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Surmounting Retraining Limits in Musicians’ Dystonia by Transcranial Stimulation

Shinichi Furuya, PhD,1 Michael A. Nitsche, MD,2 Walter Paulus, MD,2 and Eckart Altenmüller, MD1

Objective: Abnormal cortical excitability is evident in various movement disorders that compromise fine motor control. Here we tested whether skilled finger movements can be restored in musicians with focal hand dystonia through behavioral training assisted by transcranial direct current stimulation to the motor cortex of both hemispheres.

Methods: The bilateral motor cortices of 20 pianists (10 with focal dystonia, 10 healthy controls) were electrically stimulated noninvasively during bimanual mirrored finger movements.

Results: We found improvement in the rhythmic accuracy of sequential finger movements with the affected hand during and after cathodal stimulation over the affected cortex and simultaneous anodal stimulation over the unaffected cortex. The improvement was retained 4 days after intervention. Neither a stimulation with the reversed montage of electrodes nor sham stimulation yielded any improvement. Furthermore, the amount of improvement was positively correlated with the severity of the symptoms. Bihemispheric stimulation without concurrent motor training failed to improve fine motor control, underlining the importance of combined retraining and stimulation for restoring the dystonic symptoms. For the healthy pianists, none of the stimulation protocols enhanced movement accuracy.

Interpretation: These results suggest a therapeutic potential of behavioral training assisted by bihemispheric, noninvasive brain stimulation in restoring fine motor control in focal dystonia.

Various movement disorders degrade fine motor control. Focal task-specific dystonia is a nondegenerative disease (focal dystonia [FD]) characterized by involuntary movements and muscular spasms.1,2 Among musicians and writers, but also in other professions such as surgeons and golfers, it is accompanied by a reduction of inhibition in motor-related cortical regions,3 including reduced intracortical inhibition,4,5 reduced surround inhibition,6,7 and hyperactivity of the affected motor cortex.8 Thus, to improve fine motor control, a restoration of the disturbed motor cortex activation patterns could be a hypothesis-guided intervention.

Previous studies attempted in vain to use unihemispheric noninvasive excitability-diminishing transcranial direct current stimulation (tDCS) over the affected motor cortex of FD patients to reduce dystonic overactivity.9–12 The present study proposes a novel therapeutic approach that combines bimanual mirrored finger movements with bihemispheric tDCS for pianists with FD. During the bimanual mirror finger movements, motor commands issued by motor cortices spill over to the respective contralateral cortices via the corpus callosum; this is called neural crosstalk.13–15 We postulate that fine motor control of the hand affected by FD can be improved by simultaneously facilitating the neural crosstalk into the affected cortex by increasing activation in the unaffected motor cortex using anodal stimulation, and by suppressing abnormal hyperactivity in the affected motor cortex using cathodal stimulation.16–20 This stimulation can also indirectly facilitate transcallosal inhibitory input to the affected motor cortex.21
Subjects and Methods

Participants
Twenty adult pianists participated in the present experiment (10 with FD, 10 without FD). Ten pianists with FD of the right hand (4 females, 24–61 years old, mean = 39.6 years old) were recruited from the outpatient clinic of the Institute of Music Physiology and Musicians’ Medicine at Hannover University of Music, Drama, and Media (Table 1). Each pianist underwent a thorough neurological examination and was diagnosed by one of the authors (E.A.) specializing in movement disorders of musicians. Exclusion criteria were bilateral FD, generalized dystonia, epilepsy, history of any other neurological diseases, and injection of botulinum toxin A within the past 3 months. Ten pianists with no history of neurological disorders were recruited as controls (4 females, 24–37 years old; Table 2). In accordance with the Declaration of Helsinki, the experimental procedures were explained to all participants. Informed consent was obtained from all participants prior to participation in the experiment, and the experimental protocol was approved by the ethics committee at Hanover Medical School.

