Long-term treatment of bipolar disorder with a radioelectric asymmetric conveyor

Background: The bipolar spectrum disorders are considered an important and frequent psychiatric problem. The clinical complexity of these illnesses due to the coexistence of depressive and excitative phases is correlated with the global difficulty of adequate treatment; consequently, the prognosis is not optimal. For this reason, in recent years, novel nonpharmacologic physical approaches have been tested for bipolar disorders, with encouraging results. The aim of this study was to evaluate the long-term effectiveness of a radioelectric asymmetric brain stimulation device associated with lithium (REAC-lithium) versus previous treatments in subjects with bipolar disorder I or II, evaluated as the number of recurrences compared with the period of illness preceding treatment with REAC-lithium.

Methods: The charts of 56 bipolar patients attending our institute were retrospectively evaluated. Treatment with REAC-lithium was administered following the standard Rinaldi-Fontani Institute protocol. Add-on treatments were allowed in the event of manic or depressive recurrence. Eight patients (Group 1) were followed for 30.2 ± 3.0 months, 14 patients (Group 2) were followed for 25.3 ± 3.3 months, 25 patients (Group 3) were followed for 20.3 ± 1.6 months, and nine patients (Group 4) were followed for 16.2 ± 0.5 months.

Results: After REAC-lithium treatment, the number of manic and depressive episodes in Group 1 decreased from 2.1 ± 0.6 and 3.0 ± 0.7 to 0.12 ± 0.0 and 0.8 ± 0.4, respectively. In Group 2, the number of manic and depressive episodes decreased from 2.4 ± 0.6 and 3.9 ± 0.7 to 0.14 ± 0.2 and 0.0 ± 0.0, respectively. In Group 3, the number of manic and depressive episodes decreased from 2.6 ± 0.8 and 3.6 ± 0.9 to 0.04 ± 0.0 and 0.0 ± 0.0, respectively. In Group 4, the number of manic and depressive episodes decreased from 2.6 ± 1.1 and 3.7 ± 1.0 to 0.1 ± 0.0 and 0.0 ± 0.0, respectively. All results were statistically significant.

Conclusion: REAC showed good efficacy in treating both the manic and depressive phases of bipolar disorder, and in the prevention of recurrences/relapses.

Keywords: bipolar disorder, stress, radioelectric asymmetric brain stimulation

Introduction

The concept of the “bipolar spectrum” as a consequence of more accurate diagnostic assessment has progressively increased the classic epidemiologic boundary of bipolar disorder, historically affecting 0.8%–1.5%² to 15%³ of the general population. In the same way, the complexity of the illness has posed the need for dimensional (cross-sectional and longitudinal) monitoring of the patient, in order to design more personalized and phase-specific therapeutic strategies. Thus, from a therapeutic point of view, the classic drugs for bipolar disorder (lithium, carbamazepine, valproic acid, antidepressants, and neuroleptics)⁴ have been progressively supported with the new...
generation of anticonvulsants (topiramate, gabapentin, pregabalin, lamotrigine, and levetiracetam),5,6 and the atypical antipsychotics.7,8 Although the excitative phases (manic and hypomanic) are usually considered as the organizing principle of bipolar disorder,9 pooled data for the disease course show that, with exclusion of free intervals, nearly 70% of the “bipolar life” consists of depressive episodes and the remaining 30% consists of excitative phenomena.9,10 The main result of this hierarchical approach, based on the course of bipolar disorder, is a different pharmacologic treatment strategy for the manic phenomena and for bipolar depression.11,12 Despite recent efforts,13 the results of treatment of bipolar depression are not encouraging.14 The use of lamotrigine, after an initial enthusiastic introduction, has gradually reduced as an acute antidepressant in patients with bipolar disorder, and is currently used for the prevention of depressive recurrences.15–17 New pharmacologic data for atypical antipsychotics could be considered promising. These drugs seem to be useful in the depressive phase (without a manic switch) and in the manic phase, avoiding the depressive and apathetic motivational risk associated with the administration of typical antipsychotics. Unfortunately, the data available regarding the role of these drugs in the maintenance of bipolar disorder are fewer than expected.8 Moreover, it is important to emphasize that lithium remains the first-choice drug for bipolar disorder, and is currently used for the prevention of depressive recurrences.8,9

