Deviant EEG resting-state large-scale brain network dynamics in euthymic bipolar disorder patients

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19 Abstract

20 Background: Neuroimaging studies provide evidence for disrupted resting-state functional brain

21 network activity in bipolar disorder (BD). Electroencephalographic (EEG) studies found altered

22 temporal characteristics of functional EEG microstates during depressive episode within different

affective disorders. Here we investigated whether euthymic patients with BD show deviant resting-

state large-scale brain network dynamics as reflected by altered temporal characteristics of EEGmicrostates.

- 26 Methods: We used high-density EEG to explore between-group differences in duration, coverage and
- occurrence of EEG microstates in 17 euthymic adults with BD and 17 age- and gender-matched healthy
 controls.
- 29 Results: Microstate analysis revealed five microstates (A-E) in global clustering across all subjects. In
- 30 patients compared to controls, we found increased occurrence and coverage of microstate A that did
- 31 not significantly correlate with anxiety scores.
- 32 Conclusion: Our results provide neurophysiological evidence for altered large-scale brain network
- dynamics in BD patients and suggest the increased presence of A microstate to be an
 electrophysiological trait characteristic of BD.
- 34 electrophysiological trait characteristic of BL

35 1 Introduction

- 36 Bipolar disorder (BD) is a common and severe psychiatric disorder, with an important personal and
- 37 societal burden (Cloutier et al., 2018; Eaton et al., 2012). The prevalence of bipolar disorder
- 38 worldwide is considered to range between 1% and 3% (Merikangas et al., 2007). BD patients are
- 39 frequently misdiagnosed and often identified at late stages of disease progression, which can lead to
- 40 inadequate treatment (Hirschfeld, 2007) and worse functional prognosis (Vieta et al., 2018). A better

understanding of the underlying pathophysiology is needed to identify objective biomarkers of BD
 that would improve diagnostic and/or treatment stratification of patients.

43 Potential candidates for neurobiological biomarkers could arise from functional brain network

44 abnormalities in BD patients. Evidence from brain imaging studies consistently points to

45 abnormalities in circuits implicated in emotion regulation and reactivity. Particularly, attenuated

46 frontal and enhanced limbic activations are reported in BD patients (Chen et al., 2011; Houenou et

47 al., 2011; Kupferschmidt and Zakzanis, 2011). Interestingly, regions implicated in the

48 pathophysiology of the disease, such as the inferior frontal gyrus, the medial prefrontal cortex

49 (mPFC), the amygdala present altered activation patterns even in unaffected first-degree relatives of

50 BD patients (Piguet et al., 2015), pointing toward brain alterations that could underlie disease 51 vulnerability. Moreover, evidence from functional magnetic resonance imaging (fMRI) studies

51 vulnerability. Moreover, evidence from functional magnetic resonance imaging (finici) studies 52 showed aberant resting-state functional connectivity between frontal and meso-limbic areas in BD

53 when compared to healthy controls (Vargas et al., 2013). A recently developed functional

54 neuroanatomic model of BD suggests, more specifically, decreased connectivity between ventral

55 prefrontal networks and limbic brain regions including the amygdala (Strakowski et al., 2012; Chase

and Philips, 2016). The functional connectivity abnormalities in BD in brain areas associated with

57 emotion processing were shown to vary with mood state. A resting-state functional connectivity

58 study of emotion regulation networks demonstrated that subgenual anterior cingulate cortex

59 (sgACC)-amygdala coupling is critically affected during mood episodes, and that functional

60 connectivity of sgACC plays a pivotal role in mood normalization through its interactions with the

61 ventrolateral PFC and posterior cingulate cortex (Rey et al., 2016). Nevertheless, although different

62 fMRI metrics allowed to report deviant patterns of large-scale networks and altered resting-state

functional connectivity (Rey et al., 2016; Wang et al., 2016) in BD, the precise temporal dynamics of
 the functional brain networks at rest remain to be determined.

