CHAPTER 4

A collection of letters
Volume Increase of the Dentate Gyrus Induced by Electroconvulsive Therapy: Shedding Light on the Clinical Relevance of Plasticity in the Hippocampus

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Journal of ECT, 2019, 35(4), e57-e58
DEAR EDITOR:

While the underlying mechanisms of electroconvulsive therapy (ECT) remain unknown, volume increase in the hippocampus has been consistently reported (Nuninga et al., 2019; Oltedal et al., 2018; Takamiya et al., 2018, 2019) since the first study by Nordanskog et al in the Journal of ECT (Nordanskog et al., 2010). To date, the role of this volume increase, as well as whether it contributes to or is responsible for the effectiveness of ECT, is unclear. In this letter, we reanalyzed the data from a previous study (Takamiya et al., 2019), replicating a recent finding (Nuninga et al., 2019), showing that an increase in volume of the dentate gyrus (DG) is related to improvement in depressive symptoms. This replication sheds light on the relationship between volume increase and the clinical effect of ECT. Furthermore, we briefly discuss the importance of using sensitive statistical techniques to investigate the effectiveness of ECT.

The finding of ECT-induced hippocampal volume increase is consistent with multiple lines of evidence from animal studies, which have shown that electroconvulsive stimulation induced neuroplasticity in the hippocampus, including neurogenesis, synaptogenesis, gliogenesis, and angiogenesis, of which neurogenesis is the most robust finding (Bouckaert et al., 2014). Surprisingly, previous studies found no significant correlation between hippocampal volume change and clinical improvement (Oltedal et al., 2018; Takamiya et al., 2018), yet the number of ECTs, electrode placement (Oltedal et al., 2018), or even cognitive changes resulting from ECT (Bouckaert et al., 2014) have been put forward as possible explanations of ECT-related hippocampal volume change. However, the hippocampus is a multilayer structure with each substructure comprising different functions. Therefore, analyzing the whole hippocampus might not be accurate to detect a relationship between volume change and clinical improvement. Recent technological advances in neuroimage (post)processing enabled us to calculate volumes of each hippocampal subfield. Given the strong and robust finding of neurogenesis in the DG (possibly the only neurogenic region of adult human brain) after electroconvulsive stimulation in preclinical studies, several authors suggested that the volume increase of the hippocampus will selectively pertain to the DG (Nuninga et al., 2019; Takamiya et al., 2018).

In our original study (Takamiya et al., 2018), we reported that hippocampal volume increase induced by bilateral ECT was mostly driven from volume increase in the DG and that remitters showed larger volume increase in the right DG than nonremitters. However, we did not find a linear correlation between volume change in the right DG and change in Hamilton Depression Rating Scale (HAM-D) scores. Nuninga et al. (2019), however, showed that the effect of ECT was specific to the DG by using a 7-T magnetic resonance imaging (MRI) and that volume changes in the DG were significantly correlated with change in HAM-D scores. One major difference between these 2 studies is the field strength (i.e., 3 T vs 7 T) of MRI (increasing the accuracy of subfield delineation). In addition, Nuninga et al. (2019), investigated the relationship between volume change of the DG and clinical effect with a repeated measures correlation (rmcorr) (Bakdash & Marusich, 2017), whereas we used a simple linear
correlation. Rmcorr accounts for nonindependence among observation and adjust for interindiv-
didual variability. Moreover, it evaluates intraindividual association between 2 measures, and
parallel lines are fit to the data from each participant. The benefits of rmcorr include higher
statistical power than simple correlation. Because of these advantages, rmcorr seems more
suitable than simple linear correlation for data from a pre-ECT/post-ECT design. Therefore,
we reanalyzed our data in collaboration with Nuninga et al. (2019), using the statistical pack-
age R (version 3.4.3) and rmcorr.