Methods and materials

The charts of 56 patients attending the Rinaldi-Fontani Institute in Florence and the Centro di Studi Psichici in Cagliari were retrospectively evaluated. Patients diagnosed with bipolar disorder I or II according to the revised fourth edition of the Diagnostic Statistical Manual for Mental Disorders (DSM-IV-TR) criteria were included. Inclusion criteria were as follows: both genders; age 18–65 years; written informed consent obtained; history of at least 12 months of psychopharmacologically treated bipolar disorder I/II; and at least four cycles of REAC treatment. Exclusion criteria included a history of seizures, head trauma, or brain disease, and evidence of chronic or severe disease (renal or hepatic impairment, or cancer). REAC treatment was administered following the standard Rinaldi-Fontani Institute schedule of one session of neuropsychophysical optimization followed by 18 sessions of neuropsychophysical optimization. Add-on treatments were allowed in the event of manic or depressive recurrence. Patients were divided into four follow-up groups before and after REAC-lithium treatment, as follows: at least 72 months (Group 1); at least 48 months (Group 2); at least 36 months (Group 3); and at least 18 months (Group 4). Information on demographic variables, psychiatric history, concomitant treatments, and comorbid disorders were collected for each patient. The evaluation of REAC-lithium effectiveness was made by collecting the recurrence of mood episodes (manic or depressive according to DSM IV-TR). The Structured Clinical Interview for Diagnosis, the 21-item Hamilton Depression Rating Scale, and the Young Mania Rating Scale (YMRS) were used. Statistical analysis was performed using a t-test for comparative groups, and all results with a $P < 0.05$ were considered significant. Tolerability and safety were evaluated by collecting the reports of adverse events. Demographic variables and the psychiatric-psychopharmacologic history of the patients are reported in Table 1. Eight patients (Group 1) were followed for 30.2 ± 3.0 months, 14 patients (Group 2) for 25.3 ± 3.3 months, 25 patients (Group 3) for 20.3 ± 1.6 months, and nine patients (Group 4) for 16.2 ± 0.5 months. Lithium was dosed at 980.0 ± 135.6 mg/day, 825.0 ± 50.0 mg/day, 850.5 ± 100.0 mg/day, and 950.5 ± 75.5 mg/day in the four groups of patients, respectively. The demographic variables and psychiatric history of the patients are reported in Table 1.

REAC26,27 is a medical device based on innovative technology for biostimulation. REAC works with a typical range frequency of 2.4, 5.8, or 10.5 GHz, as selected by the operator for each specific protocol used. A frequency of 10.5 GHz was used in the brain stimulation protocols used in this study, with a specific absorption rate28–30 of 7 µW/kg. REAC treatments have proven efficacy in ameliorating stress-related disorders, depression, and anxiety.20–24 The REAC pulse used in this
and during the depression-free period in the remaining four patients. Mean baseline and onset ages for Group 2 (seven males [four bipolar disorder I and three bipolar disorder II], and seven females [all bipolar disorder III]) were 29.9 ± 5.0 years and 21.2 ± 4.2 years, respectively. Mean numbers of previous manic and depressive episodes per patient were 2.4 ± 0.6 and 3.9 ± 0.7, respectively, with an overall number of nine hospitalizations for depressive episodes. The onset of bipolar disorder was depressive in nine patients and manic in five patients. Patients in Group 2 were treated with lithium alone (n = 6), lithium added on to valproate (n = 3), lithium added on to olanzapine (n = 2), lithium added on to fluoxetine (n = 1), and carbamazepine (n = 2). The switch to REAC-lithium treatment occurred during a depressive phase in nine subjects and during the free period in the remaining five patients.