Large-scale neural networks dynamically and rapidly re-organize themselves to enable efficient 65 functioning (de Pasquale et al., 2018; Bressler and Menon, 2010). Fast dynamics of the resting-state 66 67 large-scale neural networks can be studied on sub-second temporal scales with EEG microstate analysis (Pascual-Marqui et al., 1995; Van de Ville et al., 2010; Michel and Koenig, 2018). EEG 68 69 microstates are defined as short periods (60-120 ms) of quasi-stable electric potential scalp 70 topography (Lehmann et al., 1987; Koenig et al., 2002). Therefore, microstate analysis can cluster the 71 scalp's topographies of the resting-state EEG activity into the set of a few microstate classes 72 including the four canonical classes A-D (Michel and Koenig, 2018) and more recent additional ones 73 (Custo et al., 2017; Bréchet et al., 2019). Since each microstate class topography reflects a coherent 74 neuronal activity (Khanna et al., 2015; Michel and Koenig, 2018), the temporal characteristics, such 75 as duration, occurrence and coverage, may be linked to the expression of spontaneous mental states 76 and be representative of the contents of consciousness (Changeux and Michel, 2004; Lehmann et al., 77 1990). Numerous studies reported abnormalities in temporal properties of resting-state EEG 78 microstates in neuropsychiatric disorders (for review see Khanna et al., 2015; Michel and Koenig, 79 2018). Evidence from microstate studies suggests that altered resting-state brain network dynamics 80 may represent a marker of risk to develop neuropsychiatric disorders (Tomescu et al., 2014, 2015; 81 Andreou et al., 2014), may predict clinical variables of an illness (Gschwind et al., 2016), or help to 82 assess the efficacy of a treatment (Atluri et al., 2018; Sverak et al., 2018). Only two studies 83 investigated resting-state EEG in BD patients (Strik et al., 1995; Damborská et al., 2019). These 84 studies examined patients during a depressive episode within different affective disorders. Adaptive

85 segmentation of resting-state EEG showed abnormal microstate topographies and reduced overall

86 average microstate duration in patients that met criteria for unipolar or bipolar mood disorders or for

87 dysthymia (Strik et al., 1995). Using a *k*-means cluster analysis, an increased occurrence of

- 88 microstate A with depression as an effect related to the symptom severity was observed during a
- 89 period of depression in unipolar and bipolar patients (Damborská et al., 2019).
- 90 Trait markers of BD based on neurobiological findings can be considered as biomarkers of illness
- 91 (Piguet et al., 2016). These trait markers of BD can be studied during the periods of remission, or
- 92 euthymia. No microstate study, however, has been performed on euthymic BD patients to the best of
- 93 our knowledge. Thus, the main goal of the current study was to explore group differences between
- 94 euthymic patients with BD and healthy controls in terms of resting-state EEG microstate dynamics.
- 95 We hypothesized that BD patients during remission will show altered temporal characteristics of
- 96 EEG microstates such as duration, coverage, and occurrence.

97 2 Materials and Methods

98 2.1 Subjects

99 Data were collected from 17 euthymic adult patients with BD and 17 healthy control (HC) subjects. 100 The patients were recruited from the Mood Disorders Unit at the Geneva University Hospital. A 101 snowball convenience sampling was used for the selection of the BD patients. Control subjects were 102 recruited by general advertisement. All subjects were clinically evaluated using clinical structured 103 interview (DIGS: Diagnostic for Genetic Studies, (Nurnberger et al., 1994). Bipolar disorder was 104 confirmed in the experimental group by the usual assessment of the specialized program, an interview 105 with a psychiatrist, and a semi-structured interview and relevant questionnaires with a psychologist. 106 Exclusion criteria for all participants were a history of head injury, current alcohol or drug abuse. 107 Additionally, a history of psychiatric or neurological illness and of any neurological comorbidity were 108 exclusion criteria for controls and bipolar patients, respectively. Symptoms of mania and depression 109 were evaluated using the Young Mania Rating Scale (YMRS) (Young et al., 1978) and the 110 Montgomery-Åsberg Depression Rating Scale (MADRS) (Williams and Kobak, 2008), respectively. 111 Participants were considered euthymic if they scored < 6 on YMRS and < 12 on MADRS at the time 112 of the experiment, and were stable for at least 4 weeks before. All patients were medicated, receiving 113 pharmacological therapy including antipsychotics, antidepressants and mood stabilizers, and had to be 114 under stable medication for at least 4 weeks. The experimental group included both BD I (n = 10) and 115 BD II (n = 7) types. Results of an event-related EEG study that was conducted on a sample partially 116 overlapping with the current dataset showed that these patients present a dysfunctional gaze processing,