Experimental Design
The pianists were asked to participate in 5 experimental sessions with different stimulation protocols (Fig 1A). Each experiment for each participant was separated by >2 weeks to minimize any carryover effect of the stimulation (see Fig 1B). The order of the stimulation protocols was balanced across participants, and the experimental design was double-blinded by asking a person different from the experimenter to set stimulation parameters so that the experimenter and participant were unaware of the ongoing stimulation protocol. However, 2 of the protocols were virtually single-blinded (i.e., unihemispheric stimulation with retraining [UniAu] and bihemispheric stimulation without retraining [NoRT]), and their order was not balanced with regard to the other 3 sessions that were performed earlier.

Each experimental session consisted of pretest, training, and post-test. During training, in 4 of 5 protocols (i.e., all except for NoRT), participants were asked to perform bimanual mirrored finger movements that consist of successive strokes on 4 adjacent keys (right, G-F-E-D; left, C-D-E-F) with the little, ring, middle, and index fingers in synchrony with a metronome (3 strokes per beat, 100 beats per minute, interstroke interval = 200 milliseconds) as evenly as possible. This movement
rate is fast enough to elicit neural crosstalk between the hemispheres. The whole training session consisted of 8 subsessions, each of which consisted of the bimanual playing for 150 seconds and subsequent rest for 30 seconds (3 minutes × 8 sessions). A sequence was played so that a key was not released until the next key was depressed at the loudness of 60% MIDI velocity (mezzo-forte). A pilot study with 3 other patients confirmed no occurrence of muscular fatigue throughout the training session with this tempo, loudness, and duration. During the training, the participants were instructed to guide their attention to the movements of the unaffected left hand for all conditions to facilitate the transfer of fine motor skill from that hand to the affected right hand, and also to facilitate plasticity of the motor cortex contralateral to the attended hand. After the training session, each participant took a rest for 5 minutes to minimize muscular fatigue. For the remaining protocol (ie, NoRT), participants did not perform the abovementioned movements during the session, thus tDCS was applied under resting conditions.

During the training session, 2 active water-soaked tDCS electrodes were put on C3 and C4 locations (primary motor cortex) in 4 of 5 protocols (ie., all except for UniAu), which were identified using the international 10–20 electroencephalogram system. The rationale behind stimulating the motor cortex was that this region represents the motor program responsible for skilled finger movement. The current montage was adopted to modulate cortical excitability of both hemispheres simultaneously. To minimize current shunt between the electrodes over the scalp, the location of the electrodes was carefully selected so that the distance between the edges of the electrodes was at least 6cm. tDCS was applied throughout the entire training session, which lasted for 24 minutes. The 4 stimulation protocols were referred to as CaAu, CuAa, sham, and NoRT. For the CaAu, sham, and NoRT conditions, the cathodal electrode was placed on the affected left hemisphere and the anodal (excitability enhancement) on the unaffected right side of the cortex, and vice versa for the CuAa condition. The UniAu condition involved bimanual training with unihemispheric anodal stimulation over the unaffected hemisphere. The stimulation duration was 24 minutes for CaAu, CuAa, UniAu, and NoRT, and 30 seconds for sham. (B) Time flow of a series of experiments. Each experiment for each participant was separated by >2 weeks.

Data Acquisition and Analysis
We recorded the time of each keystroke in pretest, training, and post-test conditions. The standard deviation of the interkey-stroke intervals across strokes was computed as an index of rhythmic variability of sequential keystrokes. High evenness of keystrokes with only very little rhythmic variability indicates a high degree of fine motor control. For the untrained task (repetitive tapping by the affected finger), the variability of keystrokes of the participants who had multiple fingers affected was evaluated by averaging rhythmic variability of the keystrokes across the affected fingers.
Statistics
To assess rhythmic variability at the pretest and post-test, and the difference for both the patient and control groups, a 2-way analysis of variance (ANOVA) with mixed design using stimulation protocol as the within-factor (CaAu, CuAa, sham, UniAu, and NoRT) and group as the between-factor (healthy pianists and pianists with FD) was designed. We excluded age-related effects by regression analysis, which yielded no significant regression between age and each of the variables (p > 0.05). Tukey post hoc tests with multiple comparison correction were performed in the case of significant results of the ANOVA. To determine factors associated with individual differences of stimulation effects on rhythmic variability of keystrokes, a correlation analysis was performed with severity of the symptom (i.e., initial rhythmic variability of keystrokes at the pretest), actual age of pianists and patients, and time period indicating the elapsed time between diagnosis of FD and participation in the study. For the training session, a 2-way repeated-measures ANOVA with session and stimulation protocol as between-factors was also performed for the patient group. To evaluate the effect of training, we subtracted rhythmic variability of the initial 5 seconds of the first training session (baseline performance) from the variability of keystrokes averaged within each of 8 training sessions of the affected right hand by the patients. This value was used as a dependent variable for the repeated-measures ANOVA, to assess improvements in motor performance over the training sessions. In addition, rhythmic variability during the 5-second baseline was assessed by 2-way ANOVA, to confirm that there was no difference across stimulation protocols.

Results
First, a 2-way mixed-design ANOVA using group and protocol as independent variables was performed, to confirm that the initial baseline performance prior to the training did not differ across the protocols. Neither main effect of protocol nor interaction effect of protocol and group was significant for rhythmic variability of the keystrokes during both the pretest of the trained sequential strokes and the baseline of the training (Table 3). Similarly, rhythmic variability of the single-finger tapping in the pretest yielded no main effect of protocol for the affected finger of the right hand in the patient group ($F_{4,36} = 2.47$, $p = 0.06$) and for all right fingers in the healthy group ($F_{4,36} = 0.79$, $p = 0.54$).

Figure 2A illustrates the group mean of the standard deviation of the interkeystroke intervals across strokes (i.e., rhythmic variability of keystrokes) after the training session (i.e., post-test) with each of the 5 stimulation protocols while FD patients and healthy individuals played a trained sequence with the affected hand. For the patient group, rhythmic variability for the CaAu protocol was smaller than for the other protocols. By contrast, none of the stimulation protocols elicited any discernible change in the healthy group. The 2-way mixed design ANOVA yielded a significant interaction effect of group and stimulation protocol ($F_{4,72} = 2.97$, $p = 0.02$). Post hoc tests confirmed significant differences between the CaAu and each of the CuAa, sham, and NoRT protocols in the patient group. In addition, a group difference was significant for the protocols except only for the CaAu. The findings indicate that CaAu stimulation elicited an improvement in rhythmic accuracy of keystrokes with the affected hand of the FD patients. Similarly, the group mean of the amount of change in rhythmic variability of keystrokes after the training test (i.e., post-test – pretest) for the CaAu, CuAa, sham, UniAu, and NoRT conditions was $-8.5 \pm 12.9$ milliseconds, $3.2 \pm 6.7$ milliseconds, $0.9 \pm 5.2$ milliseconds, $0.3 \pm 2.5$ milliseconds, and $0.6 \pm 3.6$ milliseconds for the patient group, and $-0.7 \pm 2.3$ milliseconds, $-1.5 \pm 3.0$ milliseconds, $-1.4 \pm 2.2$ milliseconds, $-0.2 \pm 2.9$ milliseconds, and $-0.3 \pm 1.6$ milliseconds for the healthy group, respectively. A 2-way mixed design ANOVA confirmed an interaction effect of group and protocol ($F_{4,72} = 3.89$, $p = 0.01$), and post hoc tests identified a significant difference between the CaAu and each of the other 4 protocols ($p < 0.01$ for CuAa and sham, $p < 0.05$ for UniAu and NoRT). None of any other protocol pairs yielded a significant difference.

By contrast, for the unaffected hand, rhythmic variability of the keystrokes in the post-test did not differ across the stimulation protocols (group × protocol: $F_{4,72} = 0.34$, $p = 0.85$ by 2-way mixed-design ANOVA; no group difference for any protocols in post hoc tests). The group mean of rhythmic accuracy of keystrokes in the post-test for the CaAu, CuAa, sham, UniAu, and NoRT protocols was $12.2 \pm 3.0$, $11.3 \pm 2.7$, $11.9 \pm 3.6$, $11.1 \pm 3.0$, and $11.1 \pm 2.8$ in the patient group, and $11.0 \pm 2.7$, $11.2 \pm 3.2$, $10.7 \pm 3.2$, $11.0 \pm 3.2$, and $10.6 \pm 2.5$ in the healthy group, respectively.

Figure 2B displays the relation between the amount of change in rhythmic variability of the keystrokes while the FD patients were playing the trained sequence with the affected hand and rhythmic variability of the keystrokes in the pretest (i.e., severity of the symptom) for the CaAu stimulation condition. A significant negative correlation was evident between these factors ($r = -0.92$, $p = 1.4 \times 10^{-4}$). By contrast, none of the other 4 conditions yielded any significant negative correlation ($r = 0.76$ and $p = 0.01$ for CuAa, $r = -0.38$ and $p = 0.28$ for sham, $r = -0.54$ and $p = 0.11$ for UniAu, $r = -0.53$ and $p = 0.12$ for NoRT). This finding indicates greater effects of the CaAu stimulation on rhythmic accuracy of keystrokes for patients with more severe symptoms of FD. In the healthy group, these variables
### TABLE 3. Rhythmic Variability of the Keystrokes at Pretest, Post-Test, and Testing Baseline

<table>
<thead>
<tr>
<th>Hand</th>
<th>Time Point</th>
<th>Group</th>
<th>CaAu</th>
<th>CuAa</th>
<th>sham</th>
<th>UniAu</th>
<th>NoRT</th>
<th>ANOVA Results</th>
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</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>SD of the inter-keystroke interval across strokes (ms)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Affected hand</td>
<td>pretest</td>
<td>Patient</td>
<td>20.7 (12.8)</td>
<td>15.0 (4.6)</td>
<td>16.7 (8.9)</td>
<td>16.6 (5.0)</td>
<td>18.1 (7.9)</td>
<td>F(1,18) 10.22 p 0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Control</td>
<td>11.4 (3.8)</td>
<td>12.1 (3.6)</td>
<td>11.9 (3.7)</td>
<td>11.7 (2.7)</td>
<td>10.9 (2.5)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>posttest</td>
<td>Patient</td>
<td>12.2 (5.0)</td>
<td>18.3 (10.6)</td>
<td>17.6 (8.4)</td>
<td>16.9 (4.3)</td>
<td>18.6 (6.7)</td>
<td>F(1,18) 8.37 p 0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Control</td>
<td>10.7 (2.3)</td>
<td>10.6 (2.9)</td>
<td>10.5 (2.5)</td>
<td>11.5 (2.8)</td>
<td>10.7 (2.0)</td>
<td></td>
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<tr>
<td>Unaffected hand</td>
<td>pretest</td>
<td>Patient</td>
<td>13.0 (9.4)</td>
<td>11.0 (2.6)</td>
<td>10.5 (2.3)</td>
<td>11.2 (2.5)</td>
<td>11.4 (2.0)</td>
<td>F(1,18) 0.82 p 0.38</td>
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<tr>
<td></td>
<td></td>
<td>Control</td>
<td>10.8 (2.7)</td>
<td>11.8 (2.6)</td>
<td>10.8 (2.8)</td>
<td>11.1 (1.7)</td>
<td>10.8 (2.7)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>posttest</td>
<td>Patient</td>
<td>12.2 (3.0)</td>
<td>11.3 (2.7)</td>
<td>11.9 (3.6)</td>
<td>11.1 (3.0)</td>
<td>11.1 (2.8)</td>
<td>F(1,18) 0.35 p 0.56</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Control</td>
<td>11.0 (2.7)</td>
<td>11.2 (3.2)</td>
<td>10.7 (3.2)</td>
<td>11.0 (3.2)</td>
<td>10.6 (2.5)</td>
<td></td>
</tr>
<tr>
<td>Affected hand</td>
<td>baseline (training)</td>
<td>Patient</td>
<td>35.9 (24.3)</td>
<td>25.7 (13.2)</td>
<td>26.3 (14.9)</td>
<td>29.9 (7.7)</td>
<td></td>
<td>F(1,18) 21.46 p 2.1×10^-4</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Control</td>
<td>11.6 (3.7)</td>
<td>12.0 (3.8)</td>
<td>12.1 (3.5)</td>
<td>12.0 (2.9)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- A number in parenthesis indicates the standard deviation within a group.
- A bold number indicates p < 0.05.
- baseline: movement variability at the affected hand during the first 5 seconds of the training session.
- A group difference was significant at both the pretest with the affected hand and baseline, but not at the pretest with the unaffected hand.
- CaAu: cathodal over the affected cortex and anodal over the unaffected cortex, CuAa: the reverted montage of CaAu,
- UniAu: unilateral anodal over the unaffected cortex, NoRT: CaAu stimulation without retraining.

