BRAINSTORMING: A study protocol for a randomised double-blind clinical trial to assess the impact of concurrent brain stimulation (tDCS) and working memory training on cognitive performance in Acquired Brain Injury (ABI)

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Abstract

Background: Acquired Brain Injury (ABI) admissions have an incidence of 385 per 100,000 of the population in the UK, and as brain injury often involves the frontal networks, cognitive domains affected are likely to be executive control, working memory, and problem-solving deficits, resulting in difficulty with everyday activities. The above observations make working memory, and related constructs such as attention and executive functioning attractive targets for neurorehabilitation. We propose a combined home-based rehabilitation protocol involving the concurrent administration of a working memory training program (adaptive N-back task) with non-invasive transcranial direct current stimulation (tDCS) of the right dorsolateral prefrontal cortex to promote long-lasting modification of brain areas underlying working memory function.

Method: Patients with a working memory deficit will be recruited and assigned to two age-matched groups receiving working memory training for 2 weeks: an active group, receiving tDCS (2 mA for 20 min), and a control group, receiving sham stimulation. After the end of the first 2 weeks, both groups will continue the working memory training for three more weeks. Outcome measures will be recorded at timepoints throughout the intervention, including baseline, after the 2 weeks of stimulation, at the end of the working memory training regimen and 1 month after the completion of the training.

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Discussion: The aim of the study is to assess if non-invasive tDCS stimulation has an impact on performance and benefits of a working memory training regimen. Specifically, we will examine the impact of brain stimulation on training gains, if changes in gains would last, and whether changes in training performance transfer to other cognitive domains. Furthermore, we will explore whether training improvements impact on everyday life activities and how the home-based training regimen is received by participants, with the view to develop an effective home healthcare tool that could enhance working memory and daily functioning.

Trial registration: This study was registered with clinicaltrials.gov: NCT04010149 on July 8, 2019.

Keywords: Brain injury, Non-invasive brain stimulation, Direct current, Working memory training, N-back, Multi-session

Background

Working memory, which involves temporary maintenance and working with information in mind, is required for cognitive skills such as problem solving, planning and auditory and reading comprehension. Working memory operates across multiple modalities (e.g., auditory and visual), and is a system that combines attentional control with temporary storage and information manipulation [1]. A core concept of Baddeley’s [2] model of working memory is that it consists of multiple components: namely, a phonological loop governing for temporary maintenance and processing of verbal information, a visuospatial sketchpad to maintain and process visual and spatial information, an episodic buffer that bridges working memory and long-term memory, and a central executive that exercises attentional control over the above-named subsystems. A network involving the dorsolateral prefrontal cortex (DLPFC) plays a fundamental role in supporting working memory [3], with spatial tasks primarily involving the right DLPFC and verbal task involving the left DLPFC [4, 5].

Working memory is important for everyday tasks such as problem solving, reasoning, and learning [1]; a deficit in working memory can lead to difficulties with many everyday activities that are necessary for work, study and general functioning [6]. Impaired working memory may consequently have a significant impact on quality of life and ability to participate in social roles, with potential for negative effects on mood and emotional wellbeing. Working memory is known to decline with aging [7] and is frequently compromised following brain injury [8].

Acquired Brain Injury (ABI) is brain damage caused by either a traumatic (e.g. a penetrating head injury due to an accident) or a non-traumatic injury (e.g., stroke, brain tumours, ischemia). ABI may be associated not only with physical but also with cognitive, emotional and behavioural impairments. Our research focuses on cognitive impairment resulting from ABI. As brain injury often involves frontal systems, cognitive domains commonly affected include: attention, executive control, working memory, and problem solving [9, 10]. Working memory impairment is highly prevalent across ABI aetiologies [11, 12] and severity levels. A recent meta-analysis of working memory deficits post traumatic brain injury revealed significant deficits in verbal short-term memory and visuospatial and verbal working memory (effect sizes ranging from .37 to .69 [11]).

The above observations make working memory, and overlapping constructs such as attention and executive functioning, important targets for neurorehabilitation, given the impact of these deficits on everyday function [6, 13]. Admissions for ABI have an incidence of 385 per 100,000 in the UK (https://www.headway.org.uk/about-brain-injury/further-information/statistics/). Individuals post ABI commonly report working memory dysfunctions [14, 15], thus solutions to improve recovery of working memory are critically needed.

Researchers have been exploring whether it is possible to train or improve working memory in clinical groups such as stroke patients [16], people with Parkinson’s disease [17], and children with attention deficit hyperactivity disorder (ADHD) [18]. For training to be considered of value, the gains in performance on the training task must transfer to other cognitive tasks or yield some benefit in daily function outside of the laboratory [19]. However, evidence for transfer to other cognitive tasks or daily activities is not yet consistently obtained [20].