- results that were reported elsewhere (Berchio et al., 2017).
- 118 To check for possible demographic or clinical differences between groups, subject characteristics such
- 119 as age, education or level of depression were compared between groups using independent *t*-tests.
- 120 Anxiety is highly associated with bipolar disorder (Simon et al., 2004; 2007) and is a potential 121 confounding variable when investigating microstate dynamics at rest. For example, decreased duration
- of EEG microstates at rest in patients with panic disorder has been reported (Wiedemann et al., 1998).
- 123 To check for possible differences in anxiety symptoms, all subjects were assessed with the State-trait
- Anxiety Inventory (STAI) (Spielberger et al., 1970) and the scores were compared between patients
- 125 and controls using independent *t*-tests.
- 126 This study was carried out in accordance with the recommendations of the Ethics Committee for
- 127 Human Research of the Geneva University Hospital, with written informed consent from all subjects.
- 128 All subjects gave written informed consent in accordance with the Declaration of Helsinki. The
- 129 protocol was approved by the Ethics Committee for Human Research of the Geneva University
- 130 Hospital, Switzerland.

131 **2.2 EEG recording and pre-processing**

132 The EEG was recorded with a high density 256-channel system (EGI System 200; Electrical Geodesic 133 Inc., OR, USA), sampling rate of 1kHz, and Cz as acquisition reference. Subjects were sitting in a 134 comfortable upright position and were instructed to stay as calm as possible, to keep their eyes closed

and to relax for 5 minutes. They were asked to stay awake.

136 To remove muscular artifacts originating in the neck and face the data were reduced to 204 channels.

137 Two to four minutes of EEG data were selected based on visual assessment of the artifacts and band-

138 pass filtered between 1 and 40 Hz. Subsequently, in order to remove ballistocardiogram and oculo-

139 motor artifacts, infomax-based Independent Component Analysis (Jung et al., 2000) was applied on all

but one or two channels rejected due to abundant artifacts. Only components related to physiological noise, such as ballistocardiogram, saccadic eve movements, and eve blinking, were removed based on

142 the waveform, topography and time course of the component. The cleaned EEG recordings were down-

sampled to 125 Hz and the previously identified noisy channels were interpolated using a three-

dimensional spherical spline (Perrin et al., 1989), and re-referenced to the average reference. All the

145 preprocessing steps were done using MATLAB and the freely available Cartool Software 3.70

146 (<u>https://sites.google.com/site/cartoolcommunity/home</u>), programmed by Denis Brunet.

147 **2.3 EEG data analysis**

148 To estimate the optimal set of topographies explaining the EEG signal, a standard microstate analysis 149 was performed using k-means clustering (see Supplementary Fig. 1). The polarity of the maps was 150 ignored in this procedure (Brunet et al., 2011; Murray et al, 2008; Pascual-Marqui et al., 1995). To 151 determine the optimal number of clusters, we applied a meta-criterion that is a combination of seven 152 independent optimization criteria (for details see Bréchet et al., 2019). In order to improve the signal-153 to-noise ratio, only the data at the time points of the local maximum of the Global Field Power (GFP) 154 were clustered (Pascual-Marqui et al., 1995; Koenig et al., 2002; Britz et al, 2010, Tomescu et al., 155 2014). The GFP is a scalar measure of the strength of the scalp potential field and is calculated as the standard deviation of all electrodes at a given time point (Michel et al., 1993; Brunet et al., 2011; 156 157 Murray et al., 2008). The cluster analysis was first computed at the individual level and then at global 158 level across all participants (patients and controls), clustering each participant's representative maps.