As a result, we now found a statistically significant negative correlation between
HAM-D score and the right DG volume \( r = -0.46; P = 0.018; 95\% \) confidence interval, \(-0.72\)
to \(-0.07\), but no significant result in the left DG \( r = -0.35; P = 0.076; 95\% \) confidence interval,
\(-0.66\) to \(0.06\). The negative correlation indicates that an increase in the DG volume is asso-
ciated with a decrease in HAM-D score. We did not find any correlations between HAM-D
score and the other subfields. Even though our data were from 3 T MRI scanner, our reana-
lyzed results are consistent with the previous study (Nuninga et al., 2019) and also support the
hypothesis that neuroplasticity underlies the efficacy of ECT.
REFERENCES


LETTER 2

The dentate gyrus in depression: directions for future research

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Molecular psychiatry, 2020
Koch et al. (2019) discuss our results regarding volume increases in the dentate gyrus (DG) (Nuniga et al., 2019) in the interesting context of research into stress-related disorders and fear generalization in combination with neurogenesis. While our research included severely depressed patients, Koch et al. (2019) raise the possibility that findings may be generalized to wider diagnostic groups, including trauma patients (i.e., PTSD) and patients with an anxiety disorder. In the following we would like to discuss this wider interpretation of our results and shed light on the further steps needed to be taken in years to come. We recognize that our work serves as a next step into understanding the molecular mechanisms behind severe depression (and perhaps other stress-related disorders) and the development of new therapies aiming to correct pathophysiological mechanisms.

In short, using 7-tesla magnetic resonance imaging (MRI) we found that after ten sessions of electroconvulsive therapy (ECT) the volume of the DG was significantly increased in severely depressed patients, leaving the other subfields of the hippocampus unaffected. These findings point in the direction of increased neurogenesis after ECT, although other functional recovery processes (such as synaptogenesis, axonal sprouting, and angiogenesis) may also contribute to the increase in volume. In healthy controls (n = 8), this increase was not present. In addition, we found that baseline DG volume could predict clinical response (measured with the 17-item Hamilton Depression Rating scale; HAM-D, where higher scores indicate more severely depressed patients) in a regression model. Furthermore, we found that the change in volume was associated to a change in HAM-D score (i.e., larger treatment responses were associated to greater increases in volume). Importantly, our technical equipment has two major advantages:

- We used ultra-high field MRI (7 tesla), which enabled us to focus on the hippocampal area with resolution of 0.286 × 0.286 × 2 mm, a considerable higher image resolution than previous work.

- We used automatic scan planning, which enabled us to rescan the exact same location using the exact same angulation on both occasions, which substantially increases sensitivity to volume changes.

As we included severely ill patients (mean baseline Hamilton score of 22.59) it was extremely challenging to motivate and engage participants to complete both scan sessions, which resulted in a small sample size (N = 23 patients). Our findings trigger other questions and together with Koch and other authors in the field, we can now design a path to further answer remaining questions in order to come to rapid, new, and better tolerated treatment options for severely ill patients.

