Role of Tobacco Smoking in Hangover Symptoms Among University Students

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dABSTRACT. Objective: Although hangover results from excessive alcohol consumption, the specific pathways through which hangover symptoms arise have not been elucidated. Research on predictors of hangover sensitivity may provide clues about such mechanisms. The present study investigated whether tobacco smoking on days of heavy drinking affects next-day hangover incidence and severity. Method: The study drew on diary data from a study on smoking and drinking among 113 students at a midwestern university in the United States. Participants completed a daily, web-based, 26-item survey for 8 weeks to assess prior-day alcohol and tobacco use as well as current-day hangover symptoms. Hierarchical linear modeling was used to test the hypothesis that amount of smoking is related to hangover, controlling for amount of alcohol consumed, sex, and other individual characteristics. Analyses were conducted after selecting only days with alcohol consumption levels that typically elicit hangover, then repeated on lighter drinking days for comparison. Validity of the hangover items was checked by comparing reports after such heavy drinking days with days of lighter drinking. Results: Across all possible person-days, 92% of daily reports were obtained. When selecting only events where an estimated blood alcohol concentration of 110 mg/dl was attained, smoking significantly increased the odds of hangover incidence and hangover severity while controlling for number of drinks consumed and sex. Additional analyses controlling for age first smoked regularly, frequency of drug use, type of drug involvement, or smoking status resulted in findings that were unchanged. Conclusions: Smoking more on heavy drinking days affects hangover sensitivity and severity, possibly because of acute pharmacological effects. (J. Stud. Alcohol Drugs, 74, 41–49, 2013)

HANGOVER REFERS TO THE CLUSTER of unpleasant symptoms of physical distress that occur as blood alcohol falls after an acute incident of drinking to intoxication. It is characterized predominantly by thirst, headache, nausea, and tiredness (Rohsenow et al., 2007) and is not to be confused with withdrawal, which occurs only after chronic administration and involves different neurological systems (Prat et al., 2009). The public health relevance of hangover is in its potential effects on occupational safety and performance (Howland et al., 2006; Rohsenow et al., 2006, 2010) and as a predictor of risk for drinking problems after the college years (Piasecki et al., 2005; Rohsenow et al., 2012). Mechanisms underlying hangover are not yet known, but proposed ones broadly include metabolic, fluid balance, hormonal changes, and toxicities resulting from metabolism of alcohol and beverage congeners (Penning et al., 2010).

Not all heavy drinkers appear to be susceptible to experiencing hangover, indicating that some drinkers have hangover insensitivity. A review of studies found that across various study designs, populations, and reference periods, around 23%–25% of people who drink enough to experience hangover report no hangover (Howland et al., 2008b). Attempts to predict who would experience hangover have had limited results so far. Predictors of frequency or severity of hangover found in some studies include having personal risk for alcoholism as measured by the MacAndrew Scale (Earleywine, 1993; MacAndrew, 1965), having a family history of alcoholism (Newlin and Pretorius, 1990; Piasecki et al., 2005), and having a certain alcohol dehydrogenase polymorphism (ADH1B*2) (Wall et al., 2005). Greater average quantity of drinking correlated with less intense hangovers in two studies (Rohsenow et al., 2012; Wall et al., 2000) and with more frequent hangovers in one study (Wall et al., 2005), whereas more frequent heavy drinking correlated with more frequent hangovers among women more than among men in another study (Piasecki et al., 2005). The only previous attempt to find individual difference predictors of hangover insensitivity combined data across three laboratory studies of heavy episodic drinkers in which participants achieved an estimated blood alcohol concentration (BAC) of 110 mg/dl (measured by breath alcohol analysis). That study found no relationship to sex, age, family history of drinking.
Alternatively, alcohol and tobacco use are positively correlated across a variety of studies, especially among heavy or problem drinkers (e.g., Burling and Ziff, 1988; Hurt et al., 1995; Kozlowski et al., 1986; Marks et al., 1997). Amounts of smoking and alcohol consumption vary on a daily basis, particularly among lighter smokers, who may drink and smoke more on social occasions (Jackson et al., 2010). Various mechanisms underlying the correlations of smoking with drinking have been proposed. Some predisposing individual differences that might result in smokers being more likely to drink more than nonsmokers include common genetic factors (Grucza and Bierut, 2006; Swan et al., 1997), negative affectivity (Hesselbrock and Hesselbrock, 2006), or childhood patterns of behavioral undercontrol that predispose both to smoking at an earlier age and to drinking more (e.g., Brown et al., 1996; Farrell et al., 1992; United States Department of Health and Human Services, 2000). In addition, smoking or nicotine use alters subjective responses to alcohol (e.g., Kouri et al., 2004; Madden et al., 1995; Piasecki et al., 2011). Some acute reasons for using nicotine with alcohol include additive effects on dopamine release in the mesolimbic system (Di Chiara and Imperato 1988; Funk et al., 2006; Pierce and Kumaresan, 2006) that could underlie the increased pleasure that is obtained from using both together (Piasecki et al., 2011; Rose et al., 2004) and the fact that nicotinic receptors are involved with effects of alcohol and may be the site of alcohol–tobacco interactions (Funk et al., 2006).

The mechanisms underlying hangover are still speculative (Penning et al., 2010); therefore, the role that cigarette smoking might play in hangover is difficult to determine. Ethanol’s acute effects on the brain include complex central nervous system changes involving γ-aminobutyric acid, dysregulated cytokine pathways, reduced acetylcholine, increased norepinephrine turnover, and other neurotransmitter changes (e.g., Kosten et al., 2005). Consequently, many brain regions may be affected by alcohol leaving the system. The effects of nicotine and alcohol on common neural systems in the brain might also increase residual effects as alcohol leaves the body. Smoking and nicotine withdrawal can acutely affect processes—such as sleep (e.g., Zhang et al., 2006), endocrine responses (Rohleder and Kirschbaum, 2006), and inflammatory responses (e.g., Fröhlich et al., 2003)—thought to play a role in hangover production or severity. Although the paucity of knowledge on neural or physiological changes that result in hangover means this area of investigation is still in the exploratory stage, the common mechanisms underlying alcohol and smoking co-occurrence suggest that it would be useful to investigate the role of smoking in hangover.

There has been little research investigating whether tobacco smoking can contribute to hangover incidence or severity. Hesse and Tutenges (2009) surveyed young adults about hangover severity and found that prior-day number of cigarettes smoked was not associated with hangover, nor was defining oneself as a “regular smoker.” Another report compared undergraduate smokers with nonsmokers and found no differences in likelihood of endorsing a liberally defined “hangover-like experience” (i.e., current hangover, “even just a little”) on waking when drinking frequency was controlled (Piasecki et al., 2010), but the effects of amount of smoking on hangover frequency or severity were not tested.

The present study drew on diary data from a study on smoking and drinking practices among university students (Jackson et al., 2010). Examination of the association between smoking and frequency of hangover must also consider consumption of alcohol in order to account for differences in both variables that might be attributable to confounds with amount of alcohol consumed, or sex differences in alcohol metabolism or in threshold for inducing hangover (Jackson, 2008). Therefore, the first aim of this study was to investigate the hypothesis that a greater amount of tobacco smoked would predict next-day sensitivity to hangover and a greater frequency of hangovers after controlling for variance in hangover that might be accounted for by prior-night number of drinks consumed and sex. Data were analyzed at the event level rather than just in aggregate. The second aim was to investigate whether the relationships are likely to be accounted for by common predisposing individual differences as outlined above. Although we could not directly assess childhood behavioral undercontrol or genetic factors, people who start smoking at an earlier age are more likely to have a pattern of behavioral undercontrol, to use alcohol earlier and more heavily, and to also use recreational drugs (e.g., Brown et al., 1996; Farrell et al., 1992). Therefore, the analyses were repeated while adding as covariates age at first daily smoking, then frequency of recreational drug use, then a categorical variable of no drug use versus only marijuana use versus other illicit drug use. Because inspection of the individual-level data from an early dose-response alcohol administration study (Chapman, 1970) shows that consistent reports of hangover require drinking to an estimated BAC of 110 mg/dl or higher (as described by Rohsenow et al., 2010, and in the review of hangover study methodology by Verster et al., 2010), analyses were conducted while selecting only nights for which estimated BAC was 110 mg/dl or greater, then on lighter drinking nights for comparison. Relationships were expected to be stronger when selecting data that involved drinking to the higher level.

**Method**

**Participants and procedure**

Participants were 113 college student drinkers enrolled in a daily diary study designed to examine alcohol use, tobacco use, and mood (Jackson et al., 2010). Participants
had to have endorsed past-month drinking and to drink during the study; smokers were oversampled (reported 100 or more lifetime cigarettes). See Table 1 for characteristics of participants. All procedures were approved by the university institutional review board, and participants signed informed consent forms.