159 In order to retrieve the temporal characteristics of the microstates, spatial correlation was calculated 160 between every map identified at the global level and the individual subject's topographical map in every instant of the pre-processed EEG recording. Each continuous time point of the subject's EEG 161 162 (not only the GFP peaks) was then assigned to the microstate class of the highest correlation, again ignoring polarity (Brunet et al., 2011; Bréchet et al., 2019; Michel and Koenig, 2018; Santarnecchi et 163 164 al., 2017). Temporal smoothing parameters (window half size = 3, strength (Besag Factor) = 10) 165 ensured that the noise during low GFP did not artificially interrupt the temporal segments of stable topography (Brunet et al., 2011; Pascual-Marqui et al., 1995). For each subject, three temporal 166 167 parameters were then calculated for each of the previously identified microstates: (i) occurrence, (ii) 168 coverage, and (iii) duration. Occurrence indicates how many times a microstate class recurs in one 169 second. The coverage in percent represents the summed amount of time spent in a given microstate 170 class as a portion of the whole analyzed period. The duration in milliseconds for a given microstate 171 class indicates the amount of time that a given microstate class is continuously present. In order to 172 assess the extent to which the representative microstate topographies explain the original EEG data, 173 the global explained variance (GEV) was calculated as the sum of the explained variances of each 174 microstate weighted by the GFP. Microstate analysis was performed using the freely available 175 Cartool Software 3.70, (https://sites.google.com/site/cartoolcommunity/home), programmed by Denis Brunet. Mann-Whitney U test was used to investigate group differences for temporal 176 177 parameters of each microstate. Multiple comparisons were corrected using the false discovery rate 178 (FDR) method (Benjamini, 2010).

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- 179 Spearman's rank correlations were calculated between the MADRS, YMRS, STAI-state, and STAI-
- 180 trait scores and significant microstate parameters to check for possible relationships between
- 181 symptoms and microstate dynamics. Statistical evaluation was performed by the routines included in
- the program package Statistica'13 (1984-2018, TIBCO, Software Inc, Version 13.4.0.14).

183 **3 Results**

- 184 There were no significant differences in age and level of education between the patient and the control
- 185 groups. In both groups, very low mean scores on depression and mania symptoms were observed,
- 186 which did not significantly differ between the two groups. BD patients showed higher scores on state
- and trait scales of the STAI. For all subject characteristics, see Table 1.
- 188 The meta-criterion used to determine the most dominant topographies revealed five resting-state 189 microstate maps explaining 82.2 % of the global variance (Fig. 1). The topographies resembled those
- previously reported as A, B, C, and D maps (Khanna et al., 2015; Michel and Koenig, 2018; Koenig et
- al., 2002; Britz et al., 2010) and one of the three recently identified additional maps (Custo et al., 2017).
- 192 We labeled these scalp maps A E in accordance with the previous literature on microstates. The scalp
- 193 topographies showed left posterior-right anterior orientation (map A), a right posterior-left anterior
- orientation (map B), an anterior-posterior orientation (map C), a fronto-central maximum (map D), and
- 195 a parieto-occipital maximum (map E).
- 196 Since some microstate parameters showed a non-homogeneity of variances in the two groups (Levene's
- tests for coverage of the C microstate and duration of the A and C microstates; p<0.01), we decided to calculate Mann-Whitney U test to investigate group differences for temporal parameters of each
- 199 microstate.
- 200 We found significant between-group differences for microstate parameters of the A and B microstates.
- 201 Both microstates showed increased presence in patients in terms of occurrence and coverage. The two
- groups did not differ in any temporal parameter of microstates C, D, or E. The results of the temporal
 characteristics of each microstate are summarized in Table 2 and Figure 2.
- The results of Spearman's rank correlation revealed a significant positive association between the coverage of the microstate B and the STAI-state (r = 0.40) and STAI-trait (r = 0.54) scores. The results
- of Spearman's rank correlation revealed a significant positive association between the occurrence of
- 207 the microstate B and the STAI-trait (r = 0.47) scores. The results of Spearman's rank correlation 208 revealed no significant associations between the STAI-state or STAI-trait scores and the occurrence or
- 209 coverage of the microstate A (all absolute r-values < 0.35).
- 210 The results of Spearman's rank correlation revealed no significant associations between the MADRS
- and YMRS scores and the occurrence or coverage of the microstate A and B (all absolute r-values < 0.30)
- 212 0.30).

213 **4 Discussion**

- Our study presents the first evidence for altered resting-state EEG microstate dynamics in euthymic patients with bipolar disorder. Patients were stable and did not significantly differ in their depressive or manic symptomatology from healthy controls at the time of experiment. Despite this fact, they showed abnormally increased presence of microstates A and B, the latter correlating with anxiety level. The key discovery in the current study is the increased occurrence and coverage of microstate A in euthymic bipolar patients compared to healthy controls. In an earlier combined fMRI-EEG study the microstate A was associated with the auditory network (Britz et al., 2010). Moreover, generators of
- 221 the functional EEG microstates were estimated in recent studies, where sources of the microstate A
- showed left-lateralized activity in the temporal lobe, insula, mPFC, and occipital gyri (Custo et al.,
- 223 2017; Bréchet et al., 2019).

