

# Effectiveness studies

transcranial Direct Current Stimulation

# tDCS



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# Guidelines on safety and application of tDCS

## Low intensity transcranial current stimulation: Safety, ethical, legal regulatory and application guidelines



Antal A, Alekseichuk I, Bikson M, Brockmüller J, Brunoni AR, Chen R, Cohen LG, Bowthwaite G, Ellrich J, Flöel A, Fregni F, Goerge MS, Hamilton R, Haueisen J et al.  
2017 - Clinical Neurophysiology, Volume 128, NO. 9

**OBJECTIVE** To review the safety and potential side effects of tDCS in humans and recommendations for safe use in clinical practice

**METHODOLOGY** Systematic hazard analysis side effects in transcranial current stimulation studies (in total 18000 stimulation sessions in healthy people, neurological and psychiatric patients). Inclusion of studies up to 2016

### RESULTS

<i>Assessment</i>	<i>Results</i>
<i>Safety</i>	<ul style="list-style-type: none"><li>- (tDCS) in humans is considered safe on the basis of this study</li><li>- side effects in the elderly, pregnant women, children are not different from those typical of adult study participants</li><li>- long-term effects in children/adolescents have not been sufficiently investigated</li></ul>
<i>Side effects</i>	<ul style="list-style-type: none"><li>- moderate side effects requiring action are rare and not clearly caused by tDCS</li><li>- mild side effects without need for action: headache, tiredness after stimulation, slight tingling or burning sensation at the stimulation site</li><li>- moderate side effects: skin burns due to improper application of the electrodes</li><li>- no reports of serious side effects</li><li>- no reported cognitive or perceptual side effects</li><li>- no reported seizures demonstrably attributable to tDCS</li></ul>
<i>Injuries</i>	<ul style="list-style-type: none"><li>- risk of injury can be minimized by:<ul style="list-style-type: none"><li>- selection and preparation of the electrode</li><li>- correct placement of the electrodes, specifically the position and contact with skin (through saline paste or sponges with sufficient saline solution)</li></ul></li></ul>

### RECOMMENDATIONS

<i>Training</i>	<ul style="list-style-type: none"><li>- sufficient training of users in safety precautions and the attaching of electrodes</li></ul>
<i>Patient Suitability</i>	<ul style="list-style-type: none"><li>- comprehensive background information on patient before therapy in an intake consultation assessing contraindications and risks</li></ul>
<i>Device technology</i>	<ul style="list-style-type: none"><li>- use devices which are verified in scientific research</li><li>- do not use homemade or uncertified equipment</li></ul>
<i>Protocol</i>	<ul style="list-style-type: none"><li>- comply with maximum stimulation levels and duration alongside correct electrode application</li></ul>

**CONCLUSION** This analysis shows that standardized applications of tDCS can be safely performed within the limits of the clinically approved and common intensity of max. 1 - 2 mA and max. 20 - 30 minutes with correct application of the electrodes. For all practitioners, instruction in handling the electrodes and the protocols is necessary.

# Depression

## Trial of Electrical Direct-Current Therapy versus Escitalopram for Depression

Brunoni AR, Moffa AH, Sampaio-Junior B, Borriero L, Moreno ML, Fernandes RA, Veronezi BP, Nogueira BS, Aparicio LVM, Razza LB, Chamorro R, Tort LC, Fraguas R, Lotufo PA, Gattaz WF, Fregni F, Benseñor IM; 2017 – The New England Journal of Medicine, Volume 376



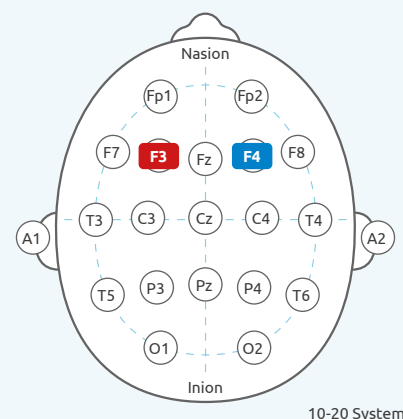
**OBJECTIVE** Influence of tDCS or escitalopram (SSRI) on depression

**PROCEDURE** Placebo group: placebo escitalopram + placebo stimulation;  
Escitalopram group: escitalopram + placebo stimulation  
tDCS group: escitalopram + active stimulation

**METHODOLOGY** Double-blind study (RCT) with 202 depressed patients (55 placebo, 75 escitalopram, 72 tDCS)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	30 minutes
<i>Anode</i>	F3
<i>Cathode</i>	F4
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	5 times a week (first 3 weeks) after that 1 x weekly (7 weeks)
<i>Scope of treatment</i>	22 sessions
<i>Assessment</i>	HDRS-17, MADRS



### RESULTS

Assessment	Placebo group	Escitalopram group	tDCS group
HDRS-17	reduction of HDRS scores compared pre-post by <b>5.8 ± 7.9 points</b>	reduction of HDRS scores compared pre-post by <b>11.3 ± 6.5 points</b>	reduction of HDRS scores compared pre-post by <b>9.0 ± 7.1 points</b>
		** Escitalopram was significantly better than placebo [Difference: 5.5, 95% CI, 3.1 zu 7.8, P<.001]	* tDCS was significantly better than placebo [Difference: 3.2, 95% CI, 0.7 zu 5.5, P=.01]
		* slight superiority of Escitalopram over tDCS [Difference: 2.3, 95% CI, -4.3 zu -0.4, <b>P=.02</b> ]	** in the 10th week, the response rate was significantly higher (reduction by >50% to baseline) in tDCS and escitalopram than in placebo stimulation
MADRS	reduction of MADRS scores compared pre-post by <b>6.6 ± 9.3 points</b>	reduction of MADRS scores compared pre-post by <b>13.4 ± 9.3 points</b>	reduction of MADRS scores compared pre-post by <b>11.0 ± 9.4 points</b>
		** Escitalopram was significantly better than placebo [Difference: 6.8, 95% CI, 5.4 zu 10.6, P<.001]	** tDCS was significantly better than placebo [Difference: 4.4, 95% CI, 2.1 zu 7.2, P=.006]
		* slight superiority of Escitalopram over tDCS [Difference: -2.4, 95% CI, -5.7 zu -1.1, <b>P=.04</b> ]	

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** This study shows that patients with moderate or severe depression (and existing anxiety disorder) can be successfully treated with tDCS. For classic pharmacological treatment, tDCS was shown to be a successful alternative in this study.

# Depression - Stroke

## Effect of transcranial direct current stimulation of stroke patients on depression and quality of life

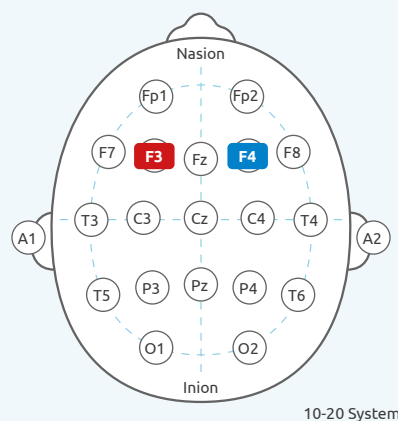
Tae-Gyu A, Soo-Han K, Ko-Un K  
2017 – The Journal of Physical Therapy Science, Volume 29



**OBJECTIVE** Effect of tDCS of DLPFC in patients with post-stroke depression  
**PROCEDURE** Placebo group: conventional occupational therapy + placebo stimulation;  
 Verum group: conventional occupational therapy + active stimulation  
**METHODOLOGY** Controlled study of 40 patients with post-stroke depression (20 control group, 20 tDCS group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	30 minutes
<i>Anode</i>	F3
<i>Cathode</i>	F4
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	1 x daily (for four weeks)
<i>Scope of treatment</i>	20 sessions
<i>Assessment</i>	BDI, SS-QOL



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
<i>BDI</i>	- no significant change in the depressive symptoms from pre 39.0 ± 4.6 to post: 37.8 ± 6.1	* significant improvement of depressive symptoms from pre 38.8 ± 4.7 to post: 16.8 ± 4.6
<i>SS-QOL</i>	- no significant change in quality of life from pre 154.3 ± 18.3 to post: 162.1 ± 18.7	* significant improvement of depressive symptoms from pre 152.1 ± 16.9 to post: 167.7 ± 25.7

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

The results of the study show the positive effect of tDCS on the DLPFC in patients after stroke. Twenty treatments reduced the depressive symptoms by half (BDI scores) and improved quality of life.

# Depression

## Transcranial direct current stimulation in depressive disorders

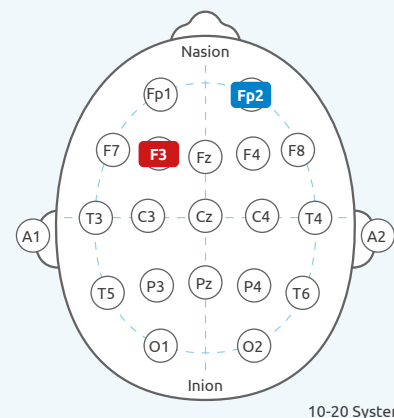
Aust S, Palm U, Padberg F, Bajbouj M  
2015 – Nervenarzt – Review, Ausgabe 12/2015

**Der Nervenarzt**

**OBJECTIVE** Efficacy of tDCS to reduce depressive disorders  
**PROCEDURE** Placebo group: placebo stimulation; active group: active stimulation  
**METHODOLOGY** Review: 6 randomized controlled studies involving 294 patients

### PARAMETER

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20-30 minutes
<i>Anode</i>	F3
<i>Cathode</i>	Fp2
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	10-15 sessions
<i>Assessment</i>	HAMD, MADRS, BDI



### RESULTS

Assessment	Placebo group	Active group
<i>HDRS-17</i>	- no significant differences	* <b>significant improvement in therapy-resistant patients by 40-47%</b>
<i>MARDS</i>	- no significant differences	* significant improvement * treatment combining combining drug and tDCS showed great success (tDCS + sertraline)

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

The review reports on the positive effect of tDCS in depressed patients - tDCS enhances activation in DLPFC. Neurological rehabilitation research points to a targeted use of tDCS to reduce depressive disorders.