Furuya et al: Noninvasive Stimulation for FD

Month 2014 5
Linear regression analysis. Negative values indicate a decrease in the data of one patient. A line was drawn based on the result of all pianists with FD in the CaAu protocol. Each point indicates rhythmic variability of keystrokes following training (measured in affected hand between pretest and post-test conditions (y-axis) and difference in rhythmic variability of keystrokes with the pretest (i.e., severity of the symptom; x-axis) over 8 sessions of training during CaAu stimulation without the retraining.

Figure 2C illustrates the group mean of variability changes in keystrokes with the affected hand relative to baseline performance over the training sessions for each of the stimulation protocols in the patient group. A decrease was evident for the CaAu condition, but not for the CuAa, UniAu, and sham conditions. A 2-way repeated-measures ANOVA revealed a significant interaction effect between session and stimulation protocol ($F_{21,189} = 2.93, p = 0.01$) and a main effect of session ($F_{4,36} = 2.30, p = 0.04$). This confirms a larger decline of movement variability over sessions for the CaAu protocol as compared to the other protocols. Post hoc tests found significant differences between the CaAu and each of the CuAa and sham protocols from the 3rd to 8th sessions, and between the CaAu and UniAu protocols from the 6th to 8th sessions, showing a smaller variability for the CaAu protocol.

To assess the duration of the aftereffect of the observed improvement of fine motor control, 4 of the 10 patients were again invited to the experiment. They underwent retraining with the CaAu condition, and 4 days after intervention played the trained sequence. The group mean of the keystroke variability was $15.3 \pm 1.6$ milliseconds prior to the stimulation (pretest), $10.1 \pm 1.7$ milliseconds immediately after the stimulation (post-test), and $11.7 \pm 1.3$ milliseconds 4 day after the retraining (retention test). The immediate effect of the retraining was therefore retained by $71.2 \pm 5.2\%$ at the retention test. One-way repeated-measures ANOVA yielded a main effect of test ($F_{2,6} = 30.6, p = 0.0007$), and Tukey post hoc tests identified a larger value at the pretest than the post-test ($p = 0.003$) and retention test ($p = 0.02$), but no difference between the post-test and retention test ($p = 0.36$). The result confirms retention of the retraining effect for at least 4 days.

For the untrained finger-tapping task, the group mean of the amount of change in rhythmic variability of the tapping by the affected finger was $-4.7 \pm 14.0$, $-7.2 \pm 8.1$, $12.1 \pm 23.5$, $-7.1 \pm 10.5$, and $-5.5 \pm 18.9$ milliseconds for the CaAu, CuAa, sham, UniAu, and NoRT protocols, respectively. One-way repeated-measures ANOVA yielded no main effect of stimulation ($F_{3,66} = 0.34, p = 0.85$), which indicates a lack of transfer effects of the training on performance of the untrained tapping.