Participants completed a daily, web-based, 26-item survey for 8 weeks to assess prior-day alcohol and tobacco use as well as current-day hangover symptoms. Each day, participants received a morning email notice prompting them to log in and complete a survey. Paper-and-pencil surveys were available in the event that participants could not access the Internet. Initially, participants were brought into the lab and were given a brief orientation to the survey, including information on how to access to the survey as well as the standard definition of a drink. At this time, they were administered a baseline survey assessing substance use, smoking history, and other psychosocial constructs not used in the present study.

Measures

Drinking. Respondents reported the number of standard drinks consumed on the prior day, ranging from 0 to 25 or more. We also assessed number of hours elapsed between first drink and last drink (in hourly intervals from 1 to 24, in addition to options for 30 minutes and 90 minutes).

Smoking. Prior-day smoking was assessed by asking number of cigarettes smoked, ranging from zero to more than three packs (with an interval of one cigarette for one pack or less, and an interval of five cigarettes for more than one pack).

Hangover. Current-day hangover symptoms were adapted empirically (based on factor loadings and inter-item correlations) from the Hangover Symptoms Scale by Slutske et al. (2003). To decrease response burden with repeated assessments, only five items were used. Although the subset does not include all valid hangover symptoms, high internal consistencies of the various published measures (e.g., \( \alpha = .79 \) for this measure [Slutske et al., 2003] and \( \alpha = .84 \) for an acute hangover measure [Rohsenow et al., 2007]) justify using a subset of items. For use in daily diaries, the questions were changed from asking about percentage of drinking occasions to “Have you felt _____ today because of yesterday’s drinking?” and included five symptoms: felt more tired than usual, had a headache, felt nauseated, felt very weak, and had difficulty concentrating on things. To provide a severity rating, item responses ranged from 1 (not at all) to 7 (extremely). A composite hangover scale was formed by taking the mean of the five items (Cronbach’s \( \alpha \) across the time points = .92; Cronbach’s \( \alpha \) aggregated across individuals = .94). We also created a dichotomous variable reflecting whether the respondent endorsed any hangover symptoms (nonzero mean score) to assess hangover insensitivity.

Blood alcohol estimation. BAC was estimated for each day based on number of drinks, body weight, and the period over which the respondent consumed alcohol, recorded in hours \( (M = 6.46 \text{ hours}, SD = 2.92) \). We used the Matthews and Miller (1979) formula: BAC estimate = \[ \frac{\text{standard drinks}}{2} \times (\text{sex constant} / \text{weight}) - (0.017 \times \text{hours}) \], where the sex constant is 9.0 for females and 7.5 for males.

Additional covariates. Age first smoked was determined by asking respondents to indicate how old they were the first time they started smoking “regularly.” Past-year frequency of use of any drugs, measured on a seven-level ordinal scale ranging from 0 (never/not in the past year) to 7 (41 or more times in the past year), was summed across nine drugs: marijuana, 3,4-methylenedioxymethamphetamine (MDMA; Ecstasy), club drugs, inhalants, stimulants, crack, psychedelics, barbiturates, and heroin. To reflect the degree of involvement with illicit drugs, a three-level drug use variable was also computed that coded no past-year drug use as 0, past-year use of marijuana only as 1, and past-year use of any drugs other than marijuana as 2.

Data analysis approach

First, Pearson’s correlations were conducted among the variables to be entered into the models to show the univariate relationships. These were calculated using all data and repeated using only data from days where estimated BAC was 110 mg/dl or greater.

Because of the clustered nature of the data, with 56 time points per respondent, all subsequent data analysis was conducted using multilevel modeling (MLM; also called hierarchical linear modeling, HLM; Raudenbush and Bryk, 2002; Snijders and Bosker, 1999). These models permit varying numbers of observations and missing observations. Each “day” consisted of 1 day’s drinking and cigarette data and the