# Depression

## Concurrent Cognitive Control Training Augments the Antidepressant Efficacy of tDCS: A Pilot Study

Segrave RA, Arnold S, Hoy K, Fitzgerald P  
2014 – Brain Stimulation, Volume 7

**BRAIN**  
STIMULATION

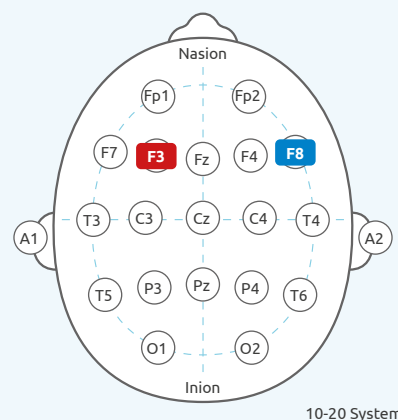
**OBJECTIVE** Clinical effects of tDCS in combination with computer-assisted cognitive training in depressive symptoms

**PROCEDURE** Placebo group: cognitive training + placebo stimulation; active group: cognitive training + active stimulation

**METHODOLOGY** **Combination of tDCS and cognitive training:** Double-blind study

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	24 minutes
<i>Anode</i>	F3
<i>Cathode</i>	F8
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	5 sessions
<i>Assessment</i>	MARDS, BDI, Two-Back



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
<i>MARDS</i>	* MARDS scores decreased [P=0.02] - after follow-up, MARDS scores decreased and were similar to the baseline	- decreased MARDS scores [P=0.06] ** after <b>follow-up, clear and increasing improvement in MARDS scores [P&lt;0.001]</b>
<i>BDI</i>	** BDI scores decreased [P=0.006] - after follow-up, BDI scores decreased and were similar to the baseline	- no significant changes [P>0.05] ** after <b>follow-up, clear and increasing improvement in the BDI scores [P=0.004]</b>
<i>Two-back (accuracy)</i>		* higher accuracy [P=0.02]
<i>Two-back (responsiveness)</i>	** responsiveness increased over time, in all [positive: $F_{(3,69)}=7.93, P=0.001$ ; neutral: $F_{(3,69)}=10.16, P<0.001$ ; negative: $F_{(3,69)}=13.25, P<0.001$ ]	

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** This study shows that tDCS can reduce depressive symptoms. Additional cognitive training (attention training) can effectively reduce depression and increase alertness in a short period of time.

# Addiction - Overview

## A Review of Brain Stimulation Methods to Treat Substance Use Disorders

Coles AS, Kozak K, George TP  
2018 – American Journal on Addictions, Volume 27

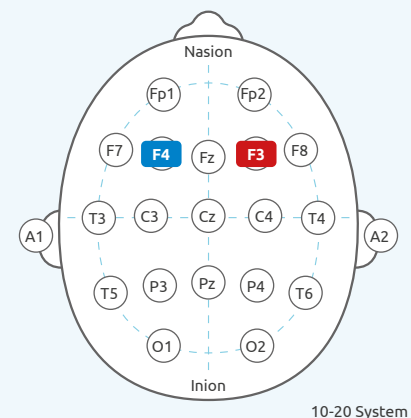
The American Journal  
on Addictions

**OBJECTIVE** Review of the clinical efficacy of various stimulation methods (rTMS, tDCS, DBS) for addiction disorders

**METHODOLOGY** Systematic analysis of studies published between 2000 and 2017 in peer-reviewed journals; test participants had diagnoses of an addiction disorder according to DSM IV and DSM V, respectively

### GENERAL RESULTS REGARDING tDCS

<i>Stim. intensity</i>	mostly 2 mA
<i>Duration</i>	20 minutes
<i>Effect sizes</i>	medium to large after several sessions
<i>Anode</i>	different positions, often lDLPFC
<i>Cathode</i>	different positions, often rDLPFC
<i>Session number</i>	various sessions (5-10) are recommended
<i>Primary / secondary objective</i>	addiction or consumption; recurrence quote
<i>Outcome</i>	effectiveness of tDCS comparable to effectiveness of rTMS



### RESULTS tDCS STUDIES WITH MULTIPLE SIMILAR SESSIONS tDCS 2 MA, 20 MINUTES

<i>Substance</i>	<i>Total number of participants</i>	<i>Results</i>
<i>Alcohol</i>	137	- craving and / or recurrence rate significantly changed with cathodal stimulation of the left DLPFC
<i>Tobacco</i>	76	- mixed results on anodal stimulation of the left DLPFC - significant reduction in craving and / or consumption in cathodal stimulation of the left DLPFC
<i>Cocaine</i>	60	- significant reduction in craving versus baseline and placebo group - no difference in anxiety, depressive symptoms and cognitive performance

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** This analysis shows that short-term treatment with tDCS leads to a decrease in craving in different types of substance-related dependencies. The recurrence rate and thus the consumption itself are in some cases significantly reduced. The best results are achieved by stimulating the DLPFC.

# Addictive disorders - Alcohol addiction

## Multiple Sessions of Transcranial Direct Current Stimulation (tDCS) Reduced Craving and Relapses for Alcohol Use: A Randomized Placebo-Controlled Trial in Alcohol Use Disorder

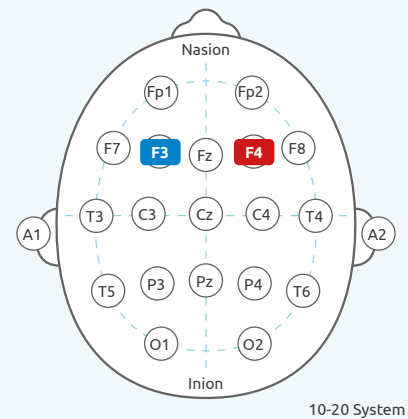
Klauss J, Anders QS, Felipe LV, Nitshe MA, Nakamura-Palacios EM  
2018 – *frontiers in Pharmacology*, Volume 9



<b>OBJECTIVE</b>	Efficacy of repetitive bilateral tDCS over the DLPFC in alcoholics
<b>PROCEDURE</b>	Placebo group: placebo stimulation; active group: active stimulation
<b>METHODOLOGY</b>	Double-blind study (RTC) with alcoholics (placebo group, active group) in a specialist hospital for addiction sufferers

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	F4 (DLPFC)
<i>Kathode</i>	F3 (DLPFC)
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	every two days
<i>Scope of treatment</i>	10 sessions
<i>Assessment</i>	OCDS (addiction pressure), abstinence



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
<i>Recurrence</i>	- 72.2% of the patients relapsed within 3 months of therapy	- 27.3% of the patients relapsed within 3 months of therapy
	difference between the groups (relapse) is significant (p=0.01)	
<i>OCDS</i>	- mean dropped from 3.82 (at the beginning) to 2.46 (in the follow-up) - small effect size (0.32)	- mean dropped from 5.25 (at the beginning) to 1.56 (in the follow-up) - big effect size (1.07)
	- no significant difference pre-post	- no significant difference pre-post
	significant difference before treatment to follow-up	
	significant group differences	

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** The study shows that bilateral stimulation of the DLPFC can significantly reduce the probability of both recurrence and craving. At the time of follow-up, three quarters of the test participants remained abstinent after the stimulation therapy. In the placebo group approximately only a quarter of the participants remained abstinent. It is therefore understood that tDCS can support the long-term success of treatment.

# Addictive disorders - Cocaine addiction

A randomized placebo-controlled trial of targeted prefrontal cortex modulation with bilateral tDCS in patients with crack-cocaine dependence

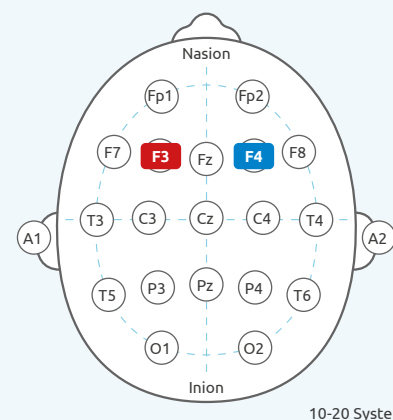
Batista E, Klauss J, Fregni F, Nitsche MA, Nakamura-Palacios EM  
2015 – International Journal of Neuropsychopharmacology, Volume 18



**OBJECTIVE** Clinical effects of repetitive bilateral tDCS on DLPFC in cocaine addiction  
**PROCEDURE** Placebo group: placebo stimulation; active group: active stimulation  
**METHODOLOGY** Double blind study (RTC) with 36 crack cocaine addicts (19 placebo group, 17 verum group)

## PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	F3 (DLPFC)
<i>Cathode</i>	F4 (DLPFC)
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	1 x daily, every second day
<i>Scope</i>	5 sessions
<i>Assessment</i>	HAM-D, HAM-A, WHOQOL, Craving



10-20 System

## RESULTS

Assessment	Placebo group	Active group
Craving	-	* active group and placebo group differ significantly in terms of craving (group and treatment: $[F(1,33)=5.29, P=.028, \text{partial } \eta^2=0.14]$ )
	-	* craving scores decreased linearly from baseline to last treatment [linear regression: $4.412 - 0.617X, r^2=0.058, F(1,66)=4.089, P=.047]$
HAM-D	-	* significant change in HAM-D values within the group ( $P=.04$ )
HAM-A	-	* opposite results: values of the active group decrease, values of the placebo group increase ( $P=.03$ )
WHOQOL	-	* significant differences between groups in WHOQOL: (Q1: $P=0.31$ & Q2: $P=0.48$ )

Significance Scores: \*  $P<0.05$ ; \*\*  $P<0.01$ ; \*\*\*  $P<0.001$

## CONCLUSION

The craving scores were lower after bilateral repetitive tDCS. The active group, compared to the placebo group, is significantly different in terms of anxiety, quality of life and health. The values of the active group improved in post and follow-up.

# Pain - Fibromyalgia

## Anodal transcranial direct current stimulation over the left dorsolateral prefrontal cortex modulates attention and pain in fibromyalgia: randomized clinical trial

Silva AF, Zortea M, Carvalho S, Leite J, da Silva Torres IL, Fregni F, Caumo W; 2017 – Nature, Volume 7



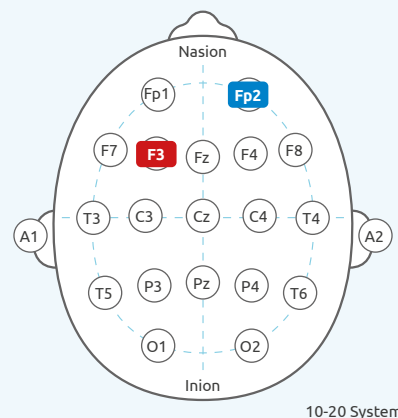
**OBJECTIVE** Efficacy of tDCS on patients with fibromyalgia and cognitive deficits

**PROCEDURE** Placebo group: Go/No-go task + placebo stimulation;  
active group: Go/No-go task + active stimulation

**METHODOLOGY** **Combination of tDCS and cognitive training:** Single blind study (RCT) with 40 patients with fibromyalgia (20 placebo group, 20 active group)

### PARAMETERS

<i>Stimulationsstärke</i>	1 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	F3
<i>Cathode</i>	Fp2
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	1 x weekly
<i>Scope</i>	2 sessions
<i>Assessment</i>	ANT, HPTH, HPTO, Go/No-go task



10-20 System

### RESULTS

<i>Assessment</i>	
	* significant group difference; active group compared to placebo group achieved higher orientation values in ANT after stimulation [F(1, 70.0)=4.189; P=0.044]
<i>ANT</i>	* significant group effect; active group compared to placebo group achieved higher levels in executive function in ANT after stimulation [F(1, 49.11)=7.94; P=0.007]
	* active group compared to placebo group performed significantly better than placebo group in congruent target condition without cue and spatial cues [P=0.040; P=0.020]
<i>HPTH</i>	* active group showed a <b>significant improvement of 4.95% of the heat pain threshold</b> compared to the placebo group [P=0.03]
<i>HPTO</i>	* active group showed a <b>significantly higher tolerance of 3.6% to heat pain</b> compared to the placebo group [P=0.03]
<i>Go/No-go Task</i>	- no significant effects

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

The results of the study show that only after a few stimulation sessions in combination with a Go/No-go training can increase the neural network for attention in orientation and executive function. Stimulation of the DLPFC can be used as an additional stimulation site adjacent to M1 to reduce pain in patients with fibromyalgia.

