Examining dose frameworks to improve aphasia rehabilitation research

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Abstract
The effect of treatment dose on recovery of post-stroke aphasia is not well understood. Inconsistent conceptualisation, measurement, and reporting of the multiple dimensions of dose hinders efforts to evaluate dose-response relationships in aphasia rehabilitation research. We review the state of dose conceptualisation in aphasia rehabilitation and compare the applicability of three existing dose frameworks to aphasia rehabilitation research – the Frequency, Intensity, Time, and Type principle (FITT), the Cumulative Intervention Intensity (CII) framework, and the Multidimensional Dose Articulation Framework (MDAF). The MDAF specifies dose in greater detail than the CII framework and the FITT principle. On this basis we selected the MDAF to be applied to three diverse examples of aphasia rehabilitation research. We next critically examined applicability of the MDAF to aphasia rehabilitation research and identified the next steps needed to systematically conceptualise, measure, and report the multiple dimensions of dose, which together can progress understanding of the effect of treatment dose on outcomes for people with aphasia following stroke. Further consideration is required to enable application of this framework to aphasia interventions that focus on participation, personal, and environmental interventions and to understand how the construct of episode difficulty applies across therapeutic activities used in aphasia interventions.

Keywords
Aphasia, rehabilitation, treatment, dose

Abbreviations
CII Cumulative Intervention Intensity
FITT Frequency, intensity, time, and type
ICF International Classification of Functioning, Disability and Health
MDAF Multidimensional Dose Articulation Framework

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Disclosures
None

Systematic investigation of treatment dose, that is, how much treatment and in what schedule, is essential for a breakthrough in stroke and aphasia recovery\textsuperscript{1}. To date, investigations of the effect of treatment dose on aphasia recovery have been exploratory, unsystematic, and hampered by issues relating to dose conceptualisation, measurement, and reporting\textsuperscript{2-4}. In this paper we examine the complexity of aphasia treatment dose exploration and consider conceptual frameworks to underpin the development of high quality dose-related aphasia rehabilitation research.

Systematic reviews and meta-analyses indicate that treatments for aphasia are, on average, effective\textsuperscript{5-7}. However, aphasia is a heterogeneous condition with highly variable treatment response across individuals and point estimates of effects may conceal important individual differences in response to a given treatment dose\textsuperscript{8}. Aphasia heterogeneity stems from differential impacts from stroke on i) diffuse neural networks that underpin language processing, and ii) the complex process of interpersonal communication. Effective communication relies on rapid interactions between multiple linguistic components (e.g., phonology, lexical-semantics, syntax, discourse processing) in multiple modalities (e.g., spoken, written, gestural) that interact with cognitive functions (e.g., attention, working memory). These are mediated by different context-dependent social norms (e.g., formality, familiarity), degrees of conversational freedom (e.g., shared referents, novel topics), and communicative goals (e.g., everyday functional transactions, group social interactions). Aphasia arises from a breakdown of one or more of the linguistic components, resulting in a variety of patholinguistic phenotypes.

Different aphasia treatments have been developed to address this heterogeneity. Impairment-focused approaches aim to target breakdown in linguistic processes. Functional approaches aim to enhance participation in personally relevant communication-related activities and may include nonverbal communication methods such as drawing, gesture, and the use of communication devices. Psychological treatments aim to address the mental health consequences of communication disability, and environmental approaches target communication partners and communication accessibility. It is common in clinical practice for people with aphasia to undertake treatments using multiple different approaches simultaneously\textsuperscript{9}. Determining the required dose of each different treatment approach to achieve the communication goals of a person with aphasia is of primary clinical and research importance. To achieve this, consistent conceptualisation and systematic measurement and reporting of treatment dose is a necessary precursor.

Aims
Following a brief review of the evidence for dose effects in aphasia treatment research, we aim to (1) identify the limitations of current dose conceptualisation, measurement, and reporting in aphasia treatment research, (2) compare and contrast the applicability of existing dose frameworks to aphasia treatment research, (3) apply the most appropriate framework to a range of aphasia rehabilitation studies, and (4) propose steps required to systematically evaluate the effect of dose on treatment outcomes for people with aphasia following stroke.

