Default Mode Network alterations in alexithymia: an EEG power spectra and connectivity study

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Recent neuroimaging studies have shown that alexithymia is characterized by functional alterations in different brain areas [e.g., posterior cingulate cortex (PCC)], during emotional/social tasks. However, only few data are available about alexithymic cortical networking features during resting state (RS). We have investigated the modifications of electroencephalographic (EEG) power spectra and EEG functional connectivity in the default mode network (DMN) in subjects with alexithymia. Eighteen subjects with alexithymia and eighteen subjects without alexithymia matched for age and gender were enrolled. EEG was recorded during 5 min of RS. EEG analyses were conducted by means of the exact Low Resolution Electric Tomography software (eLORETA). Compared to controls, alexithymic subjects showed a decrease of alpha power in the right PCC. In the connectivity analysis, compared to controls, alexithymic subjects showed a decrease of alpha connectivity between: (i) right anterior cingulate cortex and right PCC, (ii) right frontal lobe and right PCC, and (iii) right parietal lobe and right temporal lobe. Finally, mediation models showed that the association between alexithymia and EEG connectivity values was directed and was not mediated by psychopathology severity. Taken together, our results could reflect the neurophysiological substrate of some core features of alexithymia, such as the impairment in emotional awareness.

The alexithymia construct refers to a personality trait featuring several alterations in emotional processing. It has been theorized as a multidimensional construct associated with (i) difficulty in recognizing and describing one's emotions, (ii) difficulty in distinguishing between feelings and bodily sensations, (iii) impairment of affect-related fantasy and imagery activity, (iv) externally-oriented thinking (i.e. the predisposition to focus on external stimuli rather than on one's own internal experiences)1. Alexithymia is considered a crucial construct in psychosomatic medicine2 and it is related to several psychosomatic sequelae, such as medically unexplained symptoms3,4, as well as to psychopathology5.

From a neurobiological point of view, recent neuroimaging studies have shown that alexithymia is characterized by functional alterations in different brain areas (e.g., the anterior and posterior cingulate cortex), during emotional/social tasks as well as in imagery and somatosensory tasks6. On the other hand, electroencephalographic (EEG) studies in alexithymic subjects during emotional tasks (e.g. EEG connectivity and event-related potential studies) documented several abnormalities in the functional integration between brain areas (i.e. between anterior and posterior cortical regions), predominantly in the right hemisphere7-11.

As concerns cortical networking features of alexithymic subjects during resting state (RS), few data are available. RS network activity can be defined as “coherent and spontaneous fluctuations of human brain activity in distinct and spatially separate networks of varying granularity when subjects are not engaged in a particular task or superior cognitive processes”12. The default mode network (DMN) is most frequently detected during the RS condition12 and reflects the neural activity of different brain areas such as cingulate cortex, hippocampus, medial frontal lobes, inferior parietal lobes, and temporal lobes13,14. It is thought to be involved in self-consciousness, self-processing and introspection functions, including emotional awareness and processing15, which are supposed to be impaired in alexithymia16. Recently, a consistent link between alterations in emotional processing and reduced DMN connectivity has been detected16,17. Furthermore, abnormalities in DMN activity and its

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functional connectivity have been widely reported in several psychiatric disorders, such as schizophrenia and mood disorders.

Although functional magnetic resonance imaging (fMRI) is commonly used to investigate the functional connectivity of DMN, recent studies have shown that also EEG is suitable to investigate this network. Assessing changes in the synaptic synchrony of millions of neurons connected at varying time delays and frequencies. Furthermore, compared to fMRI, EEG time-series data directly relate to dynamic postsynaptic activity in the cerebral cortex with a higher temporal resolution. Conversely, MR-based methods cannot assess fast-frequency synchronized neuronal activity. Finally, the EEG offers a valuable tool to assess "real-time electrical activity in the brain and is overall a less costly, time-consuming, and complex procedure." To the best of our knowledge, only one study has directly investigated the DMN in subjects with alexithymia. Lienburg et al., in an fMRI study, showed that, compared to non-alexithymic participants, alexithymic individuals showed lower connectivity in medial frontal and temporal areas. Therefore, the main aim of the present study was to extend these previous findings by exploring the modifications of EEG power spectra and EEG functional connectivity in DMN in subjects with alexithymia. Furthermore, another aim was to investigate the association among any significantly modified connections of DMN, alexithymia severity and general psychopathology.