Rehabilitation for working memory impairments can focus on compensatory strategies, which aim to compensate for the impaired functions by using those which remain intact, or supplementary external aids [21]. In contrast, recent studies have focused on increasing working memory capacity through extensive practice and training [19, 22]. Computerized working memory training has shown promising results in healthy participants [23, 24] as well as in ABI patients [7, 16, 25–27]. Mood measures also appear to improve following working memory training [21, 25]. One of the tools that has been assessed for its working memory training potential is the n-back task, which involves determining whether a new item in a sequence is the same as the one that appeared ‘n’ times before. While results have been mixed (likely due in part to methodological inconsistencies between studies [28], a dual n-back implementation
wherein participants keep track of two streams of information (one audio and one visual), was successful in healthy participants [23, 24].

Brain plasticity, which can vary greatly by individual, plays a central role in the success of cognitive training. Transcranial direct current stimulation (tDCS) [29–32] is a non-invasive technique that is thought to modify cortical excitability which, in turn, can directly affect brain plasticity by making relevant brain networks more or less likely to fire synchronously. Synchronous firing has been suggested to modulate long-term potentiation (LTP) like plasticity at the synaptic level, a critical component for learning [33, 34].

The combination of cognitive training protocols and non-invasive tDCS has gained attention as a means for cognitive enhancement in both adults and older adults [35–37]. In young and healthy participants, anodal tDCS over the left or right dorsolateral pre-frontal cortex (DLPFC) improved working memory when administered concurrently with cognitive training using a n-back task, also showing transfer to other domains [38], as well as maintenance over time [39]. Preliminary evidence shows that the congruency between the side of stimulation, and therefore the functional specificity of the DLPFC, and the behavioural task plays an important role in training outcome [40, 41]. As many cognitive functions show a physiological decline even in healthy aging, combined tDCS/cognitive training protocols are of particular interest for older adults [37, 42, 43] but in the elderly results are mixed with some studies revealing null findings [35, 44, 45].

Despite encouraging results obtained in healthy populations, research involving brain stimulation in ABI patients is sparse. By stimulating over the left DLPFC, Kang et al. [46] obtained positive results on attention, albeit in a small sample. In a within-subject design, working memory performance in stroke patients can improve after anodal stimulation of the left DLPFC [47]. Villamar et al. [48] reviewed the potential of the therapeutic use of brain stimulation to modulate neuroplasticity in ABI patients; however, given the diversity of the nature of the lesions, more research is necessary to understand how to use brain stimulation effectively in ABI populations.

Specifically, our primary objective is to determine if the combination of tDCS and working memory training is superior to working memory training alone to improve working memory function in an n-back task in an ABI population. The key secondary objectives are to determine if the combination of brain stimulation with working memory training can:

- boost performance in other cognitive domains (attention, executive functions)
- speed training gains
- lengthen improvement duration to 1 month
- increase improvement magnitude

To address the above questions, we will use a randomized clinical trial, assessor- and patient-blinded with two parallel groups and a simple randomization with a 1:1 allocation ratio.

**Methods/design**

**Study setting**

Patients will be recruited from the Northamptonshire Healthcare NHS Foundation Trust (NHFT) by the Community Brain Injury Service, a multidisciplinary service that provides specialist neurorehabilitation to adults who have experienced an acquired brain injury. People accessing the service from which the participants will be recruited are aged 16 and over; the most common age group from an audit of last year’s referrals was 50–59, with the majority falling between the ages of 30–69. Aetiology of brain injury varies although traumatic brain injury accounted for over half of referrals last year, followed by stroke, tumours and then hypoxia and infection related ABI. The ratio of males to females was 60:40.

Training sessions with brain stimulation will be administered at home by a member of the research team (assessor), who will visit the patient at an agreed time for the first 2 weeks. During the remaining 3 weeks, patients will complete the cognitive training alone from home, via an internet-based application, while receiving a motivational catch up call every week. Weekends will be exempt from training. This home-based approach will help us reduce drop-out. Moreover, it will also increase convenience for the patient, many of whom may live some distance from the brain injury team base.

**Eligibility criteria**

The study will recruit patients who experienced an ABI and have been referred to the Community Brain Injury Service of NHFT. Patients provide written, informed consent before any study procedure occurs (see Fig. 2) and must meet eligibility criteria (Table 1) at randomization. Assessors, who will also administer the brain stimulation,
will receive training in the use of tDCS and basic life support.

