

Stimulus intensity determined by dose-titration versus age-based methods in electroconvulsive therapy in Thai patients

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Pichai Ittasakul¹
Apichaya Likitnukul¹
Umporn
Pitidhramabhorn¹
Punjaborn Waleeprakhon¹
Morris B Goldman²

¹Department of Psychiatry, Faculty of Medicine, Ramathibodi Hospital, Mahidol University, Bangkok, Thailand;

²Department of Psychiatry and Behavioral Sciences, Feinberg School of Medicine, Northwestern University, Chicago, IL, USA

Objectives: This study aimed to compare the accuracy of the stimulus intensity (SI) calculated by age-based methods with that using the dose-titration method.

Methods: The initial seizure threshold (IST) was determined using a standardized dose titration in hospitalized Thai patients treated with right unilateral (RUL, n=32) and bilateral (BL, n=57) electrode placement. The correlation between the IST and clinical variables was analyzed. The estimated SI based on the patient's age was compared with the SI determined by dose titration.

Results: Age was highly predictive of the IST for both groups (RUL, $P=0.012$; BL, $P=0.045$). Gender ($P=0.006$) and anticholinergic drug use ($P=0.025$) predicted the IST for the BL group. For the RUL group, the mean \pm SD (median) SI estimated using the half-age and age methods was 158 \pm 46 (169) mC and 315 \pm 92 (338) mC, respectively. The SI determined using the dose-titration method was higher compared with the half-age method and lower compared with the age method. For the RUL group, 31% of subjects using the half-age method and 22% of subjects using the age method would have received an SI within $\pm 20\%$ of that computed using dose titration. Additionally, 19% of subjects using the half-age method and 19% using the age method would have received unacceptably low ($<50\%$) or high ($>200\%$) intensities. For the BL group, 18% of subjects using the half-age method and 32% using the age method would have received an SI within $\pm 20\%$ of that computed using dose titration. Additionally, 39% with the half-age method and 18% with the age method would have received an unacceptably low or high SI, respectively.

Conclusion: Age strongly predicts the IST, but it does not robustly predict the SI compared with dose titration because the SI calculated using age-based methods results in an unacceptably low or high SI that is associated with a marked risk of adverse effects or inadequate response. We recommend the dose-titration method to determine the SI.

Keywords: electroconvulsive therapy, dosing, Thai patients, dose titration, age-based methods, seizure threshold, predictors

Introduction

The stimulus intensity (SI) is a critical feature of electroconvulsive therapy (ECT) because it must be substantially above the initial seizure threshold (IST) for the treatment to be effective but not so high that it unnecessarily increases the risk of cognitive side effects.^{1,2} Currently, the recommended SI is 50%–100% and 150%–500% above the IST for brief pulse bilateral (BL) and brief pulse right unilateral (RUL) ECT, respectively.³ The IST, which is generally defined as the minimum electrical charge needed to produce a generalized seizure lasting at least 20–25 s, exhibits marked variability (ie, 40–50-fold) between patients.^{1,2} Previous studies indicate that age,

Correspondence: Pichai Ittasakul
Department of Psychiatry, Faculty of Medicine, Ramathibodi Hospital, Mahidol University, 270 Rama VI Road, Ratchathewi, Bangkok 10400, Thailand
Tel +66 22 01 1478
Fax +66 22 00 3277
Email pichai.itt@mahidol.edu

sex, concurrent medication, electrode placement, anesthetic agents, past ECT, and body mass index (BMI) all contribute to this variability.^{4–8}

Two methods are typically used to calculate the SI: the dose-titration method¹ and the age-based method.^{9–11} The dose-titration method is generally considered the “gold standard” and seeks to determine the actual IST in a single session. However, the dose-titration method typically involves non-therapeutic treatment and is associated with a risk of potential adverse effects related to restimulation.⁹ The age-based methods estimate the SI based on the patient’s age.^{3,7,12} Age (age method) or one-half of the age (half-age method) is used as a percentage of the maximum output charge to calculate the ST at the first treatment. The age-based methods, however, have the potential to cause otherwise avoidable cognitive side effects and to lengthen the duration of treatment and cumulative exposure to the risk factors.^{3,11,13,14}

In a previous study in Thai patients, Chanpattana et al found that the ST for BL ECT determined using the dose-titration method was proportional to the patient’s age¹⁵ and lower than that calculated using either the half-age or age method.¹⁶ Additionally, the impact of ethnicity on the IST and SI remains unknown. In the current study, we compared the accuracy of the SI calculated using age-based methods with that of the dose-titration method in Thai subjects who received either RUL or BL ECT.

Materials and methods

Setting and study design

This study was conducted in accordance with the Declaration of Helsinki, and approved by the Ethics Committee on Human Experimentation of the Faculty of Medicine, Ramathibodi Hospital, Mahidol University, Bangkok, Thailand. Ramathibodi Hospital is a 1,300-bed university hospital with a 22-bed general psychiatric unit. There are

250–300 patients admitted per year,¹⁷ and approximately 10% of these patients receive ECT.¹⁸

The study design is retrospective. We performed chart reviews on all patients who received inpatient ECT from January 2013 to September 2016. Patients provided verbal and written informed consent before participation. Data collected were age, gender, BMI, concurrent use of psychotropic medications, history of previous ECT, psychiatric diagnosis, electrode placement, IST, and anesthetic agents.

Electroconvulsive therapy treatment

At least one psychiatrist involved in this manuscript oversaw each patient’s ECT. All patients were assessed by psychiatry and anesthesiology before receiving ECT. Benzodiazepines were discontinued at least 15 h before treatment, which was conducted in the postanesthetic care unit. A modified technique involving a brief pulse wave generated by a Mecta Spectrum 5000Q (Mecta Corp, Portland, OR, USA) was applied typically after thiopental (1–6 mg/kg intravenously (IV)) or propofol (1–2 mg/kg IV) anesthesia and succinylcholine (0.5–1 mg/kg IV) muscle relaxant were administered. ECT was performed three times per week. The titration schedule was standardized (Table 1) but other clinical decisions were made at the physician’s discretion.

Determination of the initial seizure threshold

The IST was determined at the first ECT session using the dose-titration method as shown in Table 1, except that males started at Step 2. The method of electrode placement (BL or RUL) was determined by the symptom severity, as determined by the treating psychiatrist. The IST was defined as the dose at which there was definite evidence on the electroencephalogram of generalized seizure activity for at least 25 s. None of the patients was administered more than

Table 1 Dose-titration schedule and parameter settings

Step	Right unilateral electrode placement					Bilateral electrode placement				
	Pulse width (ms)	Frequency (Hz)	Duration (s)	Current (mA)	Charge (mC)	Pulse width (ms)	Frequency (Hz)	Duration (s)	Current (mA)	Charge (mC)
1	0.5	50	0.5	800	20	1	40	0.75	800	48
2	0.5	40	1	800	32	1	40	1.25	800	80
3	0.5	40	1.5	800	48	1	40	2	800	128
4	0.5	40	1.5	800	77	1	60	2	800	192
5	1	40	2	800	128	1	60	3	800	288
6	1	60	2	800	192	1	60	4.5	800	432
7	1	60	3	800	288	1	60	6	800	576
8	1	60	4.5	800	432					
9	1	60	6	800	576					

four stimulus doses. The subsequent SI was calculated using $6 \times \text{IST}$ for the RUL electrode placement and $1.5 \times \text{IST}$ for the BL electrode placement.

Statistical analysis

All statistical analysis was performed using SPSS 21.0 for Windows (IBM Corporation, Armonk, NY, USA). Data are presented using the mean \pm SD and percentage for continuous and categorical outcomes, respectively. An assessment of the data distribution was performed to determine the appropriate statistical test. Correlation between the IST and demographic data, clinical characteristics, and ECT variables was performed using Spearman's rho because the data were not normally distributed. For the age-based IST calculation, a percentage of the maximum output charge (576 mC) determined by the patient's age or half of their age was used. Multivariate ordinal regression was used to examine the relationship of the IST to demographic and clinical variables. $P < 0.05$ was considered statistically significant.

Results

Demographic characteristics of the subjects

Ninety psychiatric inpatients had received ECT using the dose-titration method. Among these 90 patients, 57 patients received BL and the other 33 patients received RUL. Among the 33 patients who received RUL, one patient received ketamine as the anesthetic agent and was therefore not included in our study. Subjects' demographic characteristics are presented in Table 2. For the RUL group, the age range was 19.5–87.7 years. For the BL group, the age range was 18.3–80.9 years.

The initial seizure threshold and associated factors

For the RUL group, the mean \pm SD (median) IST was 42 ± 21 mC (32 mC), with a range of 19–128 mC. The IST for males was 50 ± 28 mC (48 mC) and that for females was 38 ± 15 mC (32 mC) (Mann–Whitney $U=81$, $z=-1.5$, $P=0.146$). Three outliers ($\text{IST} \geq 76$) were excluded from the analysis. Age was associated with a higher IST ($r_s=0.465$, $P=0.011$). For the multivariate ordinal regression analysis, only age predicted the IST ($P=0.012$).

For the BL group, the mean \pm SD (median) IST was 150 ± 101 mC (128 mC), with a range of 32–576 mC. For males, the IST was 190 ± 120 mC (128 mC) and for females, it was 114 ± 62 mC (104 mC) (Mann–Whitney $U=228$, $z=-2.9$, $P=0.003$). For subjects who were taking concurrent

Table 2 Demographic characteristics

Characteristics	RUL group (n=32)	BL group (n=57)
Female	21 (65.6%)	30 (52.6%)
Age (years)	54.8 \pm 16 (58.8)	43.7 \pm 13.8 (44.4)
Body mass index	23.5 \pm 5.5 (23.2)	25.8 \pm 6.1 (25.2)
Diagnosis		
Schizophrenia	1 (3.1%)	28 (49.1%)
Schizoaffective disorder	2 (6.3%)	8 (14.0%)
Bipolar disorder	7 (21.9%)	15 (26.3%)
Major depressive disorder	21 (65.6%)	2 (3.5%)
Other diagnosis	1 (3.1%)	4 (7.0%)
History of previous ECT	6 (18.8%)	17 (29.8%)
Concurrent medication		
Benzodiazepine	4 (12.5%)	2 (3.5%)
Antipsychotic	25 (78.1%)	55 (96.5%)
Antidepressant	13 (40.6%)	5 (8.8%)
Anticonvulsant	1 (3.1%)	4 (7.0%)
Anticholinergic	0	12 (21.1%)
Anesthetic agents		
Thiopental	30 (93.8%)	55 (96.5%)
Propofol	2 (6.2%)	2 (3.5%)

Note: Data are presented as the mean \pm SD (median) or n (%).

Abbreviations: BL, bilateral; ECT, electroconvulsive therapy; RUL, right unilateral.

anticholinergic medication, the IST was 105 ± 72 mC (80 mC) and for subjects who were not taking concurrent anticholinergic medications, it was 162 ± 104 mC (128 mC) (Mann–Whitney $U=155.5$, $z=-2.3$, $P=0.021$). The two outliers ($\text{IST} > 400$ mC) were excluded from the analysis. Age was associated with a higher IST ($r_s=0.271$, $P=0.045$), male gender was associated with a higher IST ($r_s=0.351$, $P=0.009$), and taking an anticholinergic medication was associated with a lower IST ($r_s=-0.295$, $P=0.029$). For other concurrent medications such as benzodiazepines, antipsychotics, antidepressants, and anticonvulsants, BMI and anesthetic agents were not associated significantly with the IST. The goodness of fit was calculated for the multivariate ordinal regression analysis ($\chi^2=318.4$, $\text{df}=1$, $P=0.531$). Age ($P=0.021$), male gender ($P=0.006$), and no concurrent anticholinergic medications ($P=0.025$) were associated with a higher IST.

