External Ear Anomalies and Minor Physical Anomalies in Depressive Disorder Patients and Healthy Controls

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ABSTRACT

Objective: To determine if external ear anomalies (EEAs) and minor physical anomalies (MPAs) are more prevalent in patients with depressive disorder than in healthy controls.

Methods: This cross-sectional study was conducted at a tertiary-level referral center between October 1, 2019, and September 30, 2020, and included 100 patients with depressive disorder (diagnosed per ICD-10 criteria) and 100 aged- and sex-matched healthy controls. The study participants were examined using the External Ear Anomalies Assessment Scale and the extended Waldrop Scale.

Results: Independent samples Mann-Whitney U test showed a higher prevalence of mean EEAs and MPAs in patients with depressive disorder. Adherent ear lobe was the most common ear anomaly in both patients (52%) and controls (41%), followed by Darwinian tubercle (21% in the patient group and 19% in the control group).

Conclusions: External ear anomalies are more prevalent in patients with depressive disorder, supporting the neurodevelopmental theory of depression. These EEAs need further description and attention for possible inclusion in scales that assess minor physical anomalies and may be used as an endophenotypic marker for depression in the future.

Depression is one of the most common mental disorders, causing major impairment1 and leading to increased mortality and morbidity.2,3 The external ear develops from the 6 auricular hillocks around the pharyngeal groove by the first and second branchial arch. These hillocks are fused by the end of 8 weeks’ gestation, leading to the characteristic shape of the ear.4,5 The external ear has a complex structure and shape that is species specific and remarkably constant in its basic normal shape.6 Yet, most of the dysmorphology literature pays poor attention to the accurate description of the ear and the definitions of abnormalities. Some of the external ear anomalies (EEAs) are known to be specific for some illnesses and are useful in their diagnosis (eg, ear lobe creases in Beckwith-Wiedemann syndrome, prominent crus of the helix in fetal alcohol syndrome and Saethre-Chotzen syndrome).4

There are various theories of development of depression and understanding of mood disorders, one of which is the “neurodevelopmental theory” that focuses on epigenetic mechanisms reflecting inherited changes in gene expression.7 The neurodevelopmental theory has been studied more in schizophrenia than in depressive disorders.8 There are a few factors that define the neurodevelopmental origin of schizophrenia, termed curious epiphenomena of schizophrenia,9 such as neuromotor anomalies and minor physical anomalies (MPAs), including EEAs.10 The similarities in origin between schizophrenia and major depressive disorder and bipolar disorder have raised questions regarding the role of neurodevelopment.11,12 Among the MPAs, the external ear is one of the most important in studying the major malformations as well as minor anomalies of phenotypic variations.4 It seems possible that a careful examination and detailed description of the ear in those with malformations and/or dysmorphic signs may prove to be of diagnostic value in specific syndromes that may have a neuroectodermal origin in the external ear and the brain.6 A study by Praharaj et al13 found that among the ear abnormalities, prominent crus of helix and abnormal anterior surface were significantly more prevalent in patients with schizophrenia compared to those with bipolar disorder, while the frequency of asymmetrical ears and auricular pits was more common in patients with bipolar disorder. They concluded that prominent crus of helix and ear lobe crease could differentiate patients with schizophrenia and bipolar disorder.13

The MPAs are clinically or cosmetically insignificant errors that occur during the development of morphologic
structures such as the mouth, ears, eyes, head, hands, and feet. MPAs develop during the early stage of gestation but persist into adulthood and can easily be detected or observed by careful examination of the defined area. As the structures that express MPAs are of the same embryonic origin as the central nervous system, MPAs are valuable biological markers of abnormal brain development.

We conducted a cross-sectional study to explore various EEAs in patients with major depressive disorder compared to a healthy control group. The aim of the study was to evaluate EEAs in patients with depression and their relation to the overall frequency of MPAs and other clinical variables.

METHODS

This cross-sectional study was conducted at a tertiary-level referral center between October 1, 2019, and September 30, 2020. The study was approved by the Institute Ethics Committee of All India Institute of Medical Sciences (AIIMS), Raipur, India (IEC no. AIIMSRPR/IEC/2019/338). Written informed consent was obtained from all participants before registering them for the study.

Participants

Patients were recruited by purposive sampling from the outpatient and inpatient services of AIIMS, Raipur. The sample included 114 patients aged 18–65 years with a diagnosis of depressive disorder (first depressive episode/bipolar/recurrent) in any phase of illness as per ICD-10 diagnostic criteria for research. Patients who were unwilling or had a history of neurologic illness, organic brain disorders, mental retardation, dementia, any substance dependence except for caffeine and tobacco, other psychiatric disorder, disruptive behavior, any physical trauma, or surgery that would affect the EEAs and MPAs were excluded from the study. The final sample included 100 patients. The healthy control group included 100 age- and gender-matched participants recruited from the hospital staff and general medicine outpatient department (attenders with patients) who were not first-degree relatives of the patients.

Clinical Assessment

All the required sociodemographic data were collected from the participants after written consent was obtained. Severity of depressive episode was assessed by the Hamilton Depression Rating Scale (HDRS). Healthy controls were screened with the General Health Questionnaire-5 (GHQ-5), only those who scored <2 were included in the study. To assess the EEAs, the External Ear Anomalies Assessment Scale developed by Praharaj et al. was used. A modified version of the Waldrop Scale, the extended Waldrop Scale, was used to assess the minor physical anomalies in all the participants. All 54 items of the extended Waldrop Scale were scored as absent or present, and all the participants were assessed for MPAs by a trained rater (P.K.S.).

Statistical Analysis

The Shapiro-Wilk test was used to check whether continuous variables were in a normal distribution. For both groups, differences for categorical and continuous variables were computed using the χ² test and independent samples Mann-Whitney U test, respectively. Comparison among types of depression (first episode/bipolar/recurrent) was done using the Kruskal-Wallis test (1-way analysis of variance). Spearman correlation coefficients were computed between MPA scores and External Ear Anomalies Assessment Scale scores. Statistical analysis was done using SPSS 23.0. The level of significance was kept at .05.

RESULTS

The mean ± SD age of the patient group was 34.4 ± 10.9 years and of the control group was 32.07 ± 11.2 years (Mann-Whitney U = 0.60). Most of the patient population were male (69%) and Hindu (93%). There was a significant difference in occupational and family income status of patients and controls (P < .00), with more of the controls being employed and having family incomes > 10,000 INR (Indian rupees)/month. For the patient group, the mean ± SD duration of illness was 2.55 ± 3.29 years, with an age at onset of 31.79 ± 10.70 years. Most of the patients had first-episode depression (64%), while 8% had bipolar disorder and 28% had recurrent depressive disorder.

When comparing ear anomalies in the 2 groups, no significant difference in the frequencies of total number of ear anomalies was seen between patients and controls. Regarding mean EEAs, the patients had significantly higher

Table 1. Comparison of Mean External Ear Anomalies and Mean Minor Physical Anomalies in Both Groups by Independent Sample Mann-Whitney U Test

<table>
<thead>
<tr>
<th>Anomaly</th>
<th>Group Statistics</th>
<th>Mean</th>
<th>Median</th>
<th>SD</th>
<th>Mean Rank</th>
<th>Mean Standard Error</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>External ear anomalies</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.027</td>
</tr>
<tr>
<td>Patients</td>
<td></td>
<td>1.53</td>
<td>1.167</td>
<td>109.21</td>
<td>0.117</td>
<td>0.107</td>
<td></td>
</tr>
<tr>
<td>Controls</td>
<td></td>
<td>1.16</td>
<td>1.070</td>
<td>91.79</td>
<td>0.107</td>
<td>0.000</td>
<td></td>
</tr>
<tr>
<td>Minor physical anomalies</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Patients</td>
<td></td>
<td>3.04</td>
<td>1.517</td>
<td>118.57</td>
<td>0.152</td>
<td>0.130</td>
<td></td>
</tr>
<tr>
<td>Controls</td>
<td></td>
<td>2.09</td>
<td>1.303</td>
<td>82.43</td>
<td>0.130</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Bolding indicates statistical significance.
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**DISCUSSION**

This pilot study revealed that among external ear anomalies, adherent earlobe, Darwinian tubercle, earlobe crease, and antverted ear lobe were more prevalent in the patient group than in the control group, although most of these anomalies were not included in the standard MPA scales. The groups had significantly different numbers of total mean EEAs and MPAs (patients = 1.53, controls = 1.16, $P < .027$) and mean total MPAs (patients = 3.04, controls = 2.09, $P < .000$).

Per these results, we can hypothesize that EEAs and MPAs are found predominantly in patients with depressive disorders compared to healthy controls. Based on earlier studies and the literature, these findings suggest a similar origin of brain and ear anomalies that point toward the neurodevelopmental origin of depression. EEAs can be assessed as an endophenotype for depressive disorder due to their higher prevalence in patients with depression. By assessing EEAs and MPAs, we may be able to identify populations who are prone to develop depression; however, this does not aid in diagnosis, as it is a nonspecific indicator for depressive disorder. The clinical relevance of EEAs and MPAs in the differential diagnosis and as a prognostic indicator is not yet clear. They can be used as an endophenotypic marker for depression or as a screening indicator for depressive disorder.

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**Table 2. Frequencies of External Ear Anomalies**

<table>
<thead>
<tr>
<th>Anomaly</th>
<th>Patients (n = 100)</th>
<th>Controls (n = 100)</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low seated ears</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Posteriorly rotated ear</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Asymmetric ear</td>
<td>3</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>Adherent ear lobe</td>
<td>52</td>
<td>41</td>
<td>93</td>
</tr>
<tr>
<td>Ear lobe crease</td>
<td>16</td>
<td>5</td>
<td>21</td>
</tr>
<tr>
<td>Cleft ear lobe</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Darwinian tubercle</td>
<td>21</td>
<td>19</td>
<td>40</td>
</tr>
<tr>
<td>Auricular pits</td>
<td>10</td>
<td>4</td>
<td>14</td>
</tr>
<tr>
<td>Indentation behind the helix</td>
<td>5</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>Cupidal ear</td>
<td>15</td>
<td>19</td>
<td>34</td>
</tr>
<tr>
<td>Prominent crus of helix</td>
<td>4</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td>Thickened ear lobe</td>
<td>5</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>Anteverted ear lobe</td>
<td>10</td>
<td>4</td>
<td>14</td>
</tr>
<tr>
<td>Abnormal anterior surface</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Protruding ear</td>
<td>12</td>
<td>10</td>
<td>22</td>
</tr>
<tr>
<td>Stahl ear</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

*Bolding indicates the most common in both groups.*

**Table 3. Comparison of the Frequency of All Minor Physical Anomalies in Both Groups by Pearson χ² Test**

<table>
<thead>
<tr>
<th>Range</th>
<th>Total</th>
<th>$\chi^2$/df</th>
<th>$P^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total no. of minor physical anomalies</td>
<td>14</td>
<td>27.097/7</td>
<td>.000</td>
</tr>
<tr>
<td>0</td>
<td>4</td>
<td>10</td>
<td>14</td>
</tr>
<tr>
<td>1</td>
<td>12</td>
<td>28</td>
<td>40</td>
</tr>
<tr>
<td>2</td>
<td>21</td>
<td>23</td>
<td>44</td>
</tr>
<tr>
<td>3</td>
<td>23</td>
<td>13</td>
<td>36</td>
</tr>
<tr>
<td>4</td>
<td>16</td>
<td>1</td>
<td>16</td>
</tr>
<tr>
<td>5</td>
<td>6</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>6</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
</tbody>
</table>

*Bolding indicates statistical significance.

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**Table 4. Comparing the Minor Physical Anomalies of Different Regions Assessed by the Extended Waldrop Scale**

<table>
<thead>
<tr>
<th>Group (n = 100)</th>
<th>Ear Region</th>
<th>Head Region</th>
<th>Mouth Region</th>
<th>Eye Region</th>
<th>Trunk Region</th>
<th>Limb Region</th>
</tr>
</thead>
<tbody>
<tr>
<td>Patients</td>
<td>0.980 (.73)</td>
<td>0.210 (.40)</td>
<td>0.590 (.69)</td>
<td>0.320 (.46)</td>
<td>0.400 (.48)</td>
<td>0.580 (.63)</td>
</tr>
<tr>
<td>Controls</td>
<td>0.939 (.89)</td>
<td>0.160 (.36)</td>
<td>0.340 (.51)</td>
<td>0.200 (.40)</td>
<td>0.320 (.60)</td>
<td>0.150 (.38)</td>
</tr>
</tbody>
</table>

*Mann-Whitney U test*

<table>
<thead>
<tr>
<th>Level of significance</th>
<th>Nonsignificant</th>
<th>Nonsignificant</th>
<th>Significant</th>
<th>Nonsignificant</th>
<th>Nonsignificant</th>
<th>Significant</th>
</tr>
</thead>
<tbody>
<tr>
<td>$P = .471$</td>
<td></td>
<td></td>
<td>$P = .011$</td>
<td></td>
<td></td>
<td>$P = .054$</td>
</tr>
<tr>
<td>$P = .364$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>$P = .180$</td>
</tr>
<tr>
<td>$P = .000$</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Values are presented as mean (SD).*
tool in high-risk populations for which further evaluation is to be done.

Figure 1 provides a comparison of the current and previous studies. The results of Sivkov and Akabaliev,22 and Culav-Sumić and Jukić25 regarding mean total MPAs are comparable to our study; however, we found no previous studies correlating the EEAs in patients with depressive disorders.

In our study, patients with depressive disorder showed an upward shift in the frequency distribution of total MPAs, which denoted a high anomaly load in this group. Correlation of the frequencies of total number of MPAs and of EEAs by Spearman correlation coefficient showed that a higher number of total MPAs were significantly (P < .000) associated with a higher number of EEAs. Hence, assessment of ear anomalies can be an easier and more efficient way to assess neurodevelopmental defects.

Our study revealed a significant difference in mean MPAs of the mouth (P < .011) and limb regions (P < .000) between patients with depressive disorders and the control group. However, we could not establish a significant difference in mean ear anomalies as per standard scales of MPAs (extended Waldrop Scale), but we could significantly differentiate the mean EEAs in both groups using the External Ear Anomalies Assessment Scale. This finding indicates that current standards are still restricted with regard to range and lack the accuracy to assess anomalies related to the ear region. These EEAs are worth including in scales that assess minor physical anomalies.

Strength and Limitations

A strength of this study is the larger sample size. We also included definitions of ear anomalies (Appendix 1) along with images of anomalies found in the study groups (Appendix 2). However, our study has some limitations, as we defined the MPAs as either “present” or “absent,” while some studies such as Akabaliev et al26 used a graded manner of assessment for some of the MPAs like fine electric hair, head circumference, epicanthus, intercanthal distance, low-seated ears, and high palate.24 Our evaluation for EEAs was not blinded and was done by only 1 investigator. A majority of the patient population had first-episode depression for which further follow-up could not be done, as some of those patients would develop a manic/hypomanic episode that would change the diagnosis to bipolar disorder. Other limitations of the study include the sample size not meeting the criteria for an effective sample size and that the evaluation was conducted by only one investigator. A design in which 2 or more raters independently evaluated subjects using a blinding protocol would be a great improvement.

CONCLUSION

External ear anomalies are more prevalent in patients with depressive disorder, supporting the neurodevelopmental theory of depression. These EEAs need further description and attention for possible inclusion in scales that assess minor physical anomalies, which will strengthen the literature and their use as an endophenotypic marker for depression in the future.

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Supplementary Material: See accompanying pages.
Supplementary Material

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**DOI Number:** https://doi.org/10.4088/PCC.22m03416

**LIST OF SUPPLEMENTARY MATERIAL FOR THE ARTICLE**

1. Appendix 1
2. Appendix 2

**DISCLAIMER**

This Supplementary Material has been provided by the author(s) as an enhancement to the published article. It has been approved by peer review; however, it has undergone neither editing nor formatting by in-house editorial staff. The material is presented in the manner supplied by the author.
Appendix 1:
List and definitions of EEAs adapted from Hunter and Yotsuyanagi (2005), Stevenson and Hall (2006), and Kumar and Burton (2008)

1. Low-seated ear: The root of the helix is below the line connecting the external occipital protuberance to the lateral angle of the eye.
2. Posteriorly rotated ears: The angle between the facial plane and the long axis of the ear exceeding 20°.
3. Asymmetrical ears: Obvious asymmetry between both ears.
5. Ear lobe creases: Linear fissures on the lobule of the ear.
7. Darwinian tubercle: A small protrusion or notch on the helix of the auricle one-third of the way beyond the upper tip of the helix.
8. Auricular pits/tags: Pit-like depressions, dimples, or fossae, or skin tags usually just at the anterior margin of the ascending limb of the helix.
9. Indentation behind the helix: Notch or indentation behind the helix.
10. Cuspidal ear: Prominent triangular form of the upper ridge of the auricle.
11. Prominent crux of helix: An unusual prominence or posterior flaring of the crus of the helix.
12. Thickened ear lobe: Thickening of the ear lobe.
14. Abnormal anterior surface: Anterior and inferior folding of the upper portion of the ear that obliterates triangular fossa and scapha.
15. Protruding ear (Bat ear): Angle relative to the mastoid bone is>40° or where the outer edge of the helix is more than 2 cm separated from the mastoid.
16. Stahl ear: It is a distinctive extra fold of crus of the antihelix that extends from the superior portion of the antihelix to the upper posterior aspect of the corner of the helix, producing a “crumpled” ear appearance.
Appendix 2:
Consent was taken from the patients for publication of the pictures of the anomalies.

Pictures showing Adherent ear lobe

Pictures showing asymmetrical ear

Picture showing cuspidal ear (Note that cuspidal ears have angled ridges instead of a round curve, at the top of auricles.)
Pictures showing double anti-helix

Pictures showing indentation behind the helix

Picture showing low-seated ear

Picture showing protruding ear
Picture showing pre-auricular tag

Picture showing ear-lobe crease

Picture A and B showing antverted ear lobe (A. also having double anti-helix and indentation behind the helix while B. showing darwinian tubercle)

Picture showing A. cup ear and B. double anti-helix
Picture showing Darwinian tubercle

Picture showing thickened ear-lobe

Picture showing auricular pits