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# Co-morbid cannabis use disorder and chronotype are associated with mood symptom onset in people with bipolar disorder

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# Abstract

Comorbid cannabis use disorder (CUD) is disproportionately high in people with bipolar disorder (BD) and has been associated with worsening of BD symptoms. However, many people with BD report regularly using cannabis to ameliorate symptoms, including sleep disturbances. Sleep and circadian rhythm disturbances are hallmark features of BD that often precede the onset of mood symptoms. Genetic studies indicate that circadian disruption may predispose individuals towards both problematic cannabis use and BD, rather than cannabis use directly impacting BD symptoms. To further disentangle these hypotheses, we aimed to investigate the relationship between chronotype, cannabis use disorder (CUD) and BD mood symptoms. Data from 212 participants with BD I from the Pharmacogenomics of Bipolar Disorder study dataset were analyzed for this study. Participants were stratified by those diagnosed with co-morbid CUD and BD symptom variables, including the mean number of mood episodes per year and age of mood symptom onset for both depression and mania symptoms. The Basic Language Morningness scale (BALM) was used to assess chronotype. There was no interaction between morningness levels and CUD on BD symptoms, however both lower morningness and CUD were independently associated with earlier age of mood symptom onset. However, patients who reported initiating cannabis use post mood symptom onset had an earlier mood symptom age of onset compared to those who reported initiating cannabis use prior to mood symptom onset. These findings could provide further evidence that circadian rhythm disruption could be an underlying factor that predisposes individuals toward both CUD and BD.

Declaration of competing interest The authors report no conflicts of interest.

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Cannabis abuse; Bipolar disorder; Mania; Depression; Chronotype

# 1. Introduction

Bipolar disorder (BD) is a neuropsychiatric disorder defined by recurrent mood shifts between depressive and manic/hypomanic episodes that contribute to poor psychosocial and health outcomes. In addition to mood episodes, people with BD are at a higher risk for co-morbid health outcomes which may further worsen their quality of life. Substance use disorders (SUD) are amongst the most prominent co-occurring conditions that present in people with BD. In fact, compared to other serious mental illnesses, BD has amongst the highest comorbidity of SUD(Gold et al., 2018). The high prevalence of co-morbid BD and SUD could indicate an underlying cause that contributes to an increased susceptibility towards developing both conditions. Circadian rhythm disturbances may increase susceptibility towards a co-morbid substance use disorder. Circadian rhythm disruptions are a hallmark feature of BD (Gold and Sylvia, 2016; Harvey et al., 2009) that have also been implicated as a contributing factor towards substance abuse(Hasler et al., 2014; Tamura et al., 2021). Notably, circadian rhythms disruptions in people with BD can worsen sleep and mood symptoms and adversely affect clinical course and treatment(Takaesu et al., 2018), including substance use outcomes(Serrano-Serrano et al., 2021). In particular, evening chronotype (i.e., a tendency to sleep and wake later and proxy for circadian phase delay) has been associated with a higher risk of BD as well as substance use and relapse (Berdynaj et al., 2016; Menculini et al., 2023; Santos et al., 2023; Serrano-Serrano et al., 2021).

Amongst other forms of substance use, cannabis use is one of the most prevalent in people with BD, with up to 70% reporting a lifetime history of regular cannabis use and up to 20% with a comorbid cannabis use disorder (CUD) diagnosis(Agrawal et al., 2011). Cannabis is also commonly used for its reputed sleep-promoting properties(Babson et al., 2017; Monti and Pandi-Perumal, 2022). In fact, 70% of people with BD reported using cannabis specifically to address sleep disturbances in one study(Miranda et al., 2023). However, the effects of cannabis on circadian regulation of sleep remain unclear; thus, there remain concerns that cannabis may interact with the compromised circadian system in BD and further exacerbate symptoms. Notably, cannabis use has been associated with a significantly younger age of schizophrenia onset (Veen et al., 2004) and age of onset at first psychosis treatment (Barrigón et al., 2010). Cannabis use has been associated with worse mania and psychosis symptoms(Sideli et al., 2019), an earlier age of BD onset (Lagerberg et al., 2011), and overall worse BD illness progression(Pinto et al., 2019). As such, cannabis use has been identified as a potential risk factor for the development of BD and the worsening of BD symptoms.