Comparison of the stimulus intensity using dose-titration and age-based methods

The SI determined using the dose-titration method and the stimulus dose used in the first treatment for both the RUL and BL groups was higher compared with that calculated using the half-age method and lower compared with that calculated using the age method (Table 3).

In the RUL group ($n=32$), using the half-age method, six (18.8%; Figure 1, left) subjects would have received at least

Table 3 Stimulus intensity determined using the dose-titration and age-based methods

	Dose-titration method	Half-age method	Age method
RUL group (n=32)	227±92 (192) (115–576)	158±46 (169) (56–253)	315±92 (338) (112–505)
BL group (n=57)	221.7±119.9 (192) (80–576)	126±39.8 (128) (53–233)	252±79.7 (256) (106–466)

Note: Data are presented as the mean±SD (median), range of stimulus intensity (mC).

Abbreviations: BL, bilateral; RUL, right unilateral.

the SI that was computed using the dose-titration method. However, using the age method, 26 (81.3%) subjects (Figure 1, right) would have received at least the SI that was computed using the dose-titration method. Figure 1 also reveals that ten (31.3%) subjects using the half-age method and seven (21.9%) subjects using the age method would have received a SI within $\pm 20\%$ of that computed using the dose-titration method. Additionally, six (18.8%) subjects using the half-age method and six (18.8%) subjects using the age method would have received a SI at least twice (200%) or less than one-half (50%) of that computed using the dose-titration method. The SI was in the range 1.5–8.6 times the subject's IST, when determined using the half-age method and in the range 2.9–17.2 times the subject's IST, when determined using the age method.

In summary, neither age-based method provided an accurate prediction of the SI (ie, $\pm 20\%$).

For the BL group (n=57), nine (15.8%) subjects using the half-age method (Figure 2, left) and 35 (61.4%) subjects

using the age method (Figure 2, right) would have received at least the SI computed using the dose-titration method. Figure 2 also shows that ten (17.5%) subjects using the half-age method and 18 (31.6%) subjects using the age method would have received an SI within $\pm 20\%$ of that computed using the dose-titration method. Additionally, 22 (38.6%) subjects using the half-age method and ten (17.5%) subjects using the age method would have received a SI at least twice (200%) or less than one-half (50%) of that computed using the dose-titration method. The SI was in the range 0.3–4.8 times the subject's IST for the half-age method and in the range 0.6–9.6 times the subject's IST for the age method.

Discussion

In our study, we found that the mean IST was 42 mC and 150 mC for the RUL and BL group, respectively. The variability in the IST was 6.7-fold for the RUL group and 18-fold for the BL group. As expected, age strongly influenced the IST for both the RUL and BL groups. Additionally, gender and concurrent anticholinergic drugs influenced the IST for the BL group. These findings are consistent with previous studies on IST variability and factors influencing the IST (such as age, sex, anticholinergic drugs).^{15,19,20} The SI determined using the dose-titration method was higher compared with that calculated using the half-age method and lower compared with that calculated using the age method (Table 3). The calculated SI for RUL using either the age or half-age method was in the range ($\pm 20\%$) of the SI obtained using the gold standard dose-titration method for only about