Evidence for dose effects in aphasia rehabilitation research
Systematic reviews have concluded that higher treatment doses may lead to better aphasia recovery\textsuperscript{3,5}. The recent network meta-analysis of individual patient data (n=959, 25 trials) by the RELEASE group\textsuperscript{7} suggested that the greatest language gains measured on standardised aphasia assessment batteries were following doses of 20-50 hours delivered at either 2-4 hours (for functional communication) or 9+ hours (for overall language and comprehension) per week. The 2016 Cochrane review of speech and language therapy in aphasia\textsuperscript{7} likewise found superiority of more
treatment hours over fewer for the recovery of functional communication and written expression but found no clear dose-response relationships for other aphasia outcomes.

The critical dose required to demonstrate clinically meaningful and statistically significant recovery of language and communication remains unknown but will likely be specific to aphasia treatment type and phase of recovery. Studies in the acute-subacute phase have found no superiority of relatively higher compared to lower treatment doses. Husak and colleagues conducted a systematic review of aphasia rehabilitation in the first four months following stroke. Of six studies meeting inclusion criteria, five studies found no significant difference in outcomes between patients provided either a lower or higher number of treatment hours, and one study reported superior findings in outcomes when participants received less treatment in the early recovery period. In contrast, dose-response relationships in the chronic phase of recovery appear to favor higher over lower doses of treatment. A recent Intensive Comprehensive Aphasia Program delivered 100 hours of treatment over 3 weeks and demonstrated medium to large effect sizes immediately following treatment and at longer term (e.g., 3-month) follow up. The dose delivered in this program was much higher than the most frequently prescribed dose of 30 hours as reported in a recent review of dose that examined 112 aphasia treatment research studies. Given these results, it is possible other trials that have published null effects have failed to deliver a sufficient dose to elicit a therapeutic effect, a phenomenon recognized across the domains of rehabilitation trials. Therefore, given the heterogeneity of aphasia and uncertainty regarding dose-response relationships, systematic investigation of treatment dose is required to progress this emerging field of research and positively influence language and communication outcomes for people living with aphasia.

Limitations associated with current dose conceptualisation, terminology, and reporting in aphasia rehabilitation research

Dose is under-specified in aphasia rehabilitation research

Consensus on dose terminology for non-pharmacological treatments has not been established within the stroke rehabilitation literature nor in aphasiology. In aphasia treatment studies, there is inconsistent use of terminology to describe dose dimensions. For example, dose, dosage, and intensity are used interchangeably to refer to divergent concepts including: the number of repetitions within a specific therapy task; the number, duration, and frequency of sessions; the overall duration of a treatment program in weeks; and, the total number of treatment hours provided over the course of an intervention. This inconsistency creates confusion and confounds attempts to examine dose-response relationships that may underpin treatment effectiveness.

Aphasia researchers and clinicians most commonly report treatment dose as the amount of time spent in the treatment environment, usually measured in hours. Hours of treatment are easy to calculate, aid comparison between studies, and can be easily interpreted by healthcare providers, recipients, policy makers, and funding bodies. However, measuring dose only in hours may be inadequate due to an inaccurate assumption that all hours of treatment are equal. Clinically, each hour of treatment may comprise a variety of different tasks, targeting different goals, each requiring the provision of a different number and combination of therapeutic tasks that might be punctuated by periods of rest or inactivity. In research reports, especially large trials, it is not always clear how often different therapeutic activities are performed within a given period of time unless treatment details are accurately defined, measured and reported.

A recurring limitation in examinations of dose effects in aphasia rehabilitation is the inconsistent measurement and reporting of treatment dose. For example, previous reviews have been constrained by a lack of comprehensive data collection in clinical trials and have suggested that improved quality of trial reporting “will further contribute to transparency, replication of findings, and subsequent meta-analyses.” Although clear reporting of intervention duration, dose, intensity, and mode of delivery is vital for interpretation of results and study replication, there are several well-accepted frameworks and guidelines to support systematic investigation of dose effects in aphasia rehabilitation. The Rehabilitation Treatment Specification System was developed with the goal of achieving consistent terminology use and reporting across disciplines, and to support the development of rehabilitation treatments. This system focuses on targets (the behaviour that is expected to change as a result of treatment), and mechanism(s) of action (why a given treatment works). It has been applied to three broad aphasia intervention approaches: neurobiological, cognitive-linguistic, and functional approaches. In each case, conceptualising and reporting “ingredients” remained a challenge because there was insufficient prescription within the Rehabilitation Treatment Specification System to guide dose reporting and no other universally agreed upon guidelines for the reporting of dose in aphasia rehabilitation.