Recruitment
Forty adults between the age of 18 and 69 years will be recruited through the Community Brain Injury Service. Patients will be randomly assigned to one of two experimental groups: an active (or right-sided tDCS) group and a control (or SHAM) group. All patients will receive WM training. As there is evidence that age modulates individual responses to both tDCS [50–52] and training regimens [53, 54], the two groups will be age-matched in two age subgroups: those aged between 18 and 45 years and those aged between 46 and 69 years, randomised within age range. Other factors may also influence the impact of the intervention on patients, including medications, time since injury, baseline performance, and these factors will be included as regressors in the analysis whenever appropriate. To reduce the variability due to different locations of lesions, patients with lesions underneath the electrodes (that could modify the induced electric field) will be excluded. Finally, as explained in the following, patients will only be included if they present with a working memory deficit, but no general intellectual impairment.

As the aim of this research is to improve working memory, only participants with a working memory impairment will be included. As we are focusing on improving a documented working memory deficit, we will not attempt to distinguish between pathologies (unless otherwise determined by the exclusion criteria for tDCS, e.g., a lesion underneath the electrodes), rather we will screen patients based on working memory performance. Final analysis will explore differences between pathologies. To identify a working memory impairment, cognitive tests from the Wechsler Adult Intelligence Test (WAIS-IV) and the Wechsler Memory Scale (WMS-IV) will be administered. These tests are administered to every patient as part of the routine care protocol. We will use the following scores:

- A WAIS-IV Full-Scale Intelligence Quotient (FSIQ) lower than 70: this score will identify any patients with significant intellectual impairment. The FSIQ will also be used as a covariate in the analysis to control for the effects of reduced overall intellectual ability rather than a specific working memory issue.
- A WAIS-IV Working Memory Index (WMI) score, obtained as a combination of the digit span score and the arithmetic score in the WAIS-IV, smaller than 85: this score identifies an auditory working memory impairment.
- A WMS-IV Visual Working Memory Index (VWMI) score, below 85 identifies weakness in visuospatial working memory.

Table 1 Eligibility criteria

| Inclusion Criteria |
|--------------------|
| 1. Referred to the service |
| 2. Are between 18 and 69 years of age, inclusive |
| 3. Have capacity and able to provide informed consent |
| 4. Normal or corrected-to-normal vision and hearing |
| 5. Having a working memory impairment (see screening procedure) |
| 6. At least 3 months between the injury and the starting of the study |
| 7. Has a computer or has access to a computer |

Exclusion criteria

1. Pre-injury psychiatric or neurological disease by self-report (e.g., anxiety disorder, ADHD, Parkinson’s disease, etc.)
2. History of diagnosed severe depression (diagnosed pre-injury)
3. History of epilepsy (diagnosed pre-injury)
4. Family history of epilepsy
5. Have had fainting spells or syncope in the last 3 years pre-injury
6. Have significant hearing loss, vision or motor impairment that would prevent them from performing the task
7. Known to be pregnant
8. Consuming medication affecting cortical excitability or recreational drugs
9. Metal (except titanium) or electronic implants in the brain/skull (e.g., splinters, fragments, clips, cochlear implant, deep brain stimulation, medication pump,...)
10. Metal (except titanium) or any electronic device at other sites in your body, such as cardiac pacemaker or traumatic metallic residual fragments
11. Have skin problems such as dermatitis, psoriasis or eczema under the stimulation sites
12. Have had brain stimulation in the past 6 months
13. Have undergone transcranial electric or magnetic stimulation in the past (more than 6 months) which resulted in adverse effects
14. Skull fractures, significant skull defects, skull plates or large vessels occlusions, no significant cortical lesion or atrophy at the site of electrode
15. Having had a seizure at the time of accident or between the injury and starting of the therapy.

We follow recommendations by McLaren et al. [49], and only exclude drugs that are shown to block tDCS effects (e.g., sodium channel blockers, calcium channel blockers, and NMDA receptor antagonist), while other medications shown to only modulate tDCS effects will be tracked and considered as covariate in the analysis. We will monitor for medication changes as a potential bias on the outcomes of the intervention and report any changes that may occur. If a patient become in need of medications shown to block tDCS effects, they will be excluded from further participation. We will monitor for medication changes as a covariate in the analysis. To identify a working memory deficit, we will use working memory indices that are a combination of auditory and visual working memory to identify an objectively defined working memory deficit. We will also run an exploratory post-hoc analysis factored by type of baseline WM deficit to ascertain if there are any modality specific benefits. Although this score reflects at least mild impairment, a weakness in working memory...
memory would be large enough to potentially improve with training.

Sample size calculation
The chosen sample size is based partly on the available literature that examines the effects of cognitive training on healthy young and older volunteers and additional pilot data collected in our laboratory from healthy young and elderly adults. Specifically, a statistical power analysis was performed for sample size estimation, based on data from Au et al. [39], comparing the ACTIVE (right-tDCS, 20 subjects) to the CONTROL (SHAM, 22 subjects) group. The effect size (ES) in this study was 0.73, considered to be medium to large using Cohen’s criteria [55]. With an alpha = .05 and power = 0.80, the projected sample size needed with this effect size (GPower 3.1) is approximately $N = 48$ for this simplest between group comparison.

Evidence has shown that individuals with low initial performance respond better to brain stimulation [56]. As we expect ABI patients in our sample to have low baseline performance, due to our selection criteria, we may also expect them to respond better to the intervention, therefore showing larger effects. Assuming, as explained above, that larger effects could reasonably be expected in a clinical population, our proposed sample size of 40 will be adequate for the main objective of this study. A post hoc power analysis will be conducted to assess if the sample size is appropriate to detect differences in a clinical population such as ABI.

Intervention
The BRAINSTORMING trial is designed as a randomized clinical trial, assessor and patient blinded with two parallel groups, and a simple randomization with a 1:1 allocation ratio. Participants identified as having reduced working memory will be randomly assigned to one of two experimental groups: an active (or right-sided tDCS) group and a control (or SHAM) group. Individuals included in the active group will receive 20 min of tDCS to the right dorsolateral prefrontal cortex while completing the WM training, whereas participants in the control group will complete the WM training while receiving sham (e.g., placebo) stimulation. Participant blindness to group membership will be achieved by the use of a SHAM protocol, which simulates current stimulation.

Randomization and blinding
A computer programme will generate forty random flags identifying the patient’s group (20 flags equal to 1 for ACTIVE, 20 flags equal to 0 for SHAM). Flag order will then be shuffled and a participant’s unique ID (sub-01, sub-02, ..., sub-40) associated to a flag, following the order of attendance. The study design is double blind: participants and the assessors will not know until the end of the intervention if the participant had received active or sham brain stimulation. A member of the research team (not the assessor) will program two stimulation protocols (active and sham) on the stimulation device, which will be set to operate into blind mode. As such, the person who administers the stimulation will be blind to the details of the protocol. Meaning they will not be able to see the details of the protocol. Patients are then randomly assigned to one or the other protocol, resulting in an intervention blind both to the assessor and the patient.

Brain stimulation protocol
Brain stimulation will be administered using a Starstim 8 brain stimulation device (Neuroelectrics®). Standardized locations derived from electroencephalography will be used to place the electrodes over the right dorsolateral prefrontal cortex (DLPFC). We will use a bipolar setup, includes two circular Ag/AgCl electrodes (area = 3.14 cm$^2$) filled with conductive gel and placed on F4 (anode) and Fp1 (cathode). Reference electrodes will be attached to the earlobe and impedances will be measured throughout the stimulation and kept below 20 kΩ. We will use a total current intensity of 2 mA for 20 min, preceded by 30 s ramping up and followed by 30 s ramping down (total stimulation time = 21 s). With these parameters and Ag/AgCl electrodes (area of 3.14 cm$^2$) we obtain a current density of approximately 0.6 mA/cm$^2$, slightly higher than the one obtained with larger electrodes, but still well below the threshold for risk of tissue damage [57–59]. The simulated current distribution obtained with such setup is shown in Fig. 1. During sham stimulation, we will use the same setup as in the active condition but after ramping up, the current will be brought back to zero and the ramp up/down process repeated 30 s before the end of the 21 min time interval (total sham stimulation time = 21 min). We have recently completed data collection with the same protocol in 28 elderly and another study on stroke patients is undergoing: in these participants, brain stimulation has been well received.

Working memory training
The working memory training task is an internet-based visuo-spatial adaptive n-back training program. The same training paradigm has recently been successfully used with stroke patients [61]. Players must attend to a visual stream of spatial information that is presented sequentially and look for matches. Participants will see a grid with eight spaces, with one space filled in on each trial and they will press a button upon seeing a “match” in position with ‘n’ trials before. Within each session, players begin at a level of $n = 1$ and can progress up to...
n = 6 based on their scores in the previous block (i.e., this program is adaptive (closely following Jaeggi et al. [24]). Each session consists of 20 blocks of 20 + n trials each (~ 1 min per block), and progression to the next block is self-paced by the participant. This training task will last for ~ 20 min.

