






BMJ Open Developing the TIDieR-Rehab checklist: a modified Delphi process to extend the Template for Intervention Description and Replication (TIDieR) for rehabilitation intervention reporting

Nada Signal , Sharon Olsen , Emeline Gomes , Caitlin McGeorge, Denise Taylor , Gemma Alder 

To cite: Signal N, Olsen S, Gomes E, *et al.* Developing the TIDieR-Rehab checklist: a modified Delphi process to extend the Template for Intervention Description and Replication (TIDieR) for rehabilitation intervention reporting. *BMJ Open* 2024;**14**:e084319. doi:10.1136/bmjopen-2024-084319

► Prepublication history and additional supplemental material for this paper are available online. To view these files, please visit the journal online (<https://doi.org/10.1136/bmjopen-2024-084319>).

Received 16 January 2024
Accepted 12 August 2024



► <http://dx.doi.org/10.1136/bmjopen-2024-084320>



© Author(s) (or their employer(s)) 2024. Re-use permitted under CC BY-NC. No commercial re-use. See rights and permissions. Published by BMJ.

Health and Rehabilitation Research Institute, Auckland University of Technology, Auckland, New Zealand

Correspondence to

Nada Signal;
nada.signal@aut.ac.nz

ABSTRACT

Objectives Rehabilitation is essential for supporting the recovery from, and management of, a range of health conditions. However, interventions are often poorly reported in rehabilitation research, hindering advancement of the field. The Template for Intervention Description and Replication (TIDieR) checklist was developed to enhance the reporting of interventions, but does not specifically address the complexities and multifaceted nature of rehabilitation interventions. This study aimed to develop an extension of the TIDieR checklist to support better reporting of rehabilitation interventions.

Design A modified Delphi study overseen by a Steering Committee.

Setting Online.

Participants Rehabilitation experts were purposively sampled for diversity in discipline, practice setting, area of expertise and geographical location.

Methods Participants (n=35) provided both quantitative and qualitative feedback on drafts of the TIDieR-Rehab through online surveys. Quantitative data was descriptively analysed by percentage of agreement, while qualitative data was analysed using conventional content analysis. Quantitative and qualitative findings were subsequently triangulated to facilitate iterative refinement of the TIDieR-Rehab.

Results Consensus was achieved after two rounds of the modified Delphi process. The TIDieR-Rehab checklist comprises seven original, three adapted and 12 new reporting items, and is supported by a supplementary manual. Specific enhancements include more detailed descriptions of the study population (*Who*) and timing of the intervention (*When*), the planned intervention dosage (*How much*, *How challenging* and *Regression/Progression*), person-centred care (*Personalisation*) and negative undesired effects (*Harms*) which were considered critical for the comprehensive reporting of rehabilitation interventions.

Conclusion The TIDieR-Rehab checklist marks a significant advancement in enhancing and standardising the reporting of rehabilitation interventions. By offering a structured format for detailing complex rehabilitation interventions, the TIDieR-Rehab supports improvements

STRENGTHS AND LIMITATIONS OF THIS STUDY

- ⇒ A modified Delphi process was used to develop the TIDieR-Rehab checklist, an extension of the original TIDieR, which supports the comprehensive reporting of rehabilitation interventions.
- ⇒ The TIDieR-Rehab extension includes seven original, three adapted and 12 new items which may facilitate the development, reporting, synthesis and translation of rehabilitation intervention research.
- ⇒ Our findings reveal ambiguity in key concepts underpinning rehabilitation, highlighting the need to develop shared conceptual understandings and a common language to describe rehabilitation interventions.
- ⇒ Survey-based modified Delphi approaches, supported by a Steering Committee, are common practice in health research and the development of reporting quality tools but may limit the depth of participant feedback.
- ⇒ Extensive efforts were made to capture a diversity of perspectives, but some relevant cultural and disciplinary perspectives may not be fully represented in this research.

in reporting quality to promote research replication and support the translation of research findings into clinical practice. Future research should validate the TIDieR-Rehab checklist across a variety of intervention types and clinical contexts.

INTRODUCTION

Beyond medical and surgical interventions, rehabilitation serves as a primary approach in the management of numerous health conditions and mitigation of disability.¹ Rehabilitation is conceptualised as a complex biopsychosocial process where multidisciplinary professionals work in collaboration with a person and their family, with the shared objective of enhancing the

person's function, autonomy and societal participation.^{1–3} Rehabilitation is commonly operationalised as a personalised, goal-directed, multicomponent package of non-pharmacological interventions, such as targeted activities, education and psychosocial support.^{1 2} The efficacy of rehabilitation interventions is often attributed to their 'essential elements' or 'active ingredients' and the underlying processes or pathways through which an intervention is expected to produce its effects.^{2 4} Understanding these essential elements and their mechanisms is essential for optimising effectiveness and minimising any negative undesired effects of an intervention.^{2 4}

The efficacy of rehabilitation interventions is also thought to be strongly influenced by the dosage of the essential elements.^{4 5} *Dosage* encompasses both amount and challenge.^{5 6} *Amount* generally refers to the total time invested in the intervention, considering session duration, frequency and overall length of the intervention programme.^{5 6} However, there is growing concern that current understandings of 'amount' do not reflect the time spent engaged in the essential elements of the intervention.⁵ Relying on session duration to determine amount, which often includes a significant proportion of time spent in non-essential or inactive elements, can potentially obscure our understanding of the efficacy of the essential elements of an intervention.^{4 5} *Challenge* has been conceptually defined as a multifaceted, multidimensional and dynamic interaction between the rehabilitation activity or task, the person's ability and their subjective experience and may reflect how 'difficult', 'demanding' or 'effortful' the task is.^{5–8} As such, the operationalisation of 'challenge' can vary widely across disciplines, interventions and tasks. Nonetheless, rehabilitation research suggests that challenge, as a critical parameter of dosage, is likely a key driver of rehabilitation efficacy.^{6 7}

Incomplete understanding of the essential elements of rehabilitation interventions and their dosage parameters has hampered advancement of the field and slowed translation of research to clinical practice.⁴ A key contributor to this gap is the incomplete reporting of rehabilitation interventions in the research literature.^{4 5 9–12} In non-pharmacological interventions, like rehabilitation, essential elements which are multidimensional, personalised and often difficult to quantify add significant complexity to intervention reporting.^{13–15} Tools such as the Standard Protocol Items: Recommendations for Interventional Trials (SPIRIT),¹⁶ Consolidated Standards of Reporting Trials (CONSORT)^{13 17} and the Template for Intervention Description and Replication (TIDieR)¹⁴ have been developed to address such concerns. Existing checklists provide a valuable foundation for improved reporting. However, they are not specific to *rehabilitation* interventions, and thus, the essential elements of rehabilitation interventions and their dosage parameters continue to be under-reported.^{4 7 9} This lack of specificity has been noted in other non-pharmacological fields and researchers have subsequently proposed extensions or adaptations to the TIDieR checklist in areas such as

public health¹⁸ and telehealth.¹⁹ The Consensus on Exercise Reporting Template (CERT)²⁰ is a recognised extension of the TIDieR that specifically addresses dosage, but its focus is limited to a single aspect of rehabilitation interventions: exercise. Therefore, current checklists and frameworks may not be applicable to the breadth and complexity of rehabilitation interventions, leading to incomplete reporting or the use of multiple checklists.^{21 22}

Without complete and readily available descriptions, rehabilitation interventions cannot be replicated for further evaluation in research or effectively translated to clinical practice. Furthermore, the synthesis and meta-analysis of rehabilitation intervention evidence relies heavily on the precise reporting of essential elements and dosage parameters.^{4 5} Inaccurate reporting can significantly undermine evidence synthesis, which is foundational to clinical practice guidelines and the consequent advancement of the field.^{5 14} This study describes the modified Delphi process used to develop the TIDieR-Rehab checklist, an extension of the original TIDieR, to enhance the reporting quality of rehabilitation interventions.

METHODS

Design

The aim of this study was to extend an existing research reporting tool (TIDieR) to promote its applicability and relevance to the field of rehabilitation. We drew on the Enhancing the Quality and Transparency of Health Research (EQUATOR) Network guidelines²³ and other relevant recommendations^{15 24} to inform the development of the TIDieR-Rehab checklist. A modified-Delphi methodology was deemed an appropriate approach to gather expert feedback in a structured manner, refine the checklist iteratively and achieve consensus.¹⁸ Through successive rounds of online surveys, drafts of the TIDieR-Rehab checklist were presented to individuals with rehabilitation expertise for feedback and iteratively refined until consensus (as defined below) was reached. The TIDieR-Rehab checklist and accompanying supplementary manual²⁵ were then piloted to inform minor refinements. A multidisciplinary Steering Committee offered guidance throughout the process. The following methods and results are described in accordance with Delphi reporting quality indicators.²⁶

Steering Committee

The Steering Committee (n=7) consisted of experts spanning a variety of rehabilitation disciplines including physiotherapy, speech-language therapy, occupational therapy and psychology, and represented different geographical locations (see Acknowledgements). Their role encompassed developing, reviewing, critiquing and piloting iterations of the TIDieR-Rehab checklist and supplementary manual.²⁴ Furthermore, they provided advice and confirmed consensus decisions, including the termination of survey rounds.

Ethics

Ethical approval for the study was granted by the Auckland University of Technology Ethics Committee (reference: 20/264).

Participants

A target of 30 participants was selected to balance expert representation, efficient consensus-building and active engagement, which aligns with literature that supports a panel size of approximately 30 participants for modified Delphi studies.^{15 24 26} Potential participants were actively involved in the design, conduct or reporting of rehabilitation intervention research. Potential participants were identified by reviewing authorship of English language peer-reviewed papers specifically addressing the reporting of rehabilitation interventions or exploring key components of rehabilitation interventions. Purposive sampling was employed to capture a breadth of perspectives, including rehabilitation discipline (eg, physiotherapy, occupational therapy, speech-language therapy, psychology, rehabilitation medicine and nursing), practice setting, area of expertise and geographical location.^{15 18 24}

Potential participants were invited to participate via their publicly available work email address and were provided an information sheet that outlined the scope of the study.¹⁸ Those interested in participating were asked to provide consent by copying a consent statement and an expression of agreement to participate into a reply email. Non-responders were sent up to two reminders.¹⁸ Invitations were dispatched in four waves of 50–70 invitations per wave. Following each wave, the research team reviewed the sample against the purposive sampling framework to inform the next wave of invitations.

TIDieR-Rehab development

The first draft of the TIDieR-Rehab checklist was developed through a multiphase process. An appraisal of the existing TIDieR¹⁴ and CERT²⁰ checklists and expert commentaries^{27–30} was undertaken to identify key issues in reporting rehabilitation interventions (refer to online supplemental file 1).^{15 24} The TIDieR¹⁴ and CERT²⁰ checklists were then applied within bodies of rehabilitation literature^{31–34} to identify limitations in their practical application. A draft of the TIDieR-Rehab checklist was then developed and presented to the Steering Committee for feedback.¹⁵ In alignment with the modified Delphi methods used in similar studies aiming to enhance established checklists,^{14 18 20} this draft was presented in the first survey round.

Survey

Delphi rounds were conducted using an online survey tool (Qualtrics, Provo, UT), where participants received personalised links to provide feedback on the TIDieR-Rehab drafts. Refer to online supplemental file 2 for copies of the surveys.

In round 1, participants reviewed the first draft of the TIDieR-Rehab checklist. For each item, participants were asked to provide quantitative and qualitative feedback. Quantitative feedback indicated agreement with the statement ‘This item should be included in the TIDieR-Rehab checklist’ via a five-point Likert scale ranging from ‘strongly disagree’ to ‘strongly agree’.¹⁸ Qualitative responses were via an optional free comments box in response to the prompt ‘Please provide an explanation of your response in the comments box of each item e.g., suitability, wording, grouping etc.’¹⁸ Items were presented in the intended checklist order to maintain the flow of reporting.^{18 20} Participants were also asked, ‘Are there any aspects of intervention reporting you believe were not adequately addressed within this draft checklist?’ and ‘Please provide any other feedback or general comments below’.

Participants who completed round 1 were invited to participate in round 2 to review revisions of the TIDieR-Rehab checklist. In round 2, participants were given a summary of the results from the preceding round.¹⁸ The summary of results indicated which items had achieved consensus and had been retained for inclusion, and which items were adapted, removed or new in response to the previous round’s feedback. For items that had achieved consensus in the previous round, participants were invited to provide qualitative feedback if they did not agree with the consensus using the method described in round 1. Items that had not yet achieved consensus and were subsequently adapted, removed or added were presented alongside a summary of the previous round’s feedback to enable consideration of other participants’ perspectives and facilitate consensus building. In response to these items, participants were asked to provide quantitative and qualitative feedback as described in round 1, except that a four-point Likert scale without the ‘neutral’ option was used to encourage decisive ratings and minimise ambiguity.³⁵ Participants were also asked if any aspects of intervention reporting were not adequately addressed and to provide general feedback about the TIDieR-Rehab checklist and the supplementary manual through free comment boxes. Survey rounds were continued until consensus was reached. Each survey took up to 30 minutes to complete.