# Pain - Migraine

## Transcranial direct current stimulation (tDCS) and its influence on analgesics effectiveness in patients suffering from migraine headache

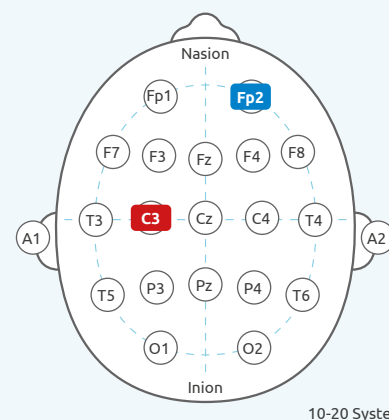
Przeklasa-Muszynska A, Kocot-Kepska M, Dobrogowski J, Wiatr M, Mika J  
2017 - Pharmacological Reports, Volume 69



<b>OBJECTIVE</b>	Effect of tDCS on drug consumption and pain in migraine patients
<b>PROCEDURE</b>	Control group: pharmacological therapy Active group: active stimulation and pharmacological therapy
<b>METHODOLOGY</b>	Single blind study (RCT) with 50 migraine patients with and without aura (20 control group, 30 active group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	C3
<i>Cathode</i>	Fp2
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	2-3 times a week
<i>Scope</i>	10 sessions
<i>Assessment</i>	duration of migraine, migraine seizures, drug use, pain intensity of migraine (NRS)



### RESULTS

Assessment	Placebo group	Active group
<i>Duration of migraine</i>	- no significant reduction - with and without aura	*** with aura: reduced from 24 hours to 10 hours *** without aura: reduced from 25 to 8.5 hours
<i>Migraine seizures</i>	- no significant reduction - with and without aura	** with aura: from 7 days a month to 4 days a month ** without aura: from 7 days a month to 4 days a month
<i>Drug consumption</i>	- no significant reduction (Analgesic) with and without aura - no significant reduction (Triptane) with and without aura	** Analgesic consumption: with aura a <b>reduction of 49%</b> * Analgesic consumption: without aura a <b>reduction of 72%</b> ** Triptans: with Aura a <b>reduction of 58%</b> * Triptans: without aura a <b>reduction of 59%</b>
*** after the 10th treatment, the groups differed quite significantly from each other		
<i>Pain intensity of migraine (NRS)</i>	reduction in pain intensity with aura by 12,5%	<b>reduction in pain intensity with aura by 36%</b>
	reduction in pain intensity without aura by 10%	<b>reduction in pain intensity without aura by 40%</b>

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** This study demonstrates the well-tolerated and effective use of tDCS in migraine patients. The pain intensity of the migraine, the migraine seizures, the migraine duration as well as drug consumption were significantly reduced by 10 treatments.

# Pain - Chronic pain

## Anodal transcranial direct current stimulation for chronic pain in the elderly: a pilot study

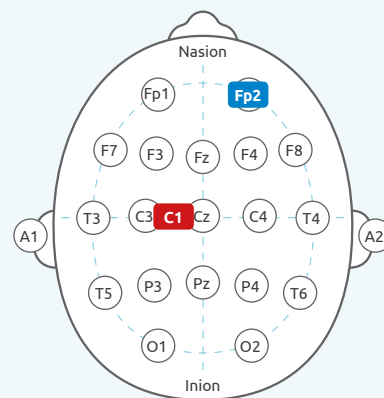
Concerto C, Al Sawah M, Chusid E, Trepal M, Taylor G, Aguglia E, Battaglia F  
2016 - Aging – Clinical and Experimental Research, Volume 28



<b>OBJECTIVE</b>	Efficacy of anodal tDCS on the motor cortex to relieve chronic pain and depression
<b>PROCEDURE</b>	Experimental group: pharmacological therapy + active stimulation
<b>METHODOLOGY</b>	Uncontrolled study: 10 patients with chronic pain

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	C1/C2 (contralateral to the pain side)
<i>Cathode</i>	Fp1/Fp2 (contralateral to the anode)
<i>Size of electrode</i>	25 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope</i>	5 sessions
<i>Assessment</i>	VAS for pain, FFI, PASS-20, drug use, HDRS



10-20 System

### RESULTS

<i>Assessment</i>	<i>Active group</i>
<i>VAS for pain</i>	*** improvement of chronic pain by 36.9% after last treatment [P<0.0001], 42,4% one week after the last stimulation [P<0.0001] ** significant improvement of chronic pain by 21.6% after last treatment four weeks after the last stimulation [P<0.006]
<i>FFI</i>	*** improved foot function by 36.8 % after the last stimulation [P<0.0001], 25.2% one week after the last stimulation [P<0.0001], 23.2% four weeks after the last stimulation [P<0.0001]
<i>PASS-20</i>	*** anxiety related to potential pain decreased by 18.2% after stimulation [P<0.0001], 15.5% one week after the last stimulation [P<0.0001], 12.7% four weeks after the last stimulation [P<0.0001]
<i>Drug consumption</i>	*** significantly reduced consumption of painkillers in follow-up [P<0.0002, P<0.0043]
<i>HDRS</i>	- no significant changes over time

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

The study shows that anodal tDCS of the motor cortex can significantly reduce pain and is long lasting. The normalization of impaired activity in the pain-processing network has increased the quality of life of chronic pain patients.

# Pain - Stroke

## Analgesic effect of transcranial direct current stimulation on central post-stroke pain

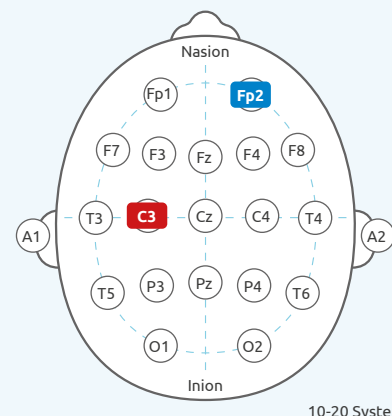
Sea-Hyun B, Gi-Do K, Kyung-Yoon K  
2014 – The Tohoku Journal of Experimental Medicine, Volume 234

Tohoku Journal of  
Experimental Medicine

<b>OBJECTIVE</b>	Effect of tDCS on the analgesic effects in patients after stroke
<b>PROCEDURE</b>	Placebo group: Placebo stimulation Active group: active stimulation
<b>METHODOLOGY</b>	Single blind study (RCT) with 14 patients after stroke with central pain (7 placebo group, 7 active group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	C3/C4-contralateral to the pain side
<i>Cathode</i>	Fp1/Fp2-contralateral to the anode
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	3 x times a week
<i>Scope of treatment</i>	3 weeks
<i>Assessment</i>	VAS for pain, skin temperature measurement, quantitative sensory measurements



### RESULTS

Assessment	Placebo group	Active group
<i>VAS for pain</i>	- no significant change pre: 4.28, post: 4.14	* significant pain reduction Pre: 4.92 and post: 3.14 [P<.05]
<i>Skin temperature</i>	- no significant change pre: 0.94, post: 0.82	* significant reduction of temperature pre: 0.96, post: 0.49 [P<.05]
<i>Quantitative sensory measurement</i>	- no significant change in cold perception: pre: 24.71, post: 25.09	* significant change in cold perception: pre: 24.71, post: 25.09 [P<.05]
	- no significant change in heat perception: pre: 7.36, post: 36.99	* significant change in heat perception: pre: 38.19, post: 35.93 [P<.05]
	- no significant change in cold pain threshold pre: 13.57, post: 13.80	** significant change in cold pain threshold pre: 12.24, post: 14.03 [P<.001]
	- no significant change in threshold for heat pain pre: 48.01, post: 48.16	* significant change in threshold for heat pain pre: 48.07, post: 47.7

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

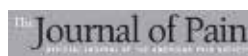
**CONCLUSION** The results demonstrate the effectiveness of tDCS in reducing pain in post stroke patients. The threshold for feeling cold and cold pain as well as for feeling warmth and heat pain changed sustainably.



# Pain - Multiple Sclerosis

## Effects of anodal transcranial direct current stimulation on chronic neuropathic pain in patients with multiple sclerosis

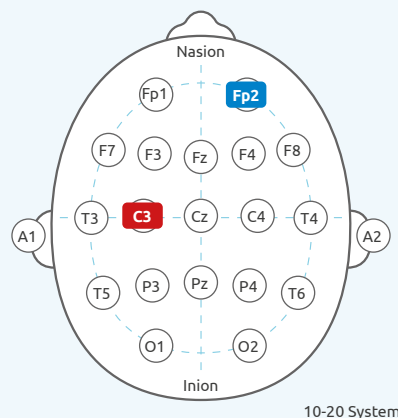
Mori F, Codeà C, Hajime K, Monteleone F, Buttari F, Fiore S, Bernardi G, Koch G; 2010 - The Journal of Pain, Volume 11



- OBJECTIVE** Efficacy of daily tDCS (on consecutive days) to reduce chronic pain in multiple sclerosis patients
- PROCEDURE** Placebo group: Placebo stimulation;  
Active group: active stimulation
- METHODOLOGY** Double-blind study (RCT) with 19 drug-resistant multiple sclerosis patients with neuropathic pain (9 placebo group, 10 active group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	C3/C4-contralateral to the pain side
<i>Cathode</i>	Fp1/Fp2-contralateral to the anode
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	5 sessions
<i>Assessment</i>	VAS for pain, SF-MPQ, MSQOL-54, BDI, VAS for anxiety



### RESULTS

Assessment	Placebo group	Active group
VAS for pain	-	** pain reduction after the fifth day compared to the first [baseline, day 1, day 2, day 3, day 4, day 5: F=9.03, P<.001] * significant differences from day three compared to baseline by 51.7% ± 10.3, day four at baseline by 43.9% ± 11,5 and day five at baseline by 45.5% ± 11,0
SF-MPQ	-	* subjective pain perception decreased after the first week and remained low [every: P<.05]
MSQOL-54	-	* significant effects time (group x time interaction) -> significant differences persisted until follow-up
BDI	-	• no significant differences
VAS for anxiety	-	• no significant differences

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

The results show the effectiveness of tDCS for the reduction of neuropathic pain in multiple sclerosis patients (significant reduction of pain in active group). A small amount of stimulation is enough to improve pain perception and quality of life in MS patients.

# Motor disorders - Stroke

## Constraint-Induced Movement Therapy Combined with Transcranial Direct Current Stimulation over Premotor Cortex Improves Motor Function in Severe Stroke: A Pilot Randomized Controlled Trial

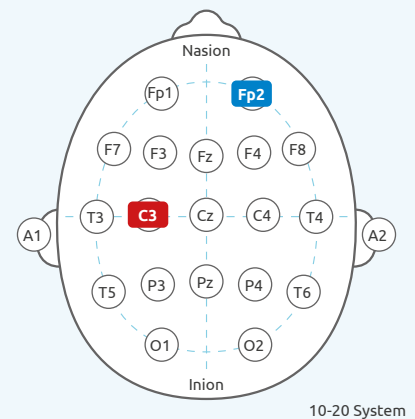
Andrade SM, Batista LM, Nogueira LLRF, de Oliveira EA, de Carvalho AGC, Lima SS, Santana JRM, de Lima ECC, Fernández-Calvo B; 2017 – Rehabilitation Research and Practice, Volume 2017



- OBJECTIVE** Effects of daily tDCS in combination with CIMT on motor rehabilitation in patients after stroke
- PROCEDURE** Placebo group: motor training + placebo stimulation;  
Active group: motor training + active stimulation
- METHODOLOGY** Double blind study (RCT) with 60 patients after stroke (20 placebo group, 20 active group (M1), 20 active group (PMC))

### PARAMETERS

<i>Stim. intensity</i>	1 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	C3 / C4 or 2.5 cm anterior to C3 / C4 (same hemisphere as lesion)
<i>Cathode</i>	Fp1 / Fp2 (contralateral to the anode)
<i>Size of electrode</i>	16 cm <sup>2</sup>
<i>Treatment</i>	5 times a week
<i>Scope of treatment</i>	10 sessions
<i>Assessment</i>	BI, UEFM, MAS, BBT, MRC



### RESULTS

Assessment	Placebo group	Active group
<b>BI</b>	<ul style="list-style-type: none"> <li>* smaller effect size in placebo group [<math>\delta = -.14, P = .01</math>]</li> <li>* significant differences between the groups [<math>\chi^2 = 11.41, df = 2, P = .01</math>]</li> </ul>	<ul style="list-style-type: none"> <li>* stronger effect size in PMC and M1 groups [PMC: <math>\delta = -.83, P = .02</math>; M1: <math>\delta = -.64, P = .01</math>]</li> </ul>
<b>UEFM</b>	<ul style="list-style-type: none"> <li>* improvement placebo group, pre-post within the group from 20 to 24 [<math>z = 0.44, P = .01</math>]</li> <li>* PMC showed greatest improvement in motor recovery compared to M1 [<math>z = -2.01; P = .04</math>] and placebo [<math>z = -2.36; P = .03</math>]</li> </ul>	<ul style="list-style-type: none"> <li>* improvement group M1, pre-post within the group from 20 to 29 [<math>z = -2.04, P = .01</math>]</li> <li>* PMC improvement, pre-post within the group from 21 to 33 [<math>z = -2.86, P = .02</math>]</li> </ul>
<b>MAS</b>	<ul style="list-style-type: none"> <li>* reduction of spasm in placebo stimulation, pre-post from 18 to 15 [<math>z = -1.16, P = .03</math>]</li> <li>* PMC greatest reduction of spasm compared to M1 [<math>z = -2.12; P = .02</math>] and placebo stimulation [<math>z = -2.51; P = .04</math>]</li> </ul>	<ul style="list-style-type: none"> <li>* reduction of spasm in M1, pre-post from 17 to 11 [<math>z = -1.94, P = .01</math>]</li> <li>* reduction of spasm in PMC, pre-post from 16 to 7 [<math>z = -2.87, P = .02</math>]</li> </ul>
<b>BBT und MRC</b>	<ul style="list-style-type: none"> <li>- no significant improvement in placebo group, pre-post [<math>p &gt; 0.05</math>]</li> </ul>	<ul style="list-style-type: none"> <li>- no significant improvement in M1, pre-post [<math>p &gt; 0.05</math>]</li> <li>* significant improvement in PMC, pre-post from 1 to 7 [<math>z = -2.98, P = .02</math>] and [<math>z = -3.01, P = .03</math>]</li> </ul>

Significance Scores: \*  $P < 0.05$ ; \*\*  $P < 0.01$ ; \*\*\*  $P < 0.001$

**CONCLUSION** The study shows that the combined use of CIMT and tDCS leads to significantly greater effects of motor rehabilitation than the isolated use of CIMT. It shows that the stimulation of the premotor cortex (PMC) can cause greater increases in performance than the stimulation of the motor cortex (M1). The stimulation of the premotor area can be a good alternative to standard stimulation of the M1.

# Motor disorders - Stroke

## Ipsilesional anodal tDCS enhances the functional benefits of rehabilitation in patients after stroke

Allman C, Amadi A, Winkler AM, Wilkins L, Filippini N, Kischka U, Stagg CJ, Johansen-Berg H; 2016 – Science Translational Medicine, Volume 8



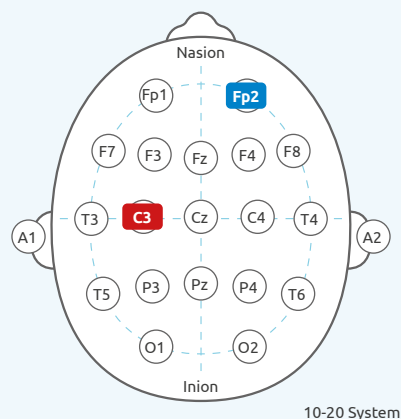
**OBJECTIVE** Effect of tDCS with Motor Training on Patients after Stroke

**PROCEDURE** Placebo group: motor training + placebo stimulation;  
Active group: motor training + active stimulation

**METHODOLOGY** **Combination of tDCS and motor training:** double-blind study (RCT) with 24 patients after stroke (13 placebo group, 11 active group)

### PARAMETERS

<i>Stim. intensity</i>	1 mA
<i>Duration</i>	20 minutes stimulation + 40 minutes motor training
<i>Anode</i>	C3 / C4 (same hemisphere as lesion)
<i>Cathode</i>	Fp1 / Fp2 (contralateral to the anode)
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	9 sessions
<i>Assessment</i>	ARAT, WMFT, UEFM und MRT



### RESULTS

<i>Assessment</i>	<i>Placebo-Gruppe</i>	<i>Active group</i>
<i>ARAT</i>	-	* significant improvement in ARAT [P=.031] compared with placebo group
<i>WMFT</i>	-	* significant improvement [P=.037] compared to the placebo group
<i>UEFM</i>	-	* UEFM showed no significance compared to the placebo group [P=.329]
<i>MRT</i>	- no significant activation patterns in the fMRI	* fMRI studies show increased activity of the active group in restricted hand movement in ipsilateral premotor and motor areas
	* MRI measurements of the active group show <b>volume increase of grey matter</b> after tDCS in the premotor and motor cortex compared to the placebo group	

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

This study shows that the combination of tDCS with motor training promotes neuronal plasticity. The significant long-term changes (follow-up) in the active group show the good possibility of combining tDCS with existing motor therapy. This suggests that motor cortex stimulation can produce plasticity and achieve greater effects than motor training without tDCS.

# Motor disorders - Stroke

## Bihemispheric-tDCS and Upper Limb Rehabilitation Improves Retention of Motor Function in Chronic Stroke: A Pilot Study

Goodwill AM, Teo W, Morgan P, Daly RM, Kidgell DJ  
2016 – Frontiers in Human Neuroscience, Volume 10



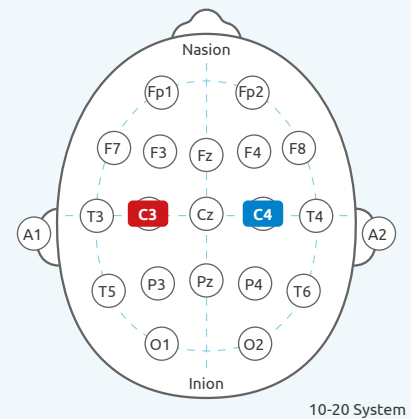
**OBJECTIVE** Effect of anodal tDCS on upper extremity motor rehabilitation and corticospinal plasticity

**PROCEDURE** Placebo group: motor training + placebo stimulation;  
Active group: motor training + active stimulation

**METHODOLOGY** Combination of tDCS and motortraining: double-blind study (RCT) with 15 chronic stroke patients (7 placebo groups, 8 active groups)

### PARAMETER

<i>Stim. intensity</i>	1,5 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	C3/C4 (same hemisphere as lesion)
<i>Cathode</i>	C3/C4 (contralateral to the anode)
<i>Size of electrode</i>	25 cm <sup>2</sup>
<i>Treatment</i>	3 times a week
<i>Scope of treatment</i>	9 sessions
<i>Assessment</i>	MAS, grip strength, tardieu, corticospinal excitability, intracortical inhibition



### RESULTS

<b>Assessment</b>	<b>Placebo group</b>	<b>Active group</b>
<b>MAS</b>	** significant improvement in pre-post in both groups: placebo group: by 43%; <b>active group: by 62%</b> [df=2, both P<0.001]	** improvements were maintained in <b>follow-up (after 3 weeks), or increased to 64%</b> [Chi-Square forest=13.25, df=2, P<0.001]
<b>Grip strength</b>		- no significant differences
<b>Tardieu Scale</b>	* significantly small increase in the Tardieu Scale in the wrist [x2 = 6.56, P = 0.02]	- no significant differences
		- no significant group differences in the elbow and spasticity
<b>Corticospinal excitability: AMT</b>		- no significant differences

<i>Corticospinal excitability: MEPs (paretic arm)</i>	- no significant changes in triggering MEPs (% M <sub>MAX</sub> ) pre-post, by 12% [Forest Chi-Square=0.86, df=2, P=0.36]	** significantly easier triggering of MEPs (% M <sub>MAX</sub> ) pre-post, by 46% [Forest Chi-Square=37.49, df=2, P<0.001]
	** in follow-up (after 3 weeks), MEPs (% M <sub>MAX</sub> ) were more easily released by 38% [Wald Chi-Square=37.49, df=2, P<0.001]	** in follow-up (after 3 weeks), MEPs (% M <sub>MAX</sub> ) were more easily released by 38% [Forest Chi-Square=37.49, df=2, P<0.001]
	- no significant changes in MEPs in the non-paretic arm	
<i>Corticospinal excitability: Laterality Index (LI)</i>	- no significant changes	** <b>increased excitability, shift of LI from 0.6 to 0.3</b> , pre-post [Forest Chi-Square = 14.80, df=2, P<0.001]
	- no significant changes	* in follow-up (after 3 weeks) the shift of the LI remained constant [Forest Chi-Square=14.80, df=2, P=0.03]
<i>Intrakortikale Inhibition: CSP, (non-paretic arm)</i>	- no significant changes	* <b>significant increase in CSP by 33%</b> , pre-post [Forest Chi-Square=8.16, df=2, P=0.01]
	- no significant changes	* in follow-up (after 3 weeks) the increase was maintained, 24% [Forest Chi-Square=8.16, df=2, P=0.04]
<i>Intracortical inhibition: SICI</i>	- no significant changes	- no significant changes
	* in follow-up (after 3 weeks), <b>SICI in the active group increased by 27%</b> compared to placebo [Forest Chi-Square=7.08, df=2, P=0.04]	

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

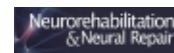
## CONCLUSION

This study demonstrates in detail the improvement of motor function of the upper limb after a tDCS stroke in combination with motor training. The increased hemispheric balance as well as improved corticospinal plasticity in the ipsilateral and contralesional M1 proves to be an important rehabilitation process for the recovery of motor functions after stroke.