Participants’ timeline
The flowchart of the procedure is shown in Fig. 2, while a schedule of enrolment, interventions, and assessments is shown in Table 2. The care team as part of the research team identifies potential participants during the cognitive assessment, which is part of the routine care, and invites them to participate. Patients expressing an interest in the study are made aware of possible side effects of brain stimulation and asked to confirm their willingness to participate. During the first in-lab session (T0), following the informed consent procedure, each participant is randomly assigned to one of the two groups (active or sham stimulation) and interviewed by the assessor with a screening questionnaire, including questions about demographic information and health history to evaluate the inclusion/exclusion criteria. We also record information about current medications and information about time since injury, handedness and colour blindness. At this point, participants who don’t meet the eligibility criteria are excluded from further participation.

Baseline session After the informed consent process is carried out, eligible participants are administered the baseline measures (T0). The baseline data are included in data analyses as potential modifiers of performance. The baseline measures involve completing a series of questionnaires (Table 5). On the same day of baseline testing, patients are given a diary for them to keep. The diary contains information on how to setup and use the training program on their computer, education on how to do the working memory training, general troubleshooting information, and also a daily table to record information about the training session and any issue they might experience. The diary also contains contact details of the researcher administering the stimulation and information about future appointments. In addition, participants are asked to refrain from excessive alcohol or caffeine intake during the testing and to maintain good sleeping habits, where possible. In total, we expect the entirety of the first in-lab session (informed consent, screening, and baseline measures (T0)) to take about 1 h.

Outcome assessment (T1) On the following day participants perform a series of pre-training outcome tasks (T1; Table 4), administered by an assessor, to measure their baseline working memory capacity and to assess, during and at the end of the training intervention, the efficacy of the training and the stimulation regime. These tests will not be used to determine eligibility. The assessor is blind to group assignment (control vs active).

On the same day as T1 testing, the assessor will familiarize them with the stimulation procedure (which is identical for sham or active) and the training game. The member of the team administering the training
answers any questions about the study. Overall, this second session lasts for about 1 h.

**Training phase 1** On the 10 consecutive training days after the T1 session, depending on the group, participants will receive active or sham brain stimulation (see section 3.5.2). As the assessor, who administer brain stimulation, is blind to the intervention, the procedure will be the same for both groups. At the same time, both groups complete the working memory training exercise for 20 min. If the WM training finishes before the stimulation, participants are asked to wait until the end of the tDCS session (20 min). Before each training session, participants are also asked to answer short questions (level of alertness, engagement, etc.). Once the training session and the stimulation are complete, the participant fills in a feedback form on the side effects of brain stimulation experienced, if any.

Participants will complete 10 consecutive training sessions (2 weeks, excluding weekends. Each session should
take about 45 min (~ 10 min setting up of tDCS and ~ 20 min of WM training). This phase will be completed at home, with the assistance of the trainer. An attempt will be made to start the intervention on a Monday, therefore aiming to have breaks in the stimulation at the same point in the 2 weeks, and to keep the training time consistent over days. Nevertheless, administration time and date will be recorded. By the end of these 10 sessions, participants will be familiar with the training game and will be able to confidently undertake the training by themselves in the next 15 sessions.

Outcome assessment (T2) When the first 10 days of training are complete, the participant will undergo time 2 (T2) assessment. As before, testing involves completing a series of computerised cognitive tasks to measure training gains and transfer (see 3.5.6). This assessment will also include the tests from the WAIS-IV and WMS-IV (Table 3).

Training phase 2 Participants will then start the second training phase, involving 3 weeks of working memory training only (no brain stimulation). During this phase, patients will access the training program via internet at home on their own. Manualised weekly phone calls using a semi-structured script will be used to monitor the patients progress and to address any issue they might experience.

Outcome assessment (T3) When the second 3-week training phase is complete, participants will undergo time 3 (T3) assessment, during which they repeat the
same series of computerised cognitive tasks as in T2 (Table 4).

**Outcome assessment (T4)** A final follow-up assessment, identical to the one at T2 and T3, will be carried out 1 month after the completion of the intervention, to assess maintenance of working memory improvements and transfer.

**Cognitive tasks and questionnaires**
A series of cognitive tasks will be administered at different time points and for a different purpose during the study: screening, training, and outcome assessment.