224 In the fMRI literature as well, resting-state functional connectivity alterations of the insula (Yin et al., 225 2018), the auditory network (Reinke et al., 2013), and the mPFC (Gong et al., 2019) were reported in 226 BD patients. Verbal episodic memory deficits and language-related symptoms in BD patients were 227 suggested to be associated with a diminished functional connectivity within the auditory/temporal 228 gyrus and to be compensated by increased fronto-temporal functional connectivity (Reinke et al., 229 2013). The mPFC was also identified as a major locus of shared abnormality in BD and schizophrenia 230 (Öngür et al., 2010), showing reduced default mode network connectivity from the mPFC to the 231 hippocampus and fusiform gyrus, as well as increased connectivity between the mPFC and primary 232 visual cortex in BD. Hypoconnectivity of the default mode network from the left posterior cingulate 233 cortex to the bilateral mPFC and bilateral precuneus, and reduced salience connectivity of the left 234 sgACC to the right inferior temporal gyrus in BD patients (Gong et al., 2019) was observed in 235 unmedicated BD patients. In euthymic BD subjects compared to healthy controls, resting-state 236 functional connectivity of the insula (Minuzzi et al., 2018) and amygdala (Li et al., 2018) to other 237 brain regions was reported to be increased and decreased, respectively. In summary, the evidence from 238 fMRI studies shows both hypoconnectivity (Gong et al., 2019; Öngür et al., 2010) and 239 hyperconnectivity (Minuzzi et al., 2018; Reinke et al., 2013; Öngür et al., 2010) pointing to complex 240 alterations of functional resting-state networks. Our findings of increased presence of the microstate A 241 in euthymic BD patients might be related to the hyperconnectivity of the underlying networks that 242 involve the temporal lobe, insula, mPFC, and occipital gyri.

243 Anxiety symptoms were previously associated with greater severity and impairment in bipolar disorder

244 (Simon et al., 2004) and euthymic bipolar patients tend to present high residual level of anxiety (Albert

- et al., 2008), as it was the case here. No significant correlation was found between the increased anxiety
- scores and the increased occurrence or coverage of the microstate A. Our results therefore indicate that this alteration of microstate dynamics might represent a characteristic feature of BD that is not affected
- by anxiety.

249 The demonstrated alterations in microstate A dynamics during clinical remission might reflect (i) an

impaired resting-state large-scale brain network dynamics as a trait characteristic of the disorder and/or
 (ii) a compensatory mechanism needed for clinical stabilization of the disorder.

252 Our study is the first to examine EEG microstate dynamics in BD patients during remission. 253 Interestingly, in our recent study we showed positive associations of depressive symptoms with the 254 occurrence of microstate A in a heterogenous group of patients with affective disorders (Damborská et 255 al., 2019). The increased microstate A occurrence with depression as an effect related to the symptom 256 severity (Damborská et al., 2019) and as a here demonstrated group difference of BD patients vs. 257 controls, is not congruent with the previously reported reduced duration of the EEG microstates during 258 a depressive episode (Strik et al., 1995). The experimental group in that study was not restricted to 259 bipolar patients, however, and included also patients who met the criteria for unipolar depression or 260 dysthymia. Moreover, authors examined the overall microstate duration and did not examine distinct 261 microstates separately. These and other aspects, such as different clustering methods used, make it 262 difficult to compare our findings with that early evidence of dirupted microstate dynamics in 263 depression.

264 The microstate B was previously associated with the visual network (Britz et al., 2010; Custo et al., 265 2014, 2017; Bréchet et al., 2019). In our group of BD patients, we found an abnormally increased 266 occurrence and coverage of microstate B that was associated with higher anxiety. Previous studies also 267 suggest that anxiety may influence visual processing (Phelps et al., 2006; Laretzaki et al., 2010) and that connections between amygdala and visual cortex might underlie enhanced visual processing of 268 269 emotionally salient stimuli in patients with social fobia (Goldin et al., 2009). Our finding of increased 270 presence of microstate B positively associated with anxiety level in euthymic BD patients is consistent 271 with these observations. To the best of our knowledge, there are no other studies that would aim to

examine in detail the relationship between anxiety and resting-state EEG microstate dynamics. An early microstate study reported decreased overall resting-state microstate duration in panic disorder (Wiedemann et al., 1998). This study, however, did not assess temporal characteristics of different microstates separately and it is therefore difficult to compare those findings with our observations. Further evidence is needed to determine, whether the increased presence of microsate B in our experimental group is a characteristic feature of BD or anxiety, or whether it is related to both conditions.