# Motor disorders - stroke

## Anodal tDCS combined with radial nerve stimulation promotes hand motor recovery in the acute phase after ischemic stroke

Sattler V, Acket B, Raposo N, Albucher JF, Thalamas C, Loubinoux I, Chollet F, Simonettra-Moreau M; 2015 – Neurorehabilitation and Neural Repair, Volume 29



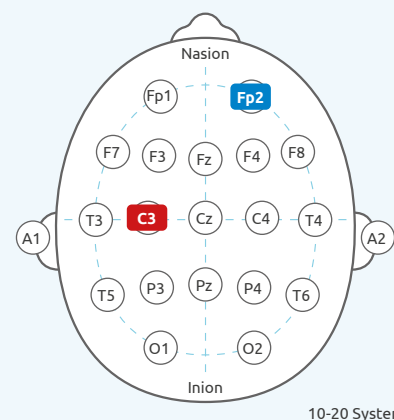
**OBJECTIVE** Effect of tDCS and FES on the functional restoration of armotory in patients after stroke in the acute phase

**PROCEDURE** Placebo group: active FES + placebo stimulation (tDCS); Active group: active FES + active stimulation (tDCS)

**METHODOLOGY** **Combination of tDCS and FES:** Double-blind study (RCT) with 20 patients after stroke with mild to moderate motor deficits (10 placebo, 10 active)

### PARAMETERS

<i>Stim. intensity</i>	1-2 mA
<i>Duration</i>	13 minutes stimulation + 13 minutes FES
<i>Anode</i>	C3/C4 (same hemisphere as lesion)
<i>Cathode</i>	Fp1/Fp2 (contralateral to the anode)
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	5 sessions
<i>Assessment</i>	JHFT, Hand Dynamometer, Nine Hole, Peg Test, Hand Tapping Test, ULFM, TMS (MEP)



### RESULTS

Assessment	Placebo group	Active group
JHFT	* significant improvement (baseline follow-up), reducing time to perform JHFT	* significant improvement (baseline follow-up), Reducing the time to perform JHFT, <b>day 5: 6 sec faster; day 15: 23 sec more quickly; day 30: 24 seconds faster</b>
	-	* <b>significantly improved motor performance after follow-up compared to the placebo group [day 15: P=.03; Day 30: P=.01]</b>
DYN		- no significant changes
9HPT		- no significant changes
HTap		- no significant changes
ULFM	*** significant temporal improvement in follow-up; improved hand and finger extension	*** <b>significant temporal improvement in follow-up, improved hand and finger extension</b>
TMS		- no significant changes regarding AMT and RMT

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** The results of the combined therapy of tDCS and FES show an improvement of the tDCS group in the follow-up (JHFT). Specifically, the functionality of the hand improved, which has a positive effect on the activities of daily life. tDCS in combination with FES can be used for motor rehabilitation.

# Motor disorders - Parkinson's

## Transcranial direct current stimulation associated with gait training in Parkinson's disease: A pilot randomized clinical trial

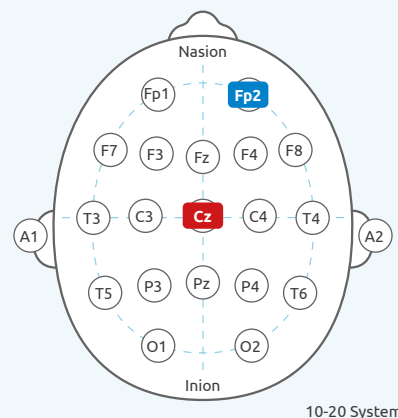
Coast-Ribeiro A, Maux A, Bosford T, Aoki Y, Castro R, Baltar A, Shirahige L, Moura Filho A, Nitsche MA, Monte-Silva K; 2016 – Developmental Neurorehabilitation, Volume 0



- OBJECTIVE** Efficacy of tDCS with gait training on functional mobility in Parkinson's patients
- PROCEDURE** Placebo group: gait training + placebo stimulation;  
Active group: gait training + active stimulation
- METHODOLOGY** **Combination of tDCS and gait training:** double blind study (RCT) with 22 Parkinson's patients (11 placebo group, 11 active group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	13 minutes stimulation + 30 minutes gait training
<i>Anode</i>	Cz (2 cm anterior)
<i>Cathode</i>	Fp2 (contralateral to the affected side)
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	3 x times a week
<i>Scope of treatment</i>	10 sessions
<i>Assessment</i>	TUG, gait speed, cadence, UPDRS, UL-MT, BBS, PDQ-39



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
<i>TUG, walking speed</i>	* significant improvements, both TUG and gait speed improved significantly <b>in post, but were no longer significant in follow-up</b>	* significant improvements, both parameters TUG and gait speed improved significantly in the post test, <b>these improvements remained significant in the follow-up</b> [gait speed: z=-2,184, P=.05; TUG: t=2.223, P=.05]
<i>Cadence</i>		* significant changes
<i>UPDRS</i>		* significant changes
<i>UL-MT</i>		* significant changes
<i>BBS</i>		* significant changes
<i>PDQ-39</i>		* significant changes

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

The result of the combined therapy of tDCS and gait training initially showed no difference to the control group, both groups improved significantly. In follow-up (1 month), only the positive effects of gait training + tDCS remain sustainable. This means that tDCS in combination with gait training prolongs and stabilizes the effects.

# Cognitive deficits - Working memory - Additional study

## tDCS-induced episodic memory enhancement and its association with functional network coupling in older adults

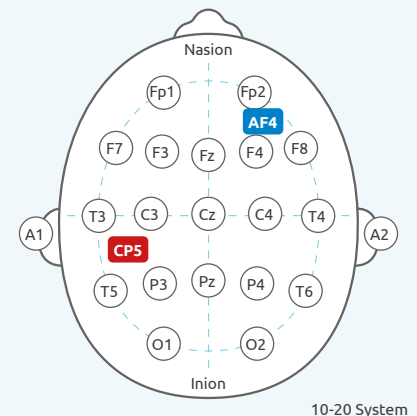
Antonenko D, Hayek D, Netzband J, Grittner U, Flöel A  
2019 – nature, Scientific Reports, Volume 9



**OBJECTIVE** Effect of tDCS over the left temporo-parietal cortex on episodic memory in the elderly  
**PROCEDURE** Learning picture-word pairs combined with placebo or active stimulation  
**METHODOLOGY** **tDCS during memory-related learning tasks:** 34 healthy older people with normal cognitive functions

### PARAMETERS

<i>Stim. intensity</i>	1 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	CP5
<i>Cathode</i>	AF4 (right supraorbital)
<i>Size of electrode</i>	anode: 35 cm <sup>2</sup> , cathode: 100 cm <sup>2</sup>
<i>Treatment</i>	5 learning blocks, stimulation during the first 4 blocks
<i>Scope of treatment</i>	2 sessions (per person 1 active / 1 placebo)
<i>Assessment</i>	accuracy, learning curve, response time, BOLD



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
<i>Accuracy of task fulfilment</i>	- significantly better performance with number of training blocks	** significantly better performance with number of training blocks (P=0.002) * significantly better performance in retrieving the learning after completing the learning blocks (P=0.014)
<i>Learning curve</i>		* steeper learning curve (P=0.014)
<i>Reaction time</i>	- response time shorter with number of training blocks	*** response time shorter with number of training blocks (P <0.001) - no significant difference placebo vs. active group
<i>BOLD</i>	- better performance with initially higher connectivity between temporo-parietal region and hippocampus	- better learning progress with initially higher connectivity

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** The study shows that tDCS also produces significant improvements in cognitive performance in the elderly. Steeper learning curves and better memory performance are achieved. The effectiveness is better with higher existing connectivity between learning-relevant brain regions.



# Cognitive deficits - Additional study

## Augmentation of working memory training by transcranial direct current stimulation (tDCS)

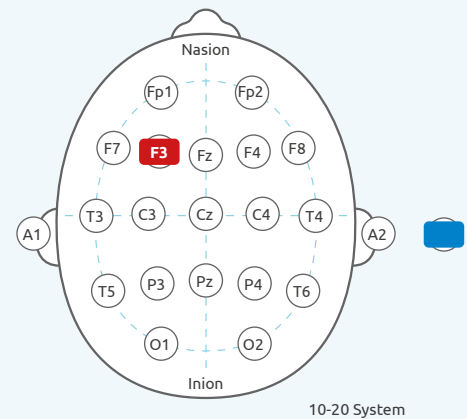
Ruf SP, Fallgatter AJ, Plewnia C  
2017 – Nature, Volume 7



- OBJECTIVE** Influence of tDCS on the working memory performance
- PROCEDURE** Placebo group: working memory training + placebo stimulation;  
Active group: working memory training + active stimulation
- METHODOLOGY** **Combination of tDCS and cognitive training:** single blind study (RCT) with 71 subjects (23 placebo group, 24 congruent group, 24 incongruent group)

### PARAMETERS

<i>Stim. intensity</i>	1 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	F3 (verbal tasks), F4 (spatial tasks)
<i>Cathode</i>	contralateral shoulder, deltoid muscle
<i>Size of electrode</i>	anode: 35 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	3 sessions
<i>Assessment</i>	verbal n-back, spatial n-back



### RESULTS

<i>Assessment</i>	
<i>Effects during training (verbal and spatial n-back)</i>	<p>*** high performance increase in active group (congruent) compared to placebo [<math>\beta_{\text{session} \times \text{congruent vs sham}} = 1.00, SE=0.27, z=3.66, P&lt;0.001</math>]</p> <p>** after two stimulations (second day) marked <b>increase in performance of the active group</b> compared active to the placebo group [<math>t(45)=3.29, P=0.002</math>]</p>
<i>Effects after follow-up (verbal and spatial n-back)</i>	<p>* active group (congruent) showed greater increase in performance in follow-up than placebo group [<math>\beta_{\text{congruent vs sham}} = 2.02, SE=1.01, t(48)=2.01, P=0.050</math>]</p> <p>** active group (congruent) showed <b>significantly better performance level</b> in follow-up than placebo group [<math>t(33)=2.96, P=0.006</math>]</p>
<i>Transfer effects</i>	<p>** patients with lower performance (baseline) benefited more from stimulation [<math>\beta_{\text{baseline} \times \text{congruent vs sham}} = 0.74, SE=0.25, t(65)=-3.00, P=0.004</math>]</p>

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** This study showed that stimulation of the DLPFC can increase working memory performance. The cognitive training accompanied by tDCS can also improve non-trained tasks through transfer effects. The improvement in working memory performance can last up to 9 months. Patients with lower initial performance benefit most from the combined use.