As an alternative hypothesis to cannabis use being a risk factor for BD or worsening of BD symptoms, genetic studies suggest that CUD and BD may instead have a shared underlying mechanism that predisposes individuals towards the development of both conditions. A

genome-wide association study on cannabis use and BD has identified shared genetic associations between CUD and BD(Pasman et al., 2018). Significant genetic associations have also been identified between chronotype and psychiatric traits, including depression symptoms(Jones et al., 2019) and lifetime cannabis use(Winiger et al., 2021), suggesting that circadian rhythm disturbances may indeed be related to both conditions.

As summarized, current literature supports two competing models of BD and cannabis use: 1) cannabis use directly contributes to the worsening of BD symptoms; and 2) a shared underlying factor predisposes individuals towards both BD and CUD. As described above, a compromised circadian system may be a predisposing factor for the latter model of BD and cannabis use, thus in this study, we aimed to further investigate the latter model 1) determine whether CUD was associated with BD symptom onset, 2) determine whether cannabis use initiation relative to BD symptom onset impacts BD symptom onset, and 3) determine whether chronotype, as a proxy for circadian rhythm disturbance, was associated with BD symptom onset, and whether co-morbid CUD moderated this relationship. Here, we used data collected from patients with BD type I, who participated in the Pharmacogenomics of Bipolar Disorder (PGBD) study that either had a diagnosed co-morbid CUD or did not. We hypothesized the following: 1) CUD would be associated with an earlier age of mood symptom onset relative to those with no CUD, 2) greater eveningness would be associated with earlier BD symptom onset and CUD would not moderate this relationship, and 3) initiation of cannabis use prior to mood symptom onset would not be associated with earlier age of BD symptom onset. If supported these hypotheses would help disentangle the two competing models and provide further support that CUD and BD share an underlying mechanism.

# 2. Methods

# 2.1. Participants

Participant data were derived from the PGBD study dataset. The Diagnostic Interview for Genetic Studies (DIGS) (Oedegaard et al., 2016) was used to assess psychiatric and substance use disorders. BD diagnosis criteria were defined using the Diagnostic and Statistical Manual of Mental Disorders IV. Only participants with BD I (n = 212) were analyzed for this study. The DIGS and other medical records were used to give as accurate descriptions of psychiatric symptoms and substance use as possible.

## 2.2. Bipolar disorder symptom variables

The Final Best Estimate method(Nurnberger et al., 1994) was used to determine mood symptom age of onset (AAO), lifetime number of mood episodes and chronicity of cannabis use. Depression and mania episode density (ED) was calculated for each participant, which measures the number of mood episodes in a given amount of time, a method described by Strejilevich et al.(Strejilevich et al., 2019, 2024). ED can be used as a proxy for lifetime burden of mood episodes, where higher ED corresponds to greater lifetime burden of mood episodes. Here, we use ED to account for differences in participant age. All BD symptom variables are listed and defined in Table 1.

# 2.3. Cannabis use disorder

Participants were stratified into two groups: those with diagnosed cannabis abuse or dependence (CUD+; n = 53) and those with no cannabis abuse or dependence (CUD-; n = 159). For CUD patients with cannabis use age of onset data available, cannabis use onset relative to mania or depression symptom onset was dichotomized as those who initiated cannabis use prior to or within the same year of mood symptom onset (Pre-onset) and those who initiated cannabis use after mood symptom onset (Post-onset). Pre- and post-onset groups were calculated relative to mania and depression onset separately.

#### 2.4. Chronotype

Chronotype was measured in units of morningness. Participants were surveyed for chronotype using the Basic Language Morningness Scale (BALM)(Brown, 1993; Smith et al., 1989). The BALM is a 13-item composite self-report scale used to measure circadian rhythms. Summary scores consisted of the sum of all 13 items, with higher summary scores indicating a higher level of morningness.

