Clinical Correlates of Alcohol Use Disorder in Depressed Patients with Unipolar and Bipolar Disorder

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Objective Alcohol use disorder (AUD) is one of the most frequent comorbid conditions in mood disorders. We aimed to examine the relationships between clinical phenotypes of acutely depressed subjects and co-occurring AUD.

Methods Clinical assessment including diagnosis of mood disorder and co-occurring AUD, the severity of depressive or manic symptoms, and affective temperaments were conducted in 137 subjects suffering from a major depressive episode. According to the presence of AUD, clinical variables were compared between the two groups. Using binary logistic regression models, the effects of mood symptoms and affective temperaments on the risk of AUD were determined.

Results Severity of manic symptoms, suicidal ideation, and childhood trauma were higher in the AUD group than in the non-AUD group. Scores for irritable and hyperthymic temperaments were higher and the score for anxious temperament was lower in the AUD group. In regression models adjusting confounders, anxious temperament was an independent protector against AUD. On the other hand, the diagnosis of bipolar disorder and the irritable manic symptom dimension increased the risk of AUD.

Conclusion Anxious temperament decreased the AUD risk, whereas irritable manic symptoms increased the risk during depression. AUD in mood disorders may be an expression of manic psychopathology.

Key Words Major depressive episode, Alcohol use disorder, Bipolar disorder, Affective temperaments, Manic symptoms dimension.

INTRODUCTION

Alcohol use disorder (AUD) is over-represented in subjects with mood disorders and has a negative impact on the course of the mood disorder. Prevalence rates of AUD among subjects with bipolar disorder (BD) range from 30% to 60%1-3 and those among subjects with major depressive disorder (MDD) from 30% to 40%.4-6 The presence of a comorbid AUD in BD has been known to be associated with an earlier onset of the mood symptoms,4 more relapses,9,10 increased suicide risks,11 violent behaviors,12 more hospitalizations,1 and treatment non-adherence.13 Similarly, comorbid AUD in MDD increases disease burden, such as increased suicide risks or other unnatural death,14 lower functioning,9 and worse outcome,15 as well.

Given the paucity of evidence supporting effective treatments,16-18 subjects with AUD and depression present a big clinical challenge to clinicians. Hypothetical explanations for the relationship between the two conditions are conflicting. The self-medication hypothesis explains substance use as an attempt to alleviate mood symptoms, while other hypothesis postulates that repeated substance administration causes brain change to mediate mood disorders.19,20 Another potential perspective assumes that both conditions share common neurobiological vulnerabilities.21 To better understand the interactive and bidirectional relationships between the two conditions, and thus to develop efficient management strategies, many questions remaining unanswered should be more extensively investigated. One pressing question is exploring phenomenological correlates of AUD in mood disorders. Though several risk indicators including male gender, earlier onset, polysubstance use and family history of AUD16,22 were found, characteristics of mood symptoms have not been fully explored yet. Distinct phenotypic expressions of mood disorders may have differential relationships with AUD, which
would allow to identify better diagnoses and treatments.

To our knowledge, few studies have analyzed mood symptoms that were related to AUD. More depressive symptoms were reported in MDD comorbid with AUD in general population and clinical samples. However, findings of these studies that used different measures are quite inconsistent to limit their generalization. Furthermore, no studies have explored the relation of manic symptoms to comorbid AUD.

Affective temperaments are a collection of biologically determined traits concept of which has been developed according to ‘basic states’ described by Kraepelin. They are postulated as a continuum of several conditions encompassing normal personality functioning, subclinical affective state, and predisposition for more severe mood episodes. Therefore, affective temperaments can serve as an important phenotype in mood disorder that complements symptom expressions of a mood episode. To date, little has focused on relations of affective temperaments to AUD in mood disorders. Only one study has assessed the relations and found that hyperthymic and irritable temperaments contributed to the association between AUD and BD. On the other hand, in AUD or other substance use disorders, cyclothymic, depressive, and irritable temperaments were implicated. The current study aimed to examine the relationships between clinical phenotypes of acute depressive episodes and co-occurring AUD, using depressive or manic symptom rating scales and measure of affective temperaments. To encompass diverse clinical features of acute depressive episodes, subjects with a major depressive episode regardless of subtypes of mood disorder were included. Childhood adversities are one of the most significant risk factors for AUD as well, and are thought to be a constitutional basis and a possible covariate of both addiction and mood symptoms. Therefore, to clarify clinical correlates of AUD in mood disorders, a measure of childhood trauma should be included as a potential confounder. We hypothesized that AUD would be more frequent in BD than in MDD and that phenotypes related to BD would predict AUD in acutely depressed subjects.