Mean baseline and onset ages for Group 3 (11 males [four bipolar disorder I and seven bipolar disorder II], and 14 females [three bipolar disorder I and 11 bipolar disorder II]) were 28.3 ± 3.7 years and 21.0 ± 3.5 years, respectively. The mean numbers of previous manic and depressive episodes per patient were 2.6 ± 0.8 and 3.6 ± 0.9, respectively, with an overall number of seven hospitalizations (four for depressive episodes and three for manic episodes). The onset of bipolar disorder was depressive in 17 patients and manic in eight patients.

Ten patients were treated with lithium alone, six with lithium added on to valproate, three with lithium added on to olanzapine, one with lithium added on to sertraline, two with lithium added on to imipramine, and three with carbamazepine. The switch to REAC-lithium treatment occurred during a depressive phase in 15 subjects and during the free period in the remaining 10 patients.

Mean baseline and onset ages for Group 4 (three males [one bipolar disorder I and two bipolar disorder II], and six females [two bipolar disorder I and four bipolar disorder II]) were 30.0 ± 3.9 years and 23.1 ± 4.0 years, respectively. The mean number of previous manic and depressive episodes were 2.6 ± 1.1 and 3.7 ± 1.0, respectively, with an overall number of two hospitalizations (one due to a depressive episode and one due to a manic episode). The onset of bipolar disorder was depressive in seven patients and manic in two patients. Previous treatments were as follows: lithium alone (n = 2); lithium added on to valproate (n = 3); olanzapine (n = 1); and lithium added on to citalopram (n = 3). The switch to REAC-lithium treatment occurred during a depressive phase in five subjects and during the free period in the remaining four patients.

Results

Before REAC-lithium therapy

Mean baseline and onset ages for Group 1 (seven males and one female; all bipolar disorder I) were 31.4 ± 2.9 years and 23.9 ± 2.7 years, respectively. The mean numbers of previous manic and depressive episodes per patient were 2.1 ± 0.6 and 3.0 ± 0.7, respectively, and the overall number of hospitalizations was seven (five due to depressive episodes and two due to a manic mood state). Bipolar disorder onset was depressive in three patients and manic in five patients.

All patients in Group 1 had comorbid pathologic gambling. Lithium was satisfactory for mood disorder, but ineffective for pathologic gambling. The switch to REAC-lithium treatment occurred during a depressive phase in three subjects and during the depression-free period in the remaining four patients. Mean baseline and onset ages for Group 2 (seven males [four bipolar disorder I and three bipolar disorder II], and seven females [all bipolar disorder III]) were 29.9 ± 5.0 years and 21.2 ± 4.2 years, respectively. Mean numbers of previous manic and depressive episodes per patient were 2.4 ± 0.6 and 3.9 ± 0.7, respectively, with an overall number of nine hospitalizations for depressive episodes. The onset of bipolar disorder was depressive in nine patients and manic in five patients. Patients in Group 2 were treated with lithium alone (n = 6), lithium added on to valproate (n = 3), lithium added on to olanzapine (n = 2), lithium added on to fluoxetine (n = 1), and carbamazepine (n = 2). The switch to REAC-lithium treatment occurred during a depressive phase in nine subjects and during the free period in the remaining five patients.

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Mean baseline and onset ages for Group 4 (three males [one bipolar disorder I and two bipolar disorder II], and six females [two bipolar disorder I and four bipolar disorder II]) were 30.0 ± 3.9 years and 23.1 ± 4.0 years, respectively. The mean number of previous manic and depressive episodes were 2.6 ± 1.1 and 3.7 ± 1.0, respectively, with an overall number of two hospitalizations (one due to a depressive episode and one due to a manic episode). The onset of bipolar disorder was depressive in seven patients and manic in two patients. Previous treatments were as follows: lithium alone (n = 2); lithium added on to valproate (n = 3); olanzapine (n = 1); and lithium added on to citalopram (n = 3). The switch to REAC-lithium treatment occurred during a depressive phase in five subjects and during the free period in the remaining four patients.