#### 2.5. Statistical analysis

Assumptions for equal variances (Levene's or Box's test of equality) and normality (Shapiro-Wilks test) were tested for all variables; variance and normality were tested across the entire sample and within each group. Non-parametric testing was used throughout to account for non-normal distributions, unequal variances and unequal sample sizes. Correlations between BALM score and BD mood symptoms were tested using Spearman's correlations. Differences (i.e., age, mood symptoms and BALM score) between CUD and CUD-groups were analyzed using Mann-Whitney U tests. Chi-square testing was used to determine group differences for nominal variables (i.e., sex, psychosis symptoms and suicidality). Group differences between no CUD, cannabis use pre-symptom onset and cannabis use post-symptom onset was first analyzed using Kruskal-Wallis tests, followed by pairwise Mann-Whitney U tests. The a level was set at 0.05 and Bonferroni multiple testing correction threshold was used (p < 0.0167). A moderated linear regression was performed using the SPSS package PROCESS (Hayes, 2022) to test whether CUD moderates the relationship between chronotype and BD symptoms. Due to the non-normal distribution of BD symptom variables, log transformation was used to achieve normality. As a sensitivity analysis, regression analyses were conducted with both non-transformed and transformed data; results were consistent thus the findings are presented using nontransformed variables BALM summary score was used as the independent variable, BD symptoms as the dependent variables, CUD as a moderating variable<sup>28</sup>. All statistical analyses were performed using SPSS 28.0 (Chicago, IL, USA).

# 3. Results

# 3.1. BD with comorbid CUD was associated with earlier onset of mood symptoms and affective psychosis

Patients with CUD were significantly younger than patients without CUD. There was a smaller proportion of females in the CUD group compared to the CUD-group (Table 2),

however there were no significant group differences between males and females on any of the BD symptom variables or BALM score. A greater proportion of CUD patients reported having psychotic symptoms in at least 2 episodes compared to patients with no CUD ( $X^2 = 8.86$ , p < 0.05; Table 2). Significantly greater percentage of CUD-patients reported never having suicidal thoughts compared to patients with CUD. ( $X^2 = 4.75$ , p < 0.05, Table 2). Consistent with previous literature, CUD + patients had significantly lower morningness as measured by BALM compared to BD patients without CUD (U = 3298, p < 0.05, Table 2). There were no significant differences in depression ED between CUD- and CUD groups. However, mania ED was significantly higher in CUD patients (U = 3236, p < 0.05, Table 3) and mania AOO was significantly earlier in CUD patients (U = 2580, p < 0.001, Table 3), compared to CUD-patients. Depression AOO was earlier in CUD patients compared to CUD-patients but the difference did not reach statistical significance (U = 3433, p = 0.086, Table 3).

#### 3.2. Chronotype was associated with mood symptom age of onset

We then tested whether CUD moderates the relationship between chronotype and BD symptom onset/ED. BALM scores were significantly associated with mania AAO and depression AAO. Greater morningness (higher BALM scores) was associated with a later AAO for mania (b = 0.23, t = 2.48, p = 0.01, Fig. 1a) and depression (b = 0.32, t = 3.19, p < 0.01, Fig. 1b). Despite the significant difference between mood symptom AAO between CUD and CUD-patients, CUD did not significantly moderate the relationship between chronotype and mood symptom AAO (Table 4). Cannabis use AAO relative to mood symptom AAO did not significantly moderate the relationship between chronotype and mood symptom AAO.

# 3.3. Cannabis use onset prior to mood symptom onset was significantly associated with later mood symptom age of onset

There was a significant difference in mania AAO between no CUD, cannabis use pre (n = 39) and post symptom onset (n = 14) groups (H = 24.73, p < 0.001, Fig. 2a). Mania AAO was significantly later in CUD-patients, compared to those who initiated cannabis use both pre (U = 2268.5, p = 0.01). and post-mania onset (U = 311.5, p < 0.001). People who initiated cannabis use post-symptom onset also had significantly earlier mania AAO compared to those who initiated cannabis use pre-symptom onset (U = 105, p < 0.001). Depression AAO was significantly different between cannabis use onset groups (H = 16.92, p < 0.001, Fig. 2b). Depression AAO was later in patients with no CUD (U = 988, p < 0.001) and in patients that reported initiating cannabis use pre-depression onset (n = 28; U = 121, p < 0.001) compared to patients who initiated cannabis use post-depression onset (n = 24). Initiation of cannabis use relative to symptom onset was not associated with depression or mania ED.