# Cognitive deficits - Traumatic brain injury

## Concomitant use of transcranial direct current stimulation and computer-assisted training for the rehabilitation of attention in traumatic brain injured patients: Behavioral and neuroimaging results

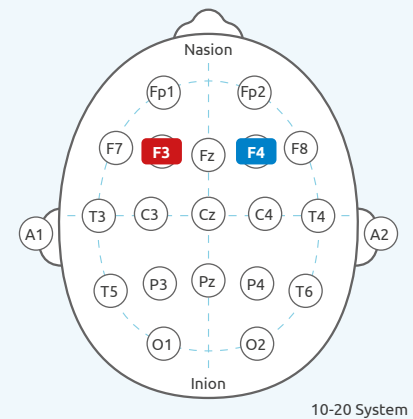
Sacco K, Galetto V, Dimitri D, Geda E, Perotti F, Zettin M, Geminiani GC  
2016 – Frontiers in Behavioral Neuroscience, Volume 10



- OBJECTIVE** Efficacy of tDCS and cognitive training on shared attention and neural activity
- PROCEDURE** Placebo group: computer-based training + placebo stimulation;  
Active group: computer-aided training + active stimulation
- METHODOLOGY** **Combination of tDCS and cognitive training:** single blind study (RCT) with 32 patients after craniocerebral trauma (16 placebo group, 16 active group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes (+30 minutes cognitive training)
<i>Anode</i>	DLPFC (Läsionsseite)
<i>Cathode</i>	DLPFC (contralateral to the anode)
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	2 x times a day
<i>Scope of treatment</i>	10 sessions
<i>Assessment</i>	TEA, rBANS, BDI, AES, MRT



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
<i>TEA</i>	- no significant differences	*** significantly better in pre-post [faster reaction time: $t=3.41, P=0.004$ ; Omission error: $t=4.49, P<0.0001$ ] <b>improvements remained until follow-up</b>
<i>rBANS</i>	-	- no significant differences
<i>BDI</i>	-	- no significant differences
<i>AES</i>	-	- no significant differences
<i>MRT</i>	-	** reduction of activity in the right cingulate gyrus

Significance Scores: \*  $P<0.05$ ; \*\*  $P<0.01$ ; \*\*\*  $P<0.001$

**CONCLUSION** After computer-assisted cognitive therapy + tDCS, abnormal cortical activity normalized only in the active group. In addition, significant, sustained improvements in shared attention and responsiveness have been demonstrated.

# Cognitive deficits - Multiple Sclerosis

## Neuroenhancement through cognitive training and anodal tDCS in multiple sclerosis

Mattioli F, Bellomi F, Stampatori C, Capra R, Miniussi C  
2016 – Multiple Sclerosis Journal, Volume 22



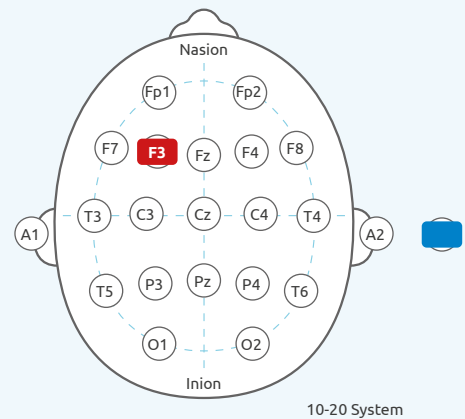
**OBJECTIVE** Effect of tDCS with cognitive training on attention performance

**PROCEDURE** Placebo group: cognitive training + placebo stimulation;  
Active group: cognitive training + active stimulation

**METHODOLOGY** **Combination of tDCS and Cognitive Training:** Double blind study (RCT) with 20 Multiple Sclerosis patients (10 placebo group, 10 verum group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	left DLPFC (F3)
<i>Cathode</i>	shoulder (right)
<i>Size of electrode</i>	25 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	10 sessions
<i>Assessment</i>	SDMT, PASAT, WCST



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
<i>SDMT</i>	* after 10 days of active group is significantly better than placebo group [P=.019]	
<i>PASAT 2</i>	-	* significant improvement
<i>PASAT 3</i>		* significant improvement
<i>WCST</i>		after 10 days active group is significantly better than placebo group: ** total error [P=.003] ** perseverative responses [P=.035] ** perseverative errors [P=.043] ** non-perseverative errors [P=.009]
<i>difficulty</i>	-	* faster achievement of higher difficulty levels [P=.02]

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** tDCS in combination with cognitive training improves attention and information processing in multiple sclerosis patients. In the active group, the significant results are sustainable (follow-up). The stimulation increases the activation of the neuron populations, which could lead to stronger response to therapy.

# Cognitive deficits - Parkinson's

Mild cognitive impairment in Parkinson's disease is improved by transcranial direct current stimulation combined with physical therapy

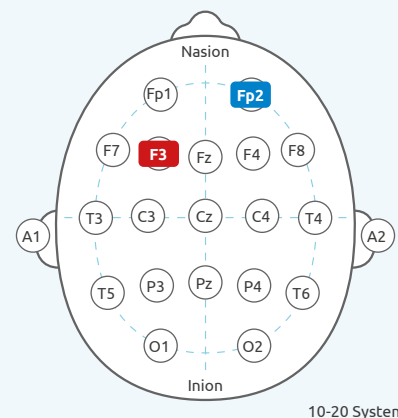
Manenti R, Brambilla M, Benussi A, Rosini S, Cobelli C, Ferrari C, Petesi M, Orizio I, Padovani A, Borroni B, Cotelli M; 2016 – Movement Disorders, Volume 31



- OBJECTIVE** Effect of tDCS and exercise training in Parkinson's patients on motor and cognitive performance
- PROCEDURE** Placebo group: exercise training + placebo stimulation;  
Active group: exercise training + active stimulation
- METHODOLOGY** **Combination of tDCS and exercise training:** double blind study (RCT) with 20 Parkinson's patients (10 placebo group, 10 active group)

## PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	25 minutes
<i>Anode</i>	DLPFC (F3/F4: contralateral to the affected side of the body)
<i>Cathode</i>	Fp2/Fp1 (contralateral to the anode)
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	10 sessions
<i>Assessment</i>	motor function tests, PD-CRS, verbal fluency, TMT



## RESULTS

Assessment	Placebo group	Active group
<i>Motor tests</i>		* both groups improved in the motor tests (e.g. balance)
<i>PD-CRS</i>	-	- results improved in post test [P = .003], <b>this effect remained stable in follow-up</b>
<i>Verbal word liquid -</i>		- improved in the post test [P=.002] and <b>remained sustainable significantly in the follow-up</b> [P=.005]
<i>TMT-B</i>	-	- less time was needed to complete the test - [P=.002], <b>also in follow-up</b>

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** The results of the study show a significant increase in cognitive performance parameters through the combination of tDCS and exercise training. The motor performance increased significantly in both groups, the memory and attention performance improved only in the active group.

# Cognitive deficits - Stroke

## The effects of transcranial direct current stimulation on cognition in stroke patients

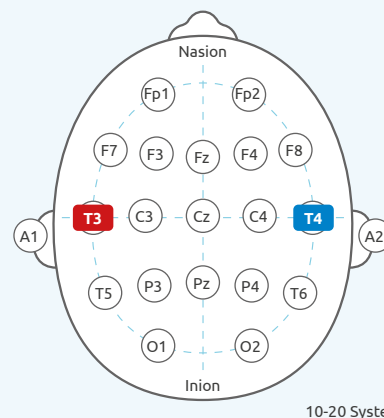
Yun GJ, Chun MH, Kim BR  
2015 – Journal of Stroke, Volume 17



- OBJECTIVE** Efficacy of tDCS on cognitive abilities in post stroke patients
- PROCEDURE** Placebo group: cognitive training + placebo stimulation;  
Active group: cognitive training + active stimulation
- METHODOLOGY** **Combination of tDCS and cognitive training:** double blind study (RCT) with 45 stroke patients (15 placebo group, 15 active group (FT-AS left), 15 active group (FT-AS right))

### PARAMETERS

Stim. intensity	2 mA
Duration	30 minutes
Anode	T3
Cathode	T4
Size of electrode	25 cm <sup>2</sup>
Treatment	1 x daily
Scope of treatment	15 sessions
Assessment	K-MBI, K-MMSE, BDST, FVST, VeLT-R



### RESULTS

Assessment	Placebo group	Active group
K-MMSE	-	* significant improvements in pre-post comparison
BDST (backwards visual)	-	* significant improvements in pre-post comparison
FVST (forward visual span)	-	* significant improvements in pre-post comparison
VeLT-R (verbal learning test - delayed recall)	-	* significant improvements in pre-post comparison
K-MBI	-	* significant improvements in pre-post comparison

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** Stimulation of the T3 region improved memory. The working memory performance only increased in the active group.

# Neglect - Stroke

## The Effect of Transcranial Direct Current Stimulation on Neglect Syndrome in Stroke Patients

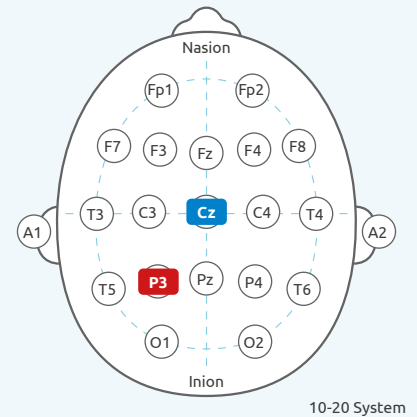
Yi YG, Chun MH, Do KH, Sung EJ, Kwon YG, Kim DY  
2016 – Annals of Rehabilitation Medicine, Volume 40



- OBJECTIVE** Effect of tDCS with conventional occupational therapy on visuo-spatial attention and post-stroke function with neurological neglect
- PROCEDURE** Placebo group: conventional therapy + placebo stimulation;  
Active group: conventional therapy + active stimulation
- METHODOLOGY** **Combination of tDCS and cognitive training:** single blind study (RCT) with 30 left-sided neglect patients (10 placebo group, 20 active group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	30 minutes
<i>Anode</i>	P3 or P4
<i>Cathode</i>	Cz
<i>Size of electrode</i>	25 cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	15 sessions
<i>Assessment</i>	MVPT, SCT, LBT, CBS, K-MBI, FAC



### RESULTS

<i>Assessment</i>	<i>Results</i>
<i>MVPT</i>	* both groups improved significantly pre-post * significantly greater improvement in the active group (from pre: 8.2 ± 6.8 to post: 14.8 ± 5.8) to the placebo group (from pre: 8.3 ± 5.2 to post: 10.3 ± 5.5) [P=0.014]
<i>SCT</i>	* both groups improved pre-post * both groups improved pre-post
<i>LBT</i>	* significantly greater improvement in the active group (from pre: 27.3 ± 18.6 to post: 12.5 ± 13.0) to the placebo group (from pre: 26.0 ± 13.2 to post: 19.0 ± 12.3) [P=0.016]
<i>CBS</i>	* both groups improved pre-post * significantly greater improvement in the active group (from pre: 16.2 ± 6.4 to post: 10.0 ± 6.2) to the placebo group (from pre: 16.0 ± 9.7 to post: 12.3 ± 10.8) [P=0.013]
<i>K-MBI</i>	* both groups improved pre-post
<i>FAC</i>	* both groups improved pre-post

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** The results of the study show that the anodal or cathodal tDCS of PPC (P3/P4) can be combined to support the existing therapy to increase visuo-spatial attention in neglect patients.