Data analysis

At the end of each round, participant responses were exported from Qualtrics to Microsoft Excel for manual deidentification and analysis.¹⁸ Quantitative responses were descriptively analysed by percentage agreement of each Likert response for each item. Qualitative responses were analysed using an inductive approach to conventional content analysis.³⁶ Researchers (EG, CM) repeatedly immersed themselves in the data item by item and independently developed inductive qualitative codes. Researchers (EG, CM guided by GA) compared and agreed on codes and then developed subcategories, categories and themes. Quantitative and qualitative data were

subsequently triangulated (EG, CM) to determine if data were convergent, complementary or dissonant.³⁷

Consensus

Item consensus was determined based on both quantitative and qualitative data. Based on quantitative data, items receiving high agreement (>80% strongly agree and agree responses) were considered for inclusion, items receiving moderate agreement (50%–80%) were considered for adaption and items receiving low agreement (<50%) were considered for exclusion from the checklist.²⁰ After each round, the research team and Steering Committee reviewed the quantitative, qualitative and triangulated findings to guide iterative development of the checklist and determine consensus for each item. Termination of survey rounds was predefined based on >70% of checklist items reaching high quantitative agreement and the Steering Committee's confirmation to conclude rounds.³⁸

Patient and public involvement

None.

RESULTS

Participants

Invitations were sent via email to 270 individuals with expertise in rehabilitation. For a summary of the potential participants invited to contribute to the study, please refer to online supplemental file 3. In total, 35 individuals with rehabilitation expertise consented and participated in round 1 (response rate=13%) and 23 of these 35 participants completed round 2 (response rate=66%). Participants self-identified as researchers (n=30), 'other' professionals (n=3) and journal editors (n=2) and were from various clinical disciplines, including neurological physiotherapy (n=14), musculoskeletal physiotherapy (n=6), other physiotherapy (n=3), rehabilitation medicine (n=4), speech-language therapy (n=4), psychology (n=2), public health (n=1) and occupational therapy (n=1). Participants were situated across 10 different geographical locations, including Australia (n=15), UK (n=6), USA (n=6), New Zealand (n=2), Brazil (n=1), Canada (n=1), France (n=1), Ireland (n=1), Netherlands (n=1) and Norway (n=1).

Modified Delphi rounds

Round 1

Round 1 proposed a first draft of the TIDieR-Rehab checklist with a total of 24 items across 11 sections. Of the 24 items, there were nine original and two adapted TIDieR items¹⁴ and 13 new TIDieR-Rehab items. The numbering of items was also updated to reflect the restructuring of the TIDieR-Rehab extension throughout. Refer to [figure 1](#) for an overview of changes through each round and online supplemental file 4 for a more comprehensive outline of the development of the TIDieR-Rehab items through each round of the modified Delphi process. The quantitative agreement from round 1 is presented in

[figure 2](#) and showed 20 out of the proposed 24 items met high agreement for inclusion (>80% strongly agree and agree). The new items that received the strongest agreement were: When and How much—Frequency (94%) and Intervention length (97%); Tailoring—Progression of dosage parameters (94%) and How well—Measurement of engagement/adherence (94%). Notably, the new items describing adverse events within the How well section also received high quantitative agreement and were convergent with positive qualitative feedback. The four items that did not meet high agreement were all newly proposed TIDieR-Rehab items and included: How—Delivery support (77%); When and How much—Work duration (69%); How challenging—Measure and set intensity (77%) and How challenging—Intensity tool (71%).

Qualitative feedback favoured the inclusion of all 24 proposed items in some form and highlighted the need for the TIDieR-Rehab checklist to enhance research and clinical practice. Suggestions and critiques were distributed across original, adapted and new items and broadly pertained to (1) terminology or wording revisions, (2) the structure of items, (3) differences in understanding, (4) relevance and application of rehabilitation concepts to particular disciplines or interventions, (5) the barriers to capturing and reporting item information and (6) the provision of item explanations or rehabilitation examples to aid clarity and usability. The four items which did not achieve high quantitative agreement were frequently accompanied by qualitative feedback which critiqued the terminology and relevance of the underlying concept.

In consultation with the Steering Committee, the triangulated quantitative and qualitative analysis from round 1 was utilised to develop a revised second draft of the TIDieR-Rehab checklist. Additionally, in response to the round 1 analysis, the TIDieR-Rehab supplementary manual was developed as an extension of the original TIDieR checklist guidance¹⁴ to provide explanations of items and rehabilitation specific examples of good reporting. These revisions and additions justified the need for further consensus building through a second survey round.

Round 2

Round 2 proposed the second draft of the TIDieR-Rehab checklist which included 22 items. Of the 22 items, the nine items from round 1 that achieved high quantitative and qualitative agreement were not presented for further quantitative feedback in round 2. The other 13 items, including the reintroduction of one original item, and nine adapted and three new TIDieR-Rehab items, were presented for quantitative and qualitative feedback in round 2. Of the 13 items presented, 11 achieved high agreement (refer to [figure 2](#)). The remaining two items bordered on high agreement. That is, the item Who received 78% agreement but received positive qualitative feedback that was dissonant to a disagreement rating and

ORIGINAL TIDieR	ROUND 1	ROUND 2	PILOTING	TIDieR-REHAB CHECKLIST
1. Brief name				1. Brief name
2. Why				# 2. Why
				* 3. Who
				* 4. When
3. What – Materials				5A. What – Materials
4. What – Procedures				5B. What – Procedures
5. Who provided				6. Who provided
6. How				7. How
7. Where				# 8. Where
8. When and How much				* 9A. How much – Session(s) duration
				* 9B. How much – Essential elements amount
				* 9C. How much – Frequency
				* 9D. How much – Intervention length
				* 10. How challenging
				* 11. Regression/Progression
9. Tailoring				* 12A. Personalisation – Needs
				* 12B. Personalisation – Preferences
10. Modifications				# 13. Protocol deviations
11. How well – Planned				14A. How well – Plan
12. How well – Actual				14B. How well – Actual
				* 15A. Harms – Plan
				* 15B. Harms – Actual

Key: **dark blue**, introduction of a new item; **blue**, major changes; **light blue**, minor changes.

Figure 1 Overview of the iterative development of the TIDieR-Rehab checklist through each stage of the modified Delphi process. *, new item unique to the TIDieR-Rehab checklist; #, adapted original TIDieR item for the TIDieR-Rehab checklist. TIDieR, Template for Intervention Description and Replication

the item Personalisation received 78% agreement but had a missing rating from one participant.

The qualitative responses in round 2 reinforced the high quantitative agreement. The qualitative analysis of rounds 1 and 2 is summarised in [table 1](#). Revisions applied to the second draft of the TIDieR-Rehab checklist as well as the addition of the supplementary manual addressed concerns that had been raised in round 1. For example, revisions to the item How challenging and its supplementary information, which included replacing the term ‘intensity’ with ‘challenging’ and providing further explanation and multidisciplinary examples of the challenge concept, were met with high agreement (87%) and largely positive qualitative feedback, supporting the changes and indicating greater shared understanding.

The round 2 responses indicated consensus and informed minor wording and ordering refinements which were reviewed and accepted by the Steering Committee. Based on participant consensus (>70% items reaching high agreement) and Steering Committee endorsement,

the modified Delphi process was terminated after the completion of round 2.

Post-survey final checklist

The TIDieR-Rehab was subsequently piloted on randomised controlled trials in two substantive areas of rehabilitation research, treadmill training and upper limb robotic rehabilitation, as part of systematic reviews and meta-analyses.^{39 40} This piloting process informed minor refinements, such as reordering of an item and minor amendments to wording and grammar in the supplementary manual (see [figure 1](#) and online supplemental file 4 for details). The 22-item TIDieR-Rehab checklist and supplementary manual were then reviewed and finalised by the Steering Committee. The final TIDieR-Rehab checklist (see [table 2](#)) retains seven original and three adapted TIDieR items and presents 12 new items unique to the TIDieR-Rehab. The checklist includes the following sections: 1. Brief name, 2. Why, 3. Who, 4. When, 5. What, 6. Who

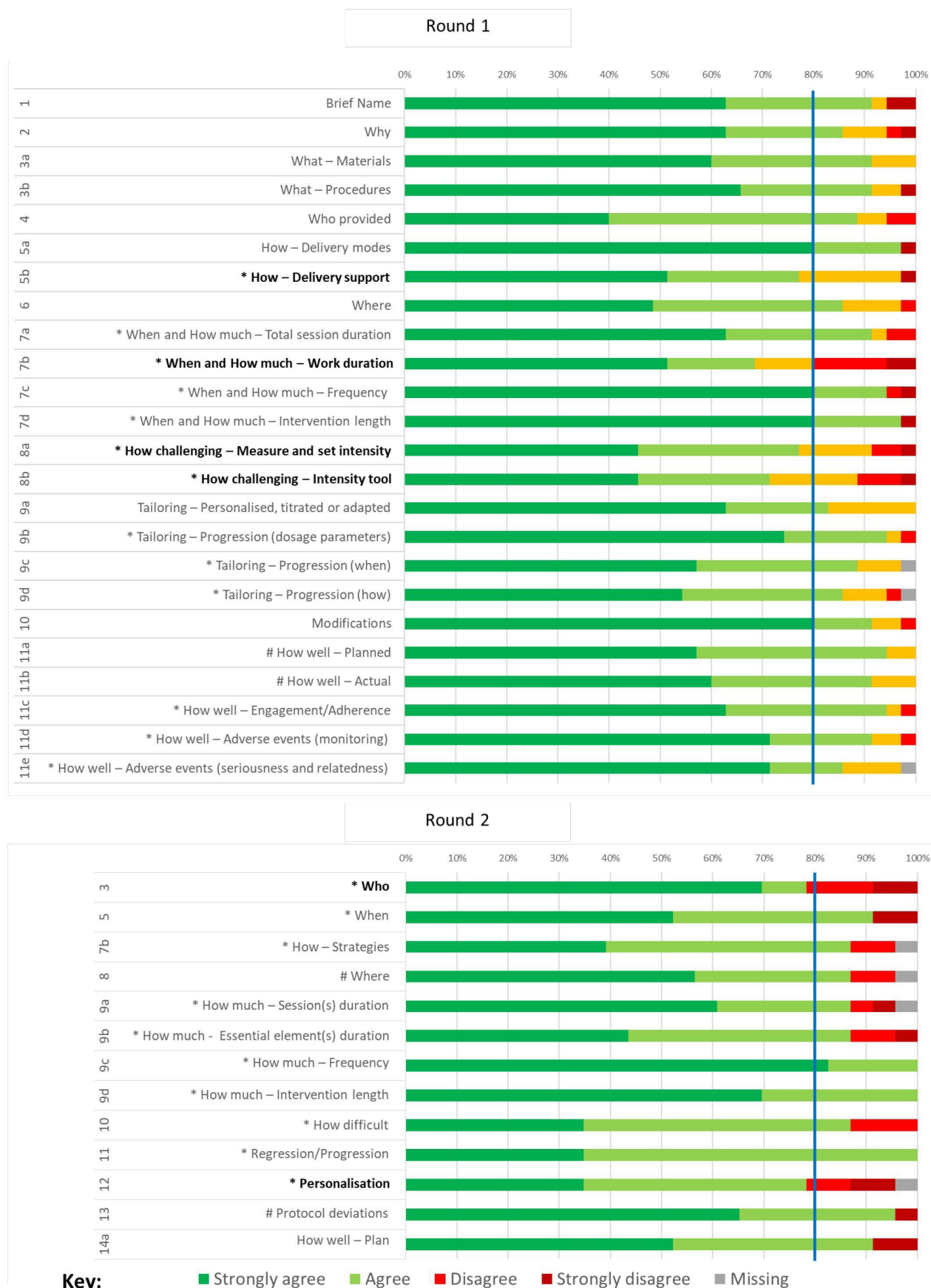


Figure 2 Rounds 1 and 2 Likert results presented in order of the TIDieR-Rehab checklist items. **Bold**, items that did not meet high quantitative agreement. *, new item unique to the TIDieR-Rehab checklist; #, adapted original TIDieR item for the TIDieR-Rehab checklist. TIDieR, Template for Intervention Description and Replication.