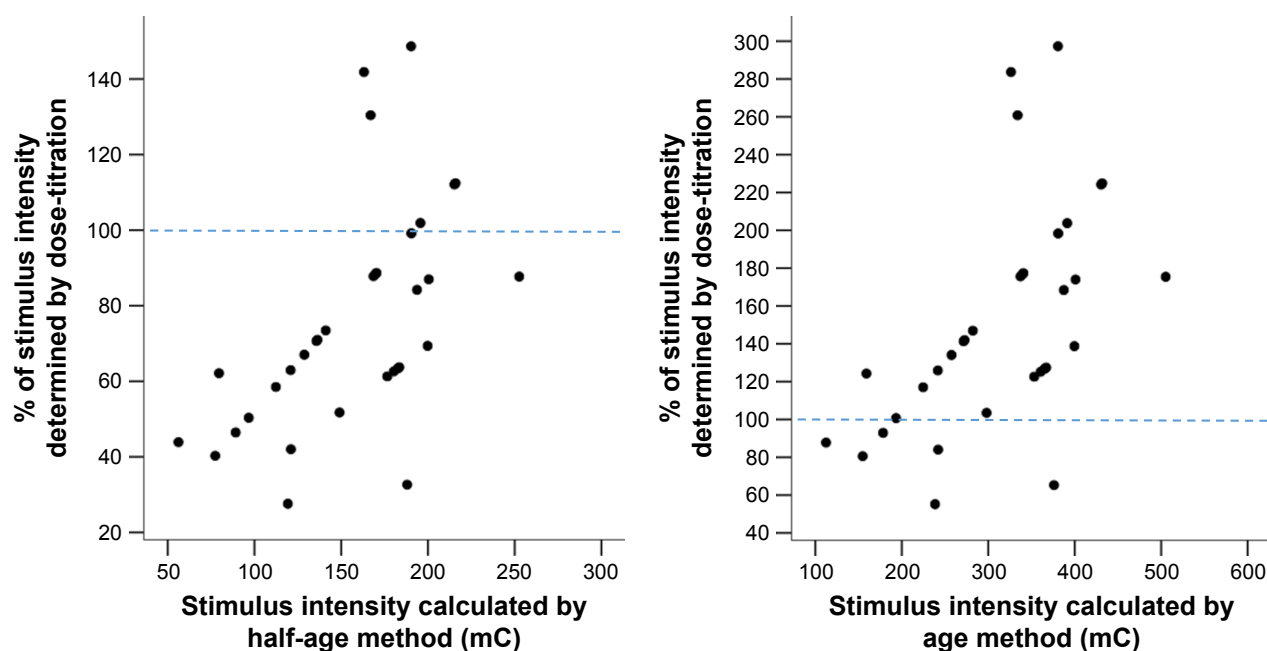


Figure 1 Comparison of the stimulus intensity determined using the dose-titration and age-based methods in the right unilateral group.