Materials and Methods

Participants. Participants were recruited from the Università Europea di Roma through advertisements posted at the university. The enrollment lasted from October 2015 to March 2016. Ninety-five subjects who agreed to participate were administered the 20-item Toronto Alexithymia Scale (TAS-20), the revised version of Symptom Check list 90 (SCL-90-R) and a checklist assessing socio-demographic data and the study inclusion criteria. Inclusion criteria were: right handedness; no history of medical, psychiatric and neurologic diseases; head trauma; no assumption of Central Nervous System active drugs in the two weeks prior to study entry. After receiving information about the aims of the study, all participants provided a written consent to participate in the study that was performed according to the Helsinki declaration standards and was approved by the Università Europea’s ethics review board. Informed consent has been obtained for all participants involved in the present study.

Eighteen consecutive subjects (three men and fifteen women, mean age: 25.00 ± 8.12 years) with TAS-20 total score ≥58 were enrolled in the alexithymia group (TAS+ group). According to a case-control design, we enrolled a control group consisting of eighteen subjects matched for sex and age (three men and fifteen women, mean age: 23.89 ± 4.85) with low alexithymia (TAS-20 total score ≤45, TAS− group). The cut-off values of TAS-20 used for selection of study groups were those adopted by Romei and coworkers. Clinical and socio-demographic characteristics of the final sample are listed in Table 1.

Questionnaires. The TAS-20 is a 20-item self-report that is one of the most commonly used measures of alexithymia. Items are rated on a 5-point Likert scale (from 1 = Strongly disagree to 5 = Strongly agree) indicating the degree to which each statement describes the respondent’s behavior. It contains the following three subscales, theoretically formulated and confirmed through factor analysis: (1) Difficulty Describing Feelings (DDF) refers to difficulty describing and communicating emotions, (2) Difficulty Identifying Feeling (DIF) refers to difficulty identifying emotions, (3) Externally-Oriented Thinking (EOT) concerns the tendency of individuals to focus their attention externally. The TAS total score (ranging from 20 to 100) corresponds to the general level of alexithymia. In the present study we used the Italian version of the TAS. The Cronbach’s α in the present sample was 0.90 for the TAS-20 total score.

The SCL-90-R is a 90-item questionnaire on 5-point Likert scale (0–4) assessing nine primary symptom dimensions: somatization (SOM), obsessive-compulsive symptoms (O-C), interpersonal sensitivity (I-S), depression (DEP), anxiety (ANX), hostility (HOS), phobic anxiety (PAR), paranoid ideation (PAR) and psychoticism (PSY). Furthermore, seven additional items assess disturbances in appetite and sleep (OTHER). The SCL-90-R also provides a global severity index (GSI) which is designed to measure overall psychological distress. Higher scores indicate more psychological symptoms in each subscale as well as a higher degree of distress, higher intensity of symptoms, and more self-reported symptoms. In the present study, a previously validated Italian version of the scale was used and the Cronbach’s alpha in the present sample was 0.92.

EEG recordings. RS recordings were performed in an EEG Laboratory, with each subject sitting in a comfortable armchair, with his/her eyes closed, in a quiet, semi-darkened silent room for 5 minutes. In order to avoid alcohol and or caffeine effects on EEG data, participants were asked to refrain from drinking alcohol and caffeine for 4 to 6 hours immediately before their EEG recordings.