# 4. Discussion

Our data support previous reports that CUD in BD is associated with an earlier onset, greater severity of symptoms and worse clinical features. Specifically, we found that people with BD and comorbid CUD had an earlier AAO for mania and depression, greater number of

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mania episodes per year and more often reported presence of affective psychosis. Our data indicate that while both chronotype and CUD were independently associated with mood symptom AAO, there was no interactive effect of CUD and chronotype on mood symptom onset. Importantly, we found that cannabis use onset prior to mood symptom onset was not associated with earlier emergence of mood symptoms; rather, mood symptom AAO was *later* in BD patients who reported initiating cannabis use prior to mood symptom onset, compared to BD patients who reporting initiating cannabis use after mood symptom onset. These findings help to clarify the temporal relationship between cannabis use and mood symptom onset, a critical step towards elucidating the causal relationship between these two psychiatric conditions.

Previous studies have similarly investigated the temporal relationship between substance dependence and BD. For example, Martínez-Ortega et al. assessed nicotine dependence onset relative to BD onset in a large cohort. The authors reported significant differences in lifetime BD symptoms and comorbid conditions between patients that reported nicotine dependence prior to BD onset (n = 135) compared to patients who reported nicotine dependence following BD onset (n = 386) (Martínez-Ortega et al., 2013). The authors also analyzed nicotine dependence relative to major depression onset using similar methods (Martínez-Ortega et al., 2017). These studies have added meaningful knowledge on the temporal relationship between mood disorders and substance use thus, we employed a similar approach by comparing cannabis use onset relative to both mania and depression onset in patients with BD. Our findings are consistent with previous literature in which CUD was associated with an earlier age of mood symptom onset and psychosis in BD(De Hert et al., 2011; Kuhns et al., 2021). For example, BD/CUD patients have been reported to have the lowest age of BD onset (mean age of cannabis use onset = 18.6 years) compared to BD patients who did not use cannabis and patients who engaged in cannabis use but did not meet diagnostic criteria for CUD(Lagerberg et al., 2014). Excessive cannabis use has also been found to predict first incidence mood episodes in BD after controlling for a number of potentially confounding factors (i.e., sociodemographic factors, childhood trauma, family psychiatric history, other substance use disorders and other mental disorders) (Cougle et al., 2015; Van Laar et al., 2007). Initiation of cannabis use pre- or post-symptom onset were both associated with an earlier age of mood symptom onset(Lagerberg et al., 2011). Our results are in part consistent with this report, such that initiation of cannabis use both pre- and post-onset of mania symptoms was associated with an early age of BD onset compared to BD patients without CUD. Our data further expand on these previous reports and demonstrate that initiation of cannabis use pre-mania onset is associated with a significantly later mania AAO compared to those initiating cannabis use post-mania onset. Further, depression AAO was statistically similar between the CUD-patients and the patients that reported initiating cannabis use pre-depression onset. These findings further strengthen the hypothesis that while CUD is associated with earlier age of onset of BD symptoms, initiation of cannabis use is not directly a major driver of early BD onset.

Our data on cannabis use onset relative to mood symptom onset indicate that other factors may contribute to BD symptom onset and severity and also predispose people towards initiation of problematic cannabis use and development of CUD. As described previously, circadian rhythm disruption could be one such factor that impacts both BD symptom onset

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and problematic cannabis use. A recent review indicated that individuals with circadian rhythm disturbances have a significantly elevated risk for BD(Scott et al., 2022). Another study supports these findings by demonstrating that higher eveningness in adolescents and young adults significantly predicted next-year depressive symptoms(Sasser et al., 2023). A longitudinal analysis of substance use in young adults and adolescents found that greater eveningness significantly predicted next-year cannabis use(Hasler et al., 2022). Lastly, mendelian randomization analyses have provided evidence that lower genetic loading for morningness significantly increases the liability of depression symptoms(Jones et al., 2019), whereas increased insomnia increases the risk of cannabis use initiation(Winiger et al., 2021). Here, we demonstrated a significant difference in chronotype between BD patients with and without co-morbid CUD. While greater morningness was associated with later mood symptom AAO, CUD did not moderate the relationship between chronotype and mood symptom AAO. The lack of interaction between CUD and chronotype on BD symptom onset indicates that CUD does not directly impact BD symptoms. These data, combined with existing literature supports the model that CUD and BD are distinct conditions similarly impacted by circadian rhythm disruption.