Table 1 Qualitative analysis of round 1 and round 2 free comments feedback

Themes	Categories (with example quote)	Subcategories	Codes (simplified)
Refining the TIDieR-Rehab checklist	Catalysts for revision 'I really like this item in some form, but I think it might be helpful to separate difficulty (as a property of the task), from challenge (how difficult a task is for a specific individual), from intensity (the effort being invested). As an analogy, playing a video game at high difficulty is less challenging for an expert than for a novice. But even then, a novice who invests more effort will likely perform better than a novice who invests less' (Participant 13).	'Correct' and clear terminology and phrasing is valued	A different name or word would be more appropriate Terms representing different rehabilitation concepts should not be used interchangeably Unclear terminology should be clarified with a definition or criteria Vague guidance may be deter people from reporting what is actually important Clarify if items report on what was 'planned' vs what 'actually' happened Unclear if challenge is 'subjective' or 'objective' Unclear boundary between 'tailoring' and 'modification' Clarify the term 'active ingredients'/'essential elements' The term 'intensity' is poorly understood within rehabilitation literature The term 'dosage' is poorly understood within rehabilitation literature Align the structure with other reporting standards Reorder words, items or sections Separate this item or section Guiding criteria could inform rationale for parameters and measurements An explanation or example would encourage better reporting An example would clarify ambiguous rehabilitation concepts Guidance is needed to understand 'when' to use the checklist The TIDieR-Rehab supplementary manual needs provide simplified guidance An item about [xxx] could be added to the TIDieR-Rehab checklist A more holistic consideration of the patient/participant experience is needed Explicit consideration of reporting on control/usual care groups is needed Reporting on this item would be insufficient or meaningless Reporting on this item would be too extensive Item is repetitive or redundant of other items Only some parts of this item are needed This aspect of research is important but may be beyond the scope of the TIDieR-Rehab checklist Not be feasible to collect this data as it would be different for every individual Fully reporting individualised rehabilitation interventions would be meaningless Concise reporting of individualised interventions requires standardisation, protocols or a word limit Overly rigorous reporting may limit the support and application of the intervention in alternate forms Emphasising the 'wrong' components of a rehabilitation intervention may attribute outcomes to incidental/non-essential elements Item risks validating a concept that is not yet well understood in rehabilitation Item should be optional as it is not applicable to all intervention types Item should be optional as it is not applicable to all disciplines Item represents a concept that is not well understood in all disciplines
		Structure supports understanding	
		Users require guidance	
		Missing items	
		Unnecessary detail or item	
		Barriers to reporting on individualised rehabilitation interventions	
		Misleading reporting	
		Questioning multidisciplinary understanding and relevance of some rehabilitation concepts	
	Tensions in complete and meaningful rehabilitation intervention reporting 'This will be highly individualised to the patients' needs, which means it will often be so broad and facile it is meaningless or so specific that is also meaningless. I think all one can do is to say (the intervention) will be given as appropriate for the patients' needs. The chances of accurately recording all these aspects of dose and encompassing the need to adapt to individual needs is low. The answer will very often be "it all depends". And if one really lists all the possibilities, it would be much too long to read' (Participant 3).		

Continued

Table 1 Continued			
Themes	Categories (with example quote)	Subcategories	Codes (simplified)
The TiDieR-Rehab checklist and supplementary manual will enhance rehabilitation	Enhancing rehabilitation research ‘[The TiDieR-Rehab is] vital to validity of results, replication from others, analysing key components of the intervention – what makes this intervention work. Thank you for doing this, a very important checklist to ensure the quality of rehabilitation interventions – much needed’ (Participant 17).	Research information gathering and evaluation	Item facilitates rehabilitation research identification
			Item facilitates literature review, synthesis and meta-analysis
		Research rigour	Supplementary manual aids clarity and usability of the checklist for exploring rehabilitation research
			Item/checklist may be used to assess or score the completeness of rehabilitation intervention reporting
			The TiDieR-Rehab checklist will clarify rehabilitation concepts
	Enhancing clinical practice ‘I think better description of rehab interventions is really important. I am also aware that filling in checklists is a pain. But I think when people are really mindful about the sorts of things captured in checklists at the start of intervention development, it can help them be really clear with some of these parameters which in rehab tend to be left at clinician’s discretion/clinical judgement, so more clarity about this is a good thing’ (Participant 24).	Future research	Item/checklist aids validity
			Item/checklist aids generalisability
		Intervention selection	Item/checklist aids transparency and fidelity
			The TiDieR-Rehab checklist may enhance the rigour of rehabilitation
			Item/checklist aids accurate replication
		Intervention translation	The TiDieR-Rehab checklist will inform the feasibility of further rehabilitation research
			The TiDieR-Rehab checklist may reduce rehabilitation research waste
			The TiDieR-Rehab checklist will inform future practice guidelines and ethics
			The TiDieR-Rehab checklist may reveal the aspects of the intervention that contributes to the outcome for further exploration
			The TiDieR-Rehab checklist may contribute to enhanced efficacy of rehabilitation
			Item/checklist will help clinicians identify which patients/groups are appropriate for the intervention
			This item/checklist informs the clinical rationale for using the intervention
			Greater attention to the multiple aspects of dosage can inform treatment planning and monitoring
			Item/checklist aids prediction of expected outcomes of the intervention
			Item is essential for responsive and adaptable translation of the intervention
			Item is essential for accurate translation of the intervention
			Clinicians need to understand all aspects of intervention delivery for effective rehabilitation
			Clinicians can use the checklist to determine and improve rehabilitation dosage
			This checklist enables clinicians to gain a better understanding of patient’s outcomes and experiences from a rehabilitation intervention
			The TiDieR-Rehab checklist may improve clinical uptake over time

Table 2 The TiDieR-Rehab checklist**SECTION 1. BRIEF NAME**

Item 1. Provide the name or a phrase that describes the intervention.

SECTION 2. WHY

Item 2. Describe any rationale, theory or goal of the elements essential to the intervention.
Essential elements, also known as 'active ingredients', are the core components of the intervention that are expected to be linked to effects or outcomes of interest.

SECTION 3. WHO

* Item 3. Describe who the intervention is intended for.

SECTION 4. WHEN

* Item 4. Describe when the intervention commenced in relation to the onset or stage of the condition and/or other relevant events.

SECTION 5. WHAT

Item 5A. Materials: Describe any physical or informational materials used in the intervention, including those provided to participants or used in intervention delivery or in training of intervention providers. Provide information on where the materials can be accessed (for example, online appendix, URL).

Item 5B. Procedures: Describe each of the procedures, activities and/or processes used in the intervention, including any enabling or support activities.

SECTION 6. WHO PROVIDED

Item 6. For each category of intervention provider (for example, psychologist, nursing assistant), describe their expertise, background and any specific training given.

SECTION 7. HOW

Item 7. Describe the modes of delivery (such as face to face or by some other mechanism, such as internet or telephone) of the intervention and whether it was provided individually or in a group.

SECTION 8. WHERE

Item 8. Describe the type(s) of environment(s) where the intervention occurred, including any necessary infrastructure or relevant features.
Environments and their features may include the relevant physical, social, cultural, economic, political and/or systemic context(s) of the intervention.

SECTION 9. HOW MUCH

* Item 9A. Session(s) duration: Specify the planned session(s) duration of the intervention.

* Item 9B. Essential elements amount: Specify the planned session(s) duration and/or repetitions of the essential elements of the intervention.
Essential elements amount refers to the time and/or repetitions, within the total duration of a single session, that is spent 'actively' participating in the core components of the intervention.

* Item 9C. Frequency: Specify the planned frequency of the intervention.

* Item 9D. Intervention length: Specify the planned overall length of the intervention.

SECTION 10. HOW CHALLENGING

* Item 10. Describe the approach(es) used to set and monitor the intervention/task challenge level.
Challenge may include the nominal, functional or perceived level of difficulty, effort, physiological intensity or cognitive load of an intervention/task at a given time and may be evaluated using subjective or objective measures.

SECTION 11. REGRESSION/PROGRESSION

* Item 11. Describe the planned regression and/or progression of dosage parameter(s), including when and how.
Dosage parameters refer to the amount (Section 9. How much) and challenge (Section 10. How challenging) of the intervention.

Continued

Table 2 Continued

SECTION 12. PERSONALISATION

* Item 12A.	Needs: If supplementary strategies were planned to enable the delivery of the essential elements of the intervention in response to specific individual or group needs, then describe what, why, when and how. <i>Supplementary strategies refer to intervention adjuncts (for example, physical assistance, verbal cueing, props) that must be used by some individuals or groups to facilitate effective participation in the essential elements.</i>
* Item 12B.	Preferences: If the intervention was planned to be adapted for personal preferences, then describe what, why, when and how.

SECTION 13. PROTOCOL DEVIATIONS

# Item 13.	If there were deviations in the intervention protocol during the course of the study, describe the changes (what, why, when and how).
------------	---

SECTION 14. HOW WELL

Item 14A.	Plan: If intervention adherence or fidelity was assessed, describe how and by whom, and if any strategies were used to maintain or improve fidelity, describe them.
Item 14B.	Actual: If intervention adherence or fidelity was assessed, describe the extent to which the intervention was delivered as planned.

SECTION 15. HARMS

* Item 15A.	Plan: Describe the monitoring of adverse consequences. <i>Adverse consequences include any negative, undesired effects related to the intervention, including physical, mental, social and/or spiritual effects.</i>
* Item 15B.	Actual: Describe any adverse consequences, including the number, seriousness and relatedness to the intervention.

*, new item unique to the TIDieR-Rehab checklist.

#, adapted original TIDieR item for the TIDieR-Rehab checklist.

TIDieR, Template for Intervention Description and Replication.

provided, 7. How, 8. Where, 9. How much, 10. How challenging, 11. Regression/Progression, 12. Personalisation (renamed from Tailoring), 13. Protocol deviations (renamed from Modifications), 14. How well and 15. Harms. The TIDieR-Rehab checklist and supplementary manual are presented in the corresponding paper.²⁵

DISCUSSION

This study developed the TIDieR-Rehab checklist and supplementary manual,²⁵ an extension of the original TIDieR, which aims to enhance the reporting of rehabilitation interventions. Rehabilitation experts reached strong consensus on the checklist items and supplementary manual. Reflecting the unique nature of rehabilitation practice, the TIDieR-Rehab extension introduces reporting requirements and guidance aligned with the current priorities and complexities of intervention reporting within rehabilitation research, emphasising concepts such as dosage, person-centred care and negative undesired effects. The new TIDieR-Rehab items within the *How much*, *How challenging* and *Regression/Progression* sections emphasised the importance experts place on understanding the role of dosage parameters in rehabilitation intervention efficacy.^{5–7 10–12} Strong agreement and qualitative feedback on these items reflected the need for structured guidance about how to describe

rehabilitation dosage parameters. Items within the *Personalisation* section underscored the centrality of person-centred care in rehabilitation.³ Furthermore, the adapted *Protocol deviations* section and the new items within the *Harms* section respond to calls for greater clarity on negative undesired effects associated with rehabilitation interventions, which is essential to evaluating their generalisability.^{20 41} The TIDieR-Rehab sets a new standard for the reporting of rehabilitation interventions, paving the way for advancements in the field.

The findings from the modified Delphi process with multidisciplinary rehabilitation experts highlight the importance of striving for a shared conceptual understanding and common language to articulate the essential elements and dosage parameters of rehabilitation interventions. Although consensus was rapidly achieved for most items based on quantitative data, the qualitative data revealed ongoing debate surrounding central rehabilitation concepts such as work duration, intensity and engagement. Such debates emerged not only between disciplines but also within them. For instance, in round 1, *How challenging* was originally framed with respect to setting and measuring the ‘intensity’ of rehabilitation. However, participants raised concerns over the terminology and its relevance. Overall, qualitative findings highlighted the ways conceptual understandings were shaped by both disciplinary perspectives and underpinning bodies

of knowledge.^{3 5 42} Surfacing these tensions was integral to revisions of the TIDieR-Rehab checklist, particularly the terminology and definitions used, which prompted the development of the TIDieR-Rehab supplementary manual. The TIDieR-Rehab supplementary manual offers nuanced explanations and multidisciplinary examples of item reporting to support a shared understanding of key concepts. Nevertheless, while there was a drive to achieve encompassing definitions of key concepts, participant recommendations informed an option for users of the checklist to determine items as ‘non-applicable’ to avoid unnecessary reporting of irrelevant items. Given the recognised variation in conceptual understandings within and across rehabilitation disciplines, the findings of this study reinforce the need for improvements in rehabilitation reporting to support the logical, systematic and coherent exploration and explanation of rehabilitation interventions and their effects.^{15 22}