EEG was recorded by means of a Micromed System Plus digital EEGraph (Micromed® S.p.A., Mogliano Veneto, TV, Italy). EEG montage included 19 standard scalp leads positioned according to the 10–20 system (recording sites: FP1, FP2, F7, F3, Fz, F4, P7, T3, C3, CZ, C4, T4, T5, P3, Pz, P4, T6, O1, O2). Electrocorticography (EOG) and Electrocardiography (EKG). The reference electrodes were placed on the linked mastoids. Impedances were kept below 5 KΩ before starting the recording and checked again at the end of the experimental recording. Sampling frequency was 256 Hz; A/D conversion was made at 16bit; pre-amplifiers amplitude range was ±3200μV and low-frequency pre-filters were set at 0.15 Hz. The following band-pass filters were used: high frequency filters (HFF) = 0.2 Hz; low frequency filters (LFF) = 128 Hz. Artifact rejection (eye movements, blinks, muscular activations, or movement artifacts) was performed visually on the raw EEG. The recordings were attended by trained technicians, and the simultaneous recording of EOG and EKG further improved the artifact recognition and removal. More details of artifact rejection procedure could be found in refs 30,31. At least 120 seconds of EEG artifact-free recording (not necessarily consecutive) were analyzed for each group, in all conditions. The average time analyzed was 276 ± 18 sec., 280 ± 21 sec.
which is a widely used procedure in signal processing\cite{35,36}. The result is a true 3-dimensional tomography, in the brain based on multichannel surface EEG recordings\cite{32}. The eLORETA software benefits from an excellent localization agreement with different multi-modal imaging techniques, also when standard 19-electrodes EEG montage was used\cite{14}.

The connectivity analysis was performed by the computation of lagged phase synchronization \cite{37,40}. This connectivity measure is widely used to assess EEG functional connectivity in both psychiatric and brain diseases\cite{38,39}. Furthermore, compared to other connectivity measure, the lagged phase synchronization has some advantages (e.g. is resistant to non-physiological artifacts and it is minimally affected by low spatial resolution)\cite{37,40}.

The eLORETA software computes lagged phase synchronization $\rho_{x,y}(\omega)$, by the formula\cite{41}:

$$\varphi^2 x, y(\omega) = \frac{\left| \text{Im} [f x, y(\omega)] \right|^2}{1 - \left| \text{Re} [f x, y(\omega)] \right|^2}$$

Details on eLORETA lagged phase synchronization formula can be found in Pascual-Marqui's studies\cite{37,41}.

In order to evaluate the connectivity in the DMN, 12 Regions of Interest (ROIs) were defined according to the review by Buckner \& colleagues\cite{13}. The 'single nearest voxel' option was chosen because eLORETA has low spatial resolution, and the single centroid voxel is considered an excellent representative of the ROI\cite{37}. In this way, each ROI consisted of a single voxel, the closest to each seed.

### Table 1. Demographic and clinical data of participants.

| Variables                     | TAS+ (n = 18)       | TAS− (n = 18)       | test     | p      |
|-------------------------------|---------------------|---------------------|----------|--------|
| Age (M (SD))                  | 25.00 ± 8.12        | 23.89 ± 4.85        | Z-test = 0.33 | 0.90   |
| Alcohol use in the last 6 months - N (%) | 8 (44.4%)          | 11 (61.1%)          | $\chi^2_1 = 1.01$ | 0.32   |
| Tobacco use in the last 6 months - N (%) | 7 (38.6%)          | 8 (44.4%)           | $\chi^2_1 = 1.11$ | 0.74   |
| TAS-20                        |                     |                     |          |        |
| TAS Total Scores - M (SD)     | 61.11 ± 5.11        | 36.83 ± 7.02        | Z-test = 3 | <0.001 |
| DDF subscale - M (SD)         | 19.33 ± 2.01        | 9.50 ± 3.13         | Z-test = 3 | <0.001 |
| DIF subscale - M (SD)         | 22.17 ± 4.06        | 12.11 ± 3.69        | Z-test = 2.5 | <0.001 |
| EOT subscale - M (SD)         | 19.61 ± 4.35        | 15.22 ± 2.46        | Z-test = 1.83 | <0.001 |
| SCL-90-R                     |                     |                     |          |        |
| GSI - M (SD)                  | 0.93 ± 0.45         | 0.47 ± 0.38         | Z-test = 1.5 | <0.05  |
| SOM - M (SD)                  | 1.07 ± 0.62         | 0.51 ± 0.46         | Z-test = 1.5 | <0.05  |
| O-C - M (SD)                  | 1.29 ± 0.65         | 0.58 ± 0.48         | Z-test = 1.68 | <0.01  |
| I-S - M (SD)                  | 0.88 ± 0.45         | 0.49 ± 0.42         | Z-test = 1 | 0.27   |
| DEP - M (SD)                  | 1.10 ± 0.69         | 0.60 ± 0.54         | Z-test = 1.5 | <0.05  |
| ANX - M (SD)                  | 1.08 ± 0.64         | 0.56 ± 0.43         | Z-test = 1.5 | <0.05  |
| HOS - M (SD)                  | 0.96 ± 0.84         | 0.35 ± 0.36         | Z-test = 1.5 | <0.05  |
| PHOB - M (SD)                 | 0.37 ± 0.53         | 0.21 ± 0.37         | Z-test = 0.68 | 0.77   |
| PAR - M (SD)                  | 0.92 ± 0.55         | 0.42 ± 0.50         | Z-test = 1.68 | <0.01  |
| PSY - M (SD)                  | 0.50 ± 0.41         | 0.21 ± 0.35         | Z-test = 1.5 | <0.05  |
| OTHER - M (SD)                | 1.04 ± 0.82         | 0.90 ± 1.03         | Z-test = 0.68 | 0.77   |

