

How Does iReadMore Therapy Change the Reading Network of Patients with Central Alexia?

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Central alexia (CA) is an acquired reading disorder co-occurring with a generalized language deficit (aphasia). The roles of perilesional and ipsilesional tissue in recovery from poststroke aphasia are unclear. We investigated the impact of reading training (using iReadMore, a therapy app) on the connections within and between the right and left hemisphere of the reading network of patients with CA. In patients with pure alexia, iReadMore increased feedback from left inferior frontal gyrus (IFG) region to the left occipital (OCC) region. We aimed to identify whether iReadMore therapy was effective through a similar mechanism in patients with CA. Participants with chronic poststroke CA ($n = 23$) completed 35 h of iReadMore training over 4 weeks. Reading accuracy for trained and untrained words was assessed before and after therapy. The neural response to reading trained and untrained words in the left and right OCC, ventral occipitotemporal, and IFG regions was examined using event-related magnetoencephalography. The training-related modulation in effective connectivity between regions was modeled at the group level with dynamic causal modeling. iReadMore training improved participants' reading accuracy by an average of 8.4% (range, -2.77 to 31.66) while accuracy for untrained words was stable. Training increased regional sensitivity in bilateral frontal and occipital regions, and strengthened feedforward connections within the left hemisphere. Our data suggest that iReadMore training in these patients modulates lower-order visual representations, as opposed to higher-order, more abstract representations, to improve word-reading accuracy.

Key words: alexia; aphasia; DCM; MEG; reading; stroke

Significance Statement

This is the first study to conduct a network-level analysis of therapy effects in participants with poststroke central alexia. When patients trained with iReadMore (a multimodal, behavioral, mass practice, computer-based therapy), reading accuracy improved by an average 8.4% on trained items. A network analysis of the magnetoencephalography data associated with this improvement revealed an increase in regional sensitivity in bilateral frontal and occipital regions and strengthening of feedforward connections within the left hemisphere. This indicates that in patients with CA iReadMore engages lower-order, intact resources within the left hemisphere (posterior to their lesion locations) to improve word reading. This provides a foundation for future research to investigate reading network modulation in different CA subtypes, or for sentence-level therapy.

Introduction

Central alexia (CA; also known as alexia with agraphia; Dejerine, 1891) is a reading disorder that occurs within the context of a generalized language disorder (aphasia). Patients with CA find

reading slow and effortful and make frequent errors (Leff and Starrfelt, 2013). There is no agreed treatment for CA, and to date there have been no group-level investigations of how neural plasticity may support reading recovery in patients with CA. In the study by Woodhead et al. (2018), we demonstrated that a computerized word-reading therapy app improved word reading in 21 patients with CA. The aim of this cross-modal training was to coactivate orthographic, phonological, and semantic representations of the word to rebuild the neuronal connections between them. The present study aimed to improve our understanding of the therapeutic mechanisms in CA, with a view to developing stratified therapy pathways in the future.

Received May 29, 2018; revised March 7, 2019; accepted March 16, 2019.

Author contributions: W.P., J.T.C., A.P.L., and Z.V.J.W. designed research; S.J.K., O.M.A., and Z.V.J.W. performed research; S.J.K. and Z.V.J.W. analyzed data; S.J.K., Z.V.J.W., and A.P.L. wrote the paper.

This trial was supported by the Medical Research Council (Grant MR/K022563/1). The trial was registered at www.clinicaltrials.gov, reference NCT02062619. We thank Gareth Barnes for his guidance regarding the MEG study design and analysis for this project.

The authors declare no competing financial interests.

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<https://doi.org/10.1523/JNEUROSCI.1426-18.2019>

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After a left hemisphere stroke, the role of spared ipsilesional regions and right hemisphere homologs in supporting aphasia recovery are unclear (Adair et al., 2000; Tsapkini et al., 2011; Crinion and Leff, 2015; Hartwigsen and Saur, 2019). There is evidence for functional reorganization in spared left hemisphere regions (Jobard et al., 2003; Fridriksson, 2010; Abel et al., 2014, 2015; van Hees et al., 2014; Bonilha et al., 2016; Pillay et al., 2018), while other studies have identified right hemisphere homologs as fulfilling this function (Meinzer et al., 2006; Richter et al., 2008; Lee et al., 2017); both accounts may be correct and aphasia recovery may rely on a combination of mechanisms (Saur et al., 2006; Kurland et al., 2008; Turkeltaub et al., 2011; Crinion and Leff, 2015; Mohr et al., 2016). We modeled a bilateral reading network in patients with CA to ascertain the effects of therapy within and between the hemispheres.

While poststroke aphasia is the result of focal damage, it is increasingly viewed as a network disorder (Hartwigsen and Saur, 2019). Neuroimaging studies of skilled readers show that word reading activates a predominantly left-lateralized network of occipitotemporal, temporal, and inferior frontal areas (Heim et al., 2005; Graves et al., 2010; Price, 2012; Carreiras et al., 2014; Hoffman et al., 2015; Perrone-Bertolotti et al., 2017; Xu et al., 2017; Zhou and Shu, 2017). The local combination detector (LCD) model of visual word recognition suggests that because neurons are tuned to progressively larger fragments of a word as their location moves anteriorly, word reading is achieved primarily through feedforward processing along the visual ventral stream (Dehaene et al., 2005). However, an alternative account suggests that efficient word recognition relies on interactive feedforward (bottom-up) and feedback (top-down) processing within this network (Cornelissen et al., 2009; Wheat et al., 2010; Price and Devlin, 2011; Woodhead et al., 2014). Dynamic causal modeling (DCM) identifies the causal influence of one region upon another, allowing us to explore the interaction between top-down and bottom-up processes.