METHODS

Subjects
The sample consisted of 89 subjects with BD and 48 subjects with MDD. Subjects were recruited from the outpatient clinic or inpatient unit of the Mood Disorders Clinic at Seoul National University Bundang Hospital and met the DSM-IV-TR criteria for BD (bipolar I disorder, bipolar II disorder, and bipolar disorder NOS) or MDD, using the Mini-International Neuropsychiatric Interview (MINI). Inclusion criteria were an absence of any lifetime history of significant neurological or medical illnesses, age between 19 and 80, the fulfillment of criteria for a major depressive episode and the score of the Hamilton Depression Rating Scale (HAMD) 8 or higher. This study was approved by the Institutional Review Board (B-1902/524-103).

Clinical assessment
The obtained clinical variables included basic demographic information such as gender, age, and duration of illness. The severity of depressive or manic symptoms was measured using the HAMD and the Young Mania Rating Scale (YMRS). The severity of suicide risk was measured using the Scale for Suicide Ideation (SSI). Experiences of childhood trauma were measured using the Childhood Trauma Questionnaire Korean version (CTQ). Diagnosis of AUD was made when DSM-IV-TR criteria for alcohol dependence and/or abuse were met within the last 12 months from the assessment, to ensure the current comorbidity. All subjects with AUD were having alcohol misuse at the time of evaluation.

Affective temperaments were measured using the short version of the Temperament Evaluation of Memphis, Pisa, Paris and San Diego-autoquestionnaire (TEMPS-A) that has 39 items assessing cyclothymic, depressive, irritable, hyperthymic and anxious temperaments. The questionnaire was translated by the authors (TH Ha & S Oh).

Statistical analysis
Clinical variables between the groups with AUD and without AUD were compared using the Student’s t-test and Chi-square test. Mixed depressive state was defined as 3 or more YMRS items were 2 or above. To find out significant predictors and to adjust possible covariates, we tested two bivariate logistic regression models. Model 1 was intended to assess the risk of diagnosis, the severity of mood symptoms and affective temperaments. Childhood trauma and suicidal ideation were added as covariates because these factors are known to relate both to mood disorders and addictive disorders. Model 2 was aimed at examining the risk of dimensions of depressive symptoms and manic symptoms. Each of the dimensions was quoted from previous factor analytic studies for HAMD and YMRS. Somatic anxiety, psychic anxiety, core depression, and anorexia were dimensions of depressive symptoms from the HAMD, and irritable mania, elated mania, and psychotic mania were dimensions of manic symptoms from the YMRS. Gender, age, and diagnosis of mood disorder were adjusted to complete these models. For all the statistical procedures, IBM SPSS Statistics ver. 25 (IBM Corp. Armonk, NY, USA) was used and significance was set as p<0.05.
RESULTS

Among 137 subjects, 29 (21.2%) had co-occurring AUD. The proportion of subjects having AUD was higher in the BD group (27.0%) than in the MDD group (10.4%) ($\chi^2=5.118$, $p=0.024$). Proportions of the BD subtypes were not different between the AUD and non-AUD groups. The AUD group had a higher YMRS mean score ($t=-3.567$, $p=0.001$), compared to the subjects without AUD. The rate of mixed depressive state, defined as 3 or more YMRS items above 1, was found to be 33.6% (12.5% of MDD and 44.9% of BD) and was higher in the AUD group ($\chi^2=7.693$, $p=0.006$). The mean scores for irritable ($t=-2.641$, $p=0.010$) and hyperthymic temperaments ($t=-2.624$, $p=0.010$) were higher, and the mean score for anxious temperament was lower ($t=2.164$, $p=0.032$) in the AUD group than in the non-AUD group. The AUD group showed more suicidal ideation ($t=-2.252$, $p=0.026$) and reported more childhood adversities ($t=-4.027$, $p<0.001$) than the non-AUD group. Gender, age, and the severity of depressive symptoms did not differ between the two groups (Table 1).

