Electroconvulsive therapy, electric field, neuroplasticity, and clinical outcomes

Zhi-De Deng1,2, Miklos Argyelan3,4,5, Jeremy Miller6, Davin K. Quinn6, Megan Lloyd6, Thomas R. Jones6, Joel Upston6, Erik Erhardt7, Shawn M. McClintock2,8 and Christopher C. Abbott8

© The Author(s) 2021

Molecular Psychiatry (2022) 27:1676–1682; https://doi.org/10.1038/s41380-021-01380-y

INTRODUCTION
Electroconvulsive therapy (ECT) remains the gold-standard treatment for patients with depressive episodes [1]. Independent of the antidepressant effect of ECT, many patients experience transient but debilitating cognitive side effects such as attention and memory impairment [2, 3]. ECT-mediated hippocampal neuroplasticity has been implicated in both antidepressant and cognitive outcomes [4, 5]. Neuroplasticity refers to the brain’s ability to restructure itself by forming new neural connections [6] and appears to be a common mechanism shared by both ECT and chemical antidepressant treatments [7]. Despite the possible relationship between hippocampal neuroplasticity and therapeutic mechanisms (neurogenesis, synaptogenesis, gliogenesis, angiogenesis), the relationship between hippocampal neuroplasticity and antidepressant response has been inconclusive. Some investigations demonstrated a direct relationship between hippocampal neuroplasticity and antidepressant response [8–13], but other investigations were negative [14–21]. The rapid hippocampal volume increase involves extensive remodeling and may also be related to procedure-related cognitive impairment [22, 23]. Consistent with this mechanism, recent ECT-imaging investigations have demonstrated a direct relationship between hippocampal neuroplasticity and impaired cognitive performance [5, 24].

Right unilateral electrode placement predominately stimulates the right hemisphere in an effort to reduce cognitive side effects [3]. The right hemisphere stimulation results in greater right hippocampal neuroplasticity that disentangles the effect from the generalized seizure [8, 10, 13, 21, 25–27]. Recent work on electric field (E-field) modeling has explored the relationship between electric field strength and neuroplasticity. Concerning ECT parameters, the electrode placement determines the geometric shape and the amplitude determines the E-field magnitude within this geometric shape [28]. Spherical head models have demonstrated that the E-field is influenced by individual anatomic variability (skull thickness, head diameter, and brain volume) [29]. The application of E-field modeling to ECT investigations has demonstrated a direct relationship between E-field and neuroplasticity, [30] but the relationship between E-field and antidepressant outcome has been mixed [30, 31]. To date, there has...
been no published examination of the E-field strength and longitudinal cognitive performance during an ECT series.

In standard ECT clinical practice, pulse amplitude tends to be fixed at 800 or 900 milliamperes (mA) without clinical or scientific rationale [28]. Lower pulse amplitudes reduce the magnitude of the induced E-field which could potentially decrease the risk of cognitive side effects. Case reports and recent clinical trials have demonstrated that 500–600 mA is sufficient to generate seizure activity with fewer cognitive side effects [32–35]. We recently completed a clinical trial of adults and older adults with depression who were randomized to right unilateral electrode placement ECT treatment with different pulse amplitude (600, 700, and 800 mA) with a bitemporal (800 mA) contingency [36]. The results of this investigation demonstrated a trade-off between cognitive safety (improved with 600 mA arm) and antidepressant response (improved with 700 and 800 mA arms). The cognitive measure most associated with amplitude mediated cognitive impairment was the Delis Kaplan Executive Function System (DKEFS) Verbal Fluency Test, which assesses frontal-temporal cognitive functions (e.g., verbal fluency, cognitive flexibility).

Based on the variability from fixed extracranial amplitude to E-field strength, we shift the focus from extracranial amplitude to hippocampal E-field strength. Here, we aim to determine the relationships between hippocampal E-field strength, hippocampal neuroplasticity, and antidepressant and cognitive outcomes. We hypothesize that E-field mediated hippocampal neuroplasticity will be associated with both antidepressant and cognitive outcomes as measured by the Hamilton Depression Rating Scale-24 item (HDRS24) and DKEFS Verbal Fluency Test, respectively. Recognizing that hippocampal neuroplasticity may be related to both antidepressant and cognitive outcomes, we further hypothesize that an optimal E-field ECT may result in sufficient hippocampal neuroplasticity for antidepressant response with cognitive safety.

METHODS
Participants, assessments, ECT, and study design
The overall study design and clinical outcomes have been previously described (ClinicalTrials.gov Identifier: NCT02999269) [36]. The University of New Mexico Human Research Protections Office approved this investigation. All subjects provided written informed consent to the research protocol and study participation. Subjects had a diagnosis of major depressive disorder (MDD; single episode or recurrent, severe with or without psychotic features), met clinical indication for ECT, were right-handed, and the age ranged between 50 and 80 years. Scheduled medication were tapered before the initiation of ECT, but as-needed medications including quetiapine (maximal cumulative dose per day: 200 mg), trazodone (200 mg), and lorazepam (3 mg) were permitted.

All subjects started the ECT series with right unilateral (RUL) electrode placement [37] and were randomized to 600, 700, and 800 mA treatment arms. Subjects commenced ECT with ultrabrief (0.3 milliseconds (ms)) pulse width until a planned interim data analysis. Because of a trend of reduced antidepressant outcomes in the 600 mA arm, subjects (n = 15) enrolled in the latter portion of this investigation received brief (1.0 ms) pulse width. The rationale of the change in pulse width from 0.3 to 1.0 ms was to improve the antidepressant outcomes in the lower amplitude arms based on the strength-duration curve (lower amplitudes may require longer pulse width). The first ECT session determined individual seizure thresholds with subsequent treatments provided at six times the seizure threshold with similar adjustments to pulse train duration and frequency across all amplitude arms [38]. Subjects received clinical, cognitive, and imaging assessments pre- (V1), mid- (after the sixth ECT treatment, V2), and post-ECT (within one week of finishing the ECT series, V3). The Hamilton Depression Rating Scale-24 item (HDRS24) was the primary antidepressant outcome measure [39]. If subjects were non-responsive to the assigned pulse amplitude (<25% reduction in from baseline HDRS24 at the second visit, n = 20), subjects then received bitemporal (BT) electrode placement (800 mA, 1.0 ms pulse width) for the remainder of the ECT series (Fig. 1) [40]. The cognitive battery included a comprehensive assessment of multiple cognitive domains. Here, we focus on the results from the DKEFS...
**Statistical analyses**

**Amplitude and hippocampal E-field.** With RUL electrode placement and the initially assigned amplitude (600, 700, or 800 mA), we assessed hippocampal E-field variability across amplitudes in the right and left hippocampi with a one-way analysis of variance and follow-up contrasts to determine amplitude differences with $E_{\text{hippo}}$.

**Hippocampal E-field and volume change.** We assessed the relationship between right hippocampal E-field strength ($E_{\text{hippo}}$) and right hippocampal volume change (percent change relative to pre-ECT HDRS), $(\Delta\text{Vol}_{\text{hippo}}/\text{Pre-ECT-Vol}_{\text{hippo}})$ with a linear regression analysis controlling for sex, age, pulse width and number of treatments. We calculated effect sizes (partial eta squared) for dependent variables of interest for all linear models.

Hippocampal E-field, volume change, and clinical outcomes. We assessed the relationships with $E_{\text{hippo}}$, right hippocampal volume change, and antidepressant outcomes (percent change in HDRS relative to pre-ECT HDRS or %ΔHDRS) with similar linear models controlling for sex, age, pulse width and treatment number. In parallel, we assessed the relationships with $E_{\text{hippo}}$, right hippocampal volume change, and cognitive outcomes (change in DKEFS Letter Fluency or DKEFS Letter Fluency controlling for sex, pulse width, treatment number, and Test of Premorbid Functioning (TOPF) to control for premorbid intelligence [51]). Electrode placement influences hippocampal E-field strength through E-field geometry and was not included as a covariate. Age was accounted for in the DKEFS Letter and Category Fluency demographic-adjusted scaled score and therefore not included as a covariate in the cognitive analyses. Using structural equation modeling, we assessed the mediation effect of hippocampal volume change ($\Delta\text{Vol}_{\text{hippo}}$) on the association between $E_{\text{hippo}}$ and %ΔHDRS. We quantified the direct effect of $E_{\text{hippo}}$ on %ΔHDRS (coefficient $c$), the indirect effect mediated by $\Delta\text{Vol}_{\text{hippo}}$ (coefficient $ab$), as well as the total effect (coefficient $c^*$, controlling for age and sex. Likewise, we assessed the mediation of the effect of ΔVol$_{\text{hippo}}$ on the association between $E_{\text{hippo}}$ and DKEFS-LF (Fig. 4F). Model fits were assessed by the root mean square error of approximation (RMSEA), the comparative fit index (CFI), and the Tucker-Lewis index (TLI). RMSEA smaller than 0.06 and a CFI and TLI larger than 0.95 indicate relatively good model fit [52]. Finally, we performed receiver operating characteristic (ROC) analysis using $E_{\text{hippo}}$ as a classifier of negative cognitive outcome. We determined the area under the ROC curve with the binary classifier of the lower bound of test-retest reliability of DKEFS Letter Fluency Scaled Score (−3) [41]. The 95% confidence interval for the area under the curve (AUC) is computed with 2000 stratified bootstrap replicates. Finally, we performed similar analysis with a control brain region, the right postcentral gyrus, which receives higher E-field dosing. This metric depends only on the electrode placement and individual head anatomy and is independent of waveform parameters including current amplitude, and thus reflects only the spatial properties of the induced E-field in the brain.

**RESULTS**

The demographic and clinical characteristics of the study sample are summarized in Supplementary Material (Supplement Material Table 1). Because of the mid-series electrode placement switch and the large difference in the left hippocampal E-field between the RUL and BT electrode placements, we focused on the right hippocampus. Because of the effect of time on hippocampal volume change (Supplementary Fig. 1), we restricted the analysis to subjects who completed the study protocol ($n = 52$). We present the results on subjects who completed the protocol with RUL electrode placement, DKEFS Category Fluency, left hippocampus and right postcentral gyrus (control region) in Supplementary Material (Supplementary Material, Sections 1–4).

**Amplitude and Hippocampal E-field**

The average (± standard deviation) $E_{\text{hippo}}$ increased across the 600, 700, and 800 mA amplitude arms: 77.5 Volts/meter (V/m) (± 11.3), 87.0 V/m (±14.8), and 101.0 V/m (±12.3) ($F_{2,50} = 14.2, p < 0.01$). The contrasts between the 600/700 mA arms have similar $E_{\text{hippo}}$ (difference: 9.5 V/m, $p = 0.11$ with Bonferroni correction). The contrasts between the 600/800 mA (difference: 23.5 V/m, $p < 0.01$) and 700/800 mA arms (difference: 14.0 V/m, $p = 0.006$) reflect amplitude differences with $E_{\text{hippo}}$ (Fig. 2A, translational implications in Fig. 2B).

**Hippocampal E-field and volume change**

$E_{\text{hippo}}$ had a direct relationship with right hippocampal volume change ($R^2 = 0.34$, $\beta = 0.04$, $t_{47} = 2.52$, $p = 0.02$, partial eta $[2] \beta^2 = 0.04$) (Fig. 3). Sex was also associated ($\beta = 1.43, t_{46} = 2.59, p = 0.01$). Post-estimation sex contrasts revealed that females had 1.43% greater hippocampal volume change relative to males (standard error ± 0.68, confidence interval: 0.06–2.79).

**Hippocampal E-field, volume change, and clinical outcomes**

$E_{\text{hippo}}$ had no relationship with %ΔHDRS ($R^2 = 0.15, \beta = -0.13, t_{46} = -0.62, p = 0.54$, effect size = 0.008).

**Fig. 2** Hippocampal E-field variability and implications for ECT dosing. A Right and left hippocampal E-field strengths across the three amplitude arms. B Individualized amplitudes have the potential to create accurate and precise E-field dosing.
Right hippocampal volume change had a relationship with %ΔHDTRS ($R^2 = 0.26$, $\beta = -5.45$, $t_{149} = -3.12$, $p = 0.003$, effect size = 0.17). ECT treatment number also contributed to %ΔHDTRS ($\beta = 4.11$, $t_{149} = 3.00$, $p = 0.004$, effect size = 0.16). Structural equation modeling showed that there was a significant direct effect of $E_{\text{hippo}}$ on %ΔHDTRS ($c = 0.39$; $z = 2.43$, $p = 0.015$) and a significant indirect effect through ΔVol$_{\text{hippo}}$ ($a = 0.042$, $b = -4.87$; $z = -2.21$, $p < 0.027$), leading to an insignificant total effect ($c' = a \times b = 0.18$; $z = 1.24$, $p = 0.22$) (Fig. 4A–C). The RMSEA for the model is $< 0.001$; the CFI and TLI for the model are 1.000 and 1.046, respectively, altogether indicating a good model fit.

**Hippocampal E-field, Hippocampal Volume Change, and Cognitive Outcomes:** $E_{\text{hippo}}$ had an inverse relationship with ΔDKEFS Letter Fluency ($R^2 = 0.24$, $\beta = -0.04$, $t_{149} = -2.44$, $p = 0.02$, effect size = 0.12). Right hippocampal volume change had no relationship with ΔDKEFS Letter Fluency ($R^2 = 0.15$, $\beta = -0.14$, $t_{149} = -0.68$, $p = 0.50$, effect size = 0.01). Structural equation modeling showed a significant direct effect of $E_{\text{hippo}}$ on ΔDKEFS Letter Fluency ($c = -0.043$; $z = -2.61$, $p = 0.009$) and significant total effect ($c = -0.042$; $z = -3.16$, $p = 0.002$) independent of changes in right hippocampal volume change (indirect effect: $z = 0.09$, $p = 0.93$). The RMSEA for the model is $< 0.001$; the CFI and TLI for the model are both 1.00, altogether indicating a good model fit. Receiver operating characteristic curves identified 112.5 V/m as the maximal right hippocampal E-field associated with cognitive safety. Fifth, the whole brain metric ($E_{\text{brain}}$) did not demonstrate the same relationships as the right hippocampal E-field suggesting anatomic specificity for these relationships.

**Recent investigations** have examined the relationships between E-field strength, neuroplasticity, and antidepressant outcomes. The Global ECT MRI Collaboration (GEMRIC) database, which included 151 subjects treated with RUL at 800 or 900 mA, revealed a direct relationship with left hippocampal E-field strength and left hippocampal neuroplasticity [30]. Our results demonstrated the direct relationship between E-field strength and neuroplasticity in both the right and left hippocampal (left hippocampal results presented in Supplementary Material Section 3). The higher amplitudes (800 and 900 mA) used in the GEMRIC database may have created a ceiling effect with the right hippocampal neuroplasticity that was not evident with the lower amplitudes (600, 700, and 800 mA) used in the present investigation. Despite the robust sample size, the GEMRIC investigation failed to establish a relationship between E-field strength and antidepressant outcome. In contrast, a recent investigation that included mixed electrode placements (both RUL and BT) at 900 mA showed that higher E-field strength in both the left and right temporal lobes was associated with worse antidepressant outcome [31]. Our investigation demonstrated that hippocampal neuroplasticity is a potential mediator in the E-field and antidepressant relationship such that increased E-field strength is associated with increased hippocampal neuroplasticity, which is associated with improved antidepressant outcomes.

While hippocampal E-field dose is related to both hippocampal neuroplasticity and cognitive outcomes, the link between hippocampal neuroplasticity and cognitive impairment is less robust. Our original cognitive hypothesis focused on hippocampal memory function as measured with the Hopkins Verbal Learning Test-Revised (HVLT-R) Percent Retention Raw Score [33], but verbal memory retention was not related to amplitude-mediated differences in cognitive performance [36]. The pathophysiology of ECT-mediated cognitive impairment may not be limited to hippocampal neuroplasticity, and thus alternative anatomic locations (e.g., prefrontal cortex) and mechanisms such as disrupted long-term potentiation and related impact on resting-state functional connectivity should be considered [54, 55]. Even if the hippocampus is not directly involved in verbal fluency performance, the hippocampus appears to be the most sensitive anatomic region for measuring the impact of E-field dosing on cognition for the brain. In contrast to antidepressant outcomes (higher E-field is better), cognitive outcomes suggest the opposite relationship (higher E-field compromises cognitive safety). Our results are demonstrated an optimal E-field strength 112.5 V/m in the right hippocampus to maximize antidepressant outcomes without compromising cognitive safety.

**DISCUSSION**

This investigation provided new insights focused on the relationships between hippocampal E-field strength, neuroplasticity, and clinical and cognitive outcomes. First, the average right hippocampal E-field strength increased with each amplitude arm. However, despite the incremental amplitude difference of 100 mA across the arms, the induced E-fields within each arm exhibit substantial variability, which is large enough to mask group differences (i.e., no E-field differences between 600 and 700-mA arms). Second, right hippocampal E-field strength is associated with hippocampal neuroplasticity, as measured by increase in hippocampal volume after ECT. Third, right hippocampal neuroplasticity mediated the relationship between right hippocampal E-field strength and antidepressant outcome. Fourth, right hippocampal E-field strength had a direct relationship with the cognitive outcome (specifically phonemic fluency). Unlike antidepressant outcomes, right hippocampal neuroplasticity did not mediate this relationship between right hippocampal E-field and cognitive outcomes. Receiver operating characteristic curves identified 112.5 V/m as the maximal right hippocampal E-field associated with cognitive safety. Fifth, the whole brain metric ($E_{\text{brain}}$) did not demonstrate the same relationships as the right hippocampal E-field suggesting anatomic specificity for these relationships.

Right hippocampal neuroplasticity ($t_{149} = 2.59$, $p = 0.01$) controlling for age, sex, pulse width and number of treatments.

**Molecular Psychiatry** (2022) 27:1676 – 1682

**Z.-D. Deng et al.**
Several limitations should be considered when interpreting these results. First, seizure is an important therapeutic component of ECT [56], but the impact of seizure on neuroplasticity and clinical outcomes was not assessed with this investigation. Seizure activity may be related to neuroplasticity with and without E-field generation (i.e., Metrazol therapy) [57, 58]. The lateralization of ictal power with RUL electrode placement suggests that E-field and ictal power may be interrelated [59]. Future work may employ similar structural equation models with topographical ictal power to disentangle the impact of E-field and ictal power on ECT’s therapeutic and iatrogenic effects. Second, our investigation included subjects who switched to BT electrode placement and 1.0 ms pulse width. BT results in a different E-field spatial distribution compared to RUL with higher E-field strength in the hippocampus. We repeated our analysis with RUL electrode placement only and maintained our relationship with E-field magnitude in the right hippocampus [61]. However, a validation study using in vivo intracranial recordings in humans demonstrated that white matter anisotropy did not significantly improve E-field modeling accuracy [62]. In recent work by Takamiya et al., E-field modeling.
models were computed in thirty depressed subjects receiving RUL ECT, a subgroup of which incorporated DTI-derived anisotropic conductivity in the models. The E-field correlations with regional brain volumetric changes were similar between both models [57]. Future work will need to validate E-field modeling accuracy (with or without DTI) prior to clinical use (“E-field-informed ECT”). Fourth, our sample included a relatively homogenous sample of older adults with MDD meeting the indication to start with RUL electrode placement. Our sample did not include subjects that required initiation of the ECT series with BT electrode placement secondary to acuity and need for rapid response. Larger samples across the adult lifespan and with different acuity levels will be needed to further assess these complex relationships [63].

CONCLUSIONS AND FUTURE DIRECTIONS

Our results have implications for ECT dosing. With a fixed extracranial current amplitude, the ECT “dose” as represented by the intracranial E-field is highly variable due to anatomic differences in skin, skull, fluid, and brain tissue [29]. This anatomic variability is prominent in older (age 50+ years) adults with MDD and can compromise both antidepressant efficacy (insufficient stimulation of mood-related circuitry) and safety (inducing cognitive impairment due to excessive stimulation of cognitive related circuitry). The E-field variability precludes the use of a fixed amplitude for older adults with MDD treated with ECT. Our results suggest a sweet spot of ECT dosing ~112.5 V/m in the right hippocampus that will maintain antidepressant benefit and maximize cognitive safety.

To improve the accuracy and precision of ECT dosing, we propose a solution towards individualized amplitude (Fig. 2B). “E-field-informed ECT” has the potential to individualize current amplitude based on the knowledge of the location (hippocampus) and strength (112.5 V/m) of the target region [64]. This method would require pre-ECT imaging (MRI T1, T2) and the capacity to perform E-field modeling before starting ECT. While achieving the goal of individualized amplitude and reducing variability of the ECT dose, the time, equipment, and expertise required for this option could have limited translational impact. Alternatively, amplitude titrated seizure thresholds will reduce the individual variability related to fixed amplitude ECT dosing [50, 65]. Amplitude titrated seizure threshold will result in consistent E-field strength despite variance in anatomic features typically associated with E-field variability, and potentially limit over-exposure of the brain to excessive stimulation [66]. Reducing the E-field variability will create a more standardized and consistent ECT dosing strategy for treatment with RUL ECT, thus improving antidepressant and cognitive outcomes.

REFERENCES

1. UK ECT Review Group. Efficacy and safety of electroconvulsive therapy in depressive disorder: a systematic review and meta-analysis. Lancet. 2003;361:799-808.
2. Semkovska M, McLoughlin DM. Objective cognitive performance associated with electroconvulsive therapy for depression: a systematic review and meta-analysis. Biol Psychiatry. 2010;68:568–77.
3. Semkovska M, Keane D, Babalola O, McLoughlin DM. Unilateral brief-pulse electroconvulsive therapy and cognition: effects of electrode placement, stimulus dosage and time. J Psychiatr Res. 2011;45:770–80.
4. Gbyl K, Rostrup E, Raghava JM, Andersen C, Rosenberg R, Larsson HBW, et al. Volume of hippocampal subregions and clinical improvement following electroconvulsive therapy in patients with depression. Prog neuro-Psychopharmacol Biol Psychiatry. 2021;104:110048.
5. Gbyl K, Stottrup MM, Mita Raghava J, Xue Jie S, Vibebeck P. Hippocampal volume and memory impairment after electroconvulsive therapy in patients with depression. Acta psychiatrca Scandinavica 2021;143:238–52.
6. Bouckaert F, Sieniawski O, Obels J, Dols A, Vandenbulcke M, Stek M, et al. ECT: its brain enabling effects: a review of electroconvulsive therapy-induced structural brain plasticity. J ECT. 2014;30:143–51.
7. Malberg JE, Eisch AJ, Nestler EJ, Duman RS. Chronic antidepressant treatment increases neurogenesis in adult rat hippocampus. J Neurosci. 2000;20:9104–10.
8. Dukart J, Regen F, Kherif F, Colla M, Bajbouj M, Heuser I, et al. Electroconvulsive therapy-induced brain plasticity determines therapeutic outcome in mood disorders. Proc Natl Acad Sci USA. 2014;111:1156–61.
9. Joshi SH, Espinoza RT, Pirnia T, Shi J, Wang Y, Ayers B, et al. Structural plasticity of the hippocampus and amygdala induced by electroconvulsive therapy in major depression. Biol Psychiatry. 2016;79:82–92.
10. Pirnia T, Joshi SH, Leaver AM, Vasavada M, Njau S, Woods RP, et al. Electroconvulsive therapy and structural neuroplasticity in neocortical, limbic and paralimbic cortex. Transl Psychiatry. 2016;6:e832.
11. Cano M, Martinez-Zalacain I, Bernabeu-Sanz A, Contreras-Rodríguez O, Hernandez-Ribas R, Via E, et al. Brain volumetric and metabolic correlates of electroconvulsive therapy for treatment-resistant depression: a longitudinal neuroimaging study. Transl Psychiatry. 2015;5:e1023.
12. Cao B, Luo Q, Fu Y, Du L, Qiu T, Yang X, et al. Predicting individual responses to the electroconvulsive therapy with hippocampal subfield volumes in major depression disorder. Sci Rep. 2018;8:5434.
13. Gryglewski G, Baldinger-Melich P, Seiger R, Godbersen GM, Michenthaler P, Klob M, et al. Structural changes in amygdala nuclei, hippocampal subfields and cortical thickness following electroconvulsive therapy in treatment-resistant depression: longitudinal analysis. Br J Psychiatry. 2019;214:59–67.
14. Nordanskog P, Larsson LR, Larsson EM, Johanson A. Hippocampal volume in relation to clinical and cognitive outcome after electroconvulsive therapy in depression. Acta Psychiatr Scand. 2014;139:303–11.
15. Ota M, Noda T, Sato N, Okazaki M, Ishikawa M, Hattori K, et al. Effect of electroconvulsive therapy on gray matter volume in major depressive disorder. J Affect Disord. 2015;186:186–91.
16. Jorgensen A, Magnusson P, Hanson LG, Kirkegaard T, Benveniste H, Lee H, et al. Regional brain volumes, diffusivity, and metabolite changes after electroconvulsive therapy for severe depression. Acta Psychiatr Scand. 2016;133:154–64.
17. Bouckaert F, Dols A, Emsell L, De Winter FL, Vansteelandt K, Claes L, et al. Relationship between hippocampal volume, Serotonin DBNF, and depression severity following electroconvulsive therapy in late-life depression. Neuropsychopharmacology. 2016;41:2741–8.
18. Sartorius A, Demirakca T, Bohringer A, Clemm von Hohenberg C, Aksay SS, Bumb JM, et al. Electroconvulsive therapy induced gray matter increase is not necessarily correlated with clinical data in depressed patients. Brain Stimul. 2019;12:335–43.
19. Yroni A, Nemmi F, Billoux S, Giron A, Spoer M, Taib S, et al. Grey Matter changes in treatment-resistant depression during electroconvulsive therapy. J Affect Disord. 2019;258:42–49.
20. Van Den Bossche MJ, Emsell L, Dols A, Vansteelandt K, De Winter FL, Van den Stock J, et al. Hippocampal volume change following ECT is mediated by rs699947 in the promoter region of VEGF. Transl Psychiatry. 2019;9:191.
21. Abbott CC, Jones T, Lemke NT, Gallegos P, McClintock SM, Mayer AR, et al. Hippocampal structural and functional changes associated with electroconvulsive therapy response. Transl Psychiatry. 2014;4:e483.
22. Akers KG, Martinez-Canabal A, Restivo L, Yiu AP, De Cristofaro A, Hsiang HL, et al. Hippocampal neurogenesis regulates forgetting during adulthood and infancy. Science. 2014;344:598–602.
23. Mongiat LA, Schinder AF. Neuroscience. A price to pay for adult neurogenesis. Science. 2014;344:594–5.
24. van Oostrom I, van Eijndhoven P, Butterbrod E, van Beek MH, Janzing J, Donders R, et al. Decreased cognitive functioning after electroconvulsive therapy is related to increased hippocampal volume: exploring the role of brain plasticity. J ECT. 2018;34:117–23.
25. Bouckaert F, De Winter FL, Emsell L, Dols A, Rhebergen D, Wampers M, et al. Grey matter volume increase following electroconvulsive therapy in patients with late life depression: a longitudinal MRI study. J Psychiatry Neurosci. 2015;40:140322.
26. Sartorius A, Demirakca T, Bohringer A, Clemm von Hohenberg C, Aksay SS, Bumb JM, et al. Electroconvulsive therapy increases temporal gray matter volume and cortical thickness. Eur Neuropsychopharmacol. 2016;26:506–17.
27. Jiang J, Wang J, Li C. Potential mechanisms underlying the therapeutic effects of electroconvulsive therapy. Neuropsychobiology. 2018;77:224–31.
28. Z.-D. Deng et al.
32. Rosa MA, Abdo GL, Lisansy SH, Peterchev A. Seizure induction with low-amplitude-current (0.5 A) electroconvulsive therapy. J ECT. 2011;27:341–2.

33. Maynar P, Harris A, Gangadhar B. 500-mA ECT-A proof of concept report. J ECT. 2015;31:e23–26.

34. Youssef NA, Sidhom E. Feasibility, safety, and preliminary efficacy of Low Amplitude Seizure Therapy (LAPT): A proof of concept clinical trial in man. J Affect Disord. 2017;222:6–10.

35. Youssef NA, Ravilla D, Patel C, Yassa M, Sadek R, Zhang LF et al. Magnitude of reduction and speed of remission of suicidality for low amplitude seizure therapy (LAPT) compared to standard right unilateral electroconvulsive therapy: a pilot double-blinded randomized clinical trial. Brain Sci 2019. https://doi.org/10.3390/brainsci9050099.

36. Abbott CC, Quinn D, Miller J, Ye E, Iqbal S, Lloyd M et al. Electroconvulsive therapy pulse amplitude and clinical outcomes. Am J Geriatr Psychiatry. 2021;29:166–78.

37. d’Elia G. Unilateral electroconvulsive therapy. Acta Psychiatr Scand Suppl. 1970;215:1–98.

38. Sackheim HA, Prudic J, Devanand DP, Nobler MS, Lisansy SH, Peyser S et al. A prospective, randomized, double-blind comparison of bilateral and right unilateral electroconvulsive therapy at different stimulus intensities. Arch Gen Psychiatry. 2000;57:425–30.

39. Hamilton M. Rating depressive patients. J Clin Psychiatry. 1980:41:21–24.

40. Kellner CH, Knapp R, Husain MM, Rasmussen K, Sampson S, Cullum M et al. Conventional Criteria Versus New Alternatives. Struct Equ Modeling. 1999;6:1–19.

41. Delis DC, Kaplan E, Kramer, JH. Delis Kaplan Executive Function System. The Psychological Corporation: San Antonio, TX, 2001.

42. Fischl B. FreeSurfer. NeuroImage. 2012;62:774–81.

43. Reuter M, Schmansky NJ, Rosas HD, Fischl B. Within-subject template estimation for unbiased longitudinal image analysis. NeuroImage. 2012;61:1402–18.

44. Reuter M, Fischl B. Avoiding asymmetry-induced bias in longitudinal image processing. NeuroImage. 2011;57:19–21.

45. Reuter M, Rosas HD, Fischl B. Highly accurate inverse consistent registration: a robust approach. NeuroImage. 2010;53:1181–96.

46. Saturnino GB, Antunes A, Thielscher A. On the importance of electrode parameters for shaping electric field patterns generated by tDCS. NeuroImage. 2015;120:25–35.

47. Smith SM, Jenkinson M, Woolrich MW, Beckmann CF, Behrens TE, Johansen-Berg H et al. Advances in functional and structural MR image analysis and implementation as FSL. NeuroImage. 2004;23:528–59.

48. Friston KJ. Statistical parametric mapping: the analysis of functional brain images. Elsevier/Academic Press: Amsterdam, 2007.

49. Nielsen JD, Madsen KH, Puonti O, Siebner HR, Bauer C, Madsen CG et al. Antidepressant efficacy of Low Amplitude Seizure Therapy (LAP-ST): A proof of concept clinical trial in man. J Affect Disord. 2017;222:6–10.

50. Lee WH, Lisansy SH, Laine AF, Peterchev AV. Regional electric field induced by electroconvulsive therapy in a realistic finite element head model: influence of white matter anisotropic conductivity. Neuroimage. 2012;59:2110–23.

51. Huang Y, Liu AA, Lafon B, Friedman D, Dayan M, Wang X et al. Measurements and models of electric fields in the in vivo human brain during transcranial electrical stimulation. Elife 2017;6. https://doi.org/10.7554/eLife.18834.

52. Olteadal L, Bartsch H, Sarhaug OJ, Kessler U, Abbott C, Dols A et al. The Global ECT-MRI Research Collaboration (GEMRIC): Establishing a multi-site investigation of the neural mechanisms underlying response to electroconvulsive therapy. NeuroImage Clin. 2017;14:422–32.

53. Saturnino GB, Siebner HR, Thielscher A, Madsen KH. Accessibility of cortical regions to focal TES: Dependence on spatial position, safety, and practical constraints. NeuroImage. 2019;203:116183.

54. Peterchev AV, Krystal AD, Rosa MA, Lisansy SH. Individualized low-amplitude seizure therapy: minimizing current for electroconvulsive therapy and magnetic seizure therapy. Neuropsychopharmacology. 2015;40:2076–84.

55. Deng Z-D, Lisansy SH, Peterchev AV. Controlling stimulation strength and focality in electroconvulsive therapy via current amplitude and electrode size and spacing: comparison with magnetic seizure therapy. J ECT. 2013;29:325–35.

**AUTHOR CONTRIBUTIONS**

CA, DQ, ZE, and SM designed the study. CA, ML, and SM completed assessments and quality assurance. CA, EE, SM, TJ, JU, ZD performed the analysis. All authors wrote, revised, approved, and agreed to be accountable for all aspects of the final manuscript.

**FUNDING INFORMATION**

This study was supported by grant from National Institute of Mental Health (U01 MH111826 Abbott, R61 MH125126 Abbott R01, MH119285 McClintock). Z-DD is supported by the National Institute of Mental Health Intramural Research Program (ZIAMM02955), and in part by Brain & Behavioral Research Foundation NARSAD Young Investigator Award (26161).

**COMPETING INTERESTS**

The authors declare no competing interests.

**ADDITIONAL INFORMATION**

Supplementary information The online version contains supplementary material available at https://doi.org/10.1038/s41380-021-01380-y.

Correspondence and requests for materials should be addressed to Christopher C. Abbott.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.