# Dysphagia - Stroke

## Randomized Trial of Transcranial Direct Current Stimulation for Poststroke Dysphagia

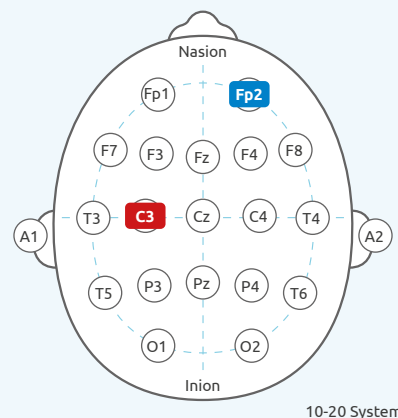
Suntrup-Krueger S, Ringmaier C, Muhle P, Wollbrink A, Kemmling A, Hanning U, Claus I, Warnecke T, Teismann I, Pantev C, Dziewas R; 2018 – Annals of Neurology, Volume 83



- OBJECTIVE** Effect of repeated tDCS in combination with swallowing on the dysphagia severity
- PROCEDURE** Placebo group: swallowing + placebo stimulation;  
Active group: swallowing + active stimulation
- METHODOLOGY** Double-blind study (RCT) with 59 patients after stroke (30 placebo group; 29 verum group)

### PARAMETERS

<i>Stim. intensity</i>	1 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	contralateral: from Cz 3.5 cm lateral and 1 cm anterior: ~ C3 or C4 (in brain stem infarction: ~ C4)
<i>Cathode</i>	Fp1 / Fp2 (contralateral to the anode)
<i>Size of electrode</i>	35 cm <sup>2</sup> anode, 100 cm <sup>2</sup> cathode
<i>Treatment</i>	4 times a week
<i>Scope of treatment</i>	4 sessions
<i>Assessment</i>	FEDSS, MEG, DSRS, FOIS, Dysphagia limit



### RESULTS

Assessment	Placebo group	Active group
<i>FEDSS</i>	* significant improvement (baseline post) [P=.027]	** significant improvement (baseline post) [P<.001]
	<b>*** significantly greater improvement, 1 FEDSS point in active group (83.3%) compared to placebo group (36.7%) average improvement 1.3 vs. 0.4 FEDSS points [η<sup>2</sup>=0.208, P&lt;.0005]</b>	
<i>MEG</i>	- no significant changes	* significant increase in swallow-related event-related desynchronization (baseline post, 13-30Hz)
<i>DSRS</i>	** significant improvement in verum group compared to placebo group [P=.001]	
<i>FOIS</i>	* significant improvement in placebo group compared to active group [P=.041]	
<i>Dysphagia limit</i>	* significant improvement in active group compared to placebo group [P=.018]	

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

The study shows that tDCS provides additional benefits over conventional dysphagia therapy currently practiced. tDCS can be used to achieve faster and greater progress in acute dysphagia patients.

# Aphasia - Stroke

## Repetitive sessions of tDCS to improve naming in post-stroke aphasia: Insights from an individual patient data (IPD) meta-analysis

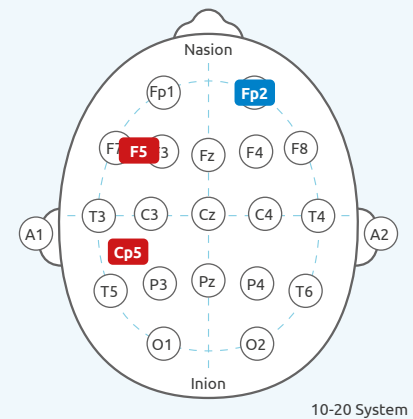
Rosso C, Arbizu C, Dhennain C, Lamy JC, Samson Y  
2018 – Restorative Neurology and Neuroscience (IOS Press), Volume 36



**OBJEKTIVE** Effectiveness of tDCS + naming training based on individual patient data  
**PROCEDURE** Placebo group: naming training + placebo stimulation;  
 tDCS group: naming training + active stimulation  
**METHODOLOGY** meta-analysis with 68 aphasics (87 active-placebo comparisons)

### PARAMETERS

<i>Stim. intensity</i>	1-2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	F5 or Cp5
<i>Cathode</i>	Fp2
<i>Size of electrode</i>	35 cm <sup>2</sup>
<i>Scope of treatment</i>	>5 sessions
<i>Assessment</i>	naming accuracy



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
	*** significant improvement in naming accuracy by 25% (± 37%) (P<.0001)	*** <b>significant improvement in naming accuracy by 35% (± 34%) (P&lt;.0001)</b>
	*** the stronger the improvement in the placebo group, the greater the improvement in the tDCS group [r=0.993, 95% CI: 0.896-0.954, R2= 87%, P<.0001]	
<i>naming accuracy</i>		* frequency of treatment: <b>repeated stimulation (more than five) increases the effects of treatment (P&lt;.02)</b>
		- anodal stimulation of the left temporoparietal flap most effective
	* for all severity grades of aphasia (mild / moderate / severe), greater enhancements can be achieved with active stimulation compared to placebo stimulation (P =.01, P=.002, P=.01)	

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

This meta-analysis shows that the combination of tDCS + naming task can lead to greater progress. The stimulation amount should exceed 5 treatments and should be done on the temporoparietal lobe.



# Aphasia - Stroke

## Electrical stimulation of the motor cortex enhances treatment outcome in post-stroke aphasia

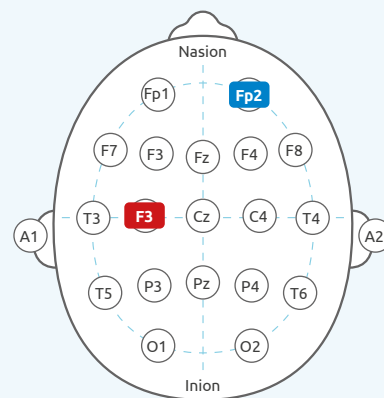
Meinzer M, Darkow R, Lindenberg R, Flöel A  
2016 – Brain – A Journal of Neurology, Volume 139



- OBJEKTIVE** Effect of tDCS on language proficiency in patients with chronic aphasia after stroke
- PROCEDURE** Placebo group: naming training + placebo stimulation;  
Active group: naming training + active stimulation
- METHODOLOGY** **Combination of tDCS and naming training:** double-blind study (RCT) with 26 patients after stroke (13 placebo group, 13 active group)

### PARAMETERS

<i>Stim. intensity</i>	1 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	C3
<i>Cathode</i>	Fp2
<i>Size of electrode</i>	Anode: 35cm <sup>2</sup> , Cathode: 100 cm <sup>2</sup>
<i>Treatment</i>	2 times a day, 4 days a week
<i>Scope of treatment</i>	2 weeks, 16 sessions
<i>Assessment</i>	Naming Test, CETI, PCQ



10-20 System

### RESULTS

Assessment	Placebo group	Active group
<i>Naming Test</i>	* significant improvement on trained objects in post and follow-up	* significant improvements in trained objects [P=.08] and <b>significantly better than the control group in follow-up (6 months)</b> [P=.01]
	* significant improvement in untrained objects, in follow-up no significant improvements (baseline follow-up)	*** significant improvements in trained objects [P=.0009] and <b>significantly better than control group in follow-up (6 months)</b> [P<.01]
<i>CETI (Communicative Effectiveness Index)</i>	* significant	* significantly <b>higher than placebo group</b> after pre-post [P=.037] and in follow-up [P=.055]
<i>PCQ (Partner Communication Questionnaire)</i>	* significant	* significantly <b>higher than placebo group</b> in pre-post [P=.06] and follow-up [P=.03]

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

The results show the positive effects of the combination of tDCS and verbal training in chronic aphasia. Progress is achieved in a short period of time. Stimulation on C3 combined with naming training improved speech ability. A transfer into the everyday life could be proven, whereby the quality of life of the patients increased.

# Aphasia - Stroke

## Differential involvement of the left frontal and temporal regions in verb naming: A tDCS treatment study

Fiori V, Di Paola M, Oliveri M, Caltagirone C  
2013 – Restorative Neurology and Neuroscience, Volume 31



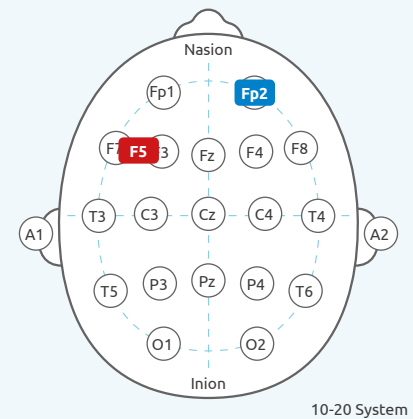
**OBJECTIVE** Effect of tDCS on verbal retrieval performance in aphasics

**PROCEDURE** Placebo setting: naming training + placebo stimulation;  
Verum setting: naming training + active stimulation

**METHODOLOGY** **Combination of tDCS and naming training:** Uncontrolled study in 7 patients with left hemispheric stroke

### PARAMETERS

<i>Stim. intensity</i>	1 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	F5
<i>Cathode</i>	Fp2
<i>Size of electrode</i>	35cm <sup>2</sup>
<i>Treatment</i>	1 x daily
<i>Scope of treatment</i>	15 sessions
<i>Assessment</i>	Naming Test



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
<i>Naming Test</i>	- no significant improvement	** significant improvement: 33% correct name [P = .004] *** significant improvement pre-post: 40% correct name [P = .000] after 5 days: 54% correctly named and recognized (baseline = 14%) in follow-up: performance at 44% (remained constant)

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** The results of the study show that tDCS + verbal training is a good therapeutic combination. The stimulation of the Broca area shows good effects in aphasia patients.