The importance of clearly defining person-centred practices as an integral part of rehabilitation is at the core of calls to improve reporting of rehabilitation interventions.²² Recognising the gap between principles of person-centred care and their implementation in practice,^{43 44} the TIDieR-Rehab checklist offers a pragmatic approach, with a dedicated *Personalisation* section, to distinguish person-centred rehabilitation practices in response to patient needs and preferences. Person-centredness was further emphasised in the *How challenging* and *Regression/Progression* sections which delineated these important dosage parameters and are often person-specific. However, the study findings recognised the inherent difficulties associated with describing and reporting personalised interventions, which influenced the items adapted and retained through the modified Delphi process. Furthermore, differences in terminology and conceptual understandings coupled with pragmatic constraints informed the removal and adaptation of proposed items. For example, in round 1, a new item related to adherence and engagement was proposed in the *How well* section; however, despite achieving high quantitative agreement (94%), participants commented on the differences in evaluating the two concepts. This was reinforced by practical concerns raised by the Steering Committee, including an expert in engagement, who questioned the items feasibility. Thus, this item was omitted. Likewise, in round 1, participants questioned the proposed new item *How much—Work duration* for its unclear distinction from *How much—Session duration*. Participants also questioned the practicality of quantifying work and rest durations, and whether this was reporting on planned or actual parameters. Such concerns led to the adaptation of this item’s terminology, definition and supplementary information to describe ‘*Essential elements amount*’, and the rewording of all items within the section *How much* to aid clarity. Describing and reporting engagement and work duration may become more feasible as conceptual understandings advance. In addition, technological advances that support the monitoring of engagement in essential elements,

such as activity or physiological monitoring, may facilitate more accurate reporting.^{42 45}

Documenting the negative undesired effects of rehabilitation is paramount to the advancement of the field. The original TIDieR guidelines did not provide explicit directives for reporting adverse consequences, likely due to coverage in other frameworks such as the CONSORT.^{17 20} Current guidelines for safety monitoring and reporting of adverse consequences are primarily focused on medical intervention-related clinical trials.⁴¹ A recent update to the CONSORT framework has proposed that all harms be clearly reported,⁴⁶ yet there remains a gap in robust reporting of adverse consequences in non-pharmacological interventions such as rehabilitation.⁴¹ This gap is addressed within the adapted item *Protocol deviations* and new item *Harms* of the TIDieR-Rehab checklist to support greater clarity of the negative undesired effects associated with rehabilitation interventions. Only by clearly defining and understanding negative undesired effects, with respect to the intervention and its proposed theoretical basis, can a fuller understanding of the potential harms be developed. Successful translation of interventions to clinical practice is dependent on such comprehensive understandings of both benefits and risks.⁴¹

Advancements in clinical trial design and reporting, coupled with a shift to online publishing, have facilitated more rigorous reporting of rehabilitation trial protocols. However, this progress has not been reflected in the reporting of interventions themselves.⁹ The TIDieR-Rehab checklist emphasises a systematic approach to the description of rehabilitation interventions. Researchers may use the TIDieR-Rehab checklist to: support the development of rehabilitation interventions, guide intervention reporting in clinical trials and facilitate the synthesis of rehabilitation research evidence. Importantly, by addressing the pitfalls of inadequate reporting, the TIDieR-Rehab checklist supports the identification of knowledge gaps in rehabilitation research, thereby minimising research wastage. In turn, the TIDieR-Rehab checklist may enhance the translation of research findings into practice by guiding the reporting of interventions in clinical practice guidelines, supporting clinicians to evaluate the potential benefits and risks of an intervention, and offering clarity for accurate and effective implementation of interventions in practice. Future research should explore the validity, reliability and usability of the TIDieR-Rehab checklist²⁵ across a range of different users to gain a better understanding of its applicability. Given the nascent stage of rehabilitation evidence, it is likely that the checklist will require revision as new knowledge emerges.

Limitations

While the TIDieR-Rehab checklist was developed through a rigorous modified Delphi process, there are several limitations which should be acknowledged. First, as opinion-based research, the modified Delphi

approach may introduce participant bias, affecting study outcomes.⁴⁷ Additionally, although online surveys enable efficient collection of diverse perspectives, this approach may introduce desirability bias and discourage in-depth feedback.^{47,48} We attempted to mitigate this issue through the inclusion of free comment boxes and open questions.¹⁸ The study concluded after two rounds of feedback based on predefined consensus criteria which considered both quantitative and qualitative data.⁴⁸ Although typical in modified Delphi studies and commonly accepted in health research,^{26,49} the absence of formal statistical analysis to assess the stability of the consensus may affect the perceived robustness of the study findings.⁴⁸ Furthermore, although consensus criteria were prespecified and documented as part of the ethical review process, the modified Delphi study protocol was not preregistered on a public platform. This would have further enhanced the transparency and trustworthiness of the research findings.⁵⁰

Although this research sought the inclusion of a broad spectrum of participants of varying nationalities, rehabilitation disciplines and professional experiences, there were limitations. While researchers may possess deep insights into rehabilitation intervention reporting, their perspectives may not wholly represent the rehabilitation community, especially clinicians. Additionally, there was a significant representation from the physiotherapy discipline. Discourses on dosage have been prevalent in physiotherapy literature, while other fields like psychology, occupational therapy and speech-language therapy have only recently engaged in such discussions, as evidenced by the under-reporting of dosage parameters in these fields.^{10–12} This limitation may have been exacerbated by reliance on literature from the physical rehabilitation field during the development^{31–34} and piloting^{39,40} of the TIDieR-Rehab checklist. The Steering Committee played a crucial role in capturing a diversity of professional and disciplinary perspectives, yet these may not be fully represented in the research. Furthermore, the recruitment strategy, which focused on authors of rehabilitation research published in the English language, likely resulted in a sample reflecting existing disparities in international research outputs, the origins of the TIDieR checklist¹⁴ and key publications in the field.^{2,4,5,9,20,51} Most participants came from high-income Organisation for Economic Co-operation and Development (OECD) countries, with a significant proportion from Australasia. This skew may limit the global applicability of our findings and potentially bias them towards Western worldviews of rehabilitation. Though the TIDieR-Rehab may risk perpetuating such dominance, the introduction of a tool that encourages more precise descriptions of rehabilitation interventions may foster a more considered examination of the underlying assumptions which shape rehabilitation practice globally.

CONCLUSION

This modified Delphi study supported the development of an extension to the original TIDieR which aimed to enhance the reporting of rehabilitation interventions. The TIDieR-Rehab checklist and supplementary manual²⁵ are grounded in the collective expertise of rehabilitation experts, and reflects critical intervention nuances specific to rehabilitation, including dosage (amount and challenge) and person-centred care. The TIDieR checklist sets a new standard for the reporting of rehabilitation interventions, emphasising systematic documentation and explication of dosage parameters and person-centred practices to promote replicability and synthesis. Recognising the vital need for better reporting of harms, the TIDieR-Rehab also emphasises the importance of documenting both the positive and negative effects of interventions. The findings exposed ongoing debates around core rehabilitation concepts and emphasised the need for a shared understanding of the multifaceted nature of rehabilitation. Furthermore, current challenges in rehabilitation intervention reporting pertaining to the personalised nature of rehabilitation and feasibility of reporting complex nonpharmacological interventions were apparent. The TIDieR-Rehab checklist offers a promising avenue for enhancing the reporting of rehabilitation interventions to promote the minimisation of research wastage, advancement of the field of rehabilitation and translation of research findings to clinical practice.

Acknowledgements We would like to thank the research participants for sharing their time and expertise to contribute to this research. Serene Boon and Sharon Lai contributed to the development of the initial tool and Delphi methodology. We also gratefully acknowledge Professor Duncan Reid, Dr Sue Lord, Associate Professor Kelly Jones, Associate Professor Felicity Bright, Dr Sarah Candy, Dr Ahmad Zamir Bin Che Daud and Elisabeth Kumar who contributed to and guided this research, as part of the Steering Committee. This research would not have been possible without your efforts.

Contributors NS, GA, SO and DT conceptualised the study. NS, GA and SO supervised the study. NS, GA, SO and EG contributed to the study design. GA, EG and CM undertook the data collection. All authors contributed to the analysis and interpretation of findings. NS, EG and CM prepared the first draft of the manuscript, which was further refined by all authors. NS and GA sourced the funding for this project. The corresponding author attests that all listed authors meet authorship criteria and that no others meeting the criteria have been omitted. The guarantor of the study is NS; NS accepts full responsibility for the finished work and/or the conduct of the study, had access to the data and controlled the decision to publish.

Funding This study was supported by a Health Research Council New Zealand Emerging Researcher First Grant (Signal-19/624) and an Auckland University of Technology Summer Research Award 2020/2021.

Competing interests None declared.

Patient and public involvement Patients and/or the public were not involved in the design, or conduct, or reporting or dissemination plans of this research.

Patient consent for publication Not applicable.

Ethics approval This study involves human participants and was approved by the Auckland University of Technology Ethics Committee (reference: 20/264). Participants gave informed consent to participate in the study before taking part.

Provenance and peer review Not commissioned; externally peer reviewed.

Data availability statement No data are available. No further data are available. All data which informed the analysis are presented in the paper and its supplementary files. Full data sets are not publicly available as they may contain

information that could compromise ethical obligation and the privacy of the participants.

Supplemental material This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

Open access This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: <http://creativecommons.org/licenses/by-nc/4.0/>.