- Screening: the Digit Span, Arithmetic, Symbol Span, and Spatial Addition subtests from the WAIS-IV and WMS-IV will be used for screening participants for cognitive impairment (see 3.5.4, 35 min to complete). A description of the task is given in Table 3.
  - Training Task: The visuo-spatial training task is a visuo-spatial adaptive n-back (see section 3.5.3).
  - Outcome: 1) the Dalhousie Computerized Attention Battery (DalCAB, [68], 35 min to complete) will be used as the outcome assessment battery. A description of the tasks in the battery is provided in Table 4. For a detailed description of the parameters used in each task see [68]; 2) In addition, a non-adaptive visuo-spatial N-back task will also be used as outcome measure, with fixed levels of N = 1, 2, 3 (~30 min to complete); 3) The Digit span, Arithmetic span, Symbol span and spatial addition will also be used as outcome measures.
- A series of questionnaires and feedback forms will also be administered at single or multiple time points throughout the intervention to record

### Table 3 Screening tasks

| Task                      | Description                                                                 |
|---------------------------|-----------------------------------------------------------------------------|
| Digit Span (WAIS-IV)      | The patient is read a sequence of digits, to repeat forwards, backwards or in ascending order of magnitude (sequence). |
| Arithmetic (WAIS-IV)      | The patient is read a number of mathematical problems increasing from very simple to complex in terms of the amount of information the person has to hold in mind. |
| Symbol Span (WMS-IV)      | Patients are asked to look at a series of symbols for 5 s then choose which symbols they saw from a multiple-choice format, pointing to them in the order they were shown from left to right. The test progresses from only one symbol with the sequence increasing depending on how well people do. |
| Spatial Addition (WMS-IV) | The patient is shown a grid for 5 s, then a second grid for 5 s. The grids contain blue and red circles. The patient is provided with a number of cards showing red, blue or white circles and asked to place them onto a cardboard grid according to the following rules: Place a blue circle in any place where you saw a blue circle on only one of the pages. Place a white circle in any place where you saw a blue circle in the same place on both pages. Ignore red circles. |

### Table 4 List of outcome measures

| TASK                              | DESCRIPTION                                                                 | Target functions          |
|-----------------------------------|-----------------------------------------------------------------------------|----------------------------|
| Simple Reaction Time              | Respond to each stimulus, with varying response-stimulus intervals.         | Vigilance                  |
| Go/No-Go                          | It employs a continuous stream of two different stimuli for which a binary decision must be made, such that one stimulus type requires a response (go) and the other stimulus type requires the participant to withhold a response (no-go). | Executive control          |
| 2-Choice Reaction Time            | Indicate the colour of each stimulus (2-choice responses; 50% each choice).  | Vigilance                  |
| Dual task                         | Complete the 2-choice reaction time task while silently counting the number of each colour of stimuli presented. Count probe for one colour at the end of each set. | Executive control          |
| Flanker                           | A central target stimulus is presented with flanking stimuli (flankers) on two sides that are either the same as (congruent) or different than (incongruent) the central target stimulus. The participant must decide and respond regarding a feature of the central stimulus (e.g., red or black) while ignoring/filtering the flanking stimuli. | Executive control          |
| Item Working memory               | Indicate whether a probe item was present or absent in a preceding study set of 2–6 items (50% present). | Working memory             |
| Location working memory           | Indicate whether a probe item was present or absent in the same position in a preceding study set of 2–6 locations (50% present). | Working memory             |
| Visual search                     | Locate and indicate orientation (upright vs. inverted; 50% each) of a target among different shape distractors that are a different colour (feature search) or the same colour (conjunction search) as the target. | Orienting and Selection    |
| Fixed visuo-spatial n-back task   | Participants must attend to a visual stream of information that is presented sequentially and look for matches, pressing a button upon seeing a “match”. Three levels of difficulty will be used (n = 1, 2, 3) | Working memory             |
variables that may impact or may be affected by performance on the task. Permission to use these instruments has been obtained for those questionnaires requiring it. A list of the questionnaires, together with a brief explanation and references is presented in Table 5.

**Outcome measures**

Our primary outcome measure will be the training gain, obtained by looking at the relative change in n-level (# items maintained in working memory) over the course of the training, as well as performance increments in the pre-/post-test outcome measures in the stimulation relative to the control (SHAM) group. Transfer of improvement to other cognitive domains will be assessed by comparing performance in a battery of cognitive tasks given at different times points throughout and after the intervention. Maintenance of improvement will be assessed by comparing outcome measures (Table 4) at different occasions during and after the conclusion of the study.

- **Training gain.** Performance in training is measured as the N level reached on every session. The training gain is defined as the difference in N levels at two time points. Accuracies and reaction times at trial level will also be collected. In addition, we will administer a non-adaptive visuo-spatial task (N = 1, N = 2, N = 3), to allow us to quantify improvements at different testing points in terms of accuracy and reaction times in the trained cognitive domain.

- **Transfer.** We will use spatial tasks that test the same cognitive process as the training task to evaluate transfer to closely related tasks. We will use the Dalhousie Computerized Attention Battery (DalCAB) [68], 8 cognitive tests to assess attentional functions in the vigilance, orienting and executive control attention network. Tasks are described in Table 4. Reaction time and accuracy will be computed for each task.