Respectively for TAS+ and TAS− subjects. All EEG analysis were performed by means of the exact Low Resolution Electric Tomography software (eLORETA) software, a validated tool for localizing the electric activity in the brain based on multichannel surface EEG recordings\cite{37}. The eLORETA software benefits from an excellent localization agreement with different multi-modal imaging techniques, also when standard 19-electrodes EEG montage was used\cite{14}.

**Frequency analysis.** Fast Fourier Transform algorithm was used to perform EEG frequency analysis, with 2 seconds interval on the EEG signal, in all scalp locations. In the present study we have considered the following frequency bands: delta (0.5–4 Hz); theta (4.5–7.5 Hz); alpha (8–12.5 Hz); beta (13–30 Hz); and gamma (30.5–60 Hz). EEG frequency analysis was performed using monopolar EEG traces (each electrode referred to joint mastoids). Topographic sources of EEG activities were determined using the eLORETA software. This software calculates the 3-dimensional current distribution throughout the brain volume by assuming that neighboring neurons are activated both simultaneously and synchronously. This is in accordance with results of single cell recordings in the brain\cite{33,34}. The computational task is to select the smoothest 3-dimensional current distribution, which is a widely used procedure in signal processing\cite{35,36}. The result is a true 3-dimensional tomography, in which the localization of brain signals is conserved with a low amount of dispersion\cite{47}.

**Connectivity analysis.** The connectivity analysis was performed by the computation of lagged phase synchronization. This connectivity measure is widely used to assess EEG functional connectivity in both psychiatric and brain diseases\cite{38,39}. Furthermore, compared to other connectivity measure, the lagged phase synchronization has some advantages (e.g. is resistant to non-physiological artifacts and it is minimally affected by low spatial resolution)\cite{37,40}.
chopathology, the GSI was included as a control variable in a partial correlation (significantly modified connections between ROIs. Due to the strong relationship between alexithymia and psy-
were reported as measures of associations among the TAS total scores, TAS sub-scales, SCL-90, along with any

Table 2. Cortical 12 regions of interest (ROIs). Adapted from Thatcher et al.\textsuperscript{14}. Note: ROI = Region of Interest; eLORETA = exact Low Resolution Electric Tomography software; MNI = Montreal Neurological Institute.

| ROI | eLORETA MNI coordinates | Anatomical regions | Brodmann areas |
|-----|-------------------------|--------------------|----------------|
| 1   | x -30 y 40 z 25         | Left Frontal Lobe  | 8-9-10         |
| 2   | x 20 y 35 z 30          | Right Frontal Lobe | 8-9-10         |
| 3   | x -45 y -15 z -25      | Left Temporal Lobe | 21-28-36       |
| 4   | x 55 y -15 z -20       | Right Temporal Lobe | 21-28-36       |
| 5   | x -5 y -5 z 35         | Left Posterior     | 23-24          |
|     | Frontal Cingulate Cortex |                    |                |
| 6   | x 5 y -10 z 30         | Right Posterior    | 23-24          |
|     | Frontal Cingulate Cortex |                    |                |
| 7   | x -5 y 30 z 20         | Left Anterior      | 32             |
|     | Cingulate Cortex       |                    |                |
| 8   | x 5 y 30 z 20          | Right Anterior     | 32             |
|     | Cingulate Cortex       |                    |                |
| 9   | x -5 y -55 z 25        | Left Hippocampus   | 29-30-31       |
| 10  | x 5 y -50 z 25         | Right Hippocampus  | 29-30-31       |
| 11  | x -45 y -50 z 40       | Left Parietal Lobe | 39-40          |
| 12  | x 45 y -50 z 35        | Right Parietal Lobe | 39-40         |