ORCID iDs

Nada Signal <http://orcid.org/0000-0001-9595-0532>

Sharon Olsen <http://orcid.org/0000-0002-7453-9127>

Emeline Gomes <http://orcid.org/0000-0002-6871-3790>

Denise Taylor <http://orcid.org/0000-0002-0955-5702>

Gemma Alder <http://orcid.org/0000-0002-8833-0375>

REFERENCES

- Wade DT. What is rehabilitation? An empirical investigation leading to an evidence-based description. *Clin Rehabil* 2020;34:571–83.
- Hart T, Ehde DM. Defining the treatment targets and active ingredients of rehabilitation: Implications for rehabilitation psychology. *Rehab Psychol* 2015;60:126–35.
- Jesus TS, Papadimitriou C, Bright FA, et al. Person-centered rehabilitation model: framing the concept and practice of person-centered adult physical rehabilitation based on a scoping review and thematic analysis of the literature. *Arch Phys Med Rehabil* 2022;103:106–20.
- Van Stan JH, Dijkers MP, Whyte J, et al. The rehabilitation treatment specification system: implications for improvements in research design, reporting, replication, and synthesis. *Arch Phys Med Rehabil* 2019;100:146–55.
- Hayward KS, Churilov L, Dalton EJ, et al. Advancing stroke recovery through improved articulation of nonpharmacological intervention dose. *Stroke* 2021;52:761–9.
- Price J, Rushton A, Tyros I, et al. Effectiveness and optimal dosage of exercise training for chronic non-specific neck pain: A systematic review with a narrative synthesis. *PLoS ONE* 2020;15:e0234511.
- Choy J, Pourkazemi F, Anderson C, et al. Dosages of swallowing exercises in stroke rehabilitation: a systematic review. *Eur Arch Otorhinolaryngol* 2023;280:1017–45.
- Gomes E, Alder G, Bright FAS, et al. Understanding task “challenge” in stroke rehabilitation: an interdisciplinary concept analysis. *Disabil Rehabil* 2024;1–11.
- Dijkers MP. Overview of reviews using the template for intervention description and replication (TIDieR) as a measure of trial intervention reporting quality. *Arch Phys Med Rehabil* 2021;102:1623–32.
- Gee BM, Lloyd K, Devine N, et al. Dosage parameters in pediatric outcome studies reported in 9 peer-reviewed occupational therapy journals from 2008 to 2014: a content analysis. *Rehabil Res Pract* 2016;2016:3580789.
- Zeng B, Law J, Lindsay G. Characterizing optimal intervention intensity: the relationship between dosage and effect size in interventions for children with developmental speech and language difficulties. *Int J Speech Lang Pathol* 2012;14:471–7.
- Hansen NB, Lambert MJ, Forman EM. The psychotherapy dose-response effect and its implications for treatment delivery services. *Clin Psychol Sci Pract* 2002;9:329–43.
- Boutron I, Altman DG, Moher D, et al. CONSORT statement for randomized trials of nonpharmacologic treatments: a 2017 update and a CONSORT extension for nonpharmacologic trial abstracts. *Ann Intern Med* 2017;167:40–7.
- Hoffmann TC, Glasziou PP, Boutron I, et al. Better reporting of interventions: template for intervention description and replication (TIDieR) checklist and guide. *BMJ* 2014;348:bmj.g1687.
- Moher D, Schulz KF, Simera I, et al. Guidance for developers of health research reporting guidelines. *PLoS Med* 2010;7:e1000217.
- Chan A-W, Tetzlaff JM, Altman DG, et al. SPIRIT 2013 statement: defining standard protocol items for clinical trials. *Ann Intern Med* 2013;158:200–7.
- Schulz KF, Altman DG, Moher D. CONSORT 2010 statement: updated guidelines for reporting parallel group randomised trials. *J Pharmacol Pharmacother* 2010;1:100–7.
- Campbell M, Katikireddi SV, Hoffmann T, et al. TIDieR-PHP: a reporting guideline for population health and policy interventions. *BMJ* 2018;361:k1079.
- Rhon DI, Fritz JM, Kerns RD, et al. TIDieR-telehealth: precision in reporting of telehealth interventions used in clinical trials - unique considerations for the Template for the Intervention Description and Replication (TIDieR) checklist. *BMC Med Res Methodol* 2022;22:161.
- Slade SC, Dionne CE, Underwood M, et al. Consensus on Exercise Reporting Template (CERT): explanation and elaboration statement. *Br J Sports Med* 2016;50:1428–37.
- Hacke C, Schreiber J, Weisser B. Application of the templates TIDieR and CERT reveal incomplete reporting and poor replicability of exercise interventions for type 2 diabetes mellitus. *Curr Diabetes Rev* 2022;18:e250821195838.
- Whyte J, Dijkers MP, Fasoli SE, et al. Recommendations for reporting on rehabilitation interventions. *Am J Phys Med Rehabil* 2021;100:5–16.
- Simera I, Moher D, Hoey J, et al. The EQUATOR Network and reporting guidelines: helping to achieve high standards in reporting health research studies. *Maturitas* 2009;63:4–6.
- Whiting P, Wolff R, Mallett S, et al. A proposed framework for developing quality assessment tools. *Syst Rev* 2017;6:204.
- Signal N, Gomes E, Olsen S, et al. Enhancing the reporting quality of rehabilitation interventions through an extension of the Template for Intervention Description and Replication (TIDieR): the TIDieR-Rehab checklist and supplementary manual. *BMJ Open* 2024;0:e084320.
- Nasa P, Jain R, Juneja D. Delphi methodology in healthcare research: how to decide its appropriateness. *World J Methodol* 2021;11:116–29.
- Page SJ, Schmid A, Harris JE. Optimizing terminology for stroke motor rehabilitation: recommendations from the American Congress of Rehabilitation Medicine Stroke Movement Interventions Subcommittee. *Arch Phys Med Rehabil* 2012;93:1395–9.
- Dijkers MP. Reporting on interventions: issues and guidelines for rehabilitation researchers. *Arch Phys Med Rehabil* 2015;96:1170–80.
- Wade DT. Describing rehabilitation interventions. *Clin Rehabil* 2005;19:811–8.
- Kleim JA, Jones TA. Principles of experience-dependent neural plasticity: implications for rehabilitation after brain damage. *J Speech Lang Hear Res* 2008;51:S225–39.
- Mehrholz J, Thomas S, Elsner B. Treadmill training and body weight support for walking after stroke. *Cochrane Database Syst Rev* 2017;8:CD002840.
- English C, Hillier SL, Lynch EA. Circuit class therapy for improving mobility after stroke. *Cochrane Database Syst Rev* 2017;2017.
- Shen X, Wong-Yu ISK, Mak MKY. Effects of exercise on falls, balance, and gait ability in Parkinson's disease: a meta-analysis. *Neurorehab Neural Repair* 2016;30:512–27.
- Hoare BJ, Wallen MA, Thorley MN, et al. Constraint-induced movement therapy in children with unilateral cerebral palsy. *Cochrane Database Syst Rev* 2019;4:CD004149.
- Trevelyan EG, Robinson PN. Delphi methodology in health research: how to do it? *Eur J Integr Med* 2015;7:423–8.
- Hsieh H-F, Shannon SE. Three approaches to qualitative content analysis. *Qual Health Res* 2005;15:1277–88.
- Cope DG. Methods and meanings: credibility and trustworthiness of qualitative research. *Oncol Nurs Forum* 2014;41:89–91.
- Diamond IR, Grant RC, Feldman BM, et al. Defining consensus: a systematic review recommends methodologic criteria for reporting of Delphi studies. *J Clin Epidemiol* 2014;67:401–9.
- Olsen S, Alder G, Rashid U, et al. Challenge level contributes to the efficacy of treadmill interventions after stroke: a systematic review and meta-analysis. *Brain Sci* 2023;13:1729.
- Charlesworth KRU, Rodriguez-Ramirez E, Browne W, et al. Device and programme requirements for effective upper-limb robotic rehabilitation following stroke: a systematic review and meta-analysis [Abstract]. In Abstracts presented at Stroke 2022 – The Annual Scientific Meeting of the Stroke Society of Australasia; Christchurch, New Zealand. *Int J Stroke* 2022.
- Papaioannou D, Cooper C, Mooney C, et al. Adverse event recording failed to reflect potential harms: a review of trial protocols of behavioral, lifestyle and psychological therapy interventions. *J Clin Epidemiol* 2021;136:64–76.

- 42 Bright FAS, Kayes NM, Worrall L, *et al.* A conceptual review of engagement in healthcare and rehabilitation. *Disabil Rehabil* 2015;37:643–54.
- 43 Fernandes JB, Vareta D, Fernandes S, *et al.* Rehabilitation workforce challenges to implement person-centered care. *Int J Environ Res Public Health* 2022;19:3199.
- 44 Yun D, Choi J. Person-centered rehabilitation care and outcomes: a systematic literature review. *Int J Nurs Stud* 2019;93:74–83.
- 45 Godecke E, Brogan E, Ciccone N, *et al.* Treatment fidelity monitoring, reporting and findings in a complex aphasia intervention trial: a substudy of the Very Early Rehabilitation in SpEEch (VERSE) trial. *Trials* 2022;23:501.
- 46 Junqueira DR, Zorzela L, Golder S, *et al.* CONSORT Harms 2022 statement, explanation, and elaboration: updated guideline for the reporting of harms in randomised trials. *BMJ* 2023;381:e073725.
- 47 Bhandari S, Hallowell MR. Identifying and controlling biases in expert-opinion research: guidelines for variations of Delphi, nominal group technique, and focus groups. *J Manage Eng* 2021;37:1–13.
- 48 Barrios M, Guilera G, Nuño L, *et al.* Consensus in the delphi method: what makes a decision change? *Technol Forecast Soc Change* 2021;163:120484.
- 49 Niederberger M, Spranger J. Delphi technique in health sciences: a map. *Front Public Health* 2020;8:457.
- 50 Grant S, Booth M, Khodyakov D. Lack of preregistered analysis plans allows unacceptable data mining for and selective reporting of consensus in Delphi studies. *J Clin Epidemiol* 2018;99:96–105.
- 51 Lohse KR, Pathania A, Wegman R, *et al.* On the reporting of experimental and control therapies in stroke rehabilitation trials: a systematic review. *Arch Phys Med Rehabil* 2018;99:1424–32.

Supplementary File 1. A synthesised appraisal of key issues in rehabilitation intervention reporting

Table 1. Key issues in rehabilitation intervention reporting.

Identified issue	Description
Lack of standardisation	There is a lack of standardisation in the language used to describe rehabilitation interventions, making it difficult to compare and replicate studies due to inconsistent definitions and interpretations of key terms.
Incomplete descriptions	Many studies do not provide a complete description of the interventions, hindering reliable implementation in clinical practice.
Inadequate theoretical rationale	Reports frequently omit detailed information on the intervention's theoretical basis and rationale.
Infrastructure reporting	Reports often omit details about the infrastructure required, including the setting and context in which the interventions are conducted.
Details on intervention providers	Information is often lacking on the intervention providers, including their qualifications, skills, and training.
Clarity of intervention ‘dosage’	There is frequently a lack of clear reporting of the 'dose' of an intervention, including its total and active duration, frequency and intensity or challenge, which is critical for evaluating efficacy.
Reporting of tailoring	Modifications or personalisation of interventions for groups or individuals are often inadequately reported, which is vital for applying findings to varied populations.
Intervention fidelity	There is often limited information available on how well the intervention was delivered according to the protocol.
Participant response	The reporting on participant response to the intervention, including engagement and adverse consequences, is frequently inadequate.

Supplementary File 2. Copies of surveys sent to modified Delphi participants.

Snapshots of the Round 1 modified Delphi survey.



STUDY SUMMARY

Project Title:

Extension of the Template for Intervention Description and Replication (TIDier) to develop a checklist to enhance the quality of reporting of rehabilitation intervention parameters: TIDier-Rehab.

Project Leaders:

Gemma Alder, Dr Sharon Olsen and Associate Professor Nada Signal
Health & Rehabilitation Research Institute, Auckland University of Technology, Auckland,
New Zealand

Thank you for agreeing to participate in our study.

This online modified-Delphi survey aims to extend the TIDier checklist to enhance the quality of reporting of rehabilitation intervention parameters, by developing the TIDier-Rehab checklist. This extension of the original TIDier checklist will enhance reporting by further addressing all parameters of intervention dosage. The original [TIDier](#) is linked below for your reference.

Link(s):

- [TIDier checklist \(original\)](#)

Approved by the Auckland University of Technology Ethics Committee on 10/12/2020,

AUTEC Reference number 20/264.

*** I wish to receive a summary of the research findings (please tick one):**

☐ Yes

☐ No

0% Survey Completion 100%





*** Please select the job description that best indicates your main role.**

☐ Researcher

☐ Clinician

☐ Journal Editor

☐ Reviewer in the Rehabilitation Field

☐ Policymaker

☐ Funder

☐ Other (please state):





INSTRUCTIONS

Please indicate the response that best characterises how you feel about the statement:

"This item should be included in the checklist."

- For each of the items, you will be offered the following responses:

Strongly disagree Disagree Neutral Agree Strongly agree

- Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc. (optional).

	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p><i>"This item should be included in the tool."</i></p> <p><i>Strongly disagree Disagree Neutral Agree Strongly agree</i></p>	<p>Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>SECTION 1 – BRIEF NAME</p> <p>1. Provide the name or phrase that describes the intervention.</p>	<p><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></p>	<div></div>

←

Survey Completion
0% ————— 100%

→



	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the tool."</p> <p>Strongly disagree Disagree Neutral Agree Strongly agree</p>	<p>Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc.</p> <p>Comments (optional):</p>
<p><u>SECTION 2 - WHY</u></p> <p>2. Describe any rationale, theory, or goal of the elements essential to the intervention.</p>	<p><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></p>	<div></div>



	<div>Please indicate the response that best characterises how you feel about the statement: "This item should be included in the tool."</div> <div>Strongly disagree Disagree Neutral Agree Strongly agree</div>	<div>Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc:</div> <div>Comments (optional):</div>
<div>SECTION 3 - WHAT</div> <div>3a. Materials: Describe any physical or informational materials used in the intervention, including those provided to participants or used in intervention delivery or in training of intervention providers. Provide information on where the materials can be accessed. <i>For example, supplementary materials, URL.</i></div> <div>3b. Procedures: Describe each of the procedures, activities, and/or processes used in the intervention, including any enabling or support activities.</div>	<div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div> <div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div>	<div></div> <div></div>



	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the tool."</p> <p>Strongly disagree Disagree Neutral Agree Strongly agree</p>	<p>Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>SECTION 4 – WHO PROVIDED</p> <p>4. For each category of intervention provider (e.g psychologist, nursing assistant) describe their expertise, background and any specific training given.</p>	<div><div></div><div></div><div></div><div></div><div></div></div>	<div></div>



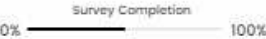
	<div>Please indicate the response that best characterises how you feel about the statement: "This item should be included in the tool."</div> <div>Strongly disagree Disagree Neutral Agree Strongly agree</div>	<div>Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc.</div> <div>Comments (optional):</div>
<div>SECTION 5 - HOW</div> <div>5a. Describe the modes of delivery of the intervention and whether it was provided individually or in a group. <i>For example, in-person or by some other mechanism, such as internet or telephone.</i></div>	<div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div>	<div></div>
<div>5b. Describe the provision of support in delivering the intervention. <i>For example, manual facilitation of stepping with physical assistance of two therapists, auditory cuing using a metronome at 2-5Hz, guidance statements for the creation of coping strategies, modelling of correct grammar during language activities.</i></div>	<div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div>	<div></div>



	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the tool."</p> <p>Strongly disagree Disagree Neutral Agree Strongly agree</p>	<p>Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>SECTION 6 - WHERE</p> <p>6. Describe the type(s) of location(s) where the intervention occurred, including any necessary infrastructure or relevant features.</p>	<p><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></p>	<div></div>



	Please indicate the response that best characterises how you feel about the statement: "This item should be included in the tool."					Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc.
	Strongly disagree	Disagree	Neutral	Agree	Strongly agree	Comments (optional):
SECTION 7 - WHEN AND HOW MUCH						
7a. Record the total session duration. Total session duration is defined as the prescribed overall length of time usually measured in minutes or hours in which a single session of therapy is administered. If minutes are not recorded, this could be recorded as number of repetitions (Page et al, 2012; Lang et al, 2015).	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	
7b. Record the work duration. Work duration is defined as the length of time, usually reported in minutes, spent participating in the task/intervention. Work duration = total session duration - rest time (Lang et al, 2015).	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	
7c. Record the frequency. Frequency is defined as the number of sessions per day or week, during the total period of time the intervention is delivered (Page et al, 2012; Lang et al, 2015). For example two sessions per day and/or two sessions per week.	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	
7d. Record the intervention length. Intervention length is defined as the total period of time, usually reported in weeks or months, an intervention is delivered for (Page et al, 2012).	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	<input type="radio"/>	





