Autistic-Like Traits in Adult Patients with Mood Disorders and Schizophrenia

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Abstract

Autism spectrum disorder often co-occurs with other psychiatric disorders. Although a high prevalence of autistic-like traits/symptoms has been identified in the pediatric psychiatric population of normal intelligence, there are no reports from adult psychiatric population. This study examined whether there is a greater prevalence of autistic-like traits/symptoms in patients with adult-onset psychiatric disorders such as major depressive disorder (MDD), bipolar disorder, or schizophrenia, and whether such an association is independent of symptom severity. The subjects were 290 adults of normal intelligence between 25 and 59 years of age (MDD, n=125; bipolar disorder, n=56; schizophrenia, n=44; healthy controls, n=65).

Almost half of the clinical subjects, except those with remitted MDD, exhibited autistic-like traits/symptoms at levels typical for sub-threshold or threshold autism spectrum disorder. Furthermore, the proportion of psychiatric patients that demonstrated high autistic-like traits/symptoms was significantly greater than that of healthy controls, and not different between that of remitted or unremitted subjects with bipolar disorder or schizophrenia. On the other hand, remitted subjects with MDD did not differ from healthy controls with regard to the prevalence or degree of high autistic-like traits/symptoms.

A substantial proportion of adults with bipolar disorder and schizophrenia showed high autistic-like traits/symptoms independent of symptom severity, suggesting a shared pathophysiology among autism spectrum disorder and these psychiatric disorders. Conversely, autistic-like traits among subjects with MDD were associated with the depressive symptom severity. These findings suggest the importance of evaluating autistic-like traits/symptoms underlying adult-onset psychiatric disorders for the best-suited treatment. Further studies with a prospective design and larger samples are needed.
Introduction

Autism spectrum disorder (ASD) is an early-onset, life-long developmental disorder characterized by persistent deficits in social reciprocity and social communication, as well as restricted, repetitive patterns of behaviors, interests, or activities [1]. The prevalence of ASD in children and adolescents is estimated to be in the range of 0.6–0.7% [autistic disorder (0.2%), pervasive developmental disorder—not otherwise specified (PDD-NOS) (0.3%), and Asperger’s disorder (0.06%)], and has continually increased over the last 15–20 years [2].

It has been recognized that youth with ASD often have co-occurring psychiatric disorders [3–5]. Such co-occurrence may negatively impact social and academic performance, even for those who have an intelligence quotient (IQ) within the normal range [6]. There have been many reports on the incidence of psychiatric comorbidity with ASD in clinical pediatric cases, although the findings vary widely [4]. There has been only one population-based study, which was conducted in the UK, for 112 children with ASD aged 10 to 14 years [3]. This study revealed that approximately 70% of children with ASD suffered from at least one comorbid Diagnostic Statistical Manual (DSM) axis-I psychiatric disorder such as social anxiety disorder, attention-deficit/hyperactivity disorder (ADHD), and oppositional defiant disorder (30% each). Approximately 10% of children with ASD had experienced a period of depression or irritability, despite the fact that major depressive disorder (MDD) or dysthymic disorder was identified in less than 1% of children with ASD. Psychotic disorders such as schizophrenia (SZ) or bipolar disorder (BPD) were not observed in this sample.

On the other hand, frequent co-occurrence of ASD among psychiatric patients has been identified in several studies of clinical youth aged 7–17 years that had been diagnosed with mood or anxiety disorders [7,8]. Pine et al. [8] observed that 57% of youth patients with BPD, 38% with MDD, and 25% with anxiety disorder exceeded the clinical cut-off for quantitative ASD scales. These findings suggest that autistic-like traits/symptoms (ALTs) as measured by quantitative ASD scales are closely associated with mood and anxiety disorders in pediatric patients, although the etiological relevance is unclear.

The psychosocial outcomes of adults who were diagnosed with ASD in childhood tend to range from poor or very poor in young adulthood (<25 years) [9], to even worse in later adulthood, although the severity of ASD decreases throughout development [10]. Reduced psychosocial quality of life (QOL) in high-functioning adults with ASD was found to be associated with concurrent behavioral and psychiatric problems [11]. Although psychiatric comorbidity, as well as core symptoms, seems to hamper the social adaptation of adults with ASD, there is only one study on the incidence of psychiatric comorbidity in adults with ASD. Hofvander et al. [12] examined patients who visited their clinic or hospital, which specialize in childhood-onset neuropsychiatric disorders, and reported that all patients of normal intelligence that were diagnosed with ASD during childhood (n = 122) had at least one life-time comorbid DSM axis-I psychiatric disorder: the most prevalent was mood disorders (53%), followed by anxiety disorders (50%), ADHD (43%), obsessive-compulsive disorder (24%), chronic tic disorders (20%), substance-related disorders (16%), and psychotic disorders (12%).