The eLORETA computed the lagged phase synchronization values between all these ROIs (total 12 × 12 = 144 connections). The eLORETA also computed the source reconstruction algorithm previously described\textsuperscript{32-42}. The EEG connectivity analysis was performed on the same blocks of EEG tracings used for power spectra analysis.

**Statistical analysis.** The power spectra analysis and EEG connectivity were compared between TAS+ and TAS− participants for each frequency band. The comparisons were performed using the statistical non-parametric mapping (SnPM) methodology supplied by the eLORETA software\textsuperscript{43}. This methodology is based on the Fisher’s permutation test (i.e., a subset of non-parametric statistics, more details could be found in ref. 43). Correction of significance for multiple testing was computed for the comparisons between TAS+ and TAS− groups for each frequency band. The non-parametric randomization procedure, available in the eLORETA program package, was performed for the correction of multiple comparison (i.e., Holmes’ non-parametric correction for multiple comparisons\textsuperscript{45,44}). The T-level thresholds were computed by the statistical software implemented in the eLORETA, which correspond to statistically significant thresholds ($p < 0.05$ and $p < 0.01$)\textsuperscript{45}. These T-level thresholds and the correspondent $p$ values were provided after applying the correction for multiple comparisons.

Two-way chi-squared and Kolmogorov-Smirnov Z test were used to analyze differences between groups, respectively, for N × N contingency tables and dimensional measures. Spearman’s $\rho$ correlation coefficients were reported as measures of associations among the TAS total scores, TAS sub-scales, SCL-90, along with any significantly modified connections between ROIs. Due to the strong relationship between alexithymia and psychopathology\textsuperscript{5}, the GSI was included as a control variable in a partial correlation ($r_p$) analyses. Furthermore, to determine whether the relationship between alexithymia and EEG connectivity data was mediated by psychopathology, multiple mediation analysis was performed using the “process” macro for SPSS\textsuperscript{46}. This tool assesses direct and indirect effects of one or more independent variables (IV) on a dependent variable (DV) through one or more mediators (M). In the present study we tested a model in which alexithymia severity (TAS total score) is the independent variable, EEG connectivity values (interconnected ROIs) are the dependent variables, and psychopathology (GSI) is a potential mediator. It is important to note that the design of this analysis is correlational in nature, which precludes a causal interpretation of the association between these variables. In the present analyses, we used standardized variables to generate standardized coefficients and corresponding $p$ values. As suggested by Preacher and Hayes\textsuperscript{47}, for indirect effects we also calculated bias-corrected and accelerated 95% CI produced using a bootstrapping method.

Two-way chi-squared test, Kolmogorov-Smirnov Z test and correlation analyses were performed using IBM SPSS Statistics for Windows, version 19.0.

**Results**

EEG recordings suitable were obtained for all participants. Qualitative visual evaluation of the EEG recordings showed no relevant modifications of the background rhythm frequency (e.g. focal abnormalities or epileptic discharges) and no evidence of drowsiness or sleep during the recordings. Differences between groups (TAS− subjects vs. TAS+ subjects) are reported in Table 1. Compared to TAS− subjects, TAS+ individuals reported higher mean scores on the GSI and on six (SOM, O-C, DEP, AXN, HOST, PSY) of the ten dimensions of the SCL-90-R. No significant differences were found for age and for substance use (alcohol and tobacco in the last 6 months) between the two groups.
significant differences were observed in the other frequency bands. eLORETA software localized these modifications in right PCC (Brodmann Area, BA 23; $T^{-} < 0.92$) compared to TAS $^+$ individuals, TAS $^+$ subjects showed a decrease of alpha power in the in right PCC (BA 23; $T^{-} = -3.28, p < 0.05$). Abbreviation: BA = Brodmann areas; PCC = Posterior Cingulate Cortex; $A =$ Anterior; $P =$ Posterior; $L =$ Left; $R =$ Right.