	<div>Please indicate the response that best characterises how you feel about the statement: "This item should be included in the tool."</div> <div>Strongly disagree Disagree Neutral Agree Strongly agree</div>	<div>Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc.</div> <div>Comments (optional):</div>
<div>SECTION 8 – HOW CHALLENGING</div> <div>8a. Describe the method used to measure and set the intervention intensity. <i>Intensity can be measured using a range of subjective and objective measures. Subjective intervention intensity could include patient-reported estimates of difficulty such as setting intensity at 13-15 on the Borg Rating of Perceived Exertion Scale. Objective intervention intensity could include %VO2max or level of language complexity such as increasing the number of steps within an instruction an individual responds too.</i></div>	<div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div>	<div></div>
<div>8b. Report a method for setting the intensity using a reliable and validated tool.</div>	<div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div>	<div></div>
<div>8c. Describe the rules for setting the starting intensity.</div>	<div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div>	<div></div>





	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the tool."</p> <p>Strongly disagree Disagree Neutral Agree Strongly agree</p>	<p>Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>SECTION 10 - MODIFICATIONS</p> <p>10. If the intervention was modified during the course of the study, describe the changes (what, why, when and how).</p>	<p><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></p>	<div></div>



	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the tool."</p> <p>Strongly disagree Disagree Neutral Agree Strongly agree</p>	<p>Please provide any comments to explain your response in the comments box of each question e.g suitability of items, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>SECTION 11 – HOW WELL</p> <p>11a. Planned: If the intervention fidelity was assessed, describe how and by whom, and if any strategies were used to maintain or improve fidelity, describe them.</p> <p><i>Fidelity refers to the extent to which the intervention was delivered as intended. For example, standardised intervention reporting forms or training of intervention providers.</i></p>	<div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div>	<div></div>
<p>11b. Actual: If intervention adherence or fidelity was assessed, describe the extent to which the intervention was delivered as planned.</p>	<div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div>	<div></div>
<p>11c. Describe how engagement/adherence was measured.</p> <p><i>For example, use of a diary, video recording of sessions, or use of a pedometer.</i></p>	<div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div>	<div></div>
<p>11d. Describe the monitoring and reporting of adverse events.</p>	<div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div>	<div></div>
<p>11e. Describe the number and seriousness of adverse events and their relatedness to the intervention.</p>	<div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div>	<div></div>



Survey Completion

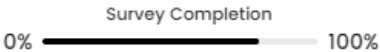
0% 100%





Are there any aspects of intervention reporting you believe were not adequately addressed within this draft checklist? (optional):

Please provide any other feedback or general comments below (optional):





Thank you for completing this survey please click SUBMIT to ensure your responses are received.

If you have any queries, please contact the project leaders:

Gemma Alder, gemma.alder@aut.ac.nz

Dr Sharon Olsen, sharon.olsen@aut.ac.nz

Do you wish to receive a copy of your responses to your email?

☐ Yes

☐ No



0% **Survey Completion** 100%

SUBMIT

Snapshots of the Round 2 modified Delphi survey.



STUDY SUMMARY

Project Title:

The TiDieR-Rehab checklist: An extension of the Template for Intervention Description and Replication (TiDieR) to enhance the reporting quality of rehabilitation intervention parameters.

Project Leaders:

Gemma Alder, Emma Gomes, Dr Sharon Olsen and Associate Professor Nada Signal
Health & Rehabilitation Research Institute, Auckland University of Technology,
Auckland, New Zealand

This survey encompasses Round Two of the modified-Delphi process which aims to refine the TiDieR-Rehab checklist. Please refer to the key documents embedded within the links below to assist with the completion of the survey.

Links:

- [TiDieR-Rehab development overview](#)
- [TiDieR-Rehab supplementary manual](#)
- [TiDieR checklist \(original\)](#)

Approved by the Auckland University of Technology Ethics Committee on 10/12/2020.

AUTEC Reference number 20/264.

0% — Survey Completion — 100%





ROUND ONE RESULTS OVERVIEW

Thank you for participating in Round One of the TiDieR-Rehab modified-Delphi survey. Your feedback is appreciated and has been rigorously analysed, triangulated and reviewed by an interdisciplinary Steering Committee to refine the TiDieR-Rehab checklist.

Round One received feedback from 35 participants from various rehabilitation backgrounds, job descriptions and geographical locations.

Quantitative results. Twenty of the 24 items presented in Round One met high quantitative agreement (>80% 'strongly agree' and/or 'agree' Likert responses for the item). Items that only met moderate agreement (50%-70% 'strongly agree' and/or 'agree') in Round One pertained to the provision of support (item 5B), work duration (item 7B), methods of measurement and setting intensity (item 8A) and the reliability and validity of the method used to set intensity (item 8B).

Qualitative results. Inductive content analysis revealed feedback generally supported the TiDieR-Rehab checklist and that items would aid intervention reporting, replication for research and clinical practice. Some critiques were made to revise the terminology and structure of items or sections, as well as suggestions to introduce new items and provide supporting information to aid clarity. Comments also highlighted potential barriers to meaningful reporting through the TiDieR-Rehab checklist, such as the individualisation of interventions or minimal relevance of some items to specific disciplines.

This survey presents Round Two of the modified-Delphi process to gain further feedback on a revised version of the TiDieR-Rehab checklist and its accompanying supplementary manual which have been developed in direct response to your feedback from Round One.



ROUND TWO SURVEY INSTRUCTIONS

All items of the revised TiDieR-Rehab checklist are chronologically presented in this Round Two survey. Items are tentatively categorised as 'accepted' or 'adapted/new' which informs the depth of feedback options.

Accepted items: These items received substantial quantitative and qualitative agreement and are mostly unchanged from Round One. 'Accepted' items will likely be included in the final TiDieR-Rehab checklist unless strong omission is received, hence, only an optional free-comments box is provided. *A summary of quantitative agreement is provided.*

Adapted/New items: These items received decent or substantial quantitative agreement but have been adapted or introduced in response to qualitative feedback. 'Adapted/New' items are seeking further feedback, hence, both a Likert scale (strongly disagree, disagree, agree and strongly agree) and an optional free-comments box have been provided. Key adaptations from Round One are highlighted by **blue text**. An overview of the iterative development of the TiDieR-Rehab checklist items through rounds is embedded within the link below for your reference. *Where applicable, a summary of the quantitative agreement and key qualitative critiques of the item (in the form it was presented in Round One) is provided.*

The TiDieR-Rehab supplementary manual: A [supplementary manual](#) has also been created for the refined Round Two items in response to feedback for item instructions, clarification and examples. ***Please refer to the corresponding item on the supplementary manual as you complete your response to each item in this survey.*** As the supplementary manual will be a supporting document to the final TiDieR-Rehab checklist, we would also appreciate any feedback on its contents within the optional free-comments box for each corresponding item, and at the end of the survey. An embedded link to the [supplementary manual](#) is provided below.

Thank you once again for participating in this research; your feedback has been invaluable to this process.

Links:

- [TiDieR-Rehab development overview](#)
- [TiDieR-Rehab supplementary manual](#)





SECTION 1: BRIEF NAME

ACCEPTED ITEM: 91% quantitative agreement (high agreement).

EXPLANATION

EXAMPLES

	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>Item 1. Provide the name or a phrase that describes the intervention.</p>	<div></div>



SECTION 2: WHY

ACCEPTED ITEM: 86% quantitative agreement (high agreement).

EXPLANATION

EXAMPLES

	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>Item 2. Describe any rationale, theory, or goal of the elements essential to the intervention.</p> <p><i>Essential elements, also known as "active ingredients", refer to the core intervention components that are hypothesised to be linked to effects or outcomes.</i></p>	<div></div>



SECTION 3: WHO PARTICIPATED

NEW ITEM:

Quantitative: nil data.

Qualitative: Who ‘received’ the intervention is missing but underpins the entire basis and needs to be explicitly reported; it should be clear if the target population is different to study population and why (*Intordisciplinary Steering Committee feedback*).

EXPLANATION

EXAMPLES

	<div>Please indicate the response that best characterises how you feel about the statement: "This item should be included in the TIDieR-Rehab checklist."</div> <div>Strongly disagreeDisagreeAgreeStrongly agree</div>	<div>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</div> <div>Comments (optional):</div>
Item 3. Provide any relevant demographic information or criteria about the study population(s).	<div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div>	<div></div>



SECTION 4: WHAT

ACCEPTED ITEM 4A – MATERIALS: 91% quantitative agreement (high agreement).

EXPLANATION

EXAMPLES

Please provide an explanation of your response in the comments box of each item
e.g. suitability, wording, grouping etc.

Comments (optional):

Item 4A. Materials: Describe any physical or informational materials used in the intervention, including those provided to participants or used in intervention delivery or in training of intervention providers. Provide information on where the materials can be accessed (e.g. online appendix, URL).

ACCEPTED ITEM 4B – PROCEDURES: 91% quantitative agreement (high agreement).

EXPLANATION

EXAMPLES

Please provide an explanation of your response in the comments box of each item
e.g. suitability, wording, grouping etc.

Comments (optional):

Item 4B. Procedures: Describe each of the procedures, activities, and/or processes used in the intervention, including any enabling or support activities.



SECTION 5: WHEN

NEW ITEM:

Quantitative: nil data.

Qualitative: "When", timing or commencement of the intervention occurred in relation to key events was not adequately addressed within the TIDieR and TIDieR - Rehab section "When and How much" which focuses on the implementation of 'amount'.

EXPLANATION

EXAMPLES

	<div>Please indicate the response that best characterises how you feel about the statement: "This item should be included in the TIDieR-Rehab checklist."</div> <div>Strongly disagree Disagree Agree Strongly agree</div>	<div>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</div> <div>Comments (optional):</div>
Item 5. Describe when the intervention was commenced in relation to the onset or stage of the conditions and/or other relevant events.	<div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div>	<div></div>



SECTION 6: WHO PROVIDED

ACCEPTED ITEM: 89% quantitative agreement (high agreement).

EXPLANATION EXAMPLES

	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>Item 6. For each category of intervention provider (e.g., psychologist, nursing assistant), describe their expertise, background and any specific training given.</p>	<div></div>

ADAPTED ITEM 7B - STRATEGIES:

Round One item:

"Describe the provision of support in delivering the intervention"

Quantitative: 77% agreement (moderate agreement);

Qualitative: the term 'support' is unclear; this may be covered in item 3B (What: Procedures); high individualisation makes this item unfeasible to report on.

EXPLANATION EXAMPLES

	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the TIDieR-Rehab checklist."</p> <p>Strongly disagree Disagree Agree Strongly agree</p>	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>Item 7B. Strategies: Describe the supplementary strategies that were required to enable the core intervention in response to specific individual or group needs.</p> <p><i>Supplementary strategies refer to the intervention adjuncts (e.g. physical assistance, verbal cueing, transportation arrangements) that must be used for some individuals or groups to facilitate their engagement or effective participation in the core intervention.</i></p>	<div><div></div><div></div><div></div><div></div></div>	<div></div>



SECTION 7: HOW

ACCEPTED ITEM 7A – MODES: 97% quantitative agreement (high agreement).

- EXPLANATION
- EXAMPLES

	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>Item 7A. Modes: Describe the modes of delivery (e.g., face-to-face or by some other mechanism, such as internet or telephone) of the intervention and whether it was provided individually or in a group.</p>	<div></div>



SECTION 8: WHERE

ADAPTED ITEM 8:

Round One item:
"Describe the type(s) of location(s) where the intervention occurred, including any necessary infrastructure or relevant features"

Quantitative: 86% agreement (high agreement).

Qualitative: Other factors beyond the physical location may influence the intervention or participants; specification of an incidental location in research may limit the application of the intervention in other viable locations; this item should be optional as it is not relevant to all disciplines or interventions.

EXPLANATION EXAMPLES

	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the TIDieR-Rehab checklist."</p> <p>Strongly disagree Disagree Agree Strongly agree</p>	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>Item 8. Describe the type(s) of environment(s) where the intervention occurred, including any necessary infrastructure or relevant features.</p> <p><i>Environments and their features may include the relevant physical, social, cultural, economic, political and/or systemic context(s) of the intervention.</i></p>	<div><div></div><div></div><div></div><div></div></div>	<div></div>



SECTION 9: HOW MUCH

ADAPTED ITEM 9A:

Round One item:

"Record the total session duration"

Total session duration is defined as the prescribed overall length of time, usually measured in minutes or hours in which a single session of therapy is administered; if minutes are not recorded, this could be recorded as number of repetitions (Page et al., 2012; Lang et al., 2015).

Quantitative: 91% agreement (high agreement).

Qualitative: 'Planned' items (parameters planned before intervention implementation) and 'actual' items (what actually occurred during the intervention) need to be clearly differentiated; repetitions should be presented with equal importance to duration.

EXPLANATION EXAMPLES

	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the TIDieR-Rehab checklist."</p> <p>Strongly disagree Disagree Agree Strongly agree</p>	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>Item 9A. Session(s) duration: provide the planned session(s) duration of the intervention.</p>	<p><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></p>	<div></div>

ADAPTED ITEM 9B:

Round One item:

"Record the work duration"

Work duration is defined as the length of time, usually reported in minutes, spent participating in the task/intervention. Work duration - total session duration minus rest time (Lang et al., 2015)

Quantitative: 69% agreement (moderate agreement).