Table 1 Overall demographic variables and psychiatric history before REAC-lithium treatment

<table>
<thead>
<tr>
<th>Gender, n (%)</th>
<th>Male</th>
<th>Female</th>
<th>Mean (SD)</th>
<th>Range</th>
<th>Mean (SD)</th>
<th>Range</th>
<th>Diagnosis, n (%)</th>
<th>Type of onset, n (%)</th>
<th>Episodes/patient mean (SD)</th>
<th>Hospitalization, n</th>
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<tbody>
<tr>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td>BD I</td>
<td>Manic</td>
<td>Depressive</td>
<td>Reason: Excitative episode</td>
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<tr>
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<td></td>
<td>BD II</td>
<td>Depressive</td>
<td></td>
<td>6</td>
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</tbody>
</table>
| Mean baseline and onset ages for Group 1 (seven males and one female; all bipolar disorder I) were 31.4 ± 2.9 years and 23.9 ± 2.7 years, respectively. The mean numbers of previous manic and depressive episodes per patient were 2.1 ± 0.6 and 3.0 ± 0.7, respectively, and the overall number of hospitalizations was seven (five due to depressive episodes and two due to a manic mood state). Bipolar disorder onset was depressive in three patients and manic in five patients. All patients in Group 1 had comorbid pathologic gambling. Lithium was satisfactory for mood disorder, but ineffective for pathologic gambling. The switch to REAC-lithium treatment occurred during a depressive phase in three subjects and during the depression-free period in the remaining four patients. Mean baseline and onset ages for Group 2 (seven males [four bipolar disorder I and three bipolar disorder II], and seven females [all bipolar disorder III]) were 29.9 ± 5.0 years and 21.2 ± 4.2 years, respectively. Mean numbers of previous manic and depressive episodes per patient were 2.4 ± 0.6 and 3.9 ± 0.7, respectively, with an overall number of nine hospitalizations for depressive episodes. The onset of bipolar disorder was depressive in nine patients and manic in five patients. Patients in Group 2 were treated with lithium alone (n = 6), lithium added on to valproate (n = 3), lithium added on to olanzapine (n = 2), lithium added on to fluoxetine (n = 1), and carbamazepine (n = 2). The switch to REAC-lithium treatment occurred during a depressive phase in nine subjects and during the free period in the remaining five patients.

Mean baseline and onset ages for Group 3 (11 males [four bipolar disorder I and seven bipolar disorder II], and 14 females [three bipolar disorder I and 11 bipolar disorder II]) were 28.3 ± 3.7 years and 21.0 ± 3.5 years, respectively. The mean numbers of previous manic and depressive episodes per patient were 2.6 ± 0.8 and 3.6 ± 0.9, respectively, with an overall number of seven hospitalizations (four for depressive episodes and three for manic episodes). The onset of bipolar disorder was depressive in 17 patients and manic in eight patients.

Ten patients were treated with lithium alone, six with lithium added on to valproate, three with lithium added on to olanzapine, one with lithium added on to sertraline, two with lithium added on to imipramine, and three with carbamazepine. The switch to REAC-lithium treatment occurred during a depressive phase in 15 subjects and during the free period in the remaining 10 patients.

Mean baseline and onset ages for Group 4 (three males [one bipolar disorder I and two bipolar disorder II], and six females [two bipolar disorder I and four bipolar disorder II]) were 30.0 ± 3.9 years and 23.1 ± 4.0 years, respectively. The mean number of previous manic and depressive episodes were 2.6 ± 1.1 and 3.7 ± 1.0, respectively, with an overall number of two hospitalizations (one due to a depressive episode and one due to a manic episode). The onset of bipolar disorder was depressive in seven patients and manic in two patients. Previous treatments were as follows: lithium alone (n = 2); lithium added on to valproate (n = 3); olanzapine (n = 1); and lithium added on to citalopram (n = 3). The switch to REAC-lithium treatment occurred during a depressive phase in five subjects and during the free period in the remaining four patients.