The results presented here add new evidence on the interrelationship between CUD, BD and circadian rhythms, however this study has several limitations. Although, participant data were clearly defined and comprehensively validated through clinical interviews and reviews, our analyses did not include current depression and mania symptom data. Importantly, we have previously reported that chronotype also predicts current depressive and mania symptoms in this cohort (McCarthy et al., 2018); thus, future analyses may investigate potential interactions between CUD and chronotype on current mood symptoms. It is also critical to note that there was no limiting eligibility criteria for mood state for this study, as such while many participants were euthymic, some presented with depression and mania/hypomania that could have impacted their memory. This recall bias may affect the participants' ability to accurately recall number of mood episodes, which may have led to an overestimation or underestimation of significance in the relationships among episode density, chronotype and CUD. Additionally, our dataset only included information on current cannabis abuse or dependence diagnoses but did not include whether participants engage in regular cannabis use that does not reach the threshold of a cannabis abuse or dependence diagnosis. This distinction is important to make for future studies considering many people with BD report regular cannabis for symptom management. In other populations, regular cannabis use has been shown to reduce symptom severity, in a dose-dependent manner, such that lighter use improves symptoms whereas heavy cannabis use worsens symptoms (Ellingson et al., 2021; Kallianpur et al., 2020; Thames et al., 2016). Thus, future studies evaluating the effects of cannabis use frequency on mood symptoms could yield important insight into manageable cannabis use practices in a population where cannabis use is highly prevalent. Multiple testing, unequal sample sizes and smaller sample sizes may have affected the statistical power of our analyses, particularly in the cannabis use relative to mania onset groups. Additionally, the significant association between earlier age of mania onset in participants and initiating cannabis use before mania onset could also indicate that some people with BD initiate cannabis use to self-medicate prodromal symptoms before full illness onset. As described previously, there is strong supporting

evidence of a shared underlying predisposition for BD and CUD. However, future studies monitoring the emergence of cannabis use and subclinical symptoms in adolescents and young adults at-risk for developing BD could further disentangle these two models of BD and cannabis use.

Here, we propose that circadian rhythm disruption contributes to both CUD and BD. Future studies could evaluate whether chronotype could be used as a biomarker to predict mood symptom onset in young adults at risk for developing BD or identify patient groups at greater risk for developing CUD. Our study is consistent with others indicating that circadian rhythm disruption could be a promising target for identifying CUD and BD risk. More research should be conducted to better understand the underlying neurobiology of this relationship, and whether circadian rhythms can be leveraged to improve BD and CUD outcomes.

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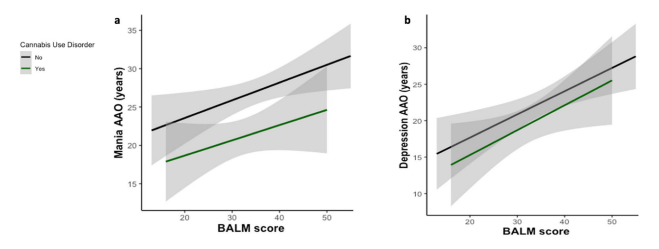
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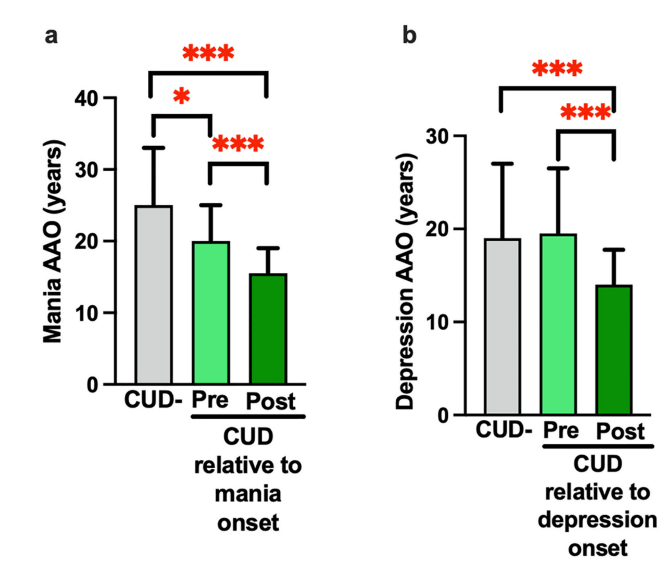
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# Fig. 1.

