Alcohol at bedtime induces minor changes in sleep stages and blood gases in stable chronic obstructive pulmonary disease

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Abstract

Purpose/background The purpose of this study is to explore the effect of a moderate dose of alcohol on sleep architecture and respiration in chronic obstructive pulmonary disease (COPD). Alcohol depresses both hypercapnic and hypoxic ventilatory drives in awake, normal individuals and reduces the amount of rapid eye movement (REM) sleep and oxygen saturation ($S_{pO_2}$) in sleeping COPD subjects.

Methods Prospectively designed, open-label interventional study in a pulmonary rehabilitation hospital. Twenty-six (nine males) stable inpatients, median forced expiratory volume first second (FEV1) 40.5 % of predicted, median age 65 years, investigated by polysomnography including transcutaneous measurement of carbon dioxide pressure increase ($\Delta P_{tcCO_2}$) in randomized order of either control sleep or intervention with 0.5 g of ethanol/kg bodyweight, taken orally immediately before lights off.

Results Alcohol induced a mean increase (95 % confidence interval, [CI]) in the mean $\Delta P_{tcCO_2}$ of 0.10 kPa (0.002–0.206, $P=0.047$) and a mean decrease (CI) in the REM-sleep percentage of total sleep time (REM % of TST) of 3.1 % (0.2–6.0), ($P=0.020$). Six subjects with apnea/hypopnea index (AHI) ≥15 had fewer apneas/hypopneas during alcohol versus control sleep (mean reduction of AHI 11 (1–20), $P=0.046$). Alcohol-sleep changes in $S_{pO_2}$ but not in $\Delta P_{tcCO_2}$, correlated with daytime arterial pressures of carbon dioxide ($P_{aCO_2}$) and oxygen ($P_{aO_2}$).

Conclusion Occasional use of a moderate, bedtime dose of alcohol has only minor respiratory depressant effects on the majority of COPD subjects, and in a minority even slightly improves respiration during sleep. However, caution is appropriate as this study is small and higher doses of alcohol may result in major respiratory depressive and additional negative health effects.

Keywords Ethanol · Transcutaneous blood gas monitoring · Hypoventilation · COPD · Carbon dioxide · Polysomnography

Introduction

Chronic obstructive pulmonary disease (COPD) has a significant morbidity and mortality [1], and chronic hypercapnic respiratory failure (CHRF) in COPD is associated with poor prognosis [2, 3]. Heavy alcohol consumption has been shown to increase the risk of developing COPD [4–6]; on the other hand, mild alcohol intake may reduce the risk both of developing severe pulmonary function abnormalities and of dying from COPD [7, 8]. Sleep is characterized by periods of unstable respiration, and in normal, awake individuals, hypoxic and hypercapnic ventilatory drives are depressed by alcohol [9]. Sleep hypoventilation (SH) has previously been shown to be frequent in severe COPD with CHRF and long-term oxygen therapy (LTOT) [10–12], and recently, we also found SH in
some normocapnic, non-LTOT individuals with stable COPD [12]. Both in a study of 20 asymptomatic men aged 20 to 65 years [13] and in a study of 20 COPD subjects (19 men) [14], alcohol increased the number of sleep apneas. However, a study of 18 asymptomatic, postmenopausal women found no difference between placebo- and alcohol-influenced sleep in the number of episodes of apnea or hypopnea or in the frequency, length, or severity of oxygen desaturation [15]. In two small studies of four normal individuals and five subjects with stable COPD, alcohol taken immediately before sleep resulted in a moderately reduced mean $S_\text{a}O_2$ and a reduced amount of rapid eye movement (REM) sleep [16, 17]. To our knowledge, no studies have explored alcohol-induced changes in carbon dioxide pressure ($PCO_2$) with polysomnography (PSG) in sleeping COPD patients. Concern is thus raised whether it is safe for individuals with COPD, especially those with CHRF or frequent apneas/hypopneas, to occasionally consume alcohol before sleep.

Hypothesizing an increase in the PCO$_2$ during sleep, the primary aim of this study was to explore to what extent a moderate dose of alcohol prior to sleep induces hypoventilation in stable COPD patients, secondary whether apneas/hypopneas and sleep architecture is altered, and finally whether the alcohol-induced changes in blood gases during sleep is correlated to daytime COPD characteristics.

### Material and methods

**Subjects**

Study participants were a randomized selection from the 166 subjects described in a previous article [12], all Caucasians with GOLD [18] defined COPD and inpatients at the Glittreklinikken Pulmonary Rehabilitation Hospital included from January 2010 through June 2011. The following exclusion criteria were used: diagnosed obstructive sleep apnea (OSA); COPD exacerbation within 3 weeks prior; other serious lung comorbidity (i.e., cancer, sarcoidosis, restrictive lung disease) or diseases affecting thoracic or abdominal movement, unstable angina pectoris, hypertension, diabetes mellitus, myocardial infarction within last 3 months, cerebral infarction, and addiction to drugs, alcohol, or narcotics. At inclusion, study subjects were randomized to receive alcohol, 5 mg of Zopiclone (a sleep drug), or supplementary oxygen (data from the latter two interventions not yet published). All subjects used prescribed medication, but no drugs known to be respiratory depressants were taken from 48 h prior to first PSG recording until end of study. The protocol was approved by the Regional Ethics Committee in south-eastern Norway. As shown in Fig. 1, 26 pairs of PSG from control and alcohol-influenced sleep were analyzed.

All subjects slept in their own bed at the hospital with full PSG for three nights; first night for acquaintance with the equipment, second and third nights randomized to either control or alcohol-influenced sleep. PSG was recorded online from Embla A101 with transcutaneously measured carbon dioxide pressure ($P_aCO_2$) from Tosca 500$^2$ to a bedside computer using Somnologica Studio Version 3.3 software. Channel setup was according to the 2007 recommendations from the American Academy of Sleep Medicine (AASM) [19]. PSGs were recorded only during weekdays. Alcohol 0.5 mg per kg bodyweight (corresponding to about three glasses of wine in a 70-kg person) was given as 96 vol% ethanol diluted in 200 ml of orange juice and ingested the last 30 min prior to sleep [20]. Immediately before lights off, breath analysis by Lion Alcometer 500$^3$ was used to check that study-alcohol was taken.

Unpublished data collected at Glittreklinikken prior to the study showed equivalence between changes in $P_aCO_2$ and $P_aCO_2$, but a delay time of 54–57 s. Details regarding delay time, $P_aCO_2$ signal stabilization, sampling frequency, and calculations according to sleep stages are given in our previous article [12].

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1. Medcare Flaga, Reykjavik, Iceland.
2. Radiometer, Basel, Switzerland.
3. Lion Laboratories Ltd., Vale of Glamorgan, UK.
Arterial blood gas samples\textsuperscript{4} were collected after a 10-min seated rest at approximately 2 p.m. prior to each PSG recording and analyzed within 10 min on the Radiometer ABL720Flex.\textsuperscript{5} At sampling, all subjects were breathing room air, with an exception for those on LTOT who used their prescribed dose of supplementary oxygen. Height, weight, and lung function tests including postbronchodilator spirometry, diffusing capacity (DLCO), and body plethysmography of total lung volumes were recorded on MasterScreen Pneumo\textsuperscript{6} as previously described [12], reference values are based on equations from the European Community for Coal and Steel [21].

Sleep scoring was done by two experienced polysomnographers according to recommendations from the AASM [12, 19]; a hypopnea was scored when nasal pressure dropped $\geq$ 30 % for $\geq$ 10 s with $\geq$ 4 % desaturation drop from baseline, with $\geq$ 90 % of the event’s duration meeting the amplitude reduction criteria for hypopnea (criterion A). Sleep hypoventilation is defined by the AASM as an increase of 1.3 kPa or more in $P_{a}CO_{2}$ during sleep, to a value exceeding 6.7 kPa for at least 10 min [22].

Statistical analysis

Sample size calculations up front showed that for $N$=28, a $P_{a}CO_{2}$ difference between alcohol and control sleep would be detected with 80 % probability at a two-sided 0.05 significance level if the true difference was 0.25 kPa, assuming a within-patient standard deviation of 0.32 kPa\textsuperscript{7}.

Data were assessed for normality of distribution and homogeneity of variance. No violation of the assumptions of normality, linearity, multicolinearity, and homoscedasticity was found preliminary to regression analysis.

Differences between groups with continuous, paired parametric data were analyzed with Student $T$, paired non-parametric data with Wilcoxon signed rank, independent non-parametric data with Mann-Whitney $U$, and non-parametric data in more than two groups by Kruskal-Wallis tests. Paired proportions were analyzed with McNemar chi-square test ($P$ value uncorrected) and correlations with Pearson’s $r$.

Two-sided $P$ values of $\leq$0.05 were considered significant in testing the main hypothesis, whereas a significance level of $P$ $\leq 0.025$ was applied in the remaining analyses. All analyses were performed using IBM SPSS Statistics version 19.

\textsuperscript{4}PICO50, Radiometer, Copenhagen, Denmark.
\textsuperscript{5}Radiometer, Copenhagen, Denmark.
\textsuperscript{6}Jaeger-Toennies, Hoechberg, Germany.
\textsuperscript{7}In our previous study of 100 COPD subjects [12], the mean increase (standard deviation) in $P_{a}CO_{2}$ during spontaneous sleep was 0.49 (0.32) kPa.

Results

The 9 men and 17 female (Table 1) were on average hyper-inflated with impaired gas exchange and severe airway obstruction, 14 (54 %) of the 26 subjects at GOLD stages III and IV. Four subjects (16 %) had daytime hypoxemia (arterial pressures of oxygen ($P_{a}O_{2}$)<8.0 kPa), whereas only one had $P_{a}O_{2}>12.0$ kPa. CHRF ($P_{a}CO_{2}$ $\geq$6.3 kPa) was found in four subjects, of which three used LTOT. The study group had a normal median BMI, five subjects (19 %) being underweight (BMI $<21$), whereas five were obese (BMI $\geq$30).

As indicated in Table 2, alcohol at bedtime reduced the REM-sleep percentage of total sleep time (REM % of TST) with a mean (SD) of 3 (7 %), and the number of awakenings with 8 (13). However, eight subjects had an increased amount of REM sleep when influenced by alcohol (Fig. 2, groups 2 and 4).

The mean $P_{a}CO_{2}$ increase from supine, resting value prior to sleep ($\Delta P_{a}CO_{2}$) was higher during alcohol versus control sleep (a mean (SD) increase of 0.10 (0.25) kPa), despite nine subjects having a decrease in the mean $\Delta P_{a}CO_{2}$ (Fig. 2, groups 3 and 4). As shown in Table 4, alcohol-sleep changes in mean $\Delta P_{a}CO_{2}$ and mean $S_{a}O_{2}$ did not correlate.

Sleep hypoventilation (SH) as defined by the AASM was found in two subjects (8 %) during control sleep and in five subjects (19 %) in alcohol sleep [22]; however, the difference was not statistically significant as one subject with SH in control sleep had no SH during alcohol sleep.

Ten subjects had a drop in alcohol-sleep mean $S_{a}O_{2}$ of $>1$ %, whereas the NREM sleep mean $S_{a}O_{2}$ for the whole study group decreased with a mean (SD) of 0.9 (2.1) % ($P$ $=$0.019). No significant change was found in REM sleep (Table 2). Interestingly, Table 2 also shows no respiratory depressant effect of alcohol during the awake periods after the initial sleep onset (N0) nor was there significant N0 changes in mean $\Delta P_{a}CO_{2}$ and mean $S_{a}O_{2}$ when analyzing only subjects with alcohol increased sleep mean $\Delta P_{a}CO_{2}$ (groups 1 and 2, Fig. 2).

The median values of apneas/hypopneas and desaturations per hour of sleep (AHI and ODI, respectively) were not changed by alcohol. Despite the exclusion of diagnosed OSA prior to the study, six subjects (four males) had an AHI of $\geq 15$ per hour in control sleep. Intriguingly, Table 3 indicates that this subgroup had alcohol sleep decreases in median AHI and ODI, all with increased mean $\Delta P_{a}CO_{2}$ and decreased REM % of TST.

As the three LTOT users were outliers in the data distribution of the mean $S_{a}O_{2}$ change, partial correlation was applied, controlling for LTOT use (yes/no). Alcohol-sleep changes in the mean $S_{a}O_{2}$ ($\Delta A_{a}O_{2}$) were highly correlated to daytime $P_{a}O_{2}$ and inversely to $P_{a}CO_{2}$ (Table 4). However, this was not the case regarding the alcohol-sleep changes in the mean $\Delta P_{a}CO_{2}$ nor did hypventilation during alcohol sleep correlate to spirometry or demographic data.
Finally, hierarchical multiple regressions were performed to determine whether the AScordO2 could be explained by daytime COPD characteristics. P02 and Pco2 were entered at step 1, explaining 34 % of the variance in AScordO2. After the entry of LTOT (yes/no) at step 2, the total variance explained by the model was 49 %, (F(3, 22)=7.09, P=0.002). Thus, LTOT (yes/no) explained an additional 15 % of the variance in the AScordO2 after controlling for P02 and Pco2 (R2 change=0.015, F change (1, 22)=6.48, P=0.018). In the full model, using LTOT recorded the highest beta value (0.70, P<0.018), over Pco2 (−0.59, P<0.031), whereas P02 did not make a unique contribution (0.31, P=0.116).

Table 1: Study population

| Demographic data | N=26 | Median (IQR) |
|------------------|------|--------------|
| Gender female    | 17 (65) |             |
| Age, years       | 65.5 (14.0) |          |
| BMI, kg/m²       | 25.0 (6.5)  |            |

| Smoking habit    | Pack years | 30.4 (22.4) |
| Current smoker   | 4 (15)  |             |

| Medication | LTOT | 3 (12)  |
|-----------|------|---------|
| SABA/LABA | 24 (92) |       |
| Statin    | 4 (15)  |         |
| ACE/AII   | 3 (12)   |         |
| Thiazide  | 3 (12)   |         |
| ASA       | 3 (12)   |         |

| Spirometry | FVC, % of pred | 81.0 (35.3) |
|------------|----------------|-------------|
| FEV1, % pred | 40.5 (30.3) |           |
| FEV1/FVC, ratio | 0.46 (0.21) |          |
| DLCO1, mmol/min/kPa | 3.92 (2.49) |         |
| RV/TLC2, ratio | 0.56 (0.20) |         |

| Clinical data | CAT, score | 18 (7) |
|---------------|------------|-------|
| MMRC, score   | 2.0 (1.0)  |       |
| 6-MWD, meter  | 420 (205)  |       |
| BODE index    | 2.0 (4.0)  |       |

| Laboratory data | P02, kPa | 9.66 (1.98) |
|-----------------|----------|-------------|
| Pco2, kPa       | 5.07 (0.96) |         |
| Sao2, %         | 96.2 (3.3)  |         |
| Alcohol, ppt    | 0.81 (0.64)  |         |

Table 2: Control versus alcohol-influenced sleep

| N=26 | Control sleep | Alcohol sleep | P value |
|------|---------------|---------------|---------|
| Sleep architecture | TST, min | 354.0 (65.3) | 354.0 (62.5) | 0.501 |
| REM % of TST         | 25 (8)   | 20 (9)  |           | 0.020 |
| NREM % of TST        | 75 (8)   | 80 (9)  |           | 0.020 |
| WASO, min            | 45.7 (74.7) | 40.9 (48.5) | 0.361 |
| Awakenings           | 28 (18)  | 19 (17) |           | 0.012 |

| Ventilation | AHI | 9.7 (10.2) | 10.4 (16.4) | 0.517 |
| H1           | 4.8 (6.2)  | 6.3 (8.6)  |           | 0.309 |
| ODIa         | 5.5 (11.6) | 8.3 (12.8) |           | 0.548 |

| Mean SaO2 % | N0 | 94.0 (5.1) | 93.9 (5.6) | 0.551 |
| NREM        | 93.7 (4.6) | 93.2 (5.4) |           | 0.019 |
| REM         | 92.5 (5.4) | 92.8 (6.4) |           | 0.258 |

| Sleep | 93.4 (4.7) | 93.2 (5.7) | 0.058 |

| Mean ΔPaCO2 kPa | N0 | 0.19 (0.23) | 0.15 (0.22) | 0.638 |
| NREM        | 0.39 (0.34) | 0.55 (0.36) |           | 0.035 |
| REM         | 0.67 (0.46) | 0.67 (0.42) |           | 0.082 |

| Sleep | 0.49 (0.35) | 0.58 (0.36) | 0.047 |

| Sleep hyperventilation | Shb | 8 (27) | 19 (40) | 0.179 |

| N=25 | Alcohol sleep vs Control sleep |
|------|--------------------------------|
| TST  | 354.0 (62.5) |
| REM  | 20 (9) |
| NREM | 80 (9) |
| WASO | 40.9 (48.5) |
| Awakenings | 19 (17) |

PaO2 and PaCO2 were entered at step 1, explaining 34 % of the variance in DeltaAsO2. After the entry of LTOT (yes/no) at step 2, the total variance explained by the model was 49 %, (F(3, 22)=7.09, P=0.002). Thus, LTOT (yes/no) explained an additional 15 % of the variance in the DeltaAsO2 after controlling for P02 and Pco2 (R2 change=0.015, F change (1, 22)=6.48, P=0.018). In the full model, using LTOT recorded the highest beta value (0.70, PaO2 and PaCO2 were entered at step 1, explaining 34 % of the variance in DeltaAsO2. After the entry of LTOT (yes/no) at step 2, the total variance explained by the model was 49 %, (F(3, 22)=7.09, P=0.002). Thus, LTOT (yes/no) explained an additional 15 % of the variance in the DeltaAsO2 after controlling for P02 and Pco2 (R2 change=0.015, F change (1, 22)=6.48, P=0.018). In the full model, using LTOT recorded the highest beta value (0.70, P<0.018), over Pco2 (−0.59, P<0.031), whereas P02 did not make a unique contribution (0.31, P=0.116).
The table 3 shows the alcohol-induced sleep changes in overlap subjects. The data includes the control and alcohol sleep measurements for the REM and NREM mean percentage of TST. The change in mean ΔPtcCO2 versus changes in REM-sleep percentage of TST is also represented in the colored figure according to group (legend in panel).

Discussion

When stable COPD subjects (2/3 female) are given a moderate dose of alcohol immediately prior to sleep, we find a very modest sleep hypoventilation as a group mean increase in PtcCO2 of about 0.1 kPa and a decrease in SpO2 of less than 1 %. This hypoventilation is not detectable when subjects are awake after the initial sleep onset. The mean REM % of TST is decreased in alcohol sleep. A novel finding is subgroups showing opposite characteristics as slight hyperventilation with decreased amount of REM sleep and hypoventilation with increased REM percentage of TST. AHI and ODI are not significantly increased by alcohol; however, in overlap subjects (AHI ≥15/h), AHI and ODI are lower. Alcohol-sleep change in SpO2, but not PtcCO2, is associated to daytime blood gases, and a model with LTOT, PtcCO2, and PtcO2 can explain about half of the variation in alcohol sleep change in SpO2.

In a much referred study from 1967, Yules et al. gave 1 g of ethanol/kg body weight to four male graduate students 4 h prior to sleep and found that the REM-sleep time reduced as a mean for the group; however, one subject showed no change in REM-sleep time [16]. Later, Easton et al. gave five COPD subjects 0.78 g of alcohol/kg bodyweight immediately prior to sleep and found a mean TST decrease of 19 %, a REM-sleep percentage of TST decrease by 12 % and a mean drop in SpO2 of about 3 % [17]. In 2007, Brander et al. studied nine males with advanced COPD using 0.5 g of alcohol/kg bodyweight albeit taken earlier in the evening, they did not record PSG but found only a trivial decrease in SpO2 (<1 %) during the first 2 h of the reported sleep [23].

In the present study, we find REM % of TST decreased as in the studies referred above, however, only in 18 (69 %) of the subjects. One possible explanation for the eight subjects showing increased REM % of TST can be alcohol abstinence. Easton et al. found the REM % of TST gradually increasing when giving alcohol repeatedly for several nights. Although none of our study subjects had a drinking problem (otherwise they would not have been included), some could nevertheless have used alcohol regularly before admission, and thus the abstinient state during the first week at the hospital might explain the increase in REM sleep when given alcohol. However, we find it unlikely that as many as 31 % of the study subjects had an undisclosed alcohol problem, and as Yules et al. also described one subject with no change in REM-sleep time [16], hitherto unknown effects of alcohol on sleep can have caused the increased mean REM % of TST in some of the subjects in the present study (Fig. 2).

In a previous article, we showed that ΔPtcCO2 increases with depth of NREM sleep in stable COPD and with the highest values recorded during REM sleep [12]. Thus, finding the REM percentage of TST being reduced in alcohol sleep, one would expect to find the mean ΔPtcCO2 to be decreased. Indeed, although not clinically impressive, the seven (27 %)

Table 3 Alcohol-induced sleep changes in overlap subjects

| N=6 (four male) | Control sleep | Alcohol sleep | P value |
|----------------|---------------|---------------|---------|
| REM % of TST   | 25 (12)       | 21 (15)       | 0.028   |
| NREM mean ΔPtcCO2, kPa | 0.48 (0.39) | 0.68 (0.27) | 0.028   |
| REM mean ΔPnCO2, kPa | 0.64 (0.63) | 0.91 (0.35) | 0.028   |
| Sleep mean ΔPnCO2, kPa | 0.51 (0.46) | 0.75 (0.28) | 0.028   |
| Sleep mean SpO2, % | 93.3 (3.8) | 93.2 (6.5) | 0.075   |
| ODI            | 32 (28)       | 24 (22)       | 0.028   |
| AHI            | 39 (40)       | 26 (22)       | 0.046   |
| REM AHI        | 49 (43)       | 36 (28)       | 0.075   |
| NREM AHI       | 31 (46)       | 30 (23)       | 0.116   |

As no record of daytime sleepiness was available, overlap was defined as having at least 15 apneas/hypopneas per hour of sleep in subjects with chronic obstructive pulmonary disease.
subjects in group 3 in Fig. 2 fit in this picture. Brief alcohol exposure is moderately bronco-dilating and increases the mucociliary motility in asthmatics [24], and thus can relieve nocturnal wheezing in reversible COPD subjects, and possibly improve gas exchange in the lungs. However, the 11 (42 %) subjects with reduced REM % of TST and increased \( P_{a}CO_{2} \) (group 1, Fig. 2) are more in accordance with previous findings in COPD [17]. The hypoventilation demonstrated in the present study with only two subjects (8 %) having a mean alcohol-sleep increase in \( \Delta P_{a}CO_{2} \geq 0.5 \) kPa and less than half having a drop in mean Sp\(_{\text{o2}}\)\( > 1 \) % was considerably less impressive than Easton et al.’s findings and more in line with the results of Brander et al.[17, 23]. One obvious explanation is the amount of alcohol and the time of ingestion related to sleep onset, as Brander et al. gave less alcohol earlier in the evening compared to Easton et al., the present study using same amount of alcohol as Brander et al., but given immediately before lights off. Thus, as REM sleep is more frequent the second half of the night when most of the alcohol is metabolized, REM-sleep hypoventilation is not significantly changed. Another explanation can be the number of study subjects; as previous studies included only up to six subjects, they could have missed the individuals having decreased \( P_{a}CO_{2} \) during alcohol sleep.

Previous studies have shown increased numbers of apneas in alcohol-sleep in asymptomatic and in snooring males, in mild to moderate OSA-males and in male, but not in female COPD subjects [13, 25, 26, 14, 15]. The present study did not find AHI, HI, and ODI significantly increased by alcohol. This can partly be explained by the majority of female subjects in the present study. Also, as apneas/hypopneas are most frequent in REM sleep, occurring at low alcohol content, the reduced REM % of TST in alcohol-sleep can result in the total AHI being unchanged. However, the subgroup of mostly men with control sleep AHI ≥ 15/h having decreased AHI and ODI during alcohol sleep seems to contradict previous findings in males. All these individuals are hypoventilating, yet they have fewer apneas, and although not statistically significant, the REM-sleep AHI tends to be lower in alcohol-sleep. A reduced tidal volume with increased frequency of respiration, shorter episodes of REM-sleep, less mucus, and reduced bronchial constriction can be possible explanations.

Interestingly, no changes are found in \( \Delta P_{a}CO_{2} \) or Sp\(_{\text{o2}}\) in N0 (awake after initial sleep onset), suggesting that moderate doses of alcohol has a significant effect on the central respiratory control only during sleep. This is in support of Sahn et al., who did not find significant alcohol-induced hypoventilation (neither as decreased PO\(_{2}\) nor increased PCO\(_{2}\)) in six awake subjects with severe COPD after drinking 0.78 g of alcohol/kg bodyweight [27].

Having COPD with daytime respiratory failure correlates with alcohol-induced sleep hypoventilation, however, only as daytime blood gases versus \( \Delta Sp_{\text{O2}}\) and not hypventilation as change in mean \( \Delta P_{a}CO_{2} \) (Table 4). As the multiple linear regressions with the variables LTOT, \( P_{a}O_{2} \), and \( P_{a}CO_{2} \) can explain only half of the decrease in Sp\(_{\text{o2}}\) in alcohol sleep, advice to COPD patients regarding the use of alcohol cannot be based on whether they have respiratory failure or not. Nor is the severity of COPD according to FEV1 as percentage of predicted, the transfer factor for carbon monoxide (DLCO) or the residual volume as percentage of total lung volume (RV/TLC) highly associated with the alcohol-induced sleep hypoventilation.

This study is to our knowledge the first to explore sleep hypoventilation as both oxygen desaturation and carbon dioxide increase in moderately alcohol-intoxicated COPD

### Table 4  Correlation between alcohol-induced blood gas changes in sleep and study population characteristics

| N=26 | Change in mean \( Sp_{\text{o2}}\)\( ^{a} \) | Change in mean \( \Delta P_{a}CO_{2} \) |
|------|----------------|----------------|
|      | Pearson’s \( r \) P value | Pearson’s \( r \) P value |
| **Demographic data** | | |
| Age, years | −0.11 | 0.616 |
| Gender | 0.08 | 0.709 |
| BMI kg/m\(^{2}\) | −0.13 | 0.543 |
| **Laboratory data** | | |
| \( P_{a}O_{2} \), kPa | 0.51 | 0.009 |
| \( P_{a}CO_{2} \), kPa | −0.58 | 0.002 |
| pH | 0.37 | 0.071 |
| **Spirometry** | | |
| FVC % of pred | 0.28 | 0.170 |
| FEV1 % of pred | 0.32 | 0.125 |
| DLCO mmol/min/kPa\(^{a}\) | 0.32 | 0.161 |
| RV/TLC ratio\(^{c}\) | −0.28 | 0.202 |
| **Sleep** | | |
| Change in REM % of TST | 0.12 | 0.554 |
| Change in awakenings | −0.11 | 0.602 |
| Change in mean \( Sp_{\text{o2}}\) | − | |

\( ^{a} \) Partial correlation controlling for LTOT use is tabulated

\( ^{b} \) Four missing from DLCO because of insufficient vital capacity or because they could not hold their breath for 10 s

\( ^{c} \) Two missing from body plethysmography because of claustrophobia

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subjects; however, it has some limitations. Many comparisons increase the risk of type I statistical errors (falsely rejecting the null hypothesis when in fact it is true) and although an alpha-correction was applied in interpreting the results of analyses other than the main hypothesis, the significance of the results must be viewed with this risk in mind. The open-label design can add a placebo bias to the alcohol intervention. However, we find it less likely that ventilation during sleep is affected by the knowledge of being drunk, and the fact that blood gases were unchanged when awake after initial sleep onset indicates little placebo impact. Another limitation is the lack of a detailed, structured alcohol consumption interview, as regular alcohol ingestion, although not being considered as a drinking problem, can result in a very different sleep and breathing patterns as discussed above. Thirdly, no records were made of whether or not the study subjects were fasting at the time of alcohol ingestion. The alcohol breath test performed immediately before lights off was not considered to be a valid measurement of blood alcohol content (BAC) as such analyses are considered to be unreliable in COPD [28–30]; nevertheless, the breath analyses showed an unexpected high variability (Table 1). Food taken together with alcohol can delay peak BAC, possibly explaining some of this variability. If BAC correlates with \( \Delta P_{\text{a}}CO_2 \), fasting individuals can have peak BACs and thus high \( \Delta P_{\text{a}}CO_2 \) values early in the night, compared to those with their BAC peak well into the night. However, supper at the hospital was at 6:30 p.m., and we find it unlikely that many study subjects ate at a later time.

**Conclusion/summary**

In this study of stable COPD patients drinking a moderate dose of alcohol immediately prior to lights off, we found a very modest sleep hypoventilation as increased mean \( \Delta P_{\text{a}}CO_2 \), a reduced REM % of TST, and no changes in AHI. However, this combination of hypoventilation and decreased REM-sleep time was evident in less than half of the subjects, as about one-fourth had a minor sleep hyperventilation with reduced REM % of TST and another one-fourth was hypoventilating with an increase in REM % of TST. Also, subjects with COPD and OSA (overlap) had lower median AHI and ODI in alcohol-sleep versus control sleep. We found an association between daytime blood gases and alcohol-sleep hypoventilation as decreased \( S_{\text{p}}O_2 \), but not increased \( \Delta P_{\text{p}}CO_2 \), and a regression model with the variables LTOT-use, \( P_{\text{p}}O_2 \), and \( P_{\text{p}}CO_2 \) could explain 49 % of the variation in alcohol sleep \( S_{\text{p}}O_2 \) change. As the blood gas changes were minor, the clinical implication of this study is that a single, moderate, bedtime dose of alcohol seems to have only minor respiratory depressant effects on the majority of COPD subjects, and in a minority even slightly improves respiration during sleep. However, caution must be applied as the study is small, and higher doses of alcohol may result in major respiratory depressive and additional negative health effects.

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