Within the domain of reading rehabilitation, in participants with pure alexia [PA; typically caused by left posterior cerebral artery (PCA) stroke], reading training was associated with stronger connectivity within the left hemisphere, and increased top-down connectivity from frontal to occipital (OCC) regions (Woodhead et al., 2013). This was interpreted as evidence that predictions from phonological and/or semantic representations in left frontal cortex facilitated visual word recognition after training. However, in CA [typically caused by left middle cerebral artery (MCA) stroke], these “central” language representations are damaged or disconnected.

As there is little in the existing literature to guide predictions of network reorganization following therapy in CA, we based our hypothesis on what is known about the reading network in healthy control subjects and pure alexia. The training used iReadMore, an adaptive word-reading training app, which improved word-reading ability for trained items in pure alexia (Woodhead et al., 2013) and CA (Woodhead et al., 2018). Using DCM of magnetoencephalography (MEG) data, we investigated how effective connectivity within the reading network changed as a result of therapy. Our speculative hypothesis was that training would strengthen feedback connections within the left hemisphere, and the self-connection of the left inferior frontal gyrus (IFG). It is anticipated that these analyses will yield predictions for future investigations of how neural network plasticity supports language recovery.

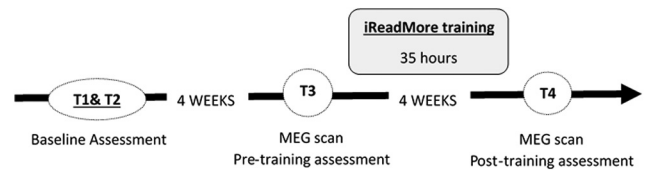


Figure 1. Study design. The baseline assessment took place over two testing sessions 1–2 weeks apart (T1 and T2). An MEG scan and behavioral assessment was conducted before (T3) and after (T4) a 4 week block of iReadMore training.

Materials and Methods

Study design

A within-subject, repeated-measures design was used. The data presented here were acquired during a larger crossover study that assessed the effects of iReadMore therapy and transcranial direct current stimulation (tDCS) on single-word reading (Woodhead et al., 2018). Participants completed an MEG scan before (T3) and after (T4) a 4 week reading therapy block (Fig. 1). Additionally, two baseline language assessments were conducted 4 weeks before training (T1 and T2) and at two time-points after training (T5 and T6).

During the therapy block, participants were asked to amass ~35 h of iReadMore training, through 40 min face-to-face sessions attended three times per week (Monday, Wednesday, and Friday; 11 sessions in total) supplemented with independent use at home.

The effect of tDCS was not analyzed in this article as (1) it was not designed to be tested using a between-subjects design, as would be required in the current analysis; and (2) the effect size of tDCS was small compared with the main effect of iReadMore.

Testing and face-to-face therapy sessions were conducted at the Institute of Cognitive Neuroscience, University College London.

Participants

Twenty-three participants with CA (15 males; mean age, 52 years; age range, 26–78 years; Table 1, demographic information), which was diagnosed by a neurologist or speech and language therapist, were recruited from either the PLORAS (Predicting Language Outcome and Recovery After Stroke) database of stroke patients held at The Wellcome Centre for Human Neuroimaging (Seghier et al., 2016), or speech and language therapy services at the National Hospital for Neurology and Neurosurgery, University College London Hospitals.

The following inclusion criteria were used: (1) left-hemisphere middle cerebral artery stroke with at least partial sparing of left IFG; (2) >12 months poststroke; (3) dominant English language use in activities of daily living; and (4) CA, operationalized as impaired word reading [Comprehensive Aphasia Test (CAT) word reading T-score, <61] and impaired spoken language (CAT naming, <63; picture description, <61). Screening and diagnoses were conducted historically in a clinical setting (data available on request from the authors), but additional baseline tests (as described by Woodhead et al., 2018) were performed at the start of the trial, including CAT Naming, nonword reading and word reading (Table 1).

Exclusion criteria included the following: (1) premorbid history of neurological or psychiatric illness; (2) history of developmental language disorder; (3) severe spoken output deficit and/or speech apraxia (CAT repetition, <44); (4) seizures in the past 12 months; (5) contraindications to MRI scanning; and (6) extensive damage to left IFG.

Participants were classified as having phonological ($n = 13$), deep ($n = 9$), or surface dyslexia ($n = 1$) according to the pattern of word and non-word-reading performance at baseline, using criteria described by Whitworth et al. (2016) (for further details, see Woodhead et al., 2018). The low proportion of patients with surface dyslexia is consistent with an opportunity sample of stroke patients described by Brookshire et al. (2014).

The participant information sheet was provided in written and auditory forms. All participants gave informed written consent in accordance with the Declaration of Helsinki. The Queen Square Research Ethics Committee approved this project.