- **Maintenance.** To identify how long benefits of training and or stimulation are maintained in time, we will compare changes between outcome measures at T3 and at T4, 1 month after the end of the intervention.

- **Everyday life improvement.** It is important to understand whether improvement in cognitive tasks targeting specific processes can transfer to improvement in everyday life activities. We will use a self-reported questionnaire (PRECIS, Patchick et al., Clinical Rehabilitation, [66]) to evaluate if and how training in cognitive tasks targeting specific processes impacts on everyday activities.

**Data collection, management, analysis, and monitoring**

Accuracies and reaction times at trial level will be collected, alongside the ‘n’ level for each block. Data will continuously be monitored for completeness and consistency. Data entry will be double-checked, and data quality will be ensured based examination of ranges. Prior to statistical analysis, we will test the distribution of the scores. We will analyse the data using appropriate statistical methods, including mixed and repeated measures ANOVAs, with factors of group and test variables. Variables potentially modulating the outcome of the intervention (such as age or time since injury) will be considered as regressors in the analysis. Scores that are non-normally distributed will be analysed with non-

### Table 5 Questionnaires

| Questionnaire                                                                 | Taken at            |
|------------------------------------------------------------------------------|---------------------|
| **Individual lifestyle**                                                     |                     |
| Participant Health History (including details on injury, medications etc.)   | T1                  |
| Epworth Sleepiness Scale (to report habitual sleepiness [62])                |                     |
| Quality of life assessment (The Whoqol Group [63]), – in four facets (Physical health, Psychological, Social relationships, Environment) |                     |
| **Individual cognitive state**                                               |                     |
| Cognitive Failures Questionnaire (CFQ, [64]) to measure self-reported failures in perception, memory, and motor function. | T1                  |
| Hospital Anxiety and Depression Scale (HADS) to measure emotional symptoms (HADS; [65]) |                     |
| Patient Reported Evaluation of Cognitive Status (PRECIS [66])                | T1/T3/T4            |
| **Feedback on the of intervention**                                          |                     |
| Side effect of the tCS [57]                                                  | Every tCS session   |
| System usability scale (SUS [67])                                            | T3                  |
| Attitude towards stimulation (to record how they feel their attitude towards the brain stimulation affected performance) |                     |
| Feedback on strategy (to record whether they use a specific strategy to complete the training task) |                     |
| Blinding questionnaire (to confirm efficacy of the blinding procedure)       | T4                  |
parametric tests. Significant main effects and interac-
tions ($p < .05$) will be followed by post-hoc tests cor-
corrected for multiple comparisons. Dependent variables on
the computerized tasks will include mean reaction time
(RT) and accuracy (% correct). The research team will
analyse the data with a combination of SPSS, JASP, R,
Python, and MATLAB.

Risk assessment and monitoring
Common side effects in tDCS experiments are mild
headache, itching, fatigue [69]. All experimenters using
these techniques will undergo thorough training about
how to deal with and minimize these risks. Clinical
members of the research team will assess seriousness,
causality (relatedness), an expectedness of any adverse
event experienced by the participant with reference to
the most recent regulatory tDCS guidelines [57]. Data
on reported side effects will be reviewed following each
treatment session by clinicians treating and supporting
the participant during treatment. Following completion
of intervention, course reported side effects will be
reviewed by the PI and clinical co-PI. If none of the par-
ticipants recruited to the study can tolerate the data col-
clection or intervention, the trial will be stopped.

Dissemination policy
Results of this study may be submitted for publication in
a peer reviewed journal. On completion of the study, the
data will be analysed and tabulated, and a final study re-
port prepared and made available on ClinicalTrials.org.
the study has been registered on the database. Results
will be published in peer-reviewed journals and pre-
sented at relevant scientific meetings. Participants will
be asked whether they would like to receive a copy of
the final report, or a simplified version of it, and will be
provided with this if they wish. Dissemination will be
achieved through conference presentations, public talks
and community talks. Upon publication, the anonymised
dataset may be made available to other researchers upon
request.

Discussion
The overarching aim of the study is to assess if a specific
form of non-invasive brain stimulation, namely direct
current, has an impact on performance during a working
memory training regimen in patients with an acquired
brain injury. The novelty of our research resides in
the combination of brain stimulation with working memory
training: while the two interventions have been used be-
fore in clinical populations, to the best of our knowledge
this is the first trial looking at the combination of both
in acquired brain injury patients.