**Power spectra analysis.** In this analysis the thresholds for significance were $T = \pm 3.23$ corresponding to $p < 0.05$, and $T = \pm 3.89$, corresponding to $p < 0.01$. A significant modification was observed in the alpha band: compared to TAS $^-$ individuals, TAS $^+$ subjects showed a decrease of alpha power in the limbic lobe (Fig. 1). The eLORETA software localized these modifications in right PCC (Brodmann Area, BA 23; $T^{-} = -3.28, p < 0.05$). No significant differences were observed in the other frequency bands.

**Connectivity analysis.** In this analysis the thresholds for significance were $T = \pm 3.72$ corresponding to $p < 0.05$, and $T = \pm 4.31$, corresponding to $p < 0.01$. Significant modifications were observed in the alpha band (Fig. 2). Compared to TAS $^-$ individuals, TAS $^+$ subjects showed a decrease of alpha lagged phase synchronization between: (i) ROI 6 (BAs 23–24) and ROI 8 (BA 32) ($T^{-} = -3.87; p < 0.05$), (ii) ROI 2 (BAs 8–9–10) and ROI 6 (BAs 23–24) ($T^{-} = -4.10; p < 0.05$), and (iii) ROI 12 (BAs 39–40) and ROI 4 (BAs 21–28–36) ($T^{-} = -3.92; p < 0.05$).

**Association among modified connections between Regions of Interest (ROIs), TAS and SCL-90-R.** TAS total scores were negatively associated with the modified connections between ROIs 2–6 ($\rho = -0.50; p < 0.01$), ROIs 8–6 ($\rho = -0.52; p < 0.01$), and ROIs 12–4 ($\rho = -0.39; p < 0.05$). Furthermore, significant correlations were also observed between TAS subscales and interconnected ROIs. The association among TAS total scores, and modified connections between ROIs 2–6 ($r_p = -0.39; p < 0.05$), ROIs 8–6 ($r_p = -0.44; p < 0.01$), and ROIs 12–4 ($r_p = -0.39; p < 0.05$) were significant after controlling for GSI. Correlations and partial correlations are respectively reported in Tables 3 and 4.

The mediation models explained between 15% ($F_{2,33} = 3.96; p < 0.05$; Fig. 3 Model 3) and 20% ($F_{2,33} = 5.31; p < 0.01$; Fig. 3 Model 2) of the data variability. All mediation models indicated that the total effect of TAS-20 on EEG functional connectivity values was significant ($b = -0.01; p < 0.01$), with more severe alexithymia associated with a greater decrease in EEG functional connectivity value. Moreover, the relationship between alexithymia and EEG functional connectivity data was directed and was not mediated by psychopathology severity [$b$ between 0.001 ($p = 0.92$) and 0.002 ($p = 0.69$)]. Finally, in all mediational models no significant direct effect was observed for GSI on EEG functional connectivity values (Fig. 3). Detailed statistics of the mediational models are listed in Table 5.

**Discussion**

The main aim of the present study was to explore the modifications of EEG power spectra and EEG functional connectivity in DMN in alexithymia. Compared to TAS $^-$ individuals, TAS $^+$ subjects showed a decrease of alpha power in the right PCC (BA 23). Furthermore, compared to TAS $^-$ individuals, TAS $^+$ subjects showed a decrease of alpha connectivity between: (i) right ACC and right PCC, (ii) right frontal lobe and right PCC, and (iii) right parietal lobe and right temporal lobe. EEG connectivity values were also negatively related to TAS total score after controlling for general psychopathology, which is known to be associated with alexithymia. Finally, our mediation models results showed that the relationship between alexithymia and EEG functional connectivity value was directed (i.e., more severe alexithymia associated with a greater decrease in EEG functional connectivity) and was not mediated by psychopathology severity.

Figure 1. Results of the eLORETA comparison of EEG power spectra in each frequency band. Threshold values ($T$) for statistical significance (corresponding to $p < 0.05$) are reported in the right side of the figure. Blue color indicates reduction of EEG power spectra. Compared to TAS $^-$ individuals, TAS $^+$ subjects showed a decrease of alpha power in the right PCC (Brodmann Area, BA 23; $T^{-} = -3.28, p < 0.05$). Abbreviation: BA = Brodmann areas; PCC = Posterior Cingulate Cortex; $A =$ Anterior; $P =$ Posterior; $L =$ Left; $R =$ Right.
Taken together, our results could reflect the neurophysiological substrate of several core features of alexithymia, such as the impairment in emotional awareness as well as the alexithymia-related imagery deficit. Our results are consistent with previous EEG and fMRI studies suggesting that alexithymia is characterized by a disruption in the integrated cortical neural network not only during emotional tasks but also during RS condition. Our study differs from and adds to previous findings by investigating EEG modifications during RS condition.
is thought to reflect intrinsic activity in the brain revealing valuable information on how different structures communicate.

Our results are in line with previous studies reporting the involvement of alpha frequency in alexithymia. Indeed, the decrease of both EEG power and EEG connectivity in the alpha frequency band was previously detected in alexithymic subjects in response to emotional visual stimuli. Due to the strong association between alpha frequency and cognitive processing related to external attention, it has been hypothesized that this neurophysiological pattern could reflect the externally-oriented thinking of alexithymic subjects.

Our data are also in accordance with previous functional imaging studies confirming the crucial role of the cingulate cortex in alexithymia. The PCC has shown to be involved in several emotional and cognitive processing. It is also considered the crucial node of the DMN, with critical relevance in maintaining a sense of self-consciousness. Moreover, PCC is engaged in self-referential thoughts during RS. In a fMRI study, Mantani et al. reported a significant decrease of PCC activation in subjects with high degrees of alexithymia during imagery tasks, suggesting a possible involvement of PCC in alexithymia-related imagery disturbance. On the other hand, the involvement of ACC in alexithymia is also well documented. A reduction of ACC activity in response to emotional stimuli (especially for emotionally negative stimuli) was observed in several neuroimaging studies. Alterations in the cingulate cortex in subjects with alexithymia were also observed by Liemburg et al. during RS condition. Therefore, the decrease of functional connectivity between ACC and PCC observed in our study could reflect both affective and cognitive disturbances in alexithymia (i.e. the impairment in emotional awareness and the externally-oriented thinking). This hypothesis is also in accordance with our correlational data (i.e., negative correlation between ACC/PCC connectivity and both DDF and EOT TAS-20 factors). Compared to Liemburg et al., we did not observe an increase of functional connectivity in brain areas involved in sensory input. The authors, using Independent Component Analysis (i.e., a data-driven method that separate the fMRI signal into spatially independent networks), showed that within the DMN, compared to

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Figure 3. Results of the multiple mediation models and related standardized β coefficients. Note. *p < 0.05; **p < 0.01; ***p < 0.001. Abbreviation: TAS-20 = Toronto Alexithymia Scale; GSI = Global Severity Index; ROIs = Regions of interest.
controls, alexithymic participant showed higher functional connectivity in the precentral gyrus and occipital areas suggesting a tendency of the alexithymic towards strong bodily expressions of emotions. Moreover, Liemburg et al. also did not detect a lateralization. The discrepancies between these results and the present research may be explained by differences in their study designs and methods (i.e., EEG vs fMRI, ROIs selections etc.).

In TAS + subjects, a decrease of alpha connectivity was also observed between right frontal lobe and right PCC. The role of frontal areas, especially of the prefrontal cortex (PFC), in alexithymia is still unclear. Although PFC plays a crucial role in several cognitive and emotional functions, such as decision-making and emotion regulation, only few studies reported the association between PFC activity and alexithymia. In a Positron Emission Tomography (PET) study of Kano et al., the alexithymic, compared to non-alexithymic subjects, showed decreased activation of right frontal areas (i.e. inferior and superior frontal cortex and the orbital PFC), in response to negative emotional stimuli. Consistently, lower connectivity within the DMN for the superior frontal gyri in alexithymic individuals was also observed. The PCC connections with PFC are thought to be crucial in the DMN13 and, as already mentioned, the decrease of connectivity between prefrontal regions and cingulate gyrus could reflect the emotional awareness disturbances, especially “the difficulties to put feelings into words”.