### Table 1. Characteristics of study participants \( (N = 113) \)

<table>
<thead>
<tr>
<th>Measure</th>
<th>M (SD) or n (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>49 (43.4%)</td>
</tr>
<tr>
<td>Age, in years ( \text{a} )</td>
<td>18.5 (0.67)</td>
</tr>
<tr>
<td>White</td>
<td>108 (96.4%)</td>
</tr>
<tr>
<td>Perceived dependence on smoking ( \text{b} )</td>
<td>79 (70%)</td>
</tr>
<tr>
<td>Daily smokers ( \text{c} )</td>
<td>59 (53%)</td>
</tr>
<tr>
<td>Reported alcohol use on weekends only during study</td>
<td>3 (3%)</td>
</tr>
<tr>
<td>Ever reached eBAC ( \geq 110 \text{ mg/dl} )</td>
<td>110 (97%)</td>
</tr>
<tr>
<td>Reached eBAC ( \geq 110 \text{ mg/dl} ) more than once</td>
<td>105 (93%)</td>
</tr>
<tr>
<td>Reported any hangover symptoms</td>
<td>110 (97%)</td>
</tr>
<tr>
<td>Reported hangover symptoms more than once</td>
<td>106 (94%)</td>
</tr>
<tr>
<td>Reported any hangover symptoms</td>
<td>108 (96%)</td>
</tr>
</tbody>
</table>

**Notes:** eBAC = estimated blood alcohol concentration. \( \approx 90\% \) were 18 or 19 years old; \( \approx \) endorsed the item “Have you ever felt that you needed tobacco or that you were dependent on it?”; ‘reported smoking “daily or almost daily” in the past month.
next morning’s hangover data. We used HLM 6.06 (Raudenbush et al., 2004) to conduct the multilevel analyses. Level 1 variables were person-centered, and Level 2 variables were grand mean-centered. Level 1 corresponds to within-person time effects, and Level 2 corresponds to between-person data. Number of cigarettes and number of drinks served as time-varying Level 1 variables. We also controlled for sex at Level 2.

Second, the hangover items were tested for validity. Although three of our items had been found valid in experimental hangover induction studies (see Rohsenow et al., 2007), “felt very weak” and “difficulty concentrating” had not been tested to see if they are rated higher after a night of drinking to an estimated BAC of 110 mg/dl or more versus drinking less. Therefore, using MLM, we compared hangover ratings on days when estimated BAC was 110 mg/dl or more the night before with days when the previous night’s estimated BAC was greater than 0 but less than 110 mg/dl. Each hangover rating and the total score were entered into the analysis after controlling for sex.

Third, we tested the effects of smoking on hangover using MLM. The equations below correspond to a model predicting hangover total scores each day from number of cigarettes and number of drinks (Level 1). The model shows how sex was included as a covariate at Level 2.

The model that was fit was as follows:

\[ \eta_{ij} = \beta_0 + \beta_1 \text{(number of cigarettes)} + \beta_2 \text{(number of drinks)} + r_{ij} \]

Level 2 model: (Participant)

\[ \beta_0 = \gamma_{00} + \gamma_{01} \text{(sex)} + u_{00} \]
\[ \beta_1 = \gamma_{10} + u_{10} \]
\[ \beta_2 = \gamma_{20} + u_{20} \]

For analyses of any hangover versus no hangover (hangover sensitivity), we used a Bernoulli (unit-specific) model for binary data. Models were estimated with robust standard errors because of the nonnormality of the outcomes. The models were first conducted while selecting only days where estimated BAC was 110 mg/dl or more to eliminate data from days when hangover was unlikely to occur, and then, for comparison, repeated while selecting days of any lighter drinking (because nondrinking days are irrelevant).

Finally, in a series of parallel models, we added as control variables age first smoked daily, frequency of drug use, and the three-level drug use variable (in separate analyses) to determine whether the relationships between smoking and hangover were attributable to these indicators of an underlying predisposition. Then, given interest in whether hangover is greater among people who routinely smoke more heavily, we sought to determine whether effects of smoking on hangover were attributable to person-level smoking status, suggesting a heavier pattern of smoking rather than the quantity smoked on a given occasion. To address this, we conducted three additional models for each hangover variable, each controlling for one of three person-level smoking status variables: a binary variable reflecting whether they endorsed ever being a dependent smoker (because we had no measure of nicotine dependence), a binary variable indicating whether they reported smoking “daily or almost daily” in the past month, and a continuous variable reflecting the mean number of cigarettes smoked over the 8-week period.

**Results**

**Descriptive information**

Across all possible person-days (56 days × 113 participants), 94% of daily drinking reports (5,930 / 6,328) were obtained. Daily participation rates, which declined over the 8-week interval, ranged from 100% to 73% (on the last survey day); the median daily retention rate was 95% (107 / 113). Table 2 presents event-level characteristics; Figure 1 portrays the number of cigarettes, number of drinks, and degree of hangover reported over the 56-day interval. A strong 7-day pattern was evident for all measures.

**Univariate correlations**

Table 3 shows the bivariate associations between measures of drinking, smoking, and hangover. The lower diagonal shows the associations across the full study period, and the upper diagonal presents associations only for events where estimated BAC was 110 mg/dl or more. These correlations were computed at the within-subjects (daily) level.
although very similar values were observed at the person-level with observations aggregated over the 8 weeks (between-subjects association). Not surprisingly, the number of drinks was bivariately highly associated with any hangover and degree of hangover across all days; the magnitude of the relationship decreased when selecting days of high estimated BAC because of the restriction of range. The number of cigarettes showed small to moderate bivariate correlations with both number of drinks and degree of hangover. There appeared to be no association between sex and hangover despite the expected sex differences in the number of drinks consumed.

**Validity of hangover ratings**

The hangover validity analyses included data on 1,643 to 1,650 events, depending on the variable. Each hangover scale item was significantly higher on mornings after estimated BAC was 110 mg/dl or greater than on mornings after drinking to a lesser estimated BAC level, $F$s (1, 111) from...
119.20 to 350.81, parameter estimates ($\gamma$) range from 0.77 (nauseated) to 1.72 (tired), as was the total score, $\gamma = 1.10$, all $p < .001$. Means and standard deviations for each item are displayed in Table 4.

### Analyses of hypotheses

The multilevel models are shown in Table 5 (estimated BAC $\geq$ 110 mg/dl in the top panel). Number of drinks predicted the hangover variables, as expected. Sex did not predict hangover and, therefore, will not be discussed again. Controlling for these variables, smoking more heavily significantly increased the odds of any hangover and predicted degree of hangover only when drinking to an estimated BAC of 110 mg/dl or greater. We conducted ancillary analyses to explore the extent to which findings were similar when using a variable reflecting whether they were smoking on the drinking day; results indicated that any smoking predicted any hangover (OR = 2.37, 95% CI [1.49, 3.78], $p < .001$) and degree of hangover, $\gamma = 0.34$, $F(1, 109) = 11.76$, $p < .001$, for an estimated BAC of 110 mg/dl or greater but was not a significant predictor and showed no trend supporting such an association for an estimated BAC less than 110 mg/dl (and, in fact, the nonsignificant coefficient was in the opposite direction when predicting any drinking, $\gamma = -0.01$).

In the models in which we added control variables—the age first smoked regularly, frequency of drug use, the three-level drug use variable, the endorsement of smoking dependence, or the average smoking rate of more than 8 weeks (in separate analyses)—gamma weights and ORs were virtually identical (data not presented).

### Table 5. Multilevel models predicting hangover incidence and degree of hangover from prior-day number of cigarettes, number of drinks, and sex

<table>
<thead>
<tr>
<th>Variable</th>
<th>Any hangover</th>
<th>Degree of hangover</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$\gamma$</td>
<td>$F$</td>
</tr>
<tr>
<td>Events $\geq$ 110 mg/dl eBAC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sex (male = 1)</td>
<td>-0.16</td>
<td>0.27</td>
</tr>
<tr>
<td>Number of drinks</td>
<td>0.32</td>
<td>65.29</td>
</tr>
<tr>
<td>Number of cigarettes</td>
<td>0.10</td>
<td>10.37</td>
</tr>
<tr>
<td>Events 0 &lt; eBAC $&lt;$ 110 mg/dl</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sex (male = 1)</td>
<td>-0.15</td>
<td>0.18</td>
</tr>
<tr>
<td>Number of drinks</td>
<td>0.68</td>
<td>66.59</td>
</tr>
<tr>
<td>Number of cigarettes</td>
<td>0.08</td>
<td>3.28</td>
</tr>
</tbody>
</table>

Notes: For events $\geq$ 110 mg/dl estimated blood alcohol concentration (eBAC), $n = 1,272$, approximate $df = 1, 108$ for sex, and approximate $df = 1, 109$ for number of drinks and number of cigarettes. For events where $0 < $eBAC $<$ 110 mg/dl, $n = 385$, approximate $df = 1, 97$ for sex, and approximate $df = 1, 98$ for number of drinks and number of cigarettes. N.S. = not significant.
Discussion

The number of cigarettes consumed on the day of a heavy drinking episode (estimated BAC ≥ 110 mg/dl) predicted both the presence and severity of hangover symptoms the following day, with heavier smoking predicting greater hangover. This is the first published study to examine daily variation in smoking as a predictor of hangover sensitivity or severity on days after very heavy drinking versus lighter drinking using event-level data. Smoking accounted for unique variance after controlling for the relationship of smoking rate to number of drinks and sex. This is important because it means the relationship is not just an artifact of the correlation between drinking rates and smoking rates each day along with the differences in drinking rates by sex. Although the relationships of drinking to hangover were significant regardless of whether data from very heavy drinking nights or lighter drinking nights were used, the effects of smoking on hangover were significant only after nights of drinking to an estimated BAC of at least 110 mg/dl.