To undertake a systematic investigation of dose, the field of aphasia research needs a conceptual framework with common terminology to guide dose articulation that can be implemented across research studies. Consistency is vital to accurately frame research questions, design studies, investigate treatment fidelity, compare across and replicate studies, and improve communication amongst all stakeholders. Although there are several well-accepted frameworks and guidelines to support systematic investigation of dose effects in aphasia rehabilitation, an ideal dose framework would provide a comprehensive and granular characterisation of the amount of treatment provided across all types of aphasia treatment. To our knowledge, there are three dose-specific frameworks for non-
Comparison of the applicability of existing dose frameworks to aphasia rehabilitation research

Frequency, Intensity, Time, and Type principle (FITT)

The American College of Sports Medicine’s Guidelines for Exercise Testing and Prescription recommend practitioners use the FITT principle – frequency (how often), intensity (how hard), time (duration or how long), and type (mode or what kind) of exercise – when designing and prescribing individualised physical exercise programs. Additional components such as volume (total amount of exercise) and progression (exercise advancement) can also be considered. The FITT principle has been applied, though not routinely, in post-stroke exercise research, and has recently been used as a framework to quantify the dose of swallowing rehabilitation exercises provided in an inpatient rehabilitation setting. It has not, to our knowledge, been applied to aphasia rehabilitation research.

Cumulative Intervention Intensity framework (CII)

Warren and colleagues’ CII framework asserts that the amount of treatment provided or received is a product of the number of times the active ingredients of treatment are applied per session and the number of sessions provided over the treatment duration. The active ingredients are the actions performed by either the treatment provider or recipient that are theoretically linked to the underlying mechanisms of that treatment. Figure 1 depicts the relationship between dose form, dose, session duration, session frequency, total intervention duration, and cumulative intervention intensity with an adaptation to the original CII framework separating dose and session duration because these two parameters can be manipulated independently (e.g., dose of 50 or 100 trials in a 30-minute session, dose of 100 trials in a 30- or 60-minute session). First developed to characterise treatments in the field of developmental disabilities, this framework has been used to report treatment dose in a small number of aphasia studies and other speech-language pathology areas such as apraxia of speech and paediatric language. For example, in a study comparing the effect of intensive versus distributed treatment, Dignam and colleagues used the CII to demonstrate non-significant differences in the average number of therapeutic inputs provided throughout treatment between groups.

Multidimensional Dose Articulation Framework (MDAF)

A group of multidisciplinary stroke researchers with expertise spanning upper limb, mobilisation, motor speech, and cognitive functions proposed the MDAF (Figure 2). It was developed to guide the specification of non-pharmacological therapeutic dose, conceptualise the multidimensional nature of, and links between, dose dimensions, and provide consistent terminology across the stroke recovery and rehabilitation fields. As Figure 2 shows, an intervention is provided over a duration (e.g., weeks, months, or years). Within that time, treatment occurs on one or more days which can vary in number and spacing (e.g., daily or weekly treatment). On any given treatment day, there will be one or more sessions which can be defined by their length in time. Sessions contain episodes of variable length that are either active (i.e., time spent on a task) or inactive (i.e., pauses or breaks). The ratio of active to inactive time in a given session renders the session density. Active episodes comprise tasks of different intensity (Figure 2 height of episode) and difficulty (colour of episode). Taken together, these multiple dimensions of dose constitute the overall amount of treatment provided or received. The MDAF emphasises it can be used to conceptually plan (methods) and actually report what dose is delivered (results). Due to the recency of publication of this framework, there are no published examples of its application in the literature.

Table 1 provides a comparison of dose dimensions and terms defined by the FITT principle, the CII framework, and the MDAF including reference to commonly used terms in the aphasia literature. There is some overlap between these conceptual frameworks but also some important structural differences. The MDAF clearly provides a more comprehensive characterisation of dose than the CII framework and the FITT principle (Table 1). In particular, the MDAF specifies more detailed temporal parameters (days, number and spacing of days and sessions) and episode-level characteristics (length, difficulty, and intensity) than the CII framework and the FITT principle. Neither the CII framework nor the FITT principle include dose dimensions that are not captured in the MDAF. The CII framework and the FITT principle include specification of the content of treatment (labelled dose form and type, respectively) whereas the MDAF does not include specification of treatment content; the MDAF was developed to examine constructs of dose irrespective of treatment type and is intended to be used in conjunction with treatment specification tools (e.g., TIDieR). Although treatment type and treatment dose are interrelated and require consideration when designing a research protocol or clinical intervention program, the current investigation is primarily concerned with dose conceptualization, not treatment specification.