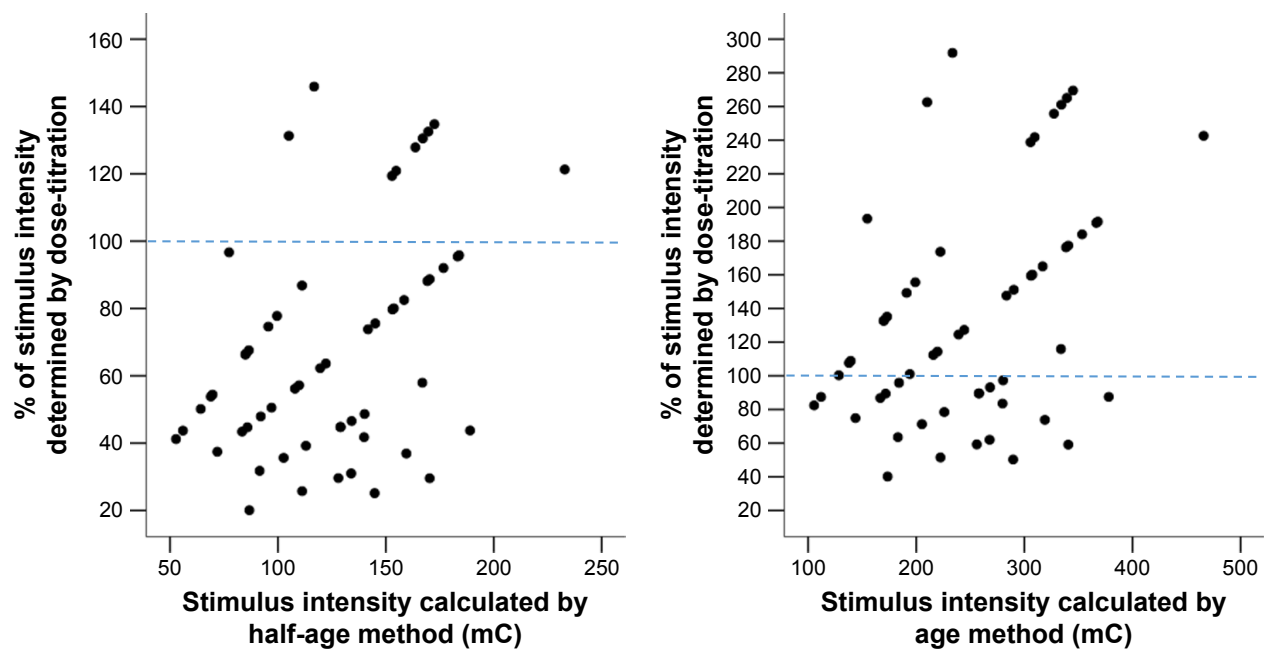


Figure 2 Comparison of the stimulus intensity determined using the dose-titration and age-based methods in the bilateral group.

20%–30% of the subjects. A slightly lower percentage for both groups was markedly outside this range (ie, $>200\%$ or $<50\%$). The overall accuracy of the two methods did not appear to be different, but it is important to note that the SI calculated using the age method is more likely to produce a therapeutic response although there is a commensurate increase in the risk of adverse effects. For the BL treatment, the age method appeared to be consistently better than the half-age method across the different indices. Fifty percent more subjects were within the range determined using the dose-titration method (32% vs 18%), and fewer than half (18% vs 39%) as many were markedly outside this range. Still fewer than half of the subjects appeared to be within the desired range using the age method.

For the RUL group, the SI may be high – up to 17.2 times higher than the subject's IST obtained using the age method. This result was consistent with a previous study from Australia which showed that the SI determined using the age method led to an excessive SI (seven or more times higher than the IST) compared with that calculated using the dose-titration method in 30% of female subjects and 8% of male subjects who were treated with RUL ECT.¹² For the BL group, the SI may be high – up to 4.8 times higher than a subject's IST calculated using the half-age method and 9.6 times higher than a subject's IST calculated using the age method. Although using the SI calculated by the age method would have resulted in successful stimulation on the first attempt in most patients by delivering an overestimated SI, this might cause cognitive side effects.

The study has several limitations. First, subjects included in this study were all inpatients at a university hospital in Thailand, so the results should be interpreted with caution in other settings. Second, this is a retrospective study so some information was missing. Third, there are other factors that might influence the IST, which were not identified in this clinical sample (eg, underlying neurological disease or substance use). Fourth, the IST was based on intervals, not a continuous charge. Fifth, the titration schedule differed by gender and electrode placement. Perhaps most important is that while the dose-titration method is the gold standard, we do not know if or how actual clinical outcomes would vary if the different methods were compared. Furthermore, our measures of adequacy (ie, $\pm 20\%$ was acceptable; $>200\%$ or $<50\%$ was highly undesirable) are somewhat arbitrary. Finally, there may also be other factors influencing the IST that should be taken into account when estimating the threshold. Thus, while we confirm the influence of age, we also confirm the influence of concurrent anticholinergic drugs, which along with BMI, concurrent psychotropic medications, and history of previous ECT may warrant further study.

Conclusion

The two age-based methods did not provide a robust substitute for the SI calculated using the dose-titration method. In particular, both methods showed equally poor results when RUL treatment was applied, and while the age-based method was superior for BL, it still appeared to expose nearly 20% of

patients to unacceptably high or low charges. Thus, the dose-titration method is still recommended to determine the SI.

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Disclosure

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