Coexistence of OSA may compensate for sleep related reduction in neural respiratory drive in patients with COPD

Bai-Ting He, Gan Lu, Si-Chang Xiao, Rui Chen, Joerg Steier, John Moxham, Michael I Polkey, Yuan-Ming Luo

ABSTRACT

Background The mechanisms underlying sleep-related hypoventilation in patients with coexisting COPD and obstructive sleep apnoea (OSA), an overlap syndrome, are incompletely understood. We compared neural respiratory drive expressed as diaphragm electromyogram (EMGdi) and ventilation during stage 2 sleep in patients with COPD alone and patients with overlap syndrome.

Methods EMGdi and airflow were recorded during full polysomnography in 14 healthy subjects, 14 patients with OSA and 39 consecutive patients with COPD. The ratio of tidal volume to EMGdi was measured to indirectly assess upper airway resistance.

Results Thirty-five patients with COPD, 12 healthy subjects and 14 patients with OSA completed the study. Of 35 patients with COPD, 19 had COPD alone (FEV1 38.5%±16.3%) whereas 16 had an overlap syndrome (FEV1 47.5%±16.2%, AH1 20.5±14.1 events/hour).

Ventilation (Vt) was lower during stage 2 sleep than wakefulness in both patients with COPD alone (8.6±2.0 to 6.5±1.5 L/min, p<0.001) and those with overlap syndrome (8.3±2.0 to 6.1±1.8 L/min). Neural respiratory drive from wakefulness to sleep decreased significantly for patients with COPD alone (29.5±13.3% to 23.0±8.9% of maximal, p<0.01) but it changed little in those with overlap syndrome. The ratio of tidal volume to EMGdi was unchanged from wakefulness to sleep in patients with COPD alone and healthy subjects but was significantly reduced in patients with OSA or overlap syndrome (p<0.05).

Conclusions Stage 2 sleep-related hypoventilation in COPD alone is due to reduction of neural respiratory drive, but in overlap syndrome it is due to increased upper airway resistance.

INTRODUCTION

COPD is a common condition and patients with COPD are subject to hypoxaemia or even respiratory failure during sleep because of hypoventilation. In prior reports we showed that this hypoventilation, in the absence of upper airway obstruction was due to a reduction in neural respiratory drive to the respiratory muscles as measured by the diaphragm electromyogram (EMGdi). Obstructive sleep apnoea (OSA) is characterised by repeated partial or complete collapse of the upper airway leading to increased upper airway resistance and arousal from sleep which are associated with increased neural respiratory drive.

What is the key question?
- The mechanisms underlying sleep-related hypoventilation in patients with coexistent COPD and obstructive sleep apnoea (OSA), an overlap syndrome, are unknown.

What is the bottom line?
- This study shows that sleep-related hypoventilation in patients with overlap syndrome is due to an increase in upper airway resistance associated with OSA rather than reduction of neural respiratory drive associated with COPD.

Why read on?
- Sleep-related hypoventilation in patients with COPD alone mainly occurs because of a decrease in neural respiratory drive whereas it is mainly a result of an increase in upper airway resistance in patients with overlap syndrome.
can be inferred by the ratio of the tidal volume ($V_t$) to EMG$_{di}$ ($V_t$/EMG$_{di}$) assuming that lung mechanics and lower airway resistance remain constant in wakefulness and sleep. Here we aimed to investigate the underlying mechanisms of hypoventilation during sleep in patients with COPD alone and patients with overlap syndrome by comparing neural respiratory drive and ventilation. The data from 10 of the patients with COPD alone and 10 healthy subjects has previously been reported; these were the participants with complete datasets from that report.

**METHODS**

**Subjects**

Thirty-nine consecutive patients with COPD from the outpatient clinic of Guangzhou Institute of Respiratory Disease were studied, including 10 patients whose data were reported previously. Exclusion criteria were a known diagnosis of OSA before the study, clinically significant coexisting diseases including cardiovascular and neuromuscular disease, an acute exacerbation of COPD in the preceding month, FEV$_1$ < 20% predicted, and use of long-term oxygen therapy. Usual medications including inhaled bronchodilators were allowed for all the patients. Fourteen healthy subjects and 14 patients with OSA were also studied; as noted 10 of the healthy subjects had appeared in our prior report. The study was approved by the Ethics Committee of the First Affiliated Hospital of Guangzhou Medical University and all subjects gave written informed consent.

**Lung function tests**

Spirometry (Cosmed Micro Quark, Cosmed, Italy) was performed on the same night as polysomnography. Measurements were repeated until maximal reproducible values of FEV$_1$ were achieved with variation of less than 150 ml between tests. The modified Medical Research Council dyspnoea scale (mMRC) was also recorded before polysomnography.

**Oesophageal electrode and its positioning**

A soft fine catheter with an external diameter of 1.6 mm was used to record the EMG$_{di}$. The catheter had 10 metal coils which provided five pairs of recording electrodes was used to record the EMG$_{di}$. The oesophageal electrode was passed through the nose into the stomach and was carefully positioned based on the EMG$_{di}$ amplitude recorded simultaneously from the five pairs of electrodes, as reported previously. Briefly the position of the electrode catheter was judged to be optimal when electrode 5 was located at the level of the diaphragm confirmed by the amplitude of EMG$_{di}$ activity being greatest in pairs 1 and 5, and smallest in pair 3 during inspiration. When the electrode catheter was in the optimal position, it was securely taped at the nose. The EMG$_{di}$ signals were amplified and band-pass filtered between 20 Hz and 1 kHz (RA-8, Yinhui Medical Technology Co., Guangzhou, China).

**Measurement of maximal EMG$_{di}$**

Maximal EMG$_{di}$ was recorded during two manoeuvres: maximal inspiration to total lung capacity (TLC) and maximal inspiration against a closed airway at functional residual capacity for 3 s. Each manoeuvre was repeatedly performed until subjects were able to master the techniques. More than three maximal efforts, with an interval of 30 s or more between them, were recorded for analysis and the largest single value was considered maximal.

**Polysonmography**

Full overnight polysomnograms including the EEG (C3-A2, C4-A1), left and right electro-oculograms (EOG), submental electromyogram (EMG$_{chin}$), airflow, snoring, body position, oxygen saturation and end-tidal CO$_2$ were recorded. Airflow was recorded with a pneumotachograph connected to a full face mask, and integrated to produce volume. All signals were recorded simultaneously using a Powerlab recording system (ADInstruments, Castle Hill, Australia). The sampling rate was 2 kHz for EMG$_{di}$ and 200 Hz for other signals. EMG$_{di}$ was recorded before and during sleep.

**Analysis of data**

Conventional polysomnography was manually analysed based on standard criteria. An obstructive apnoea event was defined as absence of airflow for longer than 10 s, while there was phasic inspiratory EMG$_{di}$. A hypopnoea event was defined as reduction of airflow of more than 30% for longer than 10 s associated with ≥3% desaturation or the event being associated with arousal. Overlap syndrome was defined as apnoea hypopnoea index (AHI) ≥ 5 events/hour in the presence of COPD. Clinically significant sleep-related desaturation was defined as SaO$_2$ < 90% which lasted longer than 5 min. The root mean square of the EMG$_{di}$ was calculated by computer with a time constant of 100 ms. The root mean square reported was that from the electrode pair with the largest EMG$_{di}$ amplitude for each breathing cycle. To avoid the influence of the electrocardiogram on the EMG$_{di}$, root mean square was measured from segments between QRS complexes. A ratio of $V_t$/EMG$_{di}$ was calculated with a pneumotachograph, therefore final data were derived from 35 patients with COPD, 12 healthy subjects and 14 patients with OSA (see table 1 and online supplementary table E-1). Using the above criteria 16 of the patients with COPD were found to have overlap syndrome (FEV$_1$ 47.5 ± 16.2%; AHI 20.5 ± 14.1 events/hour) and 19 patients had COPD alone (FEV$_1$ 38.5 ± 16.3%; AHI 1.9 ± 1.6). The body mass index (BMI) in patients with COPD alone (19.9 ± 2.7 kg/m$^2$) was significantly lower than that in patients with overlap syndrome (23.5 ± 3.8 kg/m$^2$) (p < 0.05). The FEV$_1$ in patients with COPD alone was not significantly different from that in patients with overlap syndrome (38.5% ± 16.3% vs 47.3% ± 16.2%, p > 0.05). The respiratory events in patients with overlap syndrome were predominantly hypopnoea rather than apnoea, except for subject 1 (see table 1 and online supplementary table E-2). Overall, the ratio of hypopnoea to apnoea events in patients with overlap syndrome...
was 4.4:1. COPD alone and overlap syndrome groups were similar to each other but, as expected, had significantly greater dyspnoea compared with healthy subjects and patients with OSA (p<0.001). The maximal EMGdi was similar between groups and was 176.9±72.1, 192.3±56.6, 156.1±50.3, 166.2±40.8 μV for COPD alone, patients with overlap syndrome, healthy subjects and patients with OSA (p>0.05), respectively.

The mean SaO2% over the entire sleep period was 96.8±1.4, 97.1±1.3, 98.0±1.0 and 96.9±0.7 during wakefulness and 96.0±1.6, 95.8±1.4, 97.6±0.9 and 96.1±1.4 during sleep for patients with COPD alone, patients with overlap syndrome, healthy subjects and patients with OSA, respectively. COPD alone and overlap syndrome groups were similar to each other but had a lower mean minimal SaO2% during overnight sleep compared with healthy subjects (see table 1 and online supplementary table E-2). Mean minimal SaO2% in patients with OSA was lower than those in the other three groups. Three of 19 (16%) patients with COPD alone developed SaO2<90% for longer than 5 min, whereas no subjects in both the patients with overlap syndrome and the healthy subjects developed significant oxygen desaturation (see online supplementary table E-2).

### Ventilation in patients with COPD, patients with overlap syndromes, healthy subjects and those with OSA during wakefulness and sleep

V̇E was similar between groups and was 8.6±2.0, 8.3±2.0, 8.3±1.6 and 8.0±2.7 L/min for patients with COPD alone, patients with overlap syndromes, healthy subjects and patients with OSA during wakefulness (p>0.05). V̇E decreased significantly from wakefulness to non-rapid eye movement (NREM) sleep for COPD alone (8.6±2.0 to 6.5±1.5 L/min, p<0.001), patients with overlap syndrome (8.3±2.0 to 6.1±1.8 L/min, p<0.001) and patients with OSA (8.0±2.7 to 6.3±1.9 L/min, p=0.01), although the change in V̇E failed to attain statistical significance between wakefulness and NREM sleep in healthy subjects (8.3±1.6 to 7.5±1.3 L/min, p=0.07). The decrease in V̇E from wakefulness to sleep was similar between patients with COPD and patients with overlap syndrome (24% vs 27%, p=0.05), but it was greater