Table 1. Demographic and clinical information on each patient

ID	Age (years)	Sex	Time post-stroke (months)	Lesion volume (cm ³)	CA subtype	CAT naming (%)	Pseudo-word reading (%)	Baseline word reading (%)	Reading change for trained items (%)
P01	44	Male	94	240.9	D	69	0	58.4	31.7
P02	50	Male	82	304.5	D	53	0	40.3	17.2
P03	64	Male	25	102.7	P	81	70	96.7	−2.8
P04	52	Male	66	122.7	P	66	0	71.1	18.9
P05	56	Female	93	149.8	S	5	75	63.8	8.3
P06	55	Female	75	151.2	P	93	30	91.9	3.9
P07	33	Female	59	181	P	95	2.5	90.1	2.8
P08	67	Male	107	11.7	D	72	2.5	12.5	12.5
P09	43	Female	55	399.2	D	81	0	58.2	11.7
P10	61	Male	19	195.6	D	40	0	3.4	5.0
P11	52	Male	12	31.2	P	88	75	96.3	3.9
P12	50	Female	14	59.4	P	83	25	90.6	2.2
P13	54	Male	24	149.3	P	86	65	91.5	4.4
P14	56	Male	23	45.1	P	72	0	80.3	3.3
P15	54	Male	39	189.7	P	14	2.5	47.3	6.1
P16	73	Male	158	205.2	D	71	0	20.0	5.8
P17	60	Male	16	102.6	D	33	10	28.1	10.0
P18	78	Male	22	128.5	P	43	7.5	75.4	2.2
P19	50	Female	72	141.3	P	28	5	35.9	5.0
P20	72	Male	101	243.3	D	9	0	13.4	5.8
P21	58	Female	41	297.7	P	81	0	59.5	16.1
P22	42	Male	13	43.7	P	72	27.5	74.9	12.2
P23	26	Female	81	161.9	D	79	0	75.5	6.7

Reading change (%) for trained items was calculated by subtracting pretraining (T3) word reading test accuracy (as a raw percentage) from post-training accuracy (T4) for trained words only. P, Phonological alexia; S, surface alexia; D, deep alexia.

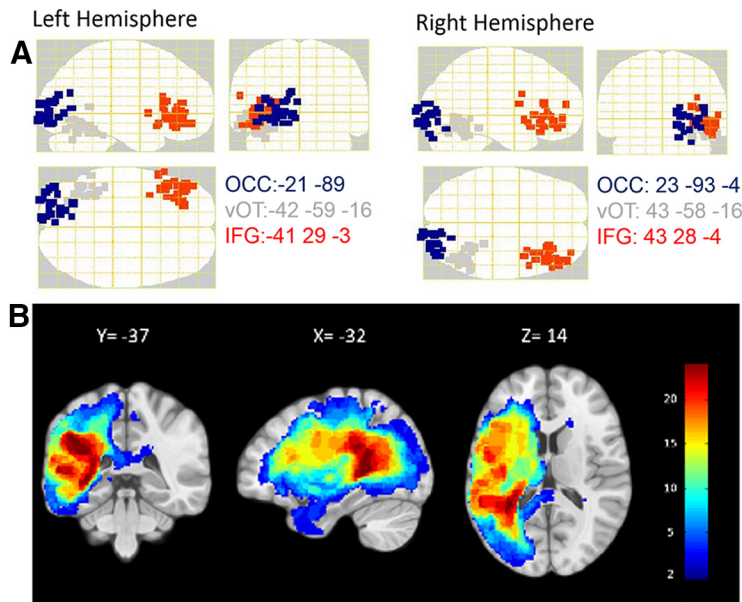


Figure 2. *A*, Optimal source locations identified using variational Bayesian equivalent current dipole modeling for each subject, plotted on a glass brain in Montreal Neurological Institute (MNI) space. Average dipole location across the group are given for the six sources: occipital (blue), ventral occipital temporal (gray), and inferior frontal gyrus (red). *B*, Lesion overlay map for the group ($n = 23$) where hotter colors indicate greater number of patients with lesions affecting that area.

Structural MRI

T1-weighted MRI scans were obtained in a 3.0 T whole-body MR system (Magnetom TIM Trio, Siemens) equipped with a standard 32 channel head coil radiofrequency (RF) receiver and RF body coil for transmission.

Data were preprocessed using Statistical Parametric Mapping 12 (SPM12; <http://www.fil.ion.ucl.ac.uk/spm/software/spm12/>) mounted in Matlab 2014b (MathWorks). Magnetic transfer (MT) maps were obtained for each participant using Voxel Based Quantification (VBQ)

toolbox in SPM12 (Weiskopf et al., 2013; Calhoun et al., 2014). The MT maps were spatially normalized into standard MNI space and segmented into tissue types (e.g., gray and white matter, CSF, atypical, or lesion). Lesions were identified using Automated Lesion Identification toolbox in SPM12 (Seghier et al., 2008). This compared a CA participant’s segmented MT maps to the MT maps of 29 healthy control subjects. A binary lesion image was created for each CA participant, upon which candidate dipole location solutions could be compared. Across our group of participants, lesion location was predominantly within the territory of the left middle cerebral artery, centered on the supramarginal gyrus (Fig. 2*B*).

iReadMore training

For a more detailed description of iReadMore training, see Woodhead et al., 2018. Briefly, iReadMore aims to retrain whole-word reading by repeatedly exposing the user to pairings of written and spoken words, and an associated picture. The aim of this cross-modal training is to coactivate orthographic, phonological, and semantic representations of the word to rebuild the neuronal connections between them. iReadMore was administered on a tablet computer. The software cycled through “training” and “challenge”

phases. During the training phase, participants were presented with 10 face-down cards. On selection, the reverse of the card revealed the written word, spoken word, and a picture of the word (all congruent with each other).

The challenge phase consisted of up to 30 trials. In each trial, a written and spoken word were presented simultaneously. In half the trials, the words were different (incongruent). Participants made the same/different judgements via a button press, and points were accrued for correct responses. If a criterion score was reached, they passed the level. The

algorithm within the iReadMore software adjusted task difficulty based on the user's performance. This modifies the following: (1) the similarity between the target spoken word and the written foils in the challenge phase (three levels); (2) the exposure duration of the written words (from a maximum of 4000 ms to a minimum of 100 ms); and (3) the criterion score required to pass a level.