Qualitative: 'Work' duration is an exercise-based term that is not relevant to some rehabilitation disciplines; 'planned' and 'actual' items should be differentiated; repetitions should be presented here (rather than Item 9A) as equally important to duration; this information may be highly individualised and impractical to record; this item should be optional.

EXPLANATION

EXAMPLES

	<div>Please indicate the response that best characterises how you feel about the statement:</div> <div>"This item should be included in the TIDieR-Rehab checklist."</div> <div>Strongly disagree Disagree Agree Strongly agree</div>	<div>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</div> <div>Comments (optional):</div>
<div>Item 9B. Essential elements duration: Provide the planned session(s) duration/repetitions for engagement with essential elements of the intervention.</div> <div>Essential element duration refers to the amount of time, within the total duration of a session, that is spent "actively" participating in the core components of the intervention.</div>	<div><div><div></div><div></div><div></div><div></div></div></div>	<div></div>

ADAPTED ITEM 9C:

Round One item:

"Record the frequency"

Frequency is defined as the number of sessions per day and/or week, during the total period of time the intervention is delivered (Page et al., 2012; Lang et al., 2015). For example, two sessions per day, three times per week.

Quantitative: 94% agreement (high agreement).

Qualitative: nil critiques.

EXPLANATION

EXAMPLES

	<div>Please indicate the response that best characterises how you feel about the statement:</div> <div>"This item should be included in the TIDieR-Rehab checklist."</div> <div>Strongly disagreeDisagreeAgreeStrongly agree</div>	<div>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</div> <div>Comments (optional):</div>
Item 9C. Frequency: Provide the planned frequency of the intervention.	<div><div></div><div></div><div></div><div></div></div>	<div></div>

ADAPTED ITEM 9D:

Round One item:

"Record the intervention length"

Intervention length is defined as the total period of time, usually reported in weeks or months, an intervention is delivered for (Page et al., 2012).

Quantitative: 97% agreement (high agreement).

Qualitative: clearly differentiate intervention length from total session duration (Round One version of item 9A).

EXPLANATION

EXAMPLES

	<div>Please indicate the response that best characterises how you feel about the statement:</div> <div>"This item should be included in the TIDieR-Rehab checklist."</div> <div>Strongly disagreeDisagreeAgreeStrongly agree</div>	<div>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</div> <div>Comments (optional):</div>
Item 9D. Intervention length: Provide the planned overall length of the intervention.	<div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div>	<div></div>



SECTION 10: HOW DIFFICULT

ADAPTED ITEM 10:

Round One item:

Item 8A. "Describe the method used to measure and set the intervention intensity"

Intensity can be measured using a range of subjective and objective measures. Subjective intensity could include patient-reported estimates of difficulty such as setting intensity at 13-16 on the Borg Rating of Perceived Exertion Scale. Objective intensity could include the level of language complexity (such as increasing the number of steps within an instruction) or %VO2max.

Quantitative: 77% agreement (moderate agreement).

Qualitative: 'intensity' is not relevant or quantifiable in some disciplines; 'intensity' described as effort or difficulty needs to be differentiated from 'amount' (addressed in Section 9, parameters); 'intensity' cannot be planned; clearer interdisciplinary examples are needed; this item should be optional.

REMOVED FROM ITEM 10:

Round One item:

Item 8B. "Report if the method for setting the intensity uses a reliable and validated tool"

Quantitative: 71% agreement (moderate agreement).

Qualitative: psychometric properties are only relevant if 'intensity' is relevant and a measure is used; this item should be optional.

- Note. Based on Round One feedback, this item has been removed as a checklist item and placed in the supplementary manual as a prompt for the new Round Two item.

EXPLANATION

EXAMPLES

	<div>Please indicate the response that best characterises how you feel about the statement: "This item should be included in the TIDieR-Rehab checklist."</div> <div>Strongly disagree Disagree Agree Strongly agree</div>	<div>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</div> <div>Comments (optional):</div>
<div>Item 10. Describe the approach(es) used to set and monitor the intervention/task difficulty level. <i>Difficulty refers to the absolute or or relative level of challenge, complexity, physiological intensity, grade, load or tolerance of an intervention/task at a given time, and may be evaluated using subjective or objective measures.</i></div>	<div><div><div></div><div></div><div></div><div></div></div></div>	<div></div>

←

Survey Completion

0%100%

→



SECTION 11: REGRESSION/PROGRESSION

ADAPTED (COMBINED) ITEM 11:

Round One item:
Item 9B: "Report the dosage parameter(s) used to progress the intervention"
Dosage parameters include the total session duration, work duration, frequency and intensity

Quantitative: 94% agreement (high agreement).
Qualitative: Round One Item's 9B, 9C and 9D should be combined into one item.

Round One item:
Item 9C: "Describe the rules for deciding when progression is implemented."

Quantitative: 91% agreement (high agreement).
Qualitative: Round One Item's 9B, 9C and 9D should be combined into one item; 'rules' is not the correct term; disambiguate progression from 'intensity'; examples would aid clarity; regression of dosage parameters should also be reported.

Round One item:
Item 9D: "Describe the rules for deciding how the progression is implemented."
For example, intervention duration was increased by two minutes or walking speed was increased by 10%.

Quantitative: 88% agreement (high agreement).

Qualitative: Round One Item's 9B, 9C and 9D should be combined into one item; 'rules' is not the correct term.

EXPLANATION

EXAMPLES

	<div>Please indicate the response that best characterises how you feel about the statement: "This item should be included in the TIDieR-Rehab checklist."</div> <div>Strongly disagree Disagree Agree Strongly agree</div>	<div>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</div> <div>Comments (optional):</div>
Item 11. Describe when and how the planned regression or progression of dosage parameter(s) was implemented.	<div><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></div>	<div></div>





SECTION 12: PERSONALISATION

ADAPTED ITEM 12:

Round One item:
"If the intervention was planned to be personalised, titrated or adapted, then describe what, why, when, and how"

Quantitative: 83% agreement (high agreement).

Qualitative: This item [as 'tailoring'] appears to be redundant of Round One Item's 9B, 9C, 9D regarding progression; description of this item should be a summary, algorithm or protocol; this item should be optional.

EXPLANATION EXAMPLES

	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the TIDieR-Rehab checklist."</p> <p>Strongly disagree Disagree Agree Strongly agree</p>	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
Item 12. Describe the way(s) in which the intervention was planned to be adapted for personal preferences.	<div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div> <div><input type="radio"/></div>	<div></div>



SECTION 13: PROTOCOL DEVIATIONS

ADAPTED ITEM 13:

Round One item:
"If the intervention was modified during the course of the study,
describe the changes (what, why, when, and how)"

Quantitative: 91% agreement (high agreement).
Qualitative: This item needs to be differentiated from items regarding "tailoring" (Round One, item 9); this will be covered by Round One, item 38 (What - Procedures); modifications will change everything else that has occurred if you are reporting what actually happened; this item should be optional.

EXPLANATION EXAMPLES

	<p>Please indicate the response that best characterises how you feel about the statement:</p> <p>"This item should be included in the TIDier-Rehab checklist."</p> <p>Strongly disagree Disagree Agree Strongly agree</p>	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
<p>Item 13. If there were deviations in the intervention protocol during the course of the study, describe the changes (what, why, when, and how).</p>	<p><input type="radio"/> <input type="radio"/> <input type="radio"/> <input type="radio"/></p>	<div></div>



SECTION 14: HOW WELL

ADAPTED ITEM 14A – PLAN:

Round One item:

Item 11A. "Planned: If intervention fidelity was assessed, describe how and by whom, and if any strategies were used to maintain or improve fidelity, describe them"

Fidelity refers to the extent to which the intervention was delivered as intended. For example, training of intervention providers or using standardised intervention reporting forms.

Quantitative: 94% agreement (high agreement).

Qualitative: Clearly differentiate 'plans' and 'actual' throughout the checklist; a rationale for fidelity strategies should be provided; examples of fidelity measures would aid clarity.

REMOVED/ADAPTED INTO ITEM 14A & 14B:

Round One item:

Item 11C. "Describe how engagement/adherence was measured"

Quantitative: 94% agreement (high agreement).

Qualitative: engagement cannot be measured; this is not always relevant or feasible to report on in some disciplines; a predefined criteria for engagement and adherence should be encouraged; this item is repetitive of fidelity questions (Round One Item 13A, 13B); engagement and adherence should not be used interchangeably.

- Note. Based on Round One qualitative feedback and Steering Committee input, this item has been removed as a checklist item and placed in the supplementary manual as a prompt for the new Round Two items within Section 14. How well.

EXPLANATION

EXAMPLES

	<div>Please indicate the response that best characterises how you feel about the statement: "This item should be included in the TIDieR-Rehab checklist."</div> <div>Strongly disagree Disagree Agree Strongly agree</div>	<div>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</div> <div>Comments (optional):</div>
Item 14A. Plan: If intervention adherence or fidelity was assessed, describe how and by whom, and if any strategies were used to maintain or improve fidelity, describe them.	<div><div></div><div></div><div></div><div></div></div>	

ACCEPTED ITEM 14B:

Round One item:

"Actual: If intervention fidelity was assessed, describe the extent to which the intervention was delivered as planned"

Quantitative: 91% agreement (high agreement).

Qualitative: An example may include the number of times a target behaviour was achieved; this may be unknown at the time of intervention reporting.

EXPLANATION

EXAMPLES

	<div>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</div> <div>Comments (optional):</div>
Item 14B. Actual: If intervention adherence or fidelity was assessed, describe the extent to which the intervention was delivered as planned.	



SECTION 15: ADVERSE EVENTS

ACCEPTED ITEM 14A – PLAN:

Round One item:
"Describe the monitoring and reporting of adverse events"

Quantitative: 91% agreement (high agreement).
Qualitative: this not important in some disciplines; this should involve a broad or summarised description.

EXPLANATION EXAMPLES

	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
Item 15A. Plan: Describe the monitoring and reporting of adverse events.	

ACCEPTED ITEM 15B - ACTUAL

Round One item:
"Describe the number and seriousness of adverse events that occur
and their relatedness to the intervention"

Quantitative: 88% agreement (high agreement).

Qualitative: differentiate this item as what actually occurred, compared to the 'plan' for adverse events (Round One, Item 12A) as this is unknown till implemented; the item should be optional; a criteria for seriousness and relatedness should be predefined for the study.

EXPLANATION EXAMPLES

	<p>Please provide an explanation of your response in the comments box of each item e.g. suitability, wording, grouping etc.</p> <p>Comments (optional):</p>
Item 15B. Actual: Describe the number and seriousness of adverse events that occur and their relatedness to the intervention.	





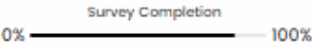
Please provide any feedback regarding the draft supplementary manual below (optional):





Are there any aspects of intervention reporting you believe were not adequately addressed within this revised checklist? (optional):

Please provide any other feedback or general comments below (optional):





This concludes the survey.
Please click **SUBMIT** to confirm your responses.

*** Do you wish to receive a PDF copy of your responses via email?**

If you select "Yes" a PDF copy of your responses will be sent to your email shortly.

☐ Yes

☐ No



0% Survey Completion 100%



Supplementary File 3. Summary of potential participants invited to contribute to the TIDieR-Rehab modified Delphi study

Table 1. Summary of potential participants’ representations of clinical disciplines.

Clinical discipline	Number of potential participants’ representation (n=270)
Psychology	58
Occupational therapy	48
Speech-language therapy	37
Neurological physiotherapy	31
Other physiotherapy (e.g., respiratory)	24
Musculoskeletal physiotherapy	22
Rehabilitation or sports medicine	23
Rehabilitation nursing	5
Exercise science	14
Public health	8

Table 2. Summary of potential participants’ representations of geographical locations.

Geographical location	Number of potential participants’ representation (n=270)
United States	80
Australia	75
United Kingdom	36
Canada	17
Netherlands	17
Denmark	8
China	6
Ireland	6
New Zealand	4
Japan	4
Germany	3
Brazil	3
Norway	3
Sweden	3
Bulgaria	1
Israel	1
France	1
Switzerland	1
Iran	1

Supplementary File 4. Iterative development of the TIDieR-Rehab checklist through each round of the modified Delphi process.

Table. Iterative development of the TIDieR-Rehab checklist through modified Delphi rounds.