279 Changes in microstate A and B have been reported in several psychiatric conditions such as dementia, 280 narcolepsy, multiple sclerosis, panic disorder, etc. (for review see Michel and Koenig, 2018). Increases 281 in duration and occurrence of microstate A and B were observed in patients with multiple sclerosis 282 (Gschwind, et al., 2016). Moreover, the changes in dynamic patterns of these two microstates, predicted 283 depression scores and other clinical variables. It was suggested that multiple sclerosis affects the 284 "sensory" (visual, auditory) rather than the higher-order (salience, central executive) functional 285 networks (Michel and Koenig, 2018). Our findings of impaired dynamics in microstates A and B 286 suggest a similar interpretation for the BD. Evidence from fMRI studies points to topographical 287 dysbalances between the default mode and sensorimotor networks in BD patients with opposing 288 patterns in depression and mania (Martino et al., 2016). Cyclothymic and depressive temperaments 289 were associated with opposite changes in the sensorimotor network variability in the resting state signal 290 measured by fractional standard deviation of Blood-Oxygen-Level Dependent signal (Conio et al., 291 2019). Our findings of altered microstates A and B dynamics is consistent with this fMRI evidence of 292 impaired sensorimotor network in affective disorders, and moreover suggests that neural correlates of 293 these deficits are prominent even during the euthymic state in BD patients.

294 In summary, results of the current study seem to indicate that dysfunctional activity of resting-state 295 brain networks underlying A microstate is a detectable impairment in BD during an euthymic state. 296 The presence of microstate A represents a measure that might be implicated in clinical practice. 297 Importantly, this parameter, whose changes were observed during remission, could be potentially 298 useful for early identification of bipolar disorder that could help better management of the disease. If 299 future studies confirm the same pattern in prodromal or vulnerable subjects, it could also help detection 300 of at-risk subjects and therefore the possiblility for early intervention. The present study has, however, 301 some limitations. Our low sample size made it impossible to examine any potential influence of 302 medication on the microstate parameters by comparing patients receiving a specific drug with those 303 not receiving it. Possible effects of medication on our results should be therefore taken into account. 304 Due to the same reason, it was not possible to examine any potential influence of subtypes of bipolar

305 disorder on microstate results.

306 **5 Conclusions**

307 Our study described altered EEG resting-state microstate temporal parameters in euthymic bipolar 308 patients. These findings provide an insight into the resting-state global brain network dynamics in 309 bipolar disorder. The increased presence of the A microstate might be considered as a candidate 310 electrophysiological non-specific trait marker of BD. Nevertheless, studies examining possible 311 interactions between microstate dynamics and BD symptoms are needed to better understand the 312 dysfunction of large-scale brain network resting-state dynamics in this affective disorder.

313 6 Table legends

Table 1. ^aEdinburgh inventory (Oldfield, 1971); ^bEducation levels: 1 = no high school, 2 = high school,

315 3 = university studies

7 Tables

317 Table 1. Subject characteristics

| Characteristic | Healthy controls $(n - 17)$ | Bipolar patients $(n - 17)$ | <i>t</i> -value | p-value |
|---|-----------------------------|-----------------------------|-----------------|---------|
| Age: mean ± SD | 36.6 ± 14.5 | 35.9 ± 11.9 | -0.17 | 0.87 |
| Gender: male, <i>n</i> | 12 | 12 | | |
| Handedness ^a : right, <i>n</i> | 14 | 14 | | |
| Education ^b : mean \pm SD | 2.3 ± 0.6 | 2.4 ± 0.5 | 0.63 | 0.53 |
| MADRS: mean \pm SD | 1.4 ± 1.6 | 2.3 ±2.9 | 1.09 | 0.29 |
| YMRS: mean \pm SD | 0.86 ± 1.4 | 0.76 ± 1.4 | -0.18 | 0.86 |
| STAI-state: mean \pm SD | 26.7 ± 4.8 | 36.9±15.2 | -2.13 | 0.04 |
| STAI-trait: mean \pm SD | 27.4±5.2 | 42.9±13.3 | -3.7 | 0.001 |