Similarly, studies of the non-clinical adult population also provided evidence of the association between ALTs and psychiatric conditions such as depression and anxiety [13, 14]. This is consistent with findings in the non-clinical child population: a sub-clinical level of ALTs was associated with emotional, behavioral, and cognitive problems [15, 16].

To our knowledge, the prevalence of ALTs among the adult psychiatric population outside ASD clinics has not been reported. In Japan, an increasing number of adults with ALTs visit general psychiatric clinics with a diverse range of chief complaints, seeking accurate diagnosis and/or treatment for concurrent psychiatric symptoms [17]. However, unlike children...
diagnosed with ASD, clinical manifestations in adulthood are often complex: core symptoms tend to become less apparent [9], or adults with ASD may use compensatory strategies to mask their deficits. For these reasons, clinical or subclinical ASD symptoms are likely to be overlooked in general psychiatric settings, which can lead to inappropriate treatment [5,18].

The aim of this study was to examine whether higher levels of ALTs are associated with adult-onset psychiatric disorders (i.e., MDD, BPD, and SZ), and whether such an association is independent of symptom severity. We hypothesized that higher ALT levels would be frequently observed among psychiatric adult patients, similar to rates observed among children/adolescents. Regarding the association between ALT degree and symptom severity of non-ASD psychiatric disorders, no specific predictions were made.

Methods
Subjects
Subjects were 290 adults (men, 48%) aged 25 to 59 years. They consisted of 44 patients with SZ (men, 46%), 125 with MDD (56%), 56 with BPD (46%), and 65 healthy controls (HC) (28%) (Table 1). They were recruited through notices posted in the National Center of Neurology and Psychiatry (NCNP) Hospital, website announcements, or advertisements in a local free paper. All subjects other than HC were outpatients who attended either the NCNP hospital or local hospitals/clinics. All subjects were interviewed by a trained psychiatrist using the Japanese version of the Mini-International Neuropsychiatric Interview [19,20], and diagnoses were confirmed based on the DSM of Mental Disorders, 4th ed., text rev. (DSM-IV-TR) [21]. Schizophrenic symptoms were corroborated by administering the Positive and Negative Symptoms Scale (PANSS) to all subjects with SZ [22]. Depression severity was assessed using the 17-item version of the Hamilton Depression Rating Scale (HDRS-17) for all subjects with MDD and BPD [23]. Manic symptoms were assessed by the Young Mania Rating Scale (YMRS) [24] for all subjects with BPD. Based on the definition of remission determined by the International Society for Bipolar Disorders Task Force [25], subjects with BPD whose YMRS total scores were 8 or over were considered to be suffering from a significant manic state and excluded from this study. Subjects with MDD and BPD were divided into remitted and unremitting subgroups based on the total score of the HDRS-17 (≤7: remitted; >7: unremitting). Likewise, subjects with SZ were divided into two subgroups based on their PANSS scores according to the criteria proposed by Andreasen et al. [26]; to meet remission criteria, a subject’s scores for all of the following items must remain under 4 for at least 6 months, and he/she must not have been hospitalized during that period: delusion, conceptual disorganization, hallucinatory behavior, blunt affect, passive apathetic social withdrawal, lack of spontaneity and flow of conversation, maniasms and posturing, unusual thought content. Those already given a clinical diagnosis of ASD were excluded. However, we did not conduct a thorough and comprehensive evaluation of ASD diagnostics for the present study. In addition, those who had intellectual disability were excluded. Full-scale IQs were assessed using the Wechsler Adult Intelligence Scale-Third edition (WAIS-III) [27] for 66 subjects with MDD (53%), 33 with BPD (59%), 36 with SZ (82%), and 58 with HC (89%); the remaining subjects were clinically judged as functioning within normal range.

This study was approved by the National Center of Neurology and Psychiatry Ethics Committee (23–185). Written informed consent was obtained from all participants prior to their inclusion in the study.