Given uncertainty regarding which dose parameters are important for recovery, greater specification may be advantageous.

Figure 1 Cumulative Intervention Intensity framework, as adapted by Baker (copyright Wiley, reproduced with permission)

Figure 2 Multidimensional Dose Articulation Framework (copyright Wolters Kluwer Health, Inc. reproduced with permission)
Table 1 Dose descriptors and common terms used in aphasia literature that are covered by the FITT principle, CII framework, and MDAF

<table>
<thead>
<tr>
<th>Conceptual descriptor (terminology used in aphasia literature)</th>
<th>FITT</th>
<th>CII</th>
<th>MDAF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Task being performed (task, activity)</td>
<td>Type</td>
<td>Dose form</td>
<td>Detailed using treatment specification tool (e.g., TIDieR)</td>
</tr>
<tr>
<td>Overall amount of treatment provided or received (dose, dosage)</td>
<td>Volume</td>
<td>Cumulative intervention intensity</td>
<td>Defined in relation to all dimensions listed below</td>
</tr>
<tr>
<td>Overall length of the intervention (total duration, intervention/treatment period/phase)</td>
<td>Total intervention duration</td>
<td>Duration</td>
<td></td>
</tr>
<tr>
<td>Number of days of intervention</td>
<td>Days (number)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>The distribution of days, number of days per week</td>
<td>Days (spacing)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Number of sessions</td>
<td>Product of total intervention duration x session frequency</td>
<td>Sessions (number)</td>
<td></td>
</tr>
<tr>
<td>The distribution of sessions, number of sessions per week (frequency, intensity&lt;sup&gt;*&lt;/sup&gt;)</td>
<td>Frequency</td>
<td>Session frequency</td>
<td>Sessions (spacing)</td>
</tr>
<tr>
<td>Timed duration of session(s)</td>
<td>Time</td>
<td>Session duration</td>
<td>Session length</td>
</tr>
<tr>
<td>Amount of time spent actively engaged in therapy activities&lt;sup&gt;2&lt;/sup&gt; (time-on-task)</td>
<td>Sum of length of active episodes</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Proportion of time spent actively engaged in therapy activities</td>
<td>Session density (i.e., sum of length of active episodes / session length)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Basic unit of treatment which contains the active ingredient(s) of a treatment</td>
<td>Teaching episode</td>
<td>Episode</td>
<td></td>
</tr>
<tr>
<td>Number of episodes administered during a session</td>
<td>Dose</td>
<td>Sum of episodes</td>
<td></td>
</tr>
<tr>
<td>How long the task is performed for, in units of time&lt;sup&gt;3&lt;/sup&gt;</td>
<td>Episode length</td>
<td></td>
<td></td>
</tr>
<tr>
<td>How hard the task is to perform&lt;sup&gt;2&lt;/sup&gt; (task difficulty, task hierarchy)</td>
<td>Episode difficulty</td>
<td></td>
<td></td>
</tr>
<tr>
<td>How much of the task is performed per episode or unit of time&lt;sup&gt;2&lt;/sup&gt; (dose rate&lt;sup&gt;15&lt;/sup&gt;)</td>
<td>Intensity</td>
<td>Episode intensity</td>
<td></td>
</tr>
</tbody>
</table>

* Commonly defined as the number of hours per week.
* Descriptors used by Hayward and colleagues<sup>15</sup>.
Table 2 Application of the MDAF to the published reports of three selected intervention studies