| Microstate | А | В | С | D | E |
|-------------------------------|----------|----------|-----------|-----------|---------------|
| Occurrence (s^{-1}) | | | | | |
| Patients (mean±s.d.) | 4.5±1.1 | 3.9±1.1 | 5.2±1.1 | 3.9±2.2 | $1.9{\pm}1.2$ |
| Controls (mean±s.d.) | 3.4±1.2 | 3.0±1.1 | 5.0±1.1 | 3.4±2.1 | 2.0±1.3 |
| Z-value | 2.51 | 2.51 | -0.03 | 0.65 | -0.14 |
| uncorrected <i>p</i> -value | 0.01 | 0.01 | 0.97 | 0.51 | 0.89 |
| FDR corrected <i>p</i> -value | 0.03 | 0.03 | 0.97 | 0.85 | 0.97 |
| Coverage (%) | | | | | |
| Patients (mean±s.d.) | 18.9±6.5 | 16.0±7.8 | 25.1±8.6 | 18.5±12.3 | 6.4±5.1 |
| Controls (mean±s.d.) | 13.3±4.9 | 11.4±6.0 | 34.7±16.8 | 17.0±12.0 | 7.0±5.3 |
| Z-value | 2.62 | 2.79 | -1.69 | 0.17 | -0.21 |
| uncorrected <i>p</i> -value | 0.009 | 0.005 | 0.09 | 0.86 | 0.84 |
| FDR corrected <i>p</i> -value | 0.02 | 0.02 | 0.15 | 0.86 | 0.86 |
| Duration (ms) | | | | | |
| Patients (mean±s.d.) | 26±4 | 25±4 | 29±4 | 26±6 | 21±4 |
| Controls (mean±s.d.) | 24±1 | 24±3 | 35±9 | 27±6 | 22±4 |
| Z-value | 1.38 | 1.03 | -2.27 | 0.17 | -0.69 |
| uncorrected <i>p</i> -value | 0.17 | 0.30 | 0.02 | 0.86 | 0.49 |
| FDR corrected <i>p</i> -value | 0.42 | 0.50 | 0.12 | 0.86 | 0.61 |

322 8 Figures



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325 Supplementary Figure 1. Microstate analysis: (A) resting-state EEG from subsample of 16 out of 204 326 electrodes; (B) global field power (GFP) curve with the GFP peaks (vertical lines) in the same EEG 327 period as shown in (A); (C) potential maps at successive GFP peaks, indicated in (B), from the first 1 328 s period of the recording; (**D**) set of five cluster maps best explaining the data as revealed by K-means 329 clustering of the maps at the GFP peaks; (E) the original EEG recording shown in (A) with 330 superimposed color-coded microstate segments. Note that each time point of the EEG recording was 331 labelled with the cluster map, shown in (D), with which the instant map correlated best. The duration 332 of segments, occurrence, and coverage for all microstates were computed on thus labeled EEG 333 recording.



Figure 1. The five microstate topographies identified in the global clustering across all subjects.
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Figure 2. Temporal dynamics of EEG microstates in patients with bipolar disorder (BD) and in healthy controls (HC). In each subplot, the raw data is plotted on top of a boxplot showing the mean (**a**), 95% confidence interval (box plot area), 1 standard deviation (whiskers), and significant differences (*). In all plots, x-axes represent the subject group; y-axes represent the occurrence (upper plots) or coverage (lower plots). Note significantly increased occurrence and coverage of the microstate A and B in the BD compared to HC group (FDR corrected p < 0.05).

347 9 Conflict of Interest

348 The authors declare that the research was conducted in the absence of any commercial or financial 349 relationships that could be construed as a potential conflict of interest.

350 10 Author Contributions

- AD designed the study, performed the analysis, and wrote the initial draft; JMA, AGD and CP were
 responsible for clinical assessment; CMM served as an advisor; CB collected the HD-EEG data
- and was responsible for the overall oversight of the study. All authors revised the manuscript.

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584 14 Data Availability Statement

585 The raw data supporting the conclusions of this manuscript will be made available by the authors, 586 without undue reservation, to any qualified researcher.