Training stimuli

High-frequency words ($\text{SUBTLEX}_{\text{WF}} > 50$) of three to six letters were drawn from the SUBTLEX database (Brysbaert and New, 2009). Two matched lists of 180 words were created. For each word on the A list, there was a corresponding word on the B list matched for letter length, syllable length, written frequency, and imageability.

Over two baseline sessions (T1 and T2), CA participants completed an assessment of the entire word corpus whereby they read each word out aloud. Based on each participant's baseline performance (word-reading accuracy and speed), a customized set of 150 matched words from the A and B word lists were selected (for further details, please see Woodhead et al., 2018, their Supplementary Materials). One list was assigned to be trained, and the other to be untrained. These word lists were individualized for each patient. The aims of this word selection process were as follows: to have no significant difference in the patient's baseline reading ability (accuracy or reaction time) between the selected A and B words; to have no significant difference in psycholinguistic variables (length, frequency, imageability, regularity, or N-size) between the selected A and B words; and to have no significant difference in reading ability (accuracy or RT) between the selected word lists and the full list of words tested at baseline. The purpose of this latter aim was to avoid the possibility of regression to the mean, which would have been an issue if we had only selected words for therapy that the participants read poorly at baseline.

At the testing sessions immediately before and after therapy (T3 and T4), participants were tested on a subset of 90 words from each word list (trained items and untrained items; for further details, see Woodhead et al., 2018). Words were presented in a random order over three blocks. E-prime software (Schneider et al., 2002) was used to present words in the center of a screen in black, lower case, size 36 Arial font on a gray background. Participants were instructed to read the words aloud as quickly and accurately as they could into a voice-key microphone. Accuracy was coded on-line as follows: 1, correct response; 0.5, self corrected errors or verbal false starts; and 0, incorrect response. Responses >4 s poststimulus onset were coded as incorrect. Reaction times were excluded for the following: (1) voice-key failures; (2) incorrect and self-corrected responses; and (3) RTs >2 SDs from the subject's mean. To identify voice-key failures, a visual cue was displayed at the bottom left corner of the screen, which informed the experimenter when the microphone had been triggered. Before inputting the accuracy of the participant's response, the experimenter coded the validity of the voice-key trigger, as follows: 1, accurate; and 2, inaccurate voice-key trigger (e.g., if the participant said "erm" or a response was not detected by the microphone).

MEG scanning procedures

Scans were acquired using a VSM MegTech Omega 275 MEG scanner with 274 axial gradiometers in software third gradient-mode at a sampling rate of 480 Hz. Fiducial markers on the nasion and left and right preauricular points were used to determine head location in the scanner. Head movements were minimized by positioning the participant in a comfortable, well supported position and using padding around the participant's head. Recordings from fiducial markers indicated that the average head movement across a run was 9.14 mm (SD, 8.18 mm).

MEG experimental paradigm and stimuli

Participants were seated upright in the scanner. Trained words ($n = 150$), untrained words ($n = 150$), "false font" symbol strings ($n = 150$); described previously in Woodhead et al., 2013), and common proper names (e.g., "Jenny," "Bob"; $n = 40$) were projected onto the screen ~ 50 cm in front of the participant. Each stimulus was presented for 1000 ms followed by a crosshair for 2000 ms with a total interstimulus interval of 3000 ms. The stimuli were presented lower case Arial font of size 50 (Fig. 3). The stimuli types were evenly distributed in a pseudorandom order across four runs and presented using Cogent software (www.vislab.ucl.

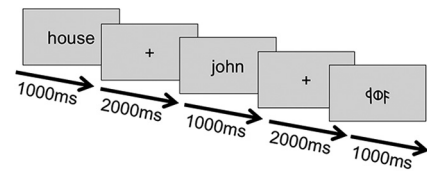


Figure 3. Stimulus presentation procedure for the MEG scans. Participants were scanned before and after training. At each session, there were 150 trials for each condition of interest (trained and untrained words), 150 trials for false fonts (omitted from this analysis) and 40 catch trials (names).

ac.uk/cogent.php). Participants were instructed to read the words silently. To ensure that participants attended to every trial, they were asked to respond via button press when they read a proper name. These catch trials were removed from the analysis. The false font condition was included to allow comparison with a dataset from healthy control participants, reported previously (Woodhead et al., 2014). The analysis of the false font trials is not reported in the current paper.

MEG preprocessing

The MEG data were preprocessed in SPM12 (<http://www.fil.ion.ucl.ac.uk/spm/software/spm12/>) using Matlab14a (<http://uk.mathworks.com/products/matlab/>). Preprocessing steps included the following: high-pass filtering at 1 Hz; removal of eye movement artifact using the Berg method (Berg and Scherg, 1994); epoching in the window -100 to 500 ms; low-pass filtering at 30 Hz; and merging the four runs. Artifact detection using a simple threshold at 2500 fT was applied, and channels with $>20\%$ of trials removed were rejected. This resulted in the removal of, on average, 40 trials (range, 0–260 trials) for each participant (of a total of 600 trials) and a total of 10 instances where channels were removed. Robust averaging across trials was conducted, and a 30 Hz low-pass filter was applied. Data from the two time-points were merged, and a single shell Boundary Element Method forward model was applied.

Source localization

Dipolar source location was performed with variational Bayes equivalent current dipole modeling (VB-ECD; Kiebel et al., 2008a), which uses a nonlinear optimization algorithm to simultaneously fit a number of dipoles with different prior distributions on their locations and moments, at a single time-point. For each participant, the M170 peak was identified in a semiautomated fashion using the average power of all trained and untrained word trials, in a time window of 0–300 ms. The sensor data at the subject-specifically identified M170 peak was used for the VB-ECD dipole modeling. The M170 peak was reliably present in all subjects and is known to represent orthographic processing (Tarkiainen et al., 1999; Marinkovic et al., 2003; Rossion et al., 2003; Pykkänen and McElree, 2007; Vartiainen et al., 2009; Zweig and Pykkänen, 2009).

The Bayesian algorithm requires the specification of a prior mean and variance for the location and moment of each dipole. The location priors were the same as those reported by Woodhead et al., 2014, who demonstrated that a six source model consisting of the left and right OCC regions (MNI coordinates: $\pm 15, -95, 2$), ventral occipital temporal (vOT) regions (MNI coordinates: $\pm 44, -58, -15$) and IFG (MNI coordinates: $\pm 48, 28, 0$) best fit the M170 peak for word reading in healthy control subjects.

Source solutions were free to move to any location. Therefore, the following restrictions were placed on the VB-ECD outputs: source locations must be (1) within the anatomically defined regions of interest, (2) >2 cm from adjacent sources, and (3) outside of the lesion. The solution with the greatest negative free energy (i.e., that best fitted the data) that met the above criteria was selected to be used in the DCM estimations.

Dynamic causal modeling

We used DCM to investigate the effective connectivity between neuronal sources within the reading network and how connection strengths were modulated in response to iReadMore therapy. For a detailed description of the methodology of DCM, the reader is directed to previous studies

(David et al., 2005; Kiebel et al., 2006, 2007, 2008b; Garrido et al., 2007; Reato et al., 2013).

Essentially, DCM uses a biologically informed neural mass model that uses the characteristic response rates and patterns of connectivity (Fellman and Van Essen, 1991) of three neuronal subpopulations (pyramidal cells, spiny stellate cells, and inhibitory interneurons) within the layers of the cortical column (Jansen and Rit, 1995) to model the connections between different sources. For example, forward connections innervate spiny stellate cells in the granular layer which results in an excitatory effect, backward connections synapse pyramidal cells and inhibitory interneurons in the supragranular and infragranular layers and hence can be excitatory or inhibitory, lateral connections can innervate all three layers of the cortical column and thus can also have an inhibitory or excitatory influence on the target region.

Self-connections are also modeled within the DCM. These quantify the maximal amplitude of the postsynaptic response in each cell population in that region (Kiebel et al., 2007). These maximal responses are modulated by gain parameters. Gain parameters greater than one increase the maximal response that can be elicited from a neuronal region. As such, the gain parameters are a measure of the sensitivity of a region to an input.

iReadMore training improved participants' word-reading accuracy for trained items only. The aim of the DCM analysis was to identify connection strengths that were significantly modulated by iReadMore training for these trained words, over and above any test–retest effects observed for untrained items. The data used for the DCM analysis were the evoked responses to trained and untrained words presented before and after therapy (Tr_Before; Un_Before; Tr_After; Un_After). We were interested in how therapy affected the early stages of word processing, so activity in the 0–300 ms time window was modeled. The sensory inputs to the model were specified as entering the left and right OCC. The A matrix modeled the connection strengths for the Tr_Before trials. Two B matrices modeled how connection strengths were modulated by therapy. The first (Matrix B1) estimated the modulation for trained words over time (Tr_Before vs Tr_After). To ensure the modulation observed in Matrix B1 did not represent a simple effect of time, rather than training per se, Matrix B2 modeled modulation for untrained items after therapy versus to-be-trained items before therapy (Tr_Before vs Un_After). It is worth noting that an alternative analysis could be to compare Un_Before vs Un_After for the B2 matrix, as this would have meant that both B1 and B2 would have compared the same items before versus after training. However, this mismatch of items in B2 is unlikely to have made a significant impact on the results because before training all items were novel and each patient's to-be-trained and never-trained word lists were matched for baseline performance and psycholinguistic properties.

Similar to other studies (Woodhead et al., 2013, 2014), and to reduce the model space to a manageable computational level, we placed the following constraints on how network connections varied among models, as follows: (1) lateral connections were only allowed within the same level of the cortical hierarchy (i.e., left OCC to right OCC) and not between levels (e.g., left OCC to right vOT); (2) lateral connections were reciprocal (e.g., a connection from the left vOT to right vOT was mirrored by a connection from the right vOT to the left vOT); and (3) forward and backward connections were symmetrical between hemispheres. This resulted in nine independently varying connections, leading to 512 models (2^9) per subject, all of which were fitted to their individual MEG data.

Bayesian model averaging

Random-effects Bayesian model averaging (BMA; Penny et al., 2010) was used to identify the average change in each connection strength across all models and all participants. BMA considers the entire model space and computes weighted averages according to the posterior probability for each model.

Experimental design and statistical analysis

Word-reading test analysis. Change in word-reading accuracy and RT were calculated over the baseline period and training block for each word list. Change was simply calculated as the difference from one time-point

to the next. Repeated-measures ANOVAs were calculated with within-subject factors of block [pretraining (T3-Baseline) vs training (T4–T3)] and Word-List (Untrained vs Trained).

MEG analysis: group-level effects of iReadMore therapy on the reading network. The DCM analysis identified the training-related modulation in effective connectivity between regions at the group level. We defined whether connections showed training-related modulation according to the following two criteria: (1) there was significant modulation in Matrix B1 (Tr_Before vs Tr_After); and (2) the therapy-specific modulation in Matrix B1 was significantly different to the nonspecific change over time in Matrix B2 (Tr_Before vs Un_After).