<table>
<thead>
<tr>
<th>Study</th>
<th>Duration</th>
<th>Days (number)</th>
<th>Days (spacing)</th>
<th>Sessions (number)</th>
<th>Sessions (spacing)</th>
<th>Session length</th>
<th>Session density</th>
<th>Episode length</th>
<th>Episode difficulty</th>
<th>Episode intensity</th>
<th>Additional dose dimensions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Harnish et al., 2014</td>
<td>2 weeks</td>
<td>8 days</td>
<td>4 days per week</td>
<td>8 sessions</td>
<td>1 per day</td>
<td>60 minutes</td>
<td>Not reported</td>
<td>Approximately 1 minute</td>
<td>Not reported</td>
<td>8 naming opportunities per episode</td>
<td>400 naming opportunities per session, 6.67 naming opportunities per minute</td>
</tr>
<tr>
<td>Rose et al., 2022</td>
<td>2 weeks</td>
<td>10 days</td>
<td>Daily</td>
<td>30 sessions</td>
<td>3 per day</td>
<td>60 minutes</td>
<td>Not reported</td>
<td>Variable depending on task and participant factors (e.g., aphasia severity)</td>
<td>The task and targets are selected based on participant performance using a prespecified rubric based on linguistic difficulty.</td>
<td>8 naming opportunities per episode</td>
<td>15-minute daily home practice Participant self-rating of fatigue and distress was measured on 100mm visual analogue scale at start and end of every day.</td>
</tr>
<tr>
<td>Attard et al., 2018</td>
<td>12 weeks</td>
<td>12 days</td>
<td>1 day per week</td>
<td>12 sessions</td>
<td>1 per day</td>
<td>120 minutes</td>
<td>Not reported</td>
<td>Variable depending on task and participant factors (e.g., aphasia severity).</td>
<td>Not reported</td>
<td>Not reported</td>
<td>Spouses involved in a parallel program: some sessions joint with person with aphasia and some sessions separate for spouses.</td>
</tr>
</tbody>
</table>
when evaluating dose-response relationships. Therefore, the MDAF is the most likely candidate of these three frameworks to progress understanding of dose-response relationships in post-stroke aphasia rehabilitation research. We will now apply the MDAF to three diverse aphasia treatments and demonstrate its potential utility.

Applications of the MDAF toaphasia rehabilitation research

Table 2 demonstrates application of the MDAF to intervention studies that we selected to cover the breadth of aphasia treatment approaches including an individual lexical-retrieval treatment15, a group-based combined lexical retrieval and syntax treatment42, and an interdisciplinary group-based intervention that targets multiple areas of the International Classification of Functioning, Disability and Health (ICF)43. Each column in Table 2 represents the overall amount of treatment provided in each study. The values in Table 2 are the doses planned to have been delivered, as reported in each study. In this study, we retrospectively applied the MDAF to published reports however, the MDAF is intended to be used prospectively to describe both the planned and actual doses delivered in a trial15. While the dimensions of duration, days, session length, and density were consistently reported, data concerning the dimensions of episode length, difficulty, and intensity were not always available. Multidimensional dose specification was comprehensive for the individual and group-based impairment-focused treatments16, 42; all dose dimensions were reported in these studies with the exception of episode difficulty in the individual therapy16. Dose specification as per the MDAF was only partially achieved in the group-based intervention targeting multiple ICF areas43 with only overall duration, number of treatment days, number of treatment sessions, and session length reported.

A key strength of the MDAF is that it provides a systematic approach for specifying and reporting dose that can be applied to a range of aphasia treatments. Lack of reporting of some dose dimensions in published studies may reflect limited attention to these dimensions in the intervention design phase. Indeed, the MDAF may be an especially useful tool in early-phase research when questions regarding dose-relationships should be addressed before scaling up to large scale definitive trials15. We will now outline challenges associated with describing and quantifying dose dimensions in complex behavioural interventions.

Challenges applying the MDAF toaphasia rehabilitation research

Aphasia treatments range in scope across the dimensions of the ICF including those focused on participation (e.g., community aphasia groups43), personal factors (e.g., mood46), and the environment (e.g., Communication Partner Training47). Table 2 demonstrates that isolating episodes within interventions focused on reducing linguistic impairment, such as naming or syntax therapies, is reasonably straightforward15 but is more complicated in group-based treatments targeting multiple areas of the ICF43. Interventions focused on participation, personal, or environmental factors are generally multifaceted in design and application, and frequently involve dyads or groups of people. Defining, isolating, and counting episodes in these multicomponent interventions requires further research. For example, within a complex group-level communication intervention that uses art making as a therapeutic activity and social interaction as the target47, what is the episode and what is episode difficulty? Can these treatment approaches be reduced to discrete episodes or do they require a less granular, more global framework that allows for flexibility and iterative adaptation as therapies develop over the course of intervention? Aphasia rehabilitation is not alone in utilising multicomponent approaches that span the entire ICF, with social work another example where alternative frameworks for conceptualising dose and ensuring treatment fidelity have been applied45. For example, Washington and colleagues47 demonstrated stronger fidelity when using a composite dose measure compared to a measure of the sum of individual elements in their study of a complex multicomponent social work intervention.