<table>
<thead>
<tr>
<th>Previous treatments, n</th>
<th>Lithium alone</th>
<th>Carbamazepine</th>
<th>Olanzapine</th>
<th>Lithium + valproate</th>
<th>Lithium + olanzapine</th>
<th>Lithium + SSRI</th>
<th>Lithium + imipramine</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>26</td>
<td>5</td>
<td>5</td>
<td>12</td>
<td>5</td>
<td>5</td>
<td>2</td>
</tr>
</tbody>
</table>

Abbreviations: BD, bipolar disorder; SD, standard deviation; SSRI, selective serotonin reuptake inhibitor.
After REAC-lithium therapy

Tables 2–5 and Figures 1–4 compare the overall mood episodes (manic and depressive) per Group before and after REAC-lithium treatment. In Group 1, three manic and one depressive episode occurred. In terms of percentages, there were 0.1 depressive episodes per patient and 0.8 ± 0.4 excitative episodes per patient, which was statistically significant in comparison with previous treatments (P < 0.001 [t-test]).

Manic episodes occurred during months 19, 22, and 23 of REAC-lithium treatment. All manic episodes were mild (YMRS scores of 24, 23, and 23, respectively), and their mean duration was 8.0 ± 0.9 weeks. The lithium dose was increased by approximately 20%, and valproic acid 1000 mg/day and zuclopenthixol 100 mg/day in one case and zuclopenthixol 125 mg/day and clonazepam 6 mg/day in the remaining two patients. The depressive episode, classified mild (Hamilton Depression Rating Scale = 21) occurred during month 23 of treatment with REAC-lithium treatment and was resolved by means of augmentation with bupropion 300 mg/day over a six-week period. In Group 2, only two manic episodes occurred during months 16 and 19 of REAC-lithium treatment. In terms of percentages, there were 0.11 ± 0.2 excitative episodes per patient, which was statistically significant in comparison with previous treatments (P < 0.001 [t-test]). Both were mild (YMRS = 22) with a mean length of 7.2 ± 0.3 weeks and were treated with valproate 1000 mg/day and zuclopenthixol (mean dose 87.5 ± 17.7 mg/day). In Group 3, the one manic episode which occurred during month 17 of REAC-lithium treatment was mild (YMRS = 25), with a length of 8.0 weeks and treatment with zuclopenthixol 75 mg/day and clonazepam 6 mg/day. In terms of percentages, there were 0.4 excitative episodes per patient, which was statistically significant in comparison with previous treatments (P = 0.000 [t-test]). In Group 4, one manic episode occurred during month 13 of REAC-lithium treatment and was mild (YMRS = 24) with a length of 10.0 weeks and treatment with zuclopenthixol 75 mg/day and valproate 1000 mg/day. In terms of percentages, there were 0.1 excitative episodes per patients, which was statistically significant in comparison with previous treatments (P = 0.000 [t-test]).

No hospitalizations occurred during REAC-lithium treatment observation periods. The safety and tolerability profiles of combined REAC-lithium treatment were favorable in all patients. The overall clinical results before and after REAC augmentation of lithium are shown in Tables 2–5 and Figures 1–4. Rates of hospitalization before and after REAC augmentation of lithium are shown in Table 6.

Discussion

The major finding of this research was the superior effectiveness of REAC-lithium treatment in comparison with the corresponding previous treatment periods, in terms of depressive rather than manic recurrences, in particular with respect to the comparison with lithium alone. It is interesting to note that this seems to be related to the REAC optimization of global brain functioning that is probably disrupted in bipolar depression as a consequence of previous manic-hypomanic episodes. The relevance of these results should be considered in relation to the well known drug resistance of this clinical disorder. The antidepressant activity of REAC-lithium treatment is the consequence of the action of the REAC device which, through the modulation of inter-intra neuronal microcurrents and microelectromagnetic fields, enables potentiation of the neurobiological activity of lithium.

Table 2 Demographic statistical, and mean values for the manic and depressive episodes, before and after REAC-lithium treatment in Group 1 patients

<table>
<thead>
<tr>
<th>Group 1 (n = 8)</th>
<th>Lithium</th>
<th>REAC-Lithium</th>
</tr>
</thead>
<tbody>
<tr>
<td>7 males [BD I]</td>
<td>2.1 ± 0.6</td>
<td>0.8 ± 0.4</td>
</tr>
<tr>
<td>1 females [BD I]</td>
<td>3 ± 0.7</td>
<td>0.1 ± 0.3</td>
</tr>
</tbody>
</table>

*t-Test: t = 11.718, DF14, P = 0.000.

Abbreviations: BD, bipolar disorder; REAC, radioelectric asymmetric brain stimulation device.

Table 3 Demographic statistical, and mean values for the manic and depressive episodes, before and after REAC-lithium treatment in Group 2 patients

<table>
<thead>
<tr>
<th>Group 2 (n = 14)</th>
<th>Lithium</th>
<th>REAC-Lithium</th>
</tr>
</thead>
<tbody>
<tr>
<td>7 males (4 BD I and 3 BD II)</td>
<td>2.4 ± 0.6</td>
<td>0.1 ± 0.2</td>
</tr>
<tr>
<td>7 females (all BD II)</td>
<td>3.9 ± 0.7</td>
<td>0 ± 0.2</td>
</tr>
</tbody>
</table>

*t-Test: t = 20.846, DF26, P = 0.000.

Abbreviations: BD, bipolar disorder; REAC, radioelectric asymmetric brain stimulation device.

Table 4 Demographic statistical, and mean values for the manic and depressive episodes, before and after REAC-lithium treatment in Group 3 patients

<table>
<thead>
<tr>
<th>Group 3 (n = 25)</th>
<th>Lithium</th>
<th>REAC-Lithium</th>
</tr>
</thead>
<tbody>
<tr>
<td>11 males [4 BD I and 7 BD II]</td>
<td>2.6 ± 0.8</td>
<td>0.4 ± 0.2</td>
</tr>
<tr>
<td>14 females [3 BD I and 11 BD II]</td>
<td>3.6 ± 0.9</td>
<td>0 ± 0.2</td>
</tr>
</tbody>
</table>

*t-Test: t = 20.000, DF48, P = 0.000.

Abbreviations: BD, bipolar disorder; REAC, radioelectric asymmetric brain stimulation device.
Radioelectric asymmetric conveyor for bipolar disorder

Other work of ours in progress conducted with functional magnetic resonance images in healthy subjects asked to perform finger tapping has demonstrated a significant reduction of the cerebral areas involved in the planning, control, execution, and stopping of this motor task (Figures 5A and 5B). Thus, according to our studies, REAC allows for better neural synchronization, such that less cerebral activity is needed to do the same movement. This deep cerebral activity of synchronization, as observed in normal subjects, is probably the main mechanism through which the REAC stabilizes the pathologic mood fluctuations in patients with bipolar disorder, so as to prevent morbid episodes, increase the duration of the so-called “free interval”, reduce the need for drug adjustments, to minimize the risk of hospitalizations. Our work shifts the focus from the biochemistry to the bioelectricity of the brain; it is important to emphasize that REAC seems synergic with lithium, which is not a drug but an ion, with a microelectric charge and a relative microelectromagnetic field. An interesting feature is that almost all patients included in the follow-up with REAC-lithium treatment experienced longer free intervals compared with previous treatment periods. REAC-lithium treatment seems to be a good strategy suitable for the treatment of free intervals in bipolar disorder. Until now, the free interval in a bipolar patient is defined as status free from depression and mania, but more recently it has been suggested that the free interval is not free. It is known that, after some time, the majority of bipolar patients show various clinical signs which correspond closely with requests for therapeutic and/or dose adjustment, and this occurs even if depressive or manic mood phases are not ongoing. This phenomenon can be explained only and exclusively with the persistence of a global neurobiological instability, for which the quantitative and qualitative boundaries are unknown. Therefore, improvement of the free interval becomes the main objective in treatment of bipolar disorder. The role of REAC-lithium treatment during intercritic periods can be considered as a therapeutic effect not related to its antidepressant or antimanic activity, but strictly related to the global optimization of brain functioning, broadening its action target so as to spread to temporal prolongation, and to qualitative improvement of the free interval. Consequently, if improvement of the state of the bipolar patient during the intercritic phases is associated with a significantly reduced risk of depressive and manic recurrences, the intercritic period should become the primary