For the first criterion, a nonparametric proportion test was used for each connection to test whether modulation in Matrix B1 (Tr_Before vs Tr_After) was significant. A Gaussian distribution based on the posterior mean and SD was generated for each connection from which 10,000 samples were obtained. A connection was deemed to be significantly stronger after therapy if >90% of samples were >1; and significantly weaker if >90% of samples were <1 (Richardson et al., 2011; Seghier, 2013; Woodhead et al., 2013).

To identify therapy-specific training effects, rather than a simple effect of time, a second analysis was performed to compare the B1 and B2 matrices. The B1 matrix provides the modulation of connections for training over time (Tr_Before vs Tr_After), whereas the B2 matrix encapsulates the main effect of time in the absence of any training (Tr_Before vs Un_After). If the experiment only induced a simple effect of time, the modulation of the two B matrices would be very similar, and not significantly different from each other. If, on the other hand, there was an additional effect of therapy over time, we would expect the modulation in the two B matrices to be different. Using a fixed-effect within-subject Bayesian model comparison, we compared the following two models: (1) Matrix B1 \neq Matrix B2; and (2) Matrix B1 = Matrix B2. Log Bayes factors of >3 indicate that connections in B1 were significantly different to those in B2 (i.e., the effect of therapy could not be simply explained as an effect of time). If both criteria are satisfied, then the connection is significantly modulated by reading therapy (criterion 1) and is not simply explained as an effect of time (criterion 2).

Results

Training effects on reading ability

Participants completed on average 33.35 h (SD = 2.65 h; range, 25.33–37.21 h) of iReadMore therapy over the training period.

A repeated-measures ANOVA revealed a significant block by word-list interaction for word-reading accuracy ($F_{(1,22)} = 11.869, p = 0.00231$; Fig. 4). Paired *t* tests showed that the change in accuracy for trained words was significantly greater during the training block compared with the pretraining block ($t_{(22)} = -3.11, p = 0.010$), and change over the training block was significantly greater for trained words compared with untrained words ($t_{(22)} = 5.89, p = 0.001$). Change in accuracy for untrained items was not significantly different between blocks ($t_{(22)} = 1.479, p = 0.153$). This indicates that therapy significantly improved word-reading accuracy for trained words only. Word-reading accuracy improved by on average 8.4% (SD = 7.36; range, -2.77% to 31.66%) for trained words compared with -0.11% (SD = 5.39; range, -13.33% to 8.36%) for untrained words. A repeated-measures ANOVA of word-reading reaction time data revealed no significant block by word list interaction ($F_{(1,21)} = 0.461, p = 0.505$) and no main effect of block ($F_{(1,21)} = 2.983, p = 0.099$) or word list $F_{(1,21)} = 0.066, p = 0.800$).

MEG scanner task results

Participants successfully completed the within-scanner name detection task. The average accuracy for name trials was 89.71% (SD = 16.01), and the average percentage of false alarms (where the button was pressed for a trial other than a name) were 3.91% (SD = 6.06).

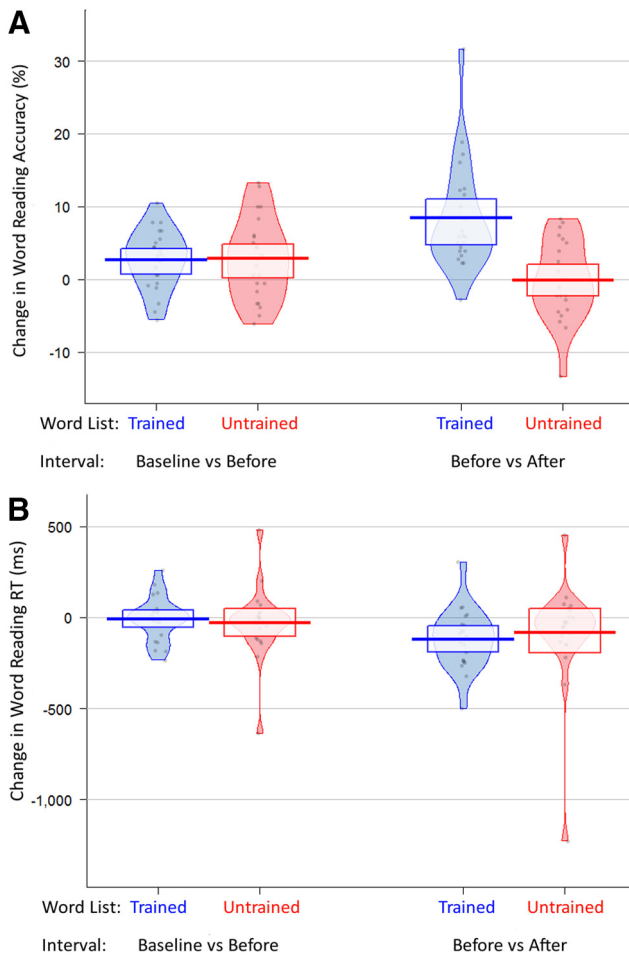


Figure 4. *A, B*, Change over time in mean word-reading accuracy (*A*; $n = 23$) and reaction times (*B*; $n = 22$) for trained words (blue) and untrained words (red). Error bars indicate 95% confidence intervals. In these pirate plots, each gray point represents a participant, the solid bars show the means and the shaded boxes indicate the 95% confidence intervals.

Cardiac artifacts

In response to a reviewer's comment, we tested whether cardiac artifacts could be confounding our results by carrying out a *post hoc* independent component analysis on the raw MEG data. A heartbeat artifact component was identifiable in $n = 18$ of 23 participants. This component was epoched according to trial onset times for the four main conditions. The "cardiac effective refractory period" data were averaged into 10 ms time bins over the 0–300 ms time window (giving 30 time bins). A 2×2 repeated-measures ANOVA at each time-point with factors time (before vs after training) and wordlist (trained vs untrained words) revealed no significant main effect of either time or wordlist in any of these 30 time bins.

Cardiac artifacts may have also added unsystematic noise to the data. This noise was, however, not related to the trial type or time from trial onset. All DCM analyses were based on averaged data (typically 150 trials), which would have significantly attenuated this confound. Additionally, we used a robust averaging procedure, which uses an iterative process to place weights on within-trial samples of data based on the degree of artifact present within the trial (Leski, 2002; Litvak et al., 2011). When the data are averaged across trials, these weights serve to down-weight outliers.

We conclude that any cardiac artifacts were unlikely to have influenced our DCM results, due to their random occurrence with respect to both stimulus onset and stimulus type allied with

Table 2. Results of the DCM analysis (group-level effects of iReadMore therapy on the reading network)

Connection	Posterior mean	Exceedance probability
Stronger with training		
LOCC to LOCC	1.02	1.00
LOCC to LvOT	1.17	1.00
LOCC to LIFG	1.16	1.00
ROCC to LOCC	1.07	0.97
ROCC to ROCC	1.07	1.00
LIFG to LIFG	1.10	1.00
RIFG to LIFG	1.08	0.96
RIFG to RIFG	1.03	0.99
Weaker with training		
LOCC to ROCC	0.86	0.00
LvOT to LOCC	0.92	0.01
RvOT to RvOT	0.97	0.01
LIFG to LvOT	0.80	0.00
RIFG to RvOT	0.91	0.00

Posterior means and exceedance probabilities from Matrix B1 (Tr_Before vs Tr_After) for the 13 connections that were shown to be significantly modulated by iReadMore therapy. L/ROCC, Left/right occipital; L/RvOT, left/right ventral occipitotemporal cortex; LIFG, left IFG; RIFG, right IFG.

the use of robust averaging to minimize any effect that they may have had on the data.

Source localization

The average latency of the M170 peak was 189.71 ms (latency range, 156.67–215.00 ms), and the average peak amplitude was 37.15 fT (amplitude range, 14.46–63.8 fT). To show that the M170 peak is related to orthographic processing a correlation was performed between baseline word-reading accuracy and M170 latency and amplitude. This revealed a significant negative correlation ($r = -0.550$, $p = 0.007$), indicating that those patients with greater word-reading accuracy had earlier M170 peaks (Fig. 2*A*, each participant's dipole location plotted on a glass brain).

MEG analysis: group-level effects of iReadMore therapy on the reading network

Table 2 displays the posterior mean and exceedance probability for connections that showed significant therapy effects (i.e., that were significantly modulated in Matrix B1; Tr_Before vs Tr_After), but this modulation was significantly different to that in Matrix B2 (Un_Before vs Tr_After). Eight connections were significantly stronger after therapy than before, and five were significantly weaker (Fig. 5).

Stronger connections for trained words after therapy

Of the eight connections significantly strengthened by iReadMore training, two were feedforward connections in the left hemisphere, two were lateral (between hemisphere) connections from right to left, and four were self-connections. More specifically, they were as follows: the feedforward connections from left OCC to left IFG and left vOT; the lateral connections between the OCCs and IFGs in the right to left direction; the self-connections in left and right OCCs and IFGs (bottom and top of the reading hierarchy, respectively). Self-connections indicate the sensitivity of a region to an input; indicating that these regions became more sensitive to trained words with therapy.

Weaker connections for trained words after therapy

Of the five connections significantly weakened by iReadMore training, three were feedback connections, two were lateral connections, and one was a self-connection. More specifically, they

Group-level effects of iReadMore therapy on the reading network

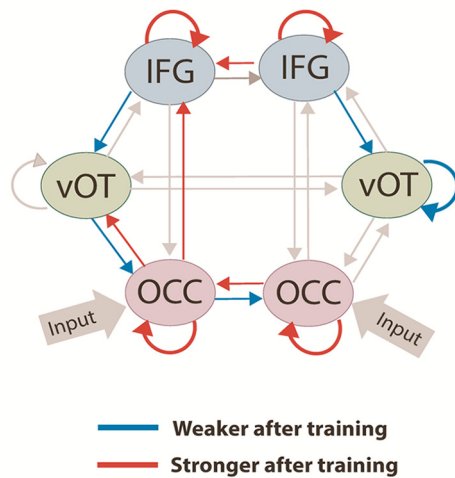


Figure 5. Results of the DCM analysis: modulated connection strengths for words trained with iReadMore after training. These are connections that met the following criteria: (1) there was significant modulation in Matrix B1 (Tr_Before vs Tr_After); and (2) the therapy-specific modulation in Matrix B1 was significantly different to the nonspecific change over time in Matrix B2 (Tr_Before vs Un_After). Connections in red became significantly stronger after training, whereas connections in blue became significantly weaker after training.

were as follows: the feedback connections from both IFGs to both vOTs and from left vOT to left OCC; the lateral connection between the OCCs in the left to right direction; the self-connection on the right vOT.

Discussion

Our analysis explored training-induced connectivity modulation within the reading network of stroke patients with CA at the group level. We observed changes distributed across the reading network. We identified increased regional sensitivity to trained words (changes in the self-connections of a region) bilaterally at the top (frontal regions) and bottom (occipital regions) of the reading network. As expected, this included the left IFG. The between-region connections modified by therapy were predominantly in the left hemisphere or, when interhemispheric, were from right to left. Contrary to our predictions, stronger connections were observed in a feedforward direction from left OCC to vOT and from left vOT to IFG. Together, these findings indicate that iReadMore training predominantly alters left hemisphere connectivity and increases the influence of bottom-up processes.

The therapy-induced inter-regional modulation of connectivity was predominantly in a feedforward direction. Stronger connections were observed between the left OCC and left IFG and left OCC and left vOT. These connections were also stronger for words compared with false fonts in the first 300 ms of reading in a group of healthy control participants (Woodhead et al., 2014). According to the LCD model (Dehaene et al., 2005; Dehaene and Cohen, 2011), neurons are tuned to progressively larger fragments of the word as their location moves along the ventral pathway. It is possible that mass exposure to the orthographic stimuli enhanced the processing of word forms within the ventral reading route. These results, when viewed with the reduced strength of feedback connections from the left IFG to left vOT and from left vOT to left OCC, suggests that iReadMore training in these patients modulates lower-order visual representations, as opposed to higher-order, more abstract ones to improve word-reading accuracy.

This finding is in contrast to patients with PA, where iReadMore training effects were driven by increased feedback from the left IFG to left OCC (Woodhead et al., 2013). It was suggested that improved predictions from the phonological and semantic representations within the IFG constrained the visual processing of trained words. This discrepancy may reflect differences in the lesion location in the two groups; with damage to the PCA territory in patients with PA and the MCA territory in patients with CA (Fig. 2B). In response to therapy, each group may have maximized their available intact resources. Therapy effects in patients with PA are likely to rely on improving feedback support from the intact phonological and semantic representation of words within their left IFG as damage affects input to the reading network. Increased IFG involvement has been identified for tasks demanding subordinate levels of semantic knowledge (Nagel et al., 2008; Whitney et al., 2011) and tasks relating to phonology (Devlin et al., 2003; Drakesmith et al., 2015). By contrast, patients with CA have damage to the central phonological and/or semantic representations (Crisp and Lambon Ralph, 2006; Robson et al., 2012; Hoffman et al., 2015) or connections to them. Therefore, therapy may increase reliance on orthographic processing to drive rebuilding or reconnecting of the phonological and/or semantic representations in a feedforward manner.

Increases in self-connection strengths were observed in the left and right OCCs and IFGs. In DCM, self-connections act as a gain control (Kiebel et al., 2007). The left IFG has been implicated in the early stages of visual word recognition (Cornelissen et al., 2009; Wheat et al., 2010; Woodhead et al., 2014) and was modulated by iReadMore therapy in patients with PA (Woodhead et al., 2013); however, we did not expect the self-connection of the right IFG in our CA patients to also become stronger. Support from the right IFG in language tasks has been reported in aphasia rehabilitation research (Crinion and Price, 2005; Naeser et al., 2011; Turkeltaub et al., 2012; Mohr et al., 2016; Nardo et al., 2017). However, it has been argued that this strategy may be ineffective compared with using perilesional left hemisphere regions (Heiss and Thiel, 2006). The stronger self-connections in both IFGs may reflect the differences in patients' progress with training. In a participant with phonological dyslexia, increased right IFG activity was observed immediately following training. However, when training continued on words read correctly immediately post-therapy, increased activation was observed in left hemisphere perilesional regions (Kurland et al., 2008). It has been suggested that the right IFG has a role in assisting with error monitoring and attention control (Hampshire et al., 2010). The increased connection strength from right IFG to left IFG may suggest that the right IFG has a different role in word reading, potentially related to error monitoring, which will have also been modulated by iReadMore.

Within the right hemisphere, the connection from right IFG to right vOT became weaker with training, as did the right vOT self-connection. This further suggests a reduced role of the right hemisphere in reading after iReadMore training.

iReadMore was designed to retrain word reading across all subtypes of CA through repeated activation of the semantic, phonological, and orthographic representations of trained words (Woodhead et al., 2018). Retraining in this omnibus manner potentially strengthened the mappings between differing cortical representations of words. It should be noted that almost all participants were classified as having either phonological or deep dyslexia (indicating a deficit in the phonological domain or sublexical reading route), which may limit our interpretations to this patient group. However, in practice we observe that few patients

have “pure” deficits of one type or another (Leff and Starrfelt, 2013), and it is an open question to what extent reading rehabilitation targets one reading route over the other. In line with previous research (Abel et al., 2015; Rueckl et al., 2015), our study suggests that therapeutic effects play out among both surviving left and right hemisphere regions, albeit with a leftward bias.

The following connections became stronger with training: (1) the right OCC self-connection; and (2) the connection from right to left OCC. This may reflect selective tuning of visual cortex to the orthographic information in trained words induced by multiple, repetitive exposure with trial-by-trial feedback. According to the split fovea theory, visual information from the front of a word is received by the right OCC as the optimal viewing position is usually just to the left of center of any given word (Nazir et al., 1992). Acceptable dipole locations were not restricted to V1 so extra-striate regions will almost certainly have contributed to the observed effects. As hemifield integration occurs above the level of V1, the changes in the right OCC self-connection, and interhemispheric connection to left OCC suggests increased sensitivity to the front part (left of fixation) of trained words (Perea and Lupker, 2003). This is consistent with the LCD reading model (Dehaene et al., 2001; Cohen et al., 2002; Perea and Lupker, 2003).

In summary, in a group of patients with CA (mainly with either phonological or deep dyslexia), improved word reading after iReadMore training was associated with distributed changes across the residual reading network. We identified a mixture of the following: (1) within-hemisphere connections (mainly left-lateralized and feedforward) that were strengthened by therapy; (2) bihemispheric connections (particularly self-connections at both the top and bottom of the reading hierarchy); and (3) between hemisphere connections (right to left pattern). The iReadMore therapy app will be available to the public in 2019 (<http://www.ucl.ac.uk/aphasialab/apps/ireadmore.html>).

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