FIGURE 2: (A) Group mean of variability of the interkeystroke interval after the training session (i.e., post-test) while pianists with focal dystonia (FD; left) and healthy pianists (right) were playing a sequence of keystrokes with the affected hand for each of the 5 stimulation protocols. Stimulation protocols (CaAu, CuAa, sham, UniAu, and NoRT) are represented on the x-axis. Error bars indicate the standard error within each of the groups. CaAu = cathodal transcranial direct current stimulation (tDCS) over the affected (left) cortex and anodal tDCS over the unaffected (right) cortex; CuAa = cathodal tDCS over the unaffected cortex and anodal tDCS over the affected cortex; UniAu = anodal tDCS over the unaffected cortex and cathodal tDCS over the contralateral orbit; NoRT = CaAu stimulation without the retraining. *$p < 0.05$, **$p < 0.01$. (B) Relation between the amount of rhythmic variability at the pretest (i.e., severity of the symptom; x-axis) and difference in rhythmic variability of keystrokes with the affected hand between pretest and post-test conditions (y-axis) of all pianists with FD in the CaAu protocol. Each point indicates the data of one patient. A line was drawn based on the result of linear regression analysis. Negative values indicate a decrease in rhythmic variability of keystrokes following training (measured in milliseconds). (C) Group mean of changes in rhythmic variability of keystrokes with the affected hand of pianists with FD over 8 sessions of training during CaAu (filled circle), CuAa (open triangle), sham (open square), and UniAu (cross) stimulation protocols. Negative values indicate a decrease in rhythmic variability of the keystrokes relative to baseline (i.e., initial 5 seconds of the first training session).
Discussion
FD is a disease extremely difficult to treat. In musicians, it frequently leads to early termination of their professional career. Retraining techniques either by repetitive movement training following the principles of induced beneficial brain plasticity by sensory–motor retuning of the affected hand, or by behavioral therapies including the slow-down technique and awareness training, remain promising therapeutic attempts. Here, we report that combining motor training with tDCS seems to be required to produce improvement on the task selected for training. This progress seems to require a combined use of bihemispheric tDCS with bimanual mirrored finger movements to elicit improvement of rhythmic accuracy of a trained sequence of finger movements. It occurs exclusively when stimulating the affected motor cortex with the cathodal and the unaffected motor cortex with the anodal electrode. So far, none of the previous studies has restored loss of inhibition at the affected motor cortex and fine motor control in FD patients using unihemispheric cathodal tDCS with the return electrode at the contralateral forehead position, even if this stimulation was combined with a retraining procedure. Bihemispheric tDCS possesses 2 possible advantages over unihemispheric stimulation. First, it is assumed to directly increase interhemispheric inhibition into the affected motor cortex during retraining and thereby might facilitate its intracortical inhibitory mechanisms. This concept was recently supported by bihemispheric tDCS with the reversed montage for stroke patients, which is assumed to decrease interhemispheric inhibition to the affected cortex. Second, bihemispheric stimulation might augment the fraction of motor commands transmitted from the unaffected to the affected cortex via the corpus callosum, and thereby normalize abnormal motor programs. This concept is supported by a lack of improvement in the fine motor control both of untrained single finger tapping following the stimulation with retraining and of trained sequential finger movements following the stimulation without retraining, making it likely that the current training specifically restored control of the trained motor skill of pianists with FD. Attention could also modulate the stimulation effect so as to suppress cortical excitation and to enhance the capacity of plastic alterations.

We acknowledge that a limitation of this study is the lack of a cortical excitability or activity assessment before and after stimulation, which would have helped to clarify mechanisms of action at the physiological level. In future studies, we will include such an assessment, which was not possible in the present design for logistical reasons. We also would like to point out that although the experimental results of our study are clear, future studies need to show if and to what extent these results translate into meaningful therapeutic improvements when combined with prolonged repeated training protocols. Nevertheless, we think that the current tDCS protocol offers qualitatively new options to improve rehabilitative efforts in occupational dystonia. Another limitation is that effects of neither unihemispheric cathodal stimulation over the affected cortex during retraining nor bihemispheric stimulation before or after retraining were evaluated due to limited feasibility in the patients.

Authorship
S.F. participated in the design of the study, carried out the experiments, analyzed data, performed the statistical analysis, and drafted the manuscript. M.A.N. participated in the design and coordination of the study and helped to interpret the data and draft the manuscript. W.P. participated in the design and coordination of the study and helped to interpret the data and draft the manuscript. E.A. participated in the design and coordination of the study, recruited and diagnosed the patients, and helped to draft the manuscript. All authors read and approved the final manuscript.

Potential Conflicts of Interest
Nothing to report.

References