The MDAF includes episode difficulty as a static quality intrinsic to the task being performed. Although the role of perceived task difficulty in aphasia treatment is being evaluated48, neither perceived nor intrinsic task difficulty have been linked to dose despite the clinical relevance. In aphasia treatments, difficulty may be operationalised in relation to choice of targets and their lexical properties, cognitive load, linguistic level, communicative context, presence or absence of distractors or cues, or speed of response. For example, in word retrieval treatment, a task requiring retrieval of an abstract, low frequency target word (e.g., justice) will be intrinsically more difficult than retrieval of a concrete, high frequency word (e.g., man)49. Using those words in sentences may be experienced as a more challenging task for individuals with syntactic processing deficits than those without. Importantly, this perceived difficulty may reduce with treatment while the intrinsic difficulty of the task remains unchanged. Clinicians may aim to pitch a given task at an appropriate challenge point by manipulating the rate that word and sentence production tasks need to be completed and the complexity of the communication environment where the task(s) take place (e.g., with the therapist in a quiet clinic room; in a group conversation; in a busy shopping centre). While reliable measurement of task difficulty within and across different therapeutic tasks is still to be established, its identification as a distinct dose parameter that is intrinsic to the tasks and could be experimentally manipulated is a necessary precursor step.

In summary, the MDAF appears well suited to dose description for treatments in which episodes are discrete and countable but more work is required to determine how best to articulate and quantify dose in complex multi-faceted aphasia treatments. Further work will be required to understand the dimension of episode difficulty across therapeutic activities used in aphasia interventions.

<table>
<thead>
<tr>
<th>Box 1 Next steps to progress systematic dose conceptualisation, measurement, and reporting inaphasia rehabilitation</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Aphasia researchers could attempt to capture the multiple dimensions of dose in treatment studies</td>
</tr>
<tr>
<td>• Aphasia researchers could use the MDAF to systematically specify dose in early-phase research protocols and reporting</td>
</tr>
<tr>
<td>• Aphasia researchers could aim to delineate and define episodes within treatments that target discourse, participation, personal factors, and the communication environment</td>
</tr>
<tr>
<td>• Aphasia researchers could conduct research that aims to understand episode difficulty within and across different therapeutic tasks. This may require development of reliable measurement tools for episode difficulty.</td>
</tr>
<tr>
<td>• Aphasia clinicians could use the MDAF to systematically capture clinical dose data</td>
</tr>
</tbody>
</table>

**Summary**

Dose is an important factor in stroke and aphasia rehabilitation. Personalised treatment prescription should consist not only of the type but also the dose of treatment required to promote long term positive change for a specific individual with aphasia. Given the heterogeneity of people with aphasia and the large variability in aphasia treatment targets and approaches, treatment doses will likely require personal calibration to a range of biological, aphasia, and psychosocial recovery factors with consideration of personal relevance, motivation, and reward. To get closer to this objective, we need to document all dose dimensions for a given treatment50. Moreover, investigation of dose prompts deep thinking into the theory behind why and how a treatment works. While the issue of what comprises a clinically practical dose to deliver is relevant, it is distinct from the question of the dose required to drive recovery. The priority for researchers is to use treatment theory and systematic investigation to determine personalised dose targets in order that aphasia outcomes can be improved for individuals. Once gold-standard treatment regimens are established, the discussion of what is feasible, practical, and economical will follow. A multidimensional dose framework is required to guide the development, implementation, and evaluation of dose-finding studies designed to determine the range of safe and tolerable doses of an intervention and dose-response relationships. Such a
framework would enable synthesis of data across studies and theoretical exploration of what drives treatment response in aphasia treatments, inform the extension of reporting guidelines, aid clinical decision-making, and guide health policy makers. We propose the Multidimensional Dose Articulation Framework13 is a first step towards this purpose and suggest further research to refine this framework to support application to aphasia interventions focused on participation, personal, and environmental intervention approaches. Aphasia researchers are urged to consider using the MDAF to describe dose prescription in research protocols and to frame the reporting of dose parameters in aphasia treatment research.

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