# Aphasia - Dementia

## Inferior parietal transcranial direct current stimulation with training improves cognition in anomic Alzheimer's disease and frontotemporal dementia

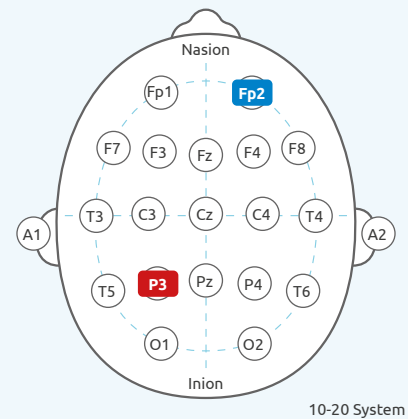
Roncero C, Kniefel H, Service E, Thiel A, Probst S, Chertkow H;  
2017 – Alzheimer's & Dementia: Translational Research & Clinical Interventions, Volume 3



- OBJECTIVE** Impact of tDCS and language training on naming performance
- PROCEDURE** Placebo group: naming training + placebo stimulation;  
Active group: naming training + active stimulation
- METHODOLOGY** **Combination of tDCS and naming training:** double blind study (RCT) with 10 dementia patients (AD or FTD); crossover design

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	30 minutes
<i>Anode</i>	P3
<i>Cathode</i>	Fp2
<i>Size of electrode</i>	35cm <sup>2</sup>
<i>Treatment</i>	once a day, every other day
<i>Scope of treatment</i>	10 sessions
<i>Assessment</i>	Naming Test, Digit Span, Interviews



### RESULTS

Assessment	Placebo group	Active group
<i>Naming Test</i>	* trained objects: significant improvement: baseline - last stimulation: 19% improvement	* trained objects: significant improvement: baseline - last stimulation: 40% improvement
	non-trained objects: * in direct comparison: baseline results are the same for both groups, after two weeks: significantly different (t (9)=3.07, P<.05)	
<i>Digit Span</i>	- baseline 11.38 (SE=1.98)	- baseline 11.75 (SE=1.01)
	- similar baselines [t(7)=0.31, P=.77]	
	- after two weeks to 9.38 (SE = 1.52)	- after two weeks to 12.75 (SE=1.39)
	* significantly different after two weeks [t (7) = 4.34, P<0.1]	
<i>Interview</i>	- no significant improvement	* <b>7/10 relatives reported significant improvements in behavior</b>

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

### CONCLUSION

The results of the study show the positive effects of parietal lobe stimulation on the naming performance. Through the combination of language training and tDCS the effect of the therapy can be strengthened. In addition, stimulated patients were better able to cope with the increased level of semantic processing.

# Schizophrenia

## Effects of transcranial direct current stimulation on working memory and negative symptoms in schizophrenia: a phase II randomized sham-controlled trial

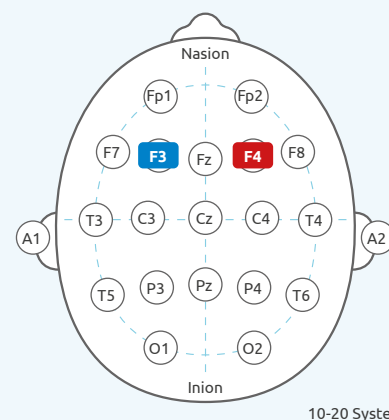
Gomes JS, Trevizol AP., Ducos DV., Gadelha A, Ortiz BB, Fonseca AO, Akiba HT, Azevedo CC, Guimaraes LSP, Shiozawa P, Cordeiro Q, Lacerda A, Dias AM; 2018 – Schizophrenia Research: Cognition, Volume 12



- OBJECTIVE** Effect of multiple tDCS in schizophrenia patients
- PROCEDURE** Placebo group: placebo stimulation;  
Active group: active stimulation
- METHODOLOGY** Double-blind study (RCT) with 24 patients with schizophrenia (12 placebo group, 12 verum group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	F3 (left DLPFC)
<i>Cathode</i>	F4 (right DLPFC)
<i>Size of electrode</i>	25cm <sup>2</sup>
<i>Treatment</i>	5 times a week
<i>Scope of treatment</i>	10 sessions
<i>Assessment</i>	PANSS, cognitive assessment



### RESULTS

<i>Assessment</i>	<i>Placebo group</i>	<i>Active group</i>
<i>PANSS-Negative symptoms</i>	reduction of baseline to last intervention: 0.17; remained constant until the FU	reduction of baseline to last intervention: 3.83; remained constant until FU
<i>PANSS-General symptoms</i>	reduction of baseline to last intervention and FU by 0.83 points	reduction of baseline to last intervention and FU by 10.75 points
<i>PANSS-Total symptoms</i>	reduction of baseline to last intervention and FU by 0.59 points	reduction of baseline to last intervention and to the FU by 6 points
<i>cognitive Assessment</i>	* time effect for working memory, processing speed, visual learning, problem solving	

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** The results of this study recommend the therapeutic use of tDCS in schizophrenia sufferers. Multiple tDCS applications significantly reduced the negative symptoms.

# Acoustic hallucinations - Schizophrenia

## Examining transcranial direct current stimulation (tDCS) as a treatment for hallucinations in schizophrenia

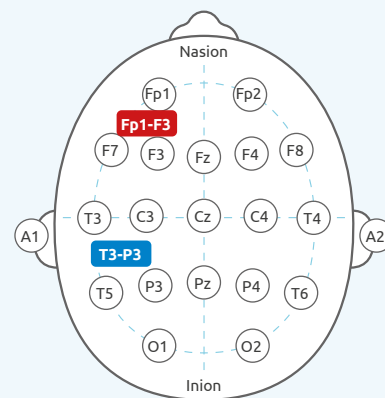
Brunelin J, Mondino M, Gassab L, Haesebaert F, Gaha L, Suaud-Chagny MF, Saoud M, Mechri A, Poulet E; 2012 – The American Journal of Psychiatry, Volume 169



- OBJECTIVE** Efficacy of tDCS in schizophrenia sufferers with severe auditory hallucinations
- PROCEDURE** Placebo group: placebo stimulation;  
Active group: active stimulation
- METHODOLOGY** Double-blind study (RCT) with 30 patients with acoustic hallucinations in schizophrenia (15 placebo group, 15 active group)

### PARAMETERS

<i>Stim. intensity</i>	2 mA
<i>Duration</i>	20 minutes
<i>Anode</i>	F1 - F3
<i>Cathode</i>	T3 - P3
<i>Size of electrode</i>	35cm <sup>2</sup>
<i>Treatment</i>	2 times a day
<i>Scope of treatment</i>	10 sessions
<i>Assessment</i>	AHRS, PANSS



10-20 System

### RESULTS

Assessment	Placebo group	Active group
AHRS	- AHRS improved by 8%	*** after 5 days, a large effect in the active group compared to the placebo group, [df=1.58, p<.0001] (6 months) relapse
	-	*** HRS improved by 31% (after 5 days)
Craving	- effect on AHRS persisted after 1 month (3%) and after 3 months (5%)	*** effect in AHRS increased after one month (36% [t=-4.48, p<.0001]) and also after three months (38% [t=-4.58, p<.0001])
	- slight reduction of negative symptoms PANSS from 82.8 to 80.5	* significant reduction in negative symptoms PANSS from 76.9 to 66.9 compared to the placebo group [P=.01]

Significance Scores: \* P<0.05; \*\* P<0.01; \*\*\* P<0.001

**CONCLUSION** The results show that auditory hallucinations in schizophrenics can be significantly reduced after 10 tDCS treatments. The improvement continued until follow-up (3 months) and improved the quality of life of patients. In addition, the stimulation caused a reduction of the negative symptoms (PANSS).



**Abbreviation test procedure**

AES	Apathy Evaluation Scale	FAC	Functional Ambulation Classification
AHRS	Auditory Hallucination Rating Scale	FEDSS	Fiber Optic Endoscopic Dysphagia Severity Scale
AMT	Active Motor Threshold	FFI	Foot Function Index
ANT	Attention Network Test	FOIS	Functional Oral Intake Scale
ARAT	Action Research Arm Test	FVST	Forward Visual Span Test
BBS	Mountain Balance Scale	HAMA	Hamilton Anxiety Scale
BBT	Box and Block Test	HAMD	Hamilton Depression Scale
BDI	Beck Depression Inventory	HDRS	Hamilton Depression Rating Scale
BDST	Backward Digit Span Test	HPTH	Heat Pain Threshold
CBS	Catherine Bergego Scale	HPTo	Heat Pain Tolerance
CATI	Communicative Effectiveness Index	HTap	Hand Tapping Test
CSP	Cortical Silent Period	JHFP J	ebesen and Taylor Hand Function Test
DSRS	Dysphagia Severity Scale	K-MBI	Korean Modified Barthel Index
DYN	Hand Dynamometer	LBT	Line Bisection Test
Dysphagia-limit	Dysphagia Maximum intake (H <sub>2</sub> O), without ingestion or aspiration	LI	Laterality Index
FAB	Frontal Assessment Battery	MARDS	Montgomery-Åsberg Depression Rating Scale
		MEG	Modified Ashworth Scale
		MEP	Motor Evoked Potential



# Learn how to apply tDCS

The neuroCademy offers one- and two-day advanced training in transcranial direct current stimulation:

- at neuroCademy training centres,
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Learn how to use tDCS in several applications, understanding:

- how tDCS works and how it is applied,
- which therapies are possible according to current research,
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MMSE	Mini-Mental State Examination	SDM	Symbol Digit Modality Test
MRC	Medical Research Council	SF-MPQ	Short-form McGill Pain Questionnaire
MSQOL-54	Multiple Sclerosis Quality of Life-54	SICI	Short-Interval Intracortical Inhibition
MVPT	Motor-Free Visual Perception Test	SS-QOL	Stroke-specific Quality of Life
NRS	Numerical Rating Scale	TEA	Test for the Examination of Attention
OCDS	Obsessive Compulsive Drinking Scale	TMT-B	Trail Making Test -Part B
PANSS	Positive and Negative Syndrome Scale	TUG	Timed Up and Go Test
PASAT 2	Paced Auditory Serial Addition Task 2	UEFM	Upper Extremity Fugl-Meyer
PASAT 3	Paced Auditory Serial Addition Task 3	ULFM	Upper Limb Fugl-Meyer
PASS-20	Pain Anxiety Symptoms Scale-20	UL-MT	Upper Limb Motor Task
PCQ	Partner Communication Questionnaire	UPDRS	Unified Parkinson's Disease Rating Scale
PD-CRS	Parkinson's Disease Cognitive Rating Scale	VAS	Visual Analog Scale
PDQ-39	Parkinson's Disease Questionnaire-39	VeLT-R	Verbal Learning Test-delayed Recall
rBANS	Repeatable Battery for the Assessment of the Neuropsychological Status	WCST	Wisconsin Card Sorting Test
RCT	Randomized Controlled Trial	WHOQOL	Quality of Life of the World Health Organization
SCT	Star Cancellation Test	WMFT	Wolf Motor Function Test
		9HPT	Nine Hole Peg Test

## Imprint

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