The results were not attributable to person-level variables associated with a predisposition to drink or use drugs more frequently because the effects were essentially the same when controlling for such variables. This suggests that the effects are more likely to be attributable to acute pharmacological effects of nicotine or other smoke constituents in the nervous system rather than being attributable to shared predisposing individual differences or traits (e.g., childhood pattern of behavioral undercontrol leading to earlier substance involvement). The results were also not attributable to a pattern of heavier smoking on average or to whether individuals smoked to a level of perceived nicotine dependence because controlling for these variables left the model results unaffected. This is consistent with others who failed to find hangover differences among smokers versus nonsmokers, although neither study modeled event-level smoking contemporaneously (Hesse and Tutenges, 2009; Piasecki et al., 2010). The results thus suggest that the amount of nicotine or smoke constituent exposure is important for affecting hangover sensitivity and severity. Although we do not know how many of the cigarettes were consumed while drinking, nicotine and other smoke constituents are well known to accumulate during the day and may play a pharmacological role while drinking. Future research would need to determine the neuropharmacological mechanisms by which smoking increases unpleasant residual effects of very heavy drinking and whether these same mechanisms play a role in harmful interactions, such as the adverse effects on brain structure, metabolites, cerebral blood flow, and neurocognition found among hazardous drinkers or alcoholics who also smoke (Durazzo et al., 2007).

Many surveys of hangover include items that have not been validated as hangover symptoms by comparing nights of drinking to a BAC of more than 110 mg/dl with nights without drinking to such levels (Rohsenow et al., 2007; Ver ster et al., 2010). Although three of the items in the measure we used had previously been validated in laboratory administration studies involving controlled heavy drinking, the other two items (feeling weak, trouble concentrating) were first validated in the present study, thus supporting preliminary evidence of these items by other studies in the field. These other studies indicated that feeling weak loaded highly on a hangover symptoms scale (Slutske et al., 2003) and was significantly elevated on mornings when college students endorsed a liberally defined hangover item in Piasecki et al. (2010). Both weakness and concentration problems were reported frequently as hangover symptoms by Dutch students (Penning et al., 2012). The ability to use event-level data to compare ratings the morning after drinking to an estimated BAC of 110 mg/dl with nights of lesser drinking makes an important methodological advance for validating hangover symptoms in the field rather than requiring controlled administration in the laboratory.

Strengths and limitations

The present study drew on a rich daily diary data set containing event-based data that permitted examination of episode-specific smoking and hangover. However, an important caveat is the restriction in temporal resolution when using daily data. With daily-level data, it was not possible to determine how much smoking occurred in the hours before reporting hangover symptoms the day after each heavy drinking episode, which could possibly influence symptom reports. However, in the natural environment, smokers will titrate to whatever nicotine level is comfortable at the time to avoid either withdrawal or too much nicotine. Therefore, this represents the naturalistic effects on any hangover rating. Interesting questions for future research may include how much smoking occurred subsequently, as a response to hangover, or as a prelude to or concomitant of additional drinking. Symptoms such as nausea might make a person tend to smoke less. Finally, it is possible that the results are not attributable to cigarettes per day per se but to some behavior correlated with higher smoking rates, such as poorer sleep.

These results are limited to a predominantly White sample of college students at one university and may be different among minority groups or among older adults with heavier smoking patterns. Smokers were oversampled in the diary study, which helps maximize variability in smoking in our analyses but somewhat reduces the generalizability of the sample.

Conclusions

These data add to what is known about interactions of smoking with alcohol use but leave questions about the
mechanisms of these effects. We encourage researchers to replicate the findings reported in the present study using different research designs, measures, and populations, in addition to exploring the mechanisms underlying these interactions in future research.

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References


Raudenbush, S. W., & Bryk, A. S. (Eds.). (2002). Hierarchical linear