We also observed a reduction of EEG connectivity between the right inferior parietal lobule (IPL) and the right temporal lobe. It has been observed that increased connectivity in these areas is associated with envisioning future tasks (i.e., imagine a future situation related to a particular cue, for a review see ref. 13). Therefore, a connectivity decrease between IPL and right temporal lobe may reflect the impairment of affect-related fantasy and imagery activity frequently observed in alexithymic individuals.

**Table 5. Statistical results of mediation models.** Abbreviation: IV = Independent variable; DV = Dependent variable; M = Mediator; TAS-20 = Toronto Alexithymia Scale; GSI = Global Severity Index; ROIs = Regions of interest; CI = Confidence interval.
Finally, we detected a decrease of EEG alpha power and EEG connectivity in the right hemisphere. It has been hypothesized that alexithymia is associated with a disturbance in right hemisphere functioning, especially in the processing of emotions\(^{(5)}\). Therefore, our results seem to be in line with both behavioral studies\(^{(6)}\) and neuroimaging data\(^{(7)}\), suggesting a right hemisphere alteration in alexithymic individuals. The decrease of EEG connectivity and EEG power in the right hemisphere observed in alexithymic subjects is also in accordance with Bucci’s model\(^{(8)}\), suggesting that this neurophysiological pattern may reflect the impairment of the symbolization of emotions, as well as the disturbances in holistic and nonverbal processing of incoming stimuli. It is important to note that our interpretations remain speculative due to the absence of imaginative/emotional tasks in the present study. However, it has been observed that RS activity may predict modifications in behavioral performance and task-evoked brain activity\(^{(9)}\).

There are some limitations in generalizing our results. One limitation that must be considered is that the sample we used is small, and this is especially relevant concerning mediation analysis. Our sample also included mostly female participants. Although conclusive EEG data on DMN gender differences are not yet available, previous studies suggested that women may show different EEG RS brain activity\(^{(10)}\). Therefore, we cannot exclude that women with alexithymia, compared to men with alexithymia, may be characterized by different EEG power and connectivity patterns during RS. Furthermore, we used scalp EEG recordings, which have an intrinsic limit in space resolution. It should be also noted that, although the SCL-90-R is considered a good tool to assess general psychopathology, it does not exclude the presence of a psychiatric disorder. Finally, we have investigated the DMN in a non-clinical sample of alexithymic subjects. Therefore, it is possible that psychiatric patients with alexithymia have different EEG patterns. Although these ideas are purely hypothetical, they might be useful in guiding future research.

Despite these limitations, to the best of our knowledge, this is the first study which simultaneously investigated EEG functional connectivity and EEG power spectra during RS condition in alexithymic subjects, controlling for psychopathology and using an accurate and validated tool (i.e., eLORETA) to localize electric activity in the brain. In conclusion, our results suggest that alexithymia is characterized by a disruption of cortical neural networks, not only during emotional processing, but also during RS condition. This is in accordance with previous hypotheses suggesting that “such distinct patterns of connectivity may be related to the diminished emotional awareness of alexithymic people”\(^{(11)}\). Therefore, our results could be of some clinical relevance both for diagnosis and therapy. On one hand, if confirmed, this EEG pattern could help to measure alexithymia in RS condition; on the other hand it highlights the possibility of developing new therapeutic approaches focused on the self neuro-modulation, such as alpha training neurofeedback.

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**Author Contributions**

C.I.: study design, data analysis, interpretation of results, manuscript writing. G.D.M.: study design, data collection, interpretation of results, manuscript writing. R.B.: study design, manuscript writing. G.C.: data collection, data analysis. C.M.: data collection, data analysis. E.M.V.: data collection, questionnaires. N.A.: data collection, questionnaires. A.C.: study design, questionnaires. B.F.: study design, data analysis, interpretation of results, manuscript writing.

**Additional Information**

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