79-0.82 (100) Stroke (inc. aphasia) population using GHQ-28: 2 month test-retest reliability with a sample of 20 individuals (r=0.90) (101)
	• 1 year test–retest interval (97), n=22 patients, r=0.992	 Chinese mandarin version; ICC=0.98 (70) Dutch ICC=0.9 (71) Greek ICC=0.96 (73) 	

	 6 months to 6.5 test-retest interval (av. 12-23 months test-retest interval; (93)), n=38 patients with chronic aphasia; WAB-AQ (r=0.968), WAB-CQ (n=9, r=0.895), WAB-LQ subtests: Spontaneous Speech – Information Content (r=0.947) and Fluency (r=0.941), Comprehension (r=0.881), Repetition (r=0.970), Naming (r=0.923), Reading (n=32; r=0.927) and Writing (n=25; r=0.956) and the Construction subtest (n=14, r=974). Test-retest reliability was adequate for the Praxis subtest (n=18, r=0.581). Danish version (98); 3.5 months test-retest interval; n=19, r=0.96. Cantonese version (99); 12 to 16 months test-retest interval; n=16 patients, Spontaneous Speech subtest – Information, Fluency and total scores (r=0.83, 0.94, 0.96 respectively), Naming subtest (r=0.91), AQ (r=0.93). 	 Hindi ICC=0.9 (74) Italian ICC=0.916 (75) (76) Japanese ICC=0.97 (77) Kannada ICC=0.8 (78) Korean ICC=0.909 (79) Malayalam ICC=0.91 (80) Persian ICC=0.93 (81) Portuguese ICC=0.927 (82) Spanish ICC=0.949 (83) Turkish ICC=0.97 (84) 	
Responsiveness	 Sub-/acute phase (up to 1 month post-onset): WAB-LQ: n=50 adults with aphasia secondary to acute stroke, who received treatment (n=42) or no treatment (n=8). Participants assessed at baseline (2-4 weeks post-onset of aphasia), 3 months, and at least 6 months post-baseline. Significant main effect for time (F=43.33, df=2.96, p<0.0001), significant differences in the mean scores for the three tests (p<0.01). (102) Very Early Rehabilitation of Speech (VERSE) trial; n=20 participants with mild-severe aphasia receiving intervention (4-5 h/wk for 5 wks) achieved 18% greater recovery on the 	 Acute to post-acute phase (up to 6 months post-onset): Generic stroke sample, n=87; people admitted to hospital with a first stroke were assessed two weeks, three months and six months post stroke. Moderate changes (d = 0.35—0.49; standardized response mean (SRM) = 0.29—0.53) from two weeks to six months support responsiveness. (55) Post-acute to chronic (3 months to 1 year) Cohort study of stroke sample with and without aphasia, n=78. Effect size r=0.22. MID estimated 0.21. (107) Chronic phase (at least 6 months post-onset): 	 Acute to post-acute phase (up to 6 months post-onset): Impact of stroke with and without aphasia across the first six months, n=87 people with stroke or stroke and aphasia; psychological distress significantly reduced with time on GHQ-12 [F (2,140) = 7.1, p=0.001] (109) Chronic phase (at least 6 months post-onset): Effects of singing in a community choir on mood; n=13 people with aphasia; 2.8 point reduction in mean GHQ-12 score was seen by week 12,