Simple slopes for the association between Basic Language Morningness Scale (BALM) score and mood symptom age of onset (AAO) in patients with and without a diagnosed cannabis use disorder (CUD). a) BALM score was significantly associated with mania AAO (b = 0.23, t = 2.48, p = 0.01), though CUD did not significantly moderate this relationship. b) BALM score was significantly associated with depression AAO (b = 0.32, t = 3.19, p = 0.002), though CUD did not significantly moderate this association. Shaded areas indicate 95% confidence intervals.

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# Fig. 2.

Cannabis use onset pre-BD symptom age of onset (AAO) is associated with a later mood symptom AAO relative to cannabis use post-BD symptom AAO. a) Mania AAO was significantly earlier in the post-mania onset group (n = 14) compared to both the pre-onset group (n = 39); U = 105, p < 0.001) and CUD-group (U = 311.5, p < 0.001). Mania AAO was also significantly later in CUD-compared to pre-onset participants (U, 2268.5, p = 0.012). b) Post depression onset cannabis use (n = 24) was associated with significantly earlier depression AAO compared to CUD- (U = 988, p < 0.001) and pre-depression onset cannabis use (n = 28; U = 121, p < 0.001). Data represents median and interquartile range; \*p < 0.05, \*\*\*p < 0.001.

BD symptom variables and definitions.

BD symptom variables	Definition
Mania age of onset (mania AOO)	Age at first diagnosed mania episode
Depression age of onset (depression AOO)	Age at first diagnosed depression episode
Mania episode density	Lifetime total number of mania episodes Age - mania age of onset
Depression episode density	Lifetime total number of depression episodes Age – depression age of onset
Affective Psychosis	Presence of psychosis during mood episodes:
	No (0-1 episodes of psychosis)
	Yes (2+ episodes with psychosis)
Suicidality	Presence of suicidality:
	Never/none
	Thoughts of suicide
	Suicide attempts

Demographic and clinical data on patients with bipolar disorder (BD), with and without comorbid cannabis use disorder (CUD). Data reported are mean (standard deviation) and percent of group.

	CUD-n = 159	<b>CUD n = 53</b>	Group differences
Age	45.6(14.3)	37.5(12.5)	CUD->CUD
Sex F/M	59%/41%	38%/62%	p < 0.05
Affective psychosis			
0–1 episodes	64%	34%	CUD- > CUD
2+ episodes	36%	66%	CUD- < CUD
Suicidality			
Never	30%	16%	CUD- > CUD
Thoughts	35%	42%	ns
Attempted	35%	42%	ns
Basic Language Morningness Scale (BALM) score	36 (11)	32 (12)	CUD->CUD

Mood symptom age of onset and episode density in patients with bipolar disorder (BD), with and without comorbid cannabis use disorder (CUD). Data reported are median (Interquartile range).

	CUD-n = 159	CUD n = 53	U	р	Group differences
Mania episode density	0.32 (0.12–0.59)	0.36 (0.19–0.98)	3236	0.021	CUD- < CUD
Depression episode density	0.33 (0.17-0.79)	0.39 (0.17–0.99)	3711	0.47	ns
Mania age of onset	25 (19–33)	19 (16–22)	2580	< 0.001	CUD->CUD
Depression age of onset	19 (15–27)	18 (13–22)	3433	0.086	ns

Summaries for linear regression models of mania and depression age of onset (AAO).

Mania AAO				
	b	SE	t	р
BALM score	0.32	0.10	3.13	0.002
CUD status	-2.84	7.15	-0.40	0.69
BALM score x CUD status	0.02	0.21	0.11	0.91
Depression AAO				
	b	SE	t	р
BALM score	0.23	0.09	2.48	0.01
CUD status	-4.25	6.62	-0.64	0.52
BALM score x CUD status	-0.03	0.20	-0.17	0.87