The focus of this study are chronic brain injury pa-
tients. These patients are no longer in a clinical setting
but have returned home and are seen by the Community
Brain Injury Service for community-based rehabilitation.
These aspects make the proposed intervention particu-
larly suitable for the population under study, as individ-
uals can undertake the long (5 weeks) training regimen
in the comfort of their home, at their own pace, minimiz-
ing the risk of drop-outs otherwise likely with long in-
terventions. Our intervention is suitable for home
healthcare, and, if proven successful, could represent a
valuable asset to the NHS to address brain injury related
cognitive impairment without putting excessive burden
on patients and carers, and the NHS. At the same time,
this trial will provide valuable knowledge to health au-
thorities and the research community on the potential of
non-invasive brain stimulation and training regimens in
addressing chronic cognitive impairment in clinical pop-
ulations. If successful, our study will provide evidence
that a restorative approach to cognitive rehabilitation is
possible, and an alternative to compensatory approaches
currently used.

By evaluating performance at 1 month after the com-
pletion of the training regimen, we will shed new light
on the long-term effects of cognitive training, also iden-
tifying potential modulatory effects of brain stimulation.
This finding will, in turn, inform future protocols, in
order to maximise improvement of cognitive functions
while minimizing the amount of training necessary. Ul-
tiltunately, patients would want these interventions to have
an impact on their everyday life: we therefore also ask
patients to self-evaluate (with the PRECIS questionnaire)
how the training regimen impacted their cognitive func-
tions and, indirectly, on their ability to cope with every-
day situations.

Finally, our intervention falls within the remit of home
healthcare medicine, with the potential to be fully self-
administered by the patient (with support of carer if re-
quired) in their home. We will therefore collect data on
the usability of the training, to inform future develop-
ment in line with homecare technologies.

Abbreviations
ABI: Acquired brain injury; tDCS: Transcranial direct current stimulation;
ADHD: Attention deficit hyperactivity disorder; NHFT: Northamptonshire
Healthcare Foundation NHS Trust; FSIQ: Full-Scale Intelligent Quotient;
WAIS: Wechsler Adult Intelligence Scale; WMS: Wechsler Memory Scale;
VWM: Visual Working Memory Index; DalCAB: Dalhousie Computerized
Attention Battery; DLFFC: Dorsolateral Pre-Frontal Cortex; CI: Chief
Investigator; REC: Research Ethics Committee; PRECIS: Patient Reported
Evaluation of Cognitive Status; HADS: Hospital Anxiety and Depression Scale;
CFQ: Cognitive Failure Questionnaire; SUS: System Usability Scale;
WHOQOL: World Health Organization quality of life assessment

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Authors’ contributions
All authors (SA, RH, GE, CG, MR, KS) were involved in developing the study design and methods. GE was responsible for designing and implementing the working memory training task. SA was responsible for writing the first draft of the manuscript. All authors read, critically revised, and approved the final manuscript.

Funding
No external funds will be used to carry out the study.

Availability of data and materials
Upon publication of the results, the datasets generated and/or analysed during the current study will be available from the corresponding author on reasonable request.

Ethics approval and consent to participate
This study protocol, informed consent forms and other relevant documents have obtained favourable opinion from the UK Health Research Authority (HRA) (ID: 238990) and by the Research Ethics Committee (REC) of the University of Birmingham (ID: RG_ 18–142), and all participants completed an informed consent process with a member of the research team. Written informed assent and consent in accordance with the Declaration of Helsinki will be obtained and the study will be conducted in accordance with the UK Policy Framework for Health and Social Care Research, the applicable UK Statutory Instruments, the General Data Protection Regulation (GDPR) / Data Protection Act 2018, and Guidelines for Good Clinical Practice (GCP). Amendments agreed between the CI and the sponsor, and approved UK Health Research Authority, will be communicated in writing to relevant stakeholders (e.g. REC, R&D). Participants may withdraw from the trial at any point without any penalty and will not receive compensation for taking part in the study. Personal information about participants collected during the consent/data collection processes are stored securely in a locked cabinet, in a locked office. Data from the software, stored electronically for the purpose of data analysis, will be password protected and participants will be identified by study code only. Data from the training task will be anonymous with only a subject code as identifier, encrypted via https during transit, and stored on a Dalhousie University-protected server. Confidentiality will be maintained by coding each participant’s data using an alphanumeric system with keys relating individuals to participant codes, kept in a separate locked place (off line) and destroyed after 1 year. Any reported adverse events and unintended effects will be discussed amongst the research team and dealt with on a case-by-case basis.

Consent for publication
Not applicable.

Competing interests
SA, KS, GE are named inventors on a patent application (1081656.PGBA) jointly submitted by University of Birmingham and Dalhousie University; RH, MR, CG have no competing interests to declare. The sponsor is not involved in study design, conduct, data analysis and interpretation, manuscript writing, and dissemination of results, and neither controls the final decision regarding any of these aspects of the study.

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