Autistic-like traits assessment
The Social Responsiveness Scale for Adults (SRS-A) [28] was distributed to either a family member or a close friend of the subject. The SRS-A is a quantitative measure of ALTs for 19 to
59-year-old adults. Similar to the original SRS [29], which was developed for 4 to 18-year-olds, the SRS-A is to be completed by a family member or a person who knows the subject well enough to provide an accurate account of his/her behaviors during the preceding 6 months. The SRS-A contains 65 Likert-scaled (0–3) items (0: never, 1: sometimes, 2: often, 3: almost always) that are divided into the following 5 subscales: social awareness, social cognition, social communication, social motivation, and autistic mannerisms. The scores are reported to be distributed widely and continuously in the general population from normal to the extreme end of the autistic spectrum [30–34]. According to the original manual [29], T-scores of 59T or less are defined as within the

Table 1. Demographics, clinical data and SRS-A total raw scores of MDD, BPD, SZ and HC.

|                          | MDD (n = 125) | BPD (n = 56) | SZ (n = 44) | HC (n = 65) | Analysis (χ²/ANOVA/t) | Significant pair-wise comparison |
|--------------------------|---------------|--------------|-------------|-------------|-----------------------|---------------------------------|
| **Demographics**         |               |              |             |             |                       |                                 |
| Men (%)                  | 70 (56%)      | 26 (46%)     | 20 (46%)    | 18 (28%)    | χ²(3) = 14, p = 0.003 | MDD, SZ, BPD > HC               |
|                          |               |              |             |             | χ²(2) = 2.2, n.s.    |                                 |
| Age (years)              | 41.5 ±9.2     | 40.4 ±7.8    | 36.9 ±7.5   | 42.2 ±8.2   | F(3, 286) = 3.9, p = 0.01 | HC, MDD > SZ                     |
| Education (years)        | 15.3 ±2.0     | 15.5 ±3.4    | 13.8 ±2.4   | 15.1 ±2.5   | F(3, 286) = 4.6, p = 0.004 | BPD, MDD > SZ                    |
| **Clinical variables**   |               |              |             |             |                       |                                 |
| Full-scale IQ³           |               |              |             |             |                       |                                 |
| M ± SD                   | 109.1 ±11.7   | 105.5 ±14.6  | 96.2 ±12.1  | 111.4 ±13.4 | F(3, 189) = 11.5, p < 0.001 | HC, MDD, BPD > SZ                |
| Range                    | 70–133        | 70–138       | 75–116      | 87–136      |                       |                                 |
| Medication (mg/day)      |               |              |             |             |                       |                                 |
| AD M ± SD²               | 155.9 ±230.5  | 161.3 ±227.3 | 54.5 ±164.9 | -           | F(2, 198) = 3.8, p = 0.024 | BPD, MDD > SZ                    |
| AP M ± SD²               | 55.8 ±130.2   | 89.3 ±169.3  | 467.3 ±530.8| -           | F(2, 198) = 35.8, p < 0.001 | SZ > BPD, MDD                   |
| Remitted N (%)           | 46 (37%)      | 20 (36%)     | 14 (32%)    | -           | χ²(2) = 0.35, n.s.   |                                 |
| HDRS-17 M ± SD           | 11.1 ±7.5     | 11.4 ±7.9    | -           | -           | t(179) = 0.3, n.s.   |                                 |
| YMRS M ± SD              | -             | 1.3 ±1.8     | -           | -           |                       |                                 |
| PANSS M ± SD             |               |              |             |             |                       |                                 |
| positive                 | -             | -            | 14.1 ±4.4   | -           | -                     |                                 |
| negative                 | -             | -            | 16.1 ±5.3   | -           | -                     |                                 |
| general psychopathol.    | -             | -            | 31.6 ±8.6   | -           | -                     |                                 |
| total                    | -             | -            | 61.8 ±15.3  | -           | -                     |                                 |
| **SRS-A total score**    |               |              |             |             |                       |                                 |
| Mean (SD)                | 48.7 (25.3)   | 55.4 (25.8)  | 59.6 (25.0) | 32.5 (19.1) | F(2, 286) = 14.0, p < 0.001 | SZ, BPD, MDD > HC**              |
| Median ± Q³              | 43.0 (15.5)   | 53.0 (18.5)  | 60.0 (21)   | 29.0 (11.5) | χ²(3) = 41.1, p < 0.001 | SZ, BPD > HC***                  |
|                          |               |              |             |             |                       | SZ > MDD*** > HC***              |

³Full-scale IQs were measured for 66 with MDD (53%), 33 with BPD (59%), 36 with SZ (82%) and 58 HC (89%).
²Imipramine equivalent dose of anti-depressant.
¹Chlorpromazine equivalent dose of anti-psychotics.
²Kruskal Wallis test was used for the effect of diagnosis on the SRS-A total raw scores; Mann-Whitney U test for between-group differences.
MDD: major depressive disorder. BPD: bipolar disorder. SZ: schizophrenia. HC: healthy control.
HDRS: Hamilton Depression Rating Score. YMRS: Young Mania Rating Scale. PANSS: Positive and Negative Symptoms Scale.
M: Mean. SD: Standard Deviation. Q: Quartile deviation. n.s. p > 0.05

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normal range, while those of 60T through 75T are defined as within the mild-to-moderate range, typical for high-functioning ASD such as Asperger’s disorder and PDD-NOS, and possibly causing substantial interference with everyday social interactions. T-scores of 76T or higher are defined as within the severe range and are typical for autistic disorder and more severe PDD-NOS, indicating severe interference with everyday social interactions. In this study using the Japanese version [34], we classified two subgroups based on SRS-A T-score [33]: one with 59T or less (low ALT: L-ALT) and another with 60T or higher (high ALT: H-ALT). A T-score of 60T (which is equivalent to raw score ≥53) derived from the Japanese standardization sample corresponds to approximately the 87th percentile in our HC sample, indicating its representativeness. These T-scores were based on combined data from both sexes, because SRS-A raw scores in the standardization sample revealed no statistically significant sex differences among adults 25–59 years old [34]. This was reconfirmed in the current study.

Statistical analysis
Statistical analyses were performed using SPSS version 22.0 (SPSS Japan, Tokyo). The proportion of categorical variables was compared using the chi-squared test. Demographic information (age, and years of education) and clinical information (IQ, and medication) were compared across diagnoses by one-way analysis of variance (ANOVA) and between remitted and unremitted subgroups within each diagnosis by the Student’s t-test. Effects of these variables on SRS-A score were examined by multiple regression analysis. Since SRS-A data were skewed, non-parametric analyses were used for the subsequent analyses. The SRS-A raw scores were compared among the 7 subgroups (HC and remitted and unremitted subgroups of MDD, BPD, and SZ) using a Kruskal-Wallis test; between-group differences were examined by a pairwise multiple comparison test using rank sums proposed by Dunn [35]. To examine the association between SRS-A raw scores and the symptom severity of each psychiatric disorder, Spearman’s correlation coefficients were used between the HDRS-17 and SRS-A raw scores in the MDD and BPD groups, and between the PANSS and SRS-A raw scores for patients with SZ. Statistical significance was set at a two-tailed p<0.05.

Results
Demographic and clinical characteristics
The demographic and clinical characteristics of the subjects are shown in Table 1. The male to female ratio was significantly higher in the MDD, SZ, and BPD groups than in the HC, but did not differ across the three clinical groups. Age, education, IQ, and medication differed significantly between the SZ group and the remaining three groups (MDD, BPD, and HC); however, multiple regression analysis in which SRS-A raw score was a dependent variable and these demographic and clinical variables were explanatory variables yielded no significant effects (p>0.05). The ratio of remitted versus unremitted subjects was not significantly different among the MDD, BPD, and SZ groups. Between remitted and unremitted subgroups within each diagnosis, there were no significant differences in demographics or clinical variables (age, years of education, IQ, age of onset, duration of illness, history of being hospitalized, hospitalized months, and recurrent episode) (S1 Table). Since the effect of sex on SRS-A raw scores was not significant in either diagnosis (S2 Table), data derived from both sexes were combined in the subsequent analyses. The distribution of SRS-A total raw scores is shown by diagnostic group in Fig 1.
Comparison of the degree of ALTs among the 7 subgroups

All the clinical subgroups, whether subjects were remitted or unremitted, except for the remitted MDD subgroup, had significantly higher total and social communication and autistic mannerisms subscale scores on the SRS-A compared to the HC group (Fig 2; S3 Table). The remitted MDD subgroup scored significantly lower overall and on the social cognition and social communication subscales relative to the unremitted SZ group, and scored lower on the social motivation subscale than did the unremitted MDD subgroup (as highlighted in S3 Table).

Comparison of the proportion of H-ALTs among the 7 subgroups

The number and percentage of H-ALTs in each subgroup are shown in S4 Table. The proportion of H-ALTs in the remitted MDD group (22%) was significantly lower compared to that in the unremitted MDD group (46%) ($\chi^2 = 7.1$, two-tailed $p = 0.012$), but did not differ significantly from that of the HC group (14%). The proportion of H-ALTs in the BPD (50%), SZ (61%) and unremitted MDD groups was significantly higher than that of the HC and remitted MDD groups, but did not significantly differ from each other. The BPD and SZ subjects did not differ regarding the proportion of H-ALTs between remitted and unremitted subgroups.
Associations between ALTs and symptom severity of psychiatric disorders

In the MDD group, including remitted and unremitted subjects, SRS-A raw scores were moderately correlated with HDRS-17 total score (\( r = 0.32, p < 0.001 \)). On the contrary, no significant correlations between the SRS-A and HDRS-17 total scores were found for the BPD group, or between the SRS-A and PANSS total scores for the SZ group. Scatter plots are shown for the MDD, BPD, and SZ groups (Fig 3).

Discussion

To our knowledge, this is the first study to examine the distribution of ALTs in the adult psychiatric population diagnosed with MDD, BPD, or SZ, and the possible association between ALTs and psychiatric symptomatology.

We found that adult subjects diagnosed with MDD, BPD, or SZ exhibited a higher degree of ALTs compared to HC. Almost half of the clinical subjects, except those with remitted MDD, fell into the mild-to-severe range for ALTs, which is typical for sub-threshold or threshold ASD. This finding suggests an aggregation of ALTs in individuals with adult-onset psychiatric disorders, and together with the findings about concurrent psychiatric disorders in clinical youths [7,8,16,29], implies a possible pathophysiological overlap between ASD and other psychiatric disorders. Furthermore, our findings may be in accordance with evidence from a recent
A genetic study examining risk loci demonstrating shared effects on ASD, ADHD, BPD, MDD, and SZ [36], with a stronger polygenic overlap between ASD and SZ and between ASD and BPD rather than between ASD and MDD. Similarly, neurocognitive and phenomenological commonalities between ASD and SZ [18, 37, 38] and ASD and BPD [39] have been noted in the literature. However, behavioral commonality between ASD and adult-onset psychiatric disorders observed in the present study does not necessarily imply etiological commonality. Interpretative caution must be taken as future research emerges.

In our study, the degree of ALTs in BPD and SZ subjects was not associated with symptom severity. Further, the proportion of subjects with high ALTs did not significantly differ between remitted and unremitted subjects in the BPD and SZ groups. Thus, ALTs as measured by the SRS-A appear independent of psychiatric symptoms in this population, and we consequently conclude that ALTs in our psychiatric population (except for MDD subjects) remain at a high level regardless of the symptom severity of the psychiatric disorder.

On the other hand, ALTs in subjects with MDD were associated with the depressive symptom severity in our study; in other words, although subjects with severe depressive symptoms tended to exhibit high ALTs, subjects with less severe depressive symptoms did not differ from healthy controls with regard to the proportion or degree of high ALTs. It is difficult to explain this finding because we did not examine intra-individual differences. The reason may be complicated because ASD and depression have several common symptoms such as social withdrawal and obsessionality. However, one possible interpretation would be that the ALTs shown by some depressive patients may decrease when depression is remitted, but resurface when depressive symptoms worsen, which would lead to misdiagnosis confounding ALTs with depressive symptoms [40]. This possibility is also mentioned in the SRS manual [28]; however, there is no evidence for this at present.

There are several limitations to this study. First, a significantly lower male to female ratio in the HC relative to the patient groups may have exaggerated the true differences between these groups. However, we confirmed that the effect of sex on SRS-A raw scores was not significant for either patient group or the HC group. Second, we did not conduct a thorough and comprehensive evaluation of ASD during our exclusion procedure. Therefore, we cannot deny the possibility that some individuals undiagnosed with ASD in the present study could have met the

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**Fig 3. Scatter plots of SRS-A total raw score and symptom severity.** Significant correlation was found between the SRS-A and HDRS-17 total in the MDD group ($r = 0.32, p < 0.001$), but not in the BPD group ($r = 0.16, n.s.$). No correlation was found between the SRS-A and PANSS total in the SZ group ($r = 0.25, n.s.$).

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ASD diagnostic criteria. This is because SRS-A scores for some patients exceeded the cutoff [34]. Although we found an overlap between high ALTs and adult-onset psychiatric disorders, irrespective of the diagnostic status for ASD in the present study, replication studies should be conducted assessing adult non-ASD psychiatric patients via more thorough diagnostic procedures. Third, the sample size was relatively small. Fourth, this was a cross-sectional study; however, we did confirm that remitted and unremitted subgroups in this study did not significantly differ in terms of demographic or clinical variables. Nonetheless, our findings need to be re-examined by performing a longitudinal study. Fifth, the demographic variables of patients with SZ were not matched to those of other diagnostic patients, although it was a natural discrepancy. However, as these variables had no significant effect on SRS-A score, it is unlikely that this discrepancy affected our results.

To conclude, the presentation of ALTs at the sub-threshold or threshold level may be closely associated with BPD and SZ. High ALTs were observed irrespective of symptom severity in a subgroup of subjects with BPD and SZ, suggesting a shared pathophysiology among BPD, SZ, and ASD. Conversely, ALTs among subjects with MDD were associated with depressive symptom severity, which should be acknowledged when assessing ALTs among MDD patients. Whether ALTs observed among subjects with MDD are trait- or state-dependent should be examined in future studies using a prospective design with a larger sample. Our results stress the importance of clinicians’ attention to underlying sub-threshold or threshold ALTs in adult patients with MDD, BPD, and SZ. Untangling persistent ALTs from childhood to adulthood and adult-onset psychiatric disorders would be helpful in the choice of an appropriate treatment plan for such patients. In general psychiatric settings, the use of validated ASD symptom scales, such as the SRS-A, represents an easy and effective way to screen for sub-threshold or threshold ASD, determine accurate diagnosis, provide the best-suited treatment, and subsequently evaluate the effects of treatment.

Supporting Information

S1 Table. Demographics and clinical data of remitted and unremitted MDD, BPD, and SZ subjects. *Hospitalized months were compared by Mann-Whitney U test. MDD: major depressive disorder. BPD: bipolar disorder. SZ: schizophrenia. PANSS: Positive and Negative Symptoms Scale. M: Mean. SD: Standard Deviation. n.s. p>0.05. (XLSX)

S2 Table. SRS-A total raw score by diagnosis and sex. aMann-Whitney U test was used to identify the effect of sex for each diagnosis. bChi-squared test was used to compare the proportion of L-ALT and H-ALT between diagnoses. M-W: Mann-Whitney U test. L-ALT: low autistic-like traits. H-ALT: high autistic-like traits. MDD: major depressive disorder. BPD: bipolar disorder. SZ: schizophrenia. HC: healthy controls. Q: quartile deviation. n.s. p>0.05. (XLSX)

S3 Table. Comparison of SRS-A total and subscale raw scores among the 7 subgroups. aKruskal Wallis test was used among the above 7 subgroups (i.e., HC and remitted/unremitted MDD, BPD, and SZ) to examine the effect of diagnosis on the SRS-A total and subscale scores. bPairwise multiple comparison test using rank sums proposed by Dunn OJ was used to further identify the between-group differences. HC: healthy control. MDD: major depressive disorder. BPD: bipolar disorder. SZ: schizophrenia. R: remitted. U: unremitted. Q: quartile deviation. * p<0.05, ** p<0.01, *** p<0.001. (XLSX)
S4 Table. Comparison of the proportion of H-ALTs among the 7 subgroups. *Chi-squared test was used to compare the proportion of L-ALT and H-ALT between diagnoses. HC: healthy control. MDD: major depressive disorder. BPD: bipolar disorder. SZ: schizophrenia. R: remitted. U: unremitted. L-ALT: low autistic-like traits. H-ALT: high autistic-like traits. ’ p<0.05, ** p<0.01, *** p<0.001.

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Author Contributions
Conceived and designed the experiments: JM TH HK. Performed the experiments: JM MO TT HH AN RT. Analyzed the data: JM. Wrote the paper: JM YK HT NM HK.

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