ORIGINAL TIDieR CHECKLIST	ROUND 1 CHECKLIST	ROUND 2 CHECKLIST	PILOTING
Item 1. Brief name Provide the name or a phrase that describes the intervention.	Section/Item 1. Brief name Provide the name or a phrase that describes the intervention.	Section/Item 1. Brief name Provide the name or a phrase that describes the intervention.	Section/Item 1. Brief name Provide the name or a phrase that describes the intervention.
	RETAINED ORIGINAL		
Item 2. Why Describe any rationale, theory, or goal of the elements essential to the intervention.	Section/Item 2. Why Describe any rationale, theory, or goal of the elements essential to the intervention.	Section/Item 2. Why Describe any rationale, theory, or goal of the elements essential to the intervention. <i>Essential elements, also known as “active ingredients”, refer to the core components of the intervention that are hypothesised to be linked to effects or outcomes.</i>	Section/Item 2. Why Describe any rationale, theory or goal of the elements essential to the intervention. <i>Essential elements, also known as “active ingredients”, are the core components of the intervention that are expected to be linked to effects or outcomes of interest.</i>
	RETAINED ORIGINAL	MINOR ADAPTATIONS TO ORIGINAL	
		Section/Item 3. Who participated Provide any relevant demographic information or criteria about the study population(s).	Section/Item 3. Who _____ Describe who the intervention is intended for.
		NEW TIDieR-REHAB	MINOR ADAPTATIONS TO TIDieR-REHAB
		Section/Item 4. When Describe when the intervention was commenced in relation to the onset or stage of the condition and/or other relevant events.	Section/Item 4. When Describe when the intervention commenced in relation to the onset or stage of the condition and/or other relevant events.
		NEW TIDieR-REHAB	
Item 3. What (materials) Materials: Describe any physical or informational materials used in the intervention, including those provided to participants or used in intervention delivery or in training of intervention providers. Provide information on where the materials can be accessed (for example, online appendix, URL).	Section 3. What Item 3A. Materials: Describe any physical or informational materials used in the intervention, including those provided to participants or used in intervention delivery or in training of intervention providers. Provide information on where the materials can be accessed (for example, online appendix, URL).	Section 4. What Item 4A. Materials: Describe any physical or informational materials used in the intervention, including those provided to participants or used in intervention delivery or in training of intervention providers. Provide information on where the materials can be accessed (for example, online appendix, URL).	Section 5. What Item 5A. Materials: Describe any physical or informational materials used in the intervention, including those provided to participants or used in intervention delivery or in training of intervention providers. Provide information on where the materials can be accessed (for example, online appendix, URL).
	RETAINED ORIGINAL		
Item 4. What (procedures) Item 4. Procedures: Describe each of the procedures, activities, and/or processes used in the intervention, including any enabling or support activities.	Item 3B. Procedures: Describe each of the procedures, activities, and/or processes used in the intervention, including any enabling or support activities.	Item 4B. Procedures: Describe each of the procedures, activities, and/or processes used in the intervention, including any enabling or support activities.	Item 5B. Procedures: Describe each of the procedures, activities and/or processes used in the intervention, including any enabling or support activities.
	RETAINED ORIGINAL		

Item 5. Who provided For each category of intervention provider (for example, psychologist, nursing assistant), describe their expertise, background and any specific training given.	Section/Item 4. Who provided For each category of intervention provider (for example psychologist, nursing assistant), describe their expertise, background and any specific training given.	Section/Item 6. Who provided For each category of intervention provider (for example psychologist, nursing assistant), describe their expertise, background and any specific training given.	Section/Item 6. Who provided For each category of intervention provider (for example, psychologist, nursing assistant), describe their expertise, background and any specific training given.
	RETAINED ORIGINAL		
Item 6. How Describe the modes of delivery (such as face to face or by some other mechanism, such as internet or telephone) of the intervention and whether it was provided individually or in a group.	Section 5. How Item 5A. Describe the modes of delivery (such as face to face or by some other mechanism, such as internet or telephone) of the intervention and whether it was provided individually or in a group.	Section 7. How Item 7A. Describe the modes of delivery (such as face to face or by some other mechanism, such as internet or telephone) of the intervention and whether it was provided individually or in a group.	Section/Item 7. How Describe the modes of delivery (such as face to face or by some other mechanism, such as internet or telephone) of the intervention and whether it was provided individually or in a group.
	RETAINED ORIGINAL Item 5B. Describe the provision of support in delivering the intervention. <i>For example, manual facilitation of stepping with physical assistance of two therapists, auditory cueing using a metronome at 2-5Hz, guidance statements for the creation of coping strategies, or modelling of correct grammar during language activities.</i>	Item 7B. Strategies: Describe the supplementary strategies that were required to enable the core intervention in response to specific individual or group needs. <i>Supplementary strategies refer to the intervention adjuncts (for example, physical assistance, verbal cueing, transportation arrangements) that must be used for some individuals or groups to facilitate their engagement or effective participation in the core intervention.</i>	
	NEW TIDieR-REHAB	MAJOR ADAPTATIONS TO TIDieR-REHAB	Moved to Section 12 (Item 12A).
Item 7. Where Describe the type(s) of location(s) where the intervention occurred, including any necessary infrastructure or relevant features.	Section/Item 6. Where Describe the type(s) of location(s) where the intervention occurred, including any necessary infrastructure or relevant features.	Section/Item 8. Where Describe the type(s) of environment(s) where the intervention occurred, including any necessary infrastructure or relevant features. <i>Environments and their features may include the relevant physical, social, cultural, economic, political and/or systemic context(s) of the intervention.</i>	Section/Item 8. Where Describe the type(s) of environment(s) where the intervention occurred, including any necessary infrastructure or relevant features. <i>Environments and their features may include the relevant physical, social, cultural, economic, political and/or systemic context(s) of the intervention.</i>
	RETAINED ORIGINAL	MINOR ADAPTATIONS TO ORIGINAL	
Item 8. When and how much Describe the number of times the intervention was delivered and over what period of time including the number of sessions, their schedule, and their duration, intensity or dose.	Section 7. When and how much Item 7A. Record the total session duration. <i>Total session duration is defined as the prescribed overall length of time, usually measured in minutes or hours in which a single session of therapy is administered; if minutes are not recorded, this could be recorded as number of repetitions (Page et al., 2012; Lang et al., 2015).</i>	Section 9. ____ How much Item 9A. Session(s) duration: Provide the planned session(s) duration of the intervention. ____	Section 9. How much Item 9A. Session(s) duration: Specify the planned session(s) duration of the intervention.
	NEW TIDieR-REHAB	MINOR ADAPTATIONS TO TIDieR-REHAB	MINOR ADAPTATIONS TO TIDieR-REHAB

	<p>Item 7B. Record the work duration. <i>Work duration is defined as the length of time, usually reported in minutes, spent participating in the task/intervention. Work duration - total session duration minus rest time (Lang et al., 2015).</i></p>	<p>Item 9B. Essential elements amount: Provide the planned session(s) duration/repetitions for engagement with essential elements of the intervention. <i>Essential elements amount refers to the time or repetitions, within the total duration of a single session, that is spent “actively” participating in the core components of the intervention.</i></p>	<p>Item 9B. Essential elements amount: Specify the planned session(s) duration and/or repetitions of the essential elements of the intervention. <i>Essential elements amount refers to the time and/or repetitions, within the total duration of a single session, that is spent “actively” participating in the core components of the intervention.</i></p>
	<p>NEW TIDieR-REHAB Item 7C. Record the frequency. <i>Frequency is defined as the number of sessions per day and/or week, during the total period of time the intervention is delivered (Page et al., 2012; Lang et al., 2015). For example, two sessions per day, three times per week.</i></p>	<p>MAJOR ADAPTATIONS TO TIDieR-REHAB Item 9C. Frequency: Provide the planned frequency of the intervention. —</p>	<p>MINOR ADAPTATIONS TO TIDieR-REHAB Item 9C. Frequency: Specify the planned frequency of the intervention.</p>
	<p>NEW TIDieR-REHAB Item 7D. Record the intervention length. <i>Intervention length is defined as the total period of time, usually reported in weeks or months, an intervention is delivered for (Page et al., 2012).</i></p>	<p>MINOR ADAPTATIONS TO TIDieR-REHAB Item 9D. Intervention length: Provide the planned overall length of the intervention. —</p>	<p>MINOR ADAPTATIONS TO TIDieR-REHAB Item 9D. Intervention length: Specify the planned overall length of the intervention.</p>
	<p>NEW TIDieR-REHAB Section 8. How challenging Item 8A. Describe the method used to measure and set the intervention intensity. <i>Intensity can be measured using a range of subjective and objective measures. Subjective intensity could include patient-reported estimates of difficulty such as setting intensity at 13-16 on the Borg Rating of Perceived Exertion Scale. Objective intensity could include the level of language complexity (such as increasing the number of steps within an instruction) or %VO2max.</i></p>	<p>MINOR ADAPTATIONS TO TIDieR-REHAB Section/Item 10. How difficult Describe the approach(es) used to set and monitor the intervention/task difficulty level. <i>Difficulty refers to the absolute or relative level of challenge, complexity, physiological intensity, grade, load or tolerance of an intervention/task at a given time and may be evaluated using subjective or objective measures.</i></p>	<p>MINOR ADAPTATIONS TO TIDieR-REHAB Section/Item 10. How challenging Describe the approach(es) used to set and monitor the intervention/task challenge level. <i>Challenge may include the nominal, functional or perceived level of difficulty, effort, physiological intensity ___ or cognitive load ___ of an intervention/task at a given time and may be evaluated using subjective or objective measures.</i></p>
	<p>NEW TIDieR-REHAB Item 8B. Report if the method for setting the intensity uses a reliable and validated tool.</p>	<p>MAJOR ADAPTATIONS TO TIDieR-REHAB</p>	<p>MINOR ADAPTATIONS TO TIDieR-REHAB</p>
Item 9. Tailoring	<p>NEW TIDieR-REHAB Section 9. Tailoring Item 9B. Report the dosage parameter(s) used to progress the intervention. <i>Dosage parameters include the total session duration, work duration, frequency and intensity.</i></p>	<p>REMOVED TIDieR-Rehab Section/Item 11. Regression/Progression Describe when and how the planned regression or progression of dosage parameter(s) was implemented. —</p>	<p>Section/Item 11. Regression/Progression Describe the planned regression and/or progression of dosage parameter(s), including when and how. <i>Dosage parameters refer to the amount (Section 9. How much) and challenge (Section 10. How challenging) of the intervention.</i></p>
	<p>NEW TIDieR-REHAB</p>	<p>MAJOR ADAPTATIONS TO TIDieR-REHAB</p>	<p>MINOR ADAPTATIONS TO TIDieR-REHAB</p>

Signal N, et al. *BMJ Open* 2024; 14:e084319. doi: 10.1136/bmjopen-2024-084319

<div>Item 12. How well – Actual</div> <div>If intervention adherence or fidelity was assessed, describe the extent to which the intervention was delivered as planned.</div>	<div>Item 11B. Actual: If intervention ___ fidelity was assessed, describe the extent to which the intervention was delivered as planned.</div> <div>MINOR ADAPTATIONS TO ORIGINAL</div> <div>Item 11C. Describe how engagement/adherence was measured. <i>For example, use of a diary, video recording of sessions or use of a pedometer.</i></div> <div>NEW TIDieR-REHAB</div> <div>Item 11D. Describe the monitoring and reporting of adverse events.</div> <div>NEW TIDieR-REHAB</div> <div>Item 11E. Describe the number and seriousness of adverse events and their relatedness to the intervention.</div> <div>NEW TIDieR-REHAB</div>	<div>Item 14B. Actual: If intervention adherence or fidelity was assessed, describe the extent to which the intervention was delivered as planned.</div> <div>REINTRODUCED ORIGINAL</div> <div>REMOVED TIDieR-REHAB</div> <div>Section 15. Adverse events</div> <div>Item 15A. Plan: Describe the monitoring ___ of adverse events.</div> <div>MINOR ADAPTATIONS TO TIDieR-REHAB</div> <div>Item 15B. Actual: Describe the number and seriousness of adverse events that occurred, and their relatedness to the intervention.</div> <div>MINOR ADAPTATIONS TO TIDieR-REHAB</div>	<div>Item 14B. Actual: If intervention adherence or fidelity was assessed, describe the extent to which the intervention was delivered as planned.</div> <div>RETAINED ORIGINAL</div> <div>Section 15. Harms</div> <div>Item 15A. Plan: Describe the monitoring of adverse consequences. <i>Adverse consequences include any negative undesired effects related to the intervention, including physical, mental, social and/or spiritual effects.</i></div> <div>MINOR ADAPTATIONS TO TIDieR-REHAB</div> <div>15B. Actual: Describe any adverse consequences, including the number, seriousness and relatedness to the intervention.</div> <div>MINOR ADAPTATIONS TO TIDieR-REHAB</div>
--	--	--	---

Key: **bold**, change from previous version; *grey*, no change from previous version; original, TIDieR item as per Hoffman et al., 2014; TIDieR-Rehab, TIDieR-Rehab extension item; RETAINED/REINTRODUCED, maintained a previous version; NEW, addition for TIDieR-Rehab; MAJOR ADAPTATIONS, revision of previous version; MINOR ADAPTATIONS, refinement of previous version; REMOVED, omitted from TIDieR-Rehab.