	 WAB-AQ compared to the usual care group (11 min/week for 3 wks) (103). Post-acute phase (2-6 months post-onset): See (102) above Prospective longitudinal study with n=75 participants with aphasia post stroke, assessments at 4, 8, 12 and 24 weeks post-stroke, significant improvement in WAB-AQ across first year post-stroke (104) Chronic phase (at least 6 months post-onset): n=10 participants with chronic aphasia. Combination of d-amphetamine, TMS, and SLT superior to control intervention of placebo with TMS and SLT; Change in AQ (from 36.13[18.23] to 38.60[19.33], P = 0.04) and LQ (from 32.41[14.93] to 35.03[15.10], P = 0.02) showed a statistically significant increase in the active experiment. Comparison of proportional changes of AQ and LQ in the placebo experiment showed a significant difference (AQ, P = 0.02; LQ, P = 0.008) (105) 	 Intensive speech and language therapy compared to a waiting list control condition; n=156; Verbal communication was significantly improved from baseline to post-treatment (mean difference 2.61 points [SD 4.94]; 95% CI 1.49 to 3.72), but not from baseline to after treatment deferral (-0.03 points [4.04]; -0.94 to 0.88; between-group difference Cohen's d=0.58; p=0.0004). F-value for the main comparison is 12.97 (df1=1, df2=153), p= 0.0004 (108) 	 suggesting a possible reduction in adverse mood symptoms that was sustained to week 20. (110) Effects of solution-focused brief therapy, n=5 people with aphasia, On GHQ-12 the mean (SD) score before therapy was 4.80 (4.60) [median (IQR) = 6.00 (0–9.00)]. This was reduced after therapy to a mean (SD) score of 2.00 (2.55) [median (IQR) = 1.00 (0–4.50)]. The effect size was large: Cohen's <i>d</i> = 0.79. (111) Caregivers of people with aphasia: Impact of a psychoeducation program on caregivers' burden and stress, n =31 caregivers of people with post stroke aphasia. Caregivers in the immediate treatment group had significant reductions in GHQ-12 measured stress (GHQ mean (SD) at baseline =6.26 (5.67), GHQ post treatment 3.21 (SD 4.20), =/0.006). (112)
Convergent	 Mixed stages n= 50 participants with aphasia (49 secondary to subacute or chronic stroke). Participants' mean scores improved significantly from pre- to post-treatment on all WAB subtests, with absolute percentages ranging from 6.5% to 13% improvement (p<0.01 to p<0.0001) (106). 	s SAQQL 20. Cood convergent well lite (c. 0.55	Convergent validity in post-stroke aphasia
validity	• Convergent validity in sample of n=15 people with aphasia (93). Comparison	• SAQOL-39: Good convergent validity (r=0.55 to 0.67)(54). Adequate correlation between	sample:

Discriminant	 with corresponding subtests of the Neurosensory Center Comprehensive Examination for Aphasia (NCCEA), using Pearson correlation coefficients Excellent correlation between: WAB Spontaneous Speech and NCCEA Description of Use and Sentence Construction (r= 0.817); WAB Comprehension and NCCEA Identification by Name and Identification by Sentence (r= 0.915); WAB Repetition and NCCEA Sentence Repetition (r= 0.880); WAB Naming and NCCEA Visual Naming and Word Fluency (r= 0.904); WAB Reading and NCCEA Reading subtests (r=0.919); WAB Writing and NCCEA Writing subtests (r=0.905); and WAB and NCCEA total scores (r=0.973). Excellent correlation between the WAB-CQ (minus the Praxis and Construction subtests) and a comparable NCCEA score (minus the Tactile Naming- Right/Left, Articulation, Digit Repetition-Forward/Backward subtests) (r=0.964). Sample of n=45 people with aphasia. Excellent correlation between the WAB and the Czech version of the Mississippi Aphasia Screening Test (MASTcz) (r= 0.933) (113) Sample of n=140 people with aphasia. 	 GHQ-12 and the SAQOL-39 mean (0.53, p<0.01). The physical, communication, and energy subscales show good convergent validity (r=0.39 to 0.67, r=0.55, r=0.32, respectively). The psychosocial subdomain shows adequate convergent (r=0.28 to 0.62) validity with only 1 correlation lower than predicted (r=0.28 with the SSS). Good correlations with Frenchay Activities Index (FAI) and ASHA Functional Assessment of Communication Skills (ASHA-FACS). SAQOL-39g: Good/excellent convergent validity for overall scale (r=0.36–0.70); and subdomains (r=0.47–0.78) (55), evidenced by moderate to high correlations with measures of stroke severity (NIHSS), activities of daily living (Barthel Index), extended activities of daily living (Frenchay Activities Index), emotional distress (GHQ-12) and language (Frenchay Aphasia Screening Test). 	 Good correlations with SAQOL 39/SAQOL-39 (English, Greek, and Turkish versions). The GHQ-12 demonstrated good convergent validity in a sample of 83 individuals with chronic stroke and aphasia, by comparison with the SAQOL-39. The study yielded an adequate correlation between the GHQ-12 and the SAQOL-39 mean (0.53, p<0.01). Correlations between the GHQ-12 and SAQOL-39 subtests were adequate (physical r=0.39, energy r=0.32, p<0.01) to excellent (psychosocial r=0.62, p<0.01). (54) Excellent discriminant validity in Swedish
validity	Comparison of WAB with Raven's	(54)	population (n=556 patient cases surveyed in specialized psychiatric care outpatient age and n=556 sex-matched controls).

AQ and the BI $(r=0.44)$ and the FAI $(r=0.50)$.

* Test-retest reliability: 1=perfect reliability; ≥ 0.9 =excellent reliability; $\geq 0.8 < 0.9$ =good reliability; $\geq 0.7 < 0.8$ =acceptable reliability; $\geq 0.6 < 0.7$ =questionable reliability; $\geq 0.5 < 0.6$ =poor reliability; < 0.5=unacceptable reliability; 0=no reliability.

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