**Table 5** Demographic statistical, and mean values for the manic and depressive episodes, before and after REAC-lithium treatment in Group 4 patients

<table>
<thead>
<tr>
<th>Group 4 (n = 9)</th>
<th>Lithium</th>
<th>REAC-Lithium</th>
</tr>
</thead>
<tbody>
<tr>
<td>3 males [1 BD I and 2 BD II]</td>
<td>6 females [2 BD I and 4 BD II]</td>
<td></td>
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<tr>
<td>Excitative episodes</td>
<td>2.6 ± 1.1</td>
<td>0.1 ± 0</td>
</tr>
<tr>
<td>Depressive episodes</td>
<td>3.7 ± 1</td>
<td>0 ± 0</td>
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*t*-Test: t = 11.100, DF16, P = 0.000.

**Abbreviations:** BD, bipolar disorder; REAC, radioelectric asymmetric brain stimulation device.

**Table 6** Rate of hospitalization before and after REAC augmentation of lithium

<table>
<thead>
<tr>
<th>Group 1</th>
<th>Group 2</th>
<th>Group 3</th>
<th>Group 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lithium</td>
<td>7 (5 D, 2 M)</td>
<td>9 (9 D)</td>
<td>7 (4 D, 3 M)</td>
</tr>
<tr>
<td>REAC-lithium</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
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</table>

**Abbreviation:** D, depressive episodes; M, excitative episodes; REAC, radioelectric asymmetric brain stimulation device.

**Figure 1** Demographic statistical, and mean values for the manic and depressive episodes, before and after REAC-lithium treatment in group 1 patients.

**Abbreviation:** REAC, radioelectric asymmetric brain stimulation device.

**Figure 2** Demographic statistical, and mean values for the manic and depressive episodes, before and after REAC-lithium treatment in group 2 patients.

**Abbreviation:** REAC, radioelectric asymmetric brain stimulation device.
objective in treatment of bipolar disorder. Furthermore, the preventive action of REAC-lithium, both in terms of manic mood and depressive recurrences, seems to be progressively stronger over time. In fact, all the mood episodes happened within 24 months from the beginning of REAC-lithium treatment. This feature seems to be in conflict with the observations drawn from olanzapine and risperidone (but not for quetiapine), for which the clinical experience shows increasing development of therapeutic tolerance phenomena. In fact, the depressive episodes of bipolar disorder seem to predispose to development of new depressive episodes, and after some time, produce a chronic hypotimic picture. Bipolar depression is a clinical dilemma because affected patients are usually poor responders to traditional antidepressant treatments, ie, tricyclic antidepressants, selective serotonin reuptake inhibitors, and selective norepinephrine reuptake inhibitors. Furthermore, with an increased risk of manic switches, cycles speed up, and there is progressive prognostic deterioration. Therefore, in light of these real difficulties of treatment, it is very important to prevent the depressive episodes of bipolar disorder, and use of REAC is a very promising therapeutic modality. Small sample size, a naturalistic and flexible dose design, maintenance of lithium, and lack of accurate assessments were the main limitations of this study. Nevertheless, we suggest that naturalistic studies and not randomized clinical trials supply a correct picture of the routine medical practice with real patients and real-life situations. Moreover, the long duration of observation can better clarify the clinical evolution of bipolar disorder. The maintenance of lithium treatment associated with REAC is due to ethical and legal considerations in relation to the fact that bipolar disorder is a severe condition (ie, carries a suicide rate >20%), and REAC is effective, albeit not well studied as yet in Italy.

Acknowledgments
The authors thank Dr Matteo Lotti Margotti for the data analysis, and Lucia Aravagli and Stefania Bini of the Rinaldi-Fontani Institute, Department of Neuro-Psycho-Physio Pathology, Florence, Italy, for their helpful discussions.

Disclosure
Salvatore Rinaldi and Vania Fontani are the inventors of the radioelectric asymmetric conveyor.
References

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