When potential confounders were adjusted, only the anxious temperament was significant in the regression model 1 (OR=0.373, 95% CI: 0.206-0.675, $p=0.001$). Among confounding variables, age and childhood trauma were significant in this model (Table 2).

In regression model 2 to test the relationships between specific mood symptom dimensions and AUD, the irritable mania dimension significantly predicted AUD (OR=1.412, 95% CI: 1.162-1.717, $p=0.001$). None of the dimensions of depressive symptoms were significant in this model. Among confounders, the diagnosis of BD was significant (OR=3.677, 95% CI: 1.031–13.119, $p=0.045$) (Table 3).

DISCUSSION

To our knowledge, the current study is the first to explore specific mood symptoms and affective temperaments in depressed subjects comorbid with AUD. The main findings of this study were that a diagnosis of BD, specific mood symptoms and affective temperaments were risk factors for co-occurring AUD in depressed subjects. Specifically, in subjects with depression, irritable manic symptoms increased the risk of AUD while the anxious temperament decreased it.

27% of BD and 10% of MDD subjects in this study had a co-occurring AUD. Compared to previous studies, those rates were lower, which may be related to the characteristics of our sample. Subjects in our study were likely to be in their mid-thirties and approximately 10 years younger than previous studies. AUD may be a complication during the course of mood disorder, given the cumulative elevation of the comor-bidity rates. In line with this assumption, our regression model revealed that age increased the risk of AUD. Male gender was usually considered as a risk factor for AUD in mood disorders, which was not replicated in our study. Female predominance (74%) in our sample may explain this inconsistency.

The irritable and hyperthymic temperaments were higher in the AUD group than in the non-AUD group. However, when adjusting diagnosis and other confounding variables, these did not predict AUD. The predominance of BD subjects in the AUD group (83%) may explain the group differences. Only one study by Singh et al. explored affective temperaments in comorbid AUD and BD. In their study, irritable and hyperthymic temperaments independently predicted AUD in subjects with BD. Possible explanations for this inconsistency include characteristics of the sample and study design. Sub-

| Table 1. Demographic and clinical characteristics of the subjects |
|---------------------------------------------------------------|
| Without AUD (N=108)                                           |
| With AUD (N=29)                                               |
| Statistics p                                                 |
| Demographic characteristics                                   |
| Female (%)                                                    |
| 81 (75.0)                                                     |
| 21 (72.4)                                                     |
| $\chi^2=0.080$                                               |
| 0.777                                                        |
| Age                                                           |
| 35.9±12.4                                                     |
| 34.2±11.0                                                    |
| t=0.682                                                      |
| 0.497                                                        |
| Duration of illness                                           |
| 9.0±9.1                                                      |
| 9.9±10.0                                                     |
| t=0.479                                                      |
| 0.633                                                        |
| Diagnosis of BD (%)                                           |
| 65 (60.2)                                                     |
| 24 (82.8)                                                     |
| $\chi^2=5.118$                                               |
| 0.024                                                        |
| Subtypes of BD (%)                                            |
| BD I                                                         |
| 5 (4.6)                                                      |
| 4 (13.8)                                                     |
| $\chi^2=1.633$                                               |
| 0.442                                                        |
| BD II                                                        |
| 43 (39.8)                                                    |
| 15 (51.7)                                                    |
| BD NOS                                                       |
| 17 (15.7)                                                    |
| 5 (17.2)                                                     |
| Mixed depressive state                                       |
| 30 (27.8)                                                    |
| 16 (55.2)                                                    |
| $\chi^2=7.693$                                               |
| 0.006                                                        |
| Severity of mood symptoms                                    |
| HAMD                                                          |
| 17.7±6.4                                                     |
| 19.7±7.1                                                     |
| t=1.427                                                      |
| 0.156                                                        |
| YMRS                                                          |
| 5.6±4.5                                                      |
| 9.1±5.6                                                      |
| t=3.567                                                      |
| 0.001                                                        |
| Affective temperament                                         |
| Cyclothymic                                                  |
| 6.4±3.5                                                      |
| 7.7±3.2                                                      |
| t=1.773                                                      |
| 0.079                                                        |
| Depressive                                                   |
| 3.7±2.5                                                      |
| 4.0±2.4                                                      |
| t=-0.751                                                     |
| 0.454                                                        |
| Irritable                                                    |
| 2.4±2.3                                                      |
| 3.7±2.4                                                      |
| t=-2.641                                                     |
| 0.009                                                        |
| Hyperthymic                                                  |
| 2.7±2.3                                                      |
| 4.0±2.7                                                      |
| t=-2.624                                                     |
| 0.010                                                        |
| Anxious                                                      |
| 1.7±1.1                                                      |
| 1.1±1.2                                                      |
| t=2.164                                                      |
| 0.032                                                        |
| Suicidal ideation, childhood trauma                           |
| SSI                                                          |
| 12.8±9.1                                                     |
| 17.0±8.5                                                     |
| t=-2.252                                                     |
| 0.026                                                        |
| CTQ                                                           |
| 59.3±18.6                                                    |
| 75.7±22.4                                                    |
| t=-4.027                                                     |
| 0.000                                                        |

AUD: alcohol use disorder; BD: bipolar disorder; BD I: bipolar I disorder, BD II: bipolar II disorder, BD NOS: bipolar disorder NOS, HAMD: Hamilton Depression Rating Scale, YMRS: Young Mania Rating Scale, SSI: Scale for Suicide Ideation, CTQ: Childhood Trauma Questionnaire.
jects in our study were in a depressive episode, while mood states of subjects in the study by Singh et al.\textsuperscript{29} were not uniform. The severity of depressive symptoms in their study was much lower than that in our study. Reports by subjects may be influenced by their mood states. In depressed subjects, subjective reports of ‘up’ signs may be attenuated and those of ‘down’ signs may be exaggerated. The trends of differences in raw scores are quite similar between the two results. However, the scores of depressive temperament were 4.0 vs. 3.7 in our sample and were 2.7 vs. 2.2 in Singh et al.\textsuperscript{29} Another explanation may be related to statistical power. The sample size in our study was much smaller than that in Singh et al.\textsuperscript{29} study.

The protective effect of anxious temperament was a unique and unexpected finding. There are few studies on the role of anxious temperament in AUD in mood disorders. In previous studies on substance use disorders, anxious temperament was a risk factor,\textsuperscript{30,48} which ran contrary to our findings. However, we can assume that the functional roles of anxious temperament may be distinct from different conditions. Anxious temperament correlates positively with harm avoidance and negatively with novelty seeking,\textsuperscript{41} which is exactly opposite to hyperthymic temperament. High novelty seeking is related to increased alcohol use while high harm avoidance with abstinence.\textsuperscript{49} Therefore, it seems logical to say that anxious temperament is related to low drinking and hyperthymic temperament is related to high drinking, according to Cloninger’s temperament dimensions. In this context, we may hypothesize that, in a depressed state, hyperthymic temperament may

| Table 2. Regression model 1 including sex, age, severity of mood symptoms, affective temperaments, childhood trauma, and suicidal ideation as covariates |
|---------------------------------|-----------------|-----------------|-----------------|
|                                | OR              | 95% CI of OR    | p               | SE              |
|                                | Lower limit     | Upper limit     |                 |                 |
| Female                         | 0.415           | 0.119           | 1.446           | 0.167           |
| Age                            | 1.080           | 1.010           | 1.154           | 0.023           |
| Diagnosis of BD               | 1.521           | 0.390           | 5.926           | 0.546           |
| HAMD                           | 1.040           | 0.946           | 1.143           | 0.415           |
| YMRS                           | 1.110           | 0.987           | 1.248           | 0.081           |
| Cyclothymic temperament        | 1.092           | 0.886           | 1.346           | 0.411           |
| Depressive temperament         | 0.946           | 0.711           | 1.258           | 0.704           |
| Irritable temperament          | 1.275           | 0.914           | 1.779           | 0.152           |
| Hyperthymic temperament        | 1.236           | 0.974           | 1.569           | 0.082           |
| Anxious temperament            | 0.373           | 0.206           | 0.675           | 0.001           |
| CTQ                            | 1.048           | 1.017           | 1.080           | 0.002           |
| SSI                            | 1.062           | 0.990           | 1.138           | 0.095           |

BD: bipolar disorder, HAMD: Hamilton Depression Rating Scale, YMRS: Young Mania Rating Scale, SSI: Scale for Suicide Ideation, CTQ: Childhood Trauma Questionnaire

| Table 3. Regression model 2 including sex, age, diagnosis, and dimensions of mood symptoms as covariates |
|---------------------------------|-----------------|-----------------|-----------------|
|                                | OR              | 95% CI of OR    | p               | SE              |
|                                | Lower limit     | Upper limit     |                 |                 |
| Female                         | 0.629           | 0.220           | 1.798           | 0.387           |
| Age                            | 0.986           | 0.943           | 1.030           | 0.518           |
| Diagnosis of BD               | 3.677           | 1.031           | 13.119          | 0.045           |
| Somatic anxiety of HAMD        | 0.948           | 0.774           | 1.161           | 0.607           |
| Psychic anxiety of HAMD        | 1.155           | 0.882           | 1.514           | 0.295           |
| Core depression of HAMD        | 1.032           | 0.814           | 1.308           | 0.798           |
| Anorexia of HAMD               | 1.318           | 0.817           | 2.126           | 0.257           |
| Irritable mania of YMRS        | 1.412           | 1.162           | 1.717           | 0.001           |
| Elated mania of YMRS           | 0.960           | 0.667           | 1.381           | 0.825           |
| Psychotic Mania of YMRS        | 0.759           | 0.467           | 1.234           | 0.266           |

BD: bipolar disorder, HAMD: Hamilton Depression Rating Scale, YMRS: Young Mania Rating Scale
play a less significant role and anxious temperament play a more important role than in euthymic or euphoric states. The significant effect of anxious temperament was independent of childhood trauma that was another risk for AUD. Childhood trauma may cause a range of mental disorders including depression, alcohol misuse and suicidal attempts. Our results add further evidence that childhood adversities may complicate the course of mood disorder.

Regarding mood symptoms, none of the depressive symptom dimensions were found to be associated with AUD. Combined with findings of previous studies, it seems that there are little connections between specific depressive symptoms and AUD in mood disorder. The severity of depression was not associated with AUD as well. On the other hand, mixed depressive state was more frequent in the AUD group and the severity of manic symptoms and irritable manic symptoms were associated with AUD. In particular, the irritable manic symptom dimension predicted AUD independently of BD diagnosis. These findings may imply that AUD in mood disorders may relate to manic components but not to depressive components. Evidence from previous studies supports our findings, such as more frequent addictive disorders in the mixed episodes rather than the pure depressive episodes, drinkings resulting from manic severity, and increased predisposition to impulsivity in BD comorbid with substance use.

Whilst the current study has strengths over previous studies in that we have focused on depressive state and assessed systematic confounders, several limitations should be noted. Firstly, the sample size was not large enough to ensure subgroup analyses and to show sufficient statistical power. In future studies, subtypes of BD, specifiers of depressive episodes or courses need to be considered. Secondly, our analyses were limited to co-occurring AUD to magnify the phenomenon during a depressive episode. Lifetime comorbidity, temporal sequence of the onset, and longitudinal courses should be further explored. Thirdly, data were collected from a single-center, which limits the generalizability of the findings. Characteristics of subjects such as younger age and female predominance make a direct comparison with findings from previous studies difficult. Other limitations include the reliability of TEMPS-A. The authors did not validate the translated version. However, authors who are experienced in this field confirmed the face validity and empirically found the usefulness of the measure. Despite these limitations, our findings offer an important insight into understanding AUD found in depressed patients.

In conclusion, co-occurring AUD during depression was associated with a diagnosis of BD, the severity of manic symptoms and irritability symptoms. Among affective temperaments, anxious temperament played a protective role. Our findings suggest that, even during depression, alcohol use may be an expression of manic symptom dimension.

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Conflicts of Interest
The authors have no potential conflicts of interest to disclose.

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Conceptualization: Minseok Hong, Tae Hyon Ha. Data curation: Minseok Hong, Suyeon Lee, Sunghee Oh. Formal analysis: Minseok Hong. Funding acquisition: Tae Hyon Ha. Investigation: Minseok Hong, Suyeon Lee, Woojae Myung. Methodology: Minseok Hong, Tae Hyon Ha. Project administration: Tae Hyon Ha, Suyeon Lee, Sunghee Oh. Resources: Tae Hyon Ha, Woojae Myung. Software: Minseok Hong. Supervision: Tae Hyon Ha. Validation: Tae Hyon Ha. Visualization & Writing—original draft: Minseok Hong. Writing—review & editing: Tae Hyon Ha.

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