In answer to the first point raised by Koch et al. (2019), regarding the difference of
DG baseline volume between patients and controls, we did not find a significant difference of baseline DG volumes (left and right) between patients and controls (difference left DG patients - controls = -33.03 mm$^3$, Cohen's d = -0.32, difference test t = -0.69, p = 0.51; difference right DG patients - controls = -19.90 mm$^3$, Cohen's d = -0.26, difference test t = -0.5, p = 0.63). The volumes at post treatment of the left and right DG of patients are not statistically different from that of the controls (difference left DG patients - controls = -1.38 mm$^3$, Cohen's d = -0.012, difference test t = -0.03, p = 0.97; difference right DG patients - controls = -0.45 mm$^3$, Cohen's d = -0.004, difference test t = -0.01, p = 0.99). A second question raised by Koch et al. (2019), concerns the association between DG volumes and depression severity at baseline. Koch et al. state that baseline volumes of the DG could be associated to depression severity and that this association could explain the predictive effect of DG volumes. However, we do not find an association between left or right DG and depression severity at baseline in the patient group (left r = -0.21, p = 0.37, right r = 0.28 p = 0.24). Moreover, including depression severity at baseline in our regression model predicting clinical change, did not change our results: baseline DG volume (left/right) still predict treatment response (in the patient group). Therefore, predicting treatment response based on baseline DG volumes cannot be explained by depression severity at baseline in our sample (although the effect of depression severity could be missed due to possible type II errors). Further, left and right DG have opposite effects in the linear regression model predicting clinical response. At baseline a smaller left DG is associated with better response, while for the right DG the inverse seems true. Interestingly, when computing the widely used asymmetry index ((left - right)/ (left + right) (Kurth, Gaser, & Luders, 2015; Postema et al., 2019) for the DG, the index is able to predict the response of ECT ($t = -3.44$, $p = 0.004$). This prediction model, with age and gender as covariates, is significant and explains 45.8% of the variance ($F(3,15) = 4.23$, $p = 0.02$). Again, inclusion of baseline Hamilton scores does not significantly change the results (the asymmetry index remains a significant predictor: $t = -2.68$, $p = 0.018$). Whether this observation will hold in larger samples, and especially at conventional (3 tesla) MRI, will be a valuable clinical question. Specifically, when this observation holds at 3 tesla MRI it could be more easily implemented in the clinic to help predict clinical response for individual patients (in combination with the help of other predictors, e.g., DG related tasks such as pattern separation).

For further research, a first important question is to answer whether or not the findings (i.e., a significant increase in volume of the DG after ECT in severely depressed patients) from our previous study (Nuninga et al., 2019) using ultra-high field MRI can be replicated with standard clinical MRI scanners operating at 3 tesla. To answer this first question, we collaborated with a group from Tokyo University. Together with Takamiya et al. we reanalyzed their previously published data (Takamiya, Plitman, et al., 2019), now reporting a significant correlation between changes in DG volume and clinical response after bilateral ECT in an independent sample using 3 tesla MRI (Takamiya, Nuninga, et al., 2019). In the aforementioned previously published study (Takamiya, Plitman, et al., 2019), Takamiya et al. reported
volume changes in DG volumes but did not find a simple linear correlation between difference scores in DG volume and HAM-D. However, when they implemented the same repeated measures correlation (Bakdash & Marusich, 2017) we used in our study (Nuninga et al., 2019), a significant correlation was found between a decrease in HAM-D score and an increase in right DG volume [yet not significant in left DG (Takamiya, Nuninga, et al., 2019)]. This finding again highlights the importance of the DG and neuroplastic changes in the DG in response to ECT treatment and suggests feasibility of replicating our findings using conventional 3 tesla MRI.

The next questions to answer regards the generalizability of our findings as well as a confirmation that they are related to neurogenesis. Now that we have a replicable method to assess DG volume and changes in that volume during recovery, we and others can set out to assess whether volumetric changes are related to plastic changes of the DG during remission. Second, animal research using both MRI and post-mortem quantification of neurogenesis is needed to confirm our theory that DG changes are caused by neurogenesis. If volumetric changes reflect plastic changes of the DG during recovery and if decreased plasticity of the DG can be confirmed to underlie the broader category of stress related disorders, this would be a major aid to develop new treatments targeting this mechanism.

In terms of treatment, while ECT is highly effective, its tolerability is low, which restricts its use. Previous animal research has delivered a wealth of information regarding processes that can positively impact neurogenesis, which include: fasting for at least 24 h (Lee, Duan, & Mattson, 2002) physical exercise [especially running (van Praag, Kempermann, & Gage, 1999)], sleep (Hairston et al., 2005) and demanding cognitive tasks (Shohayeb, Diab, Ahmed, & Ng, 2018). We envision a treatment with intensive use of these four elements as an effective and noninvasive new treatment for depression and perhaps other stress-related disorders. An extra challenge will be the motivation of patients for such a combined intervention. To this end, we may use knowledge from the gaming industry to develop an attractive and engaging program that motivates even apathetic participants to continue their practice in order to stimulate neurogenesis and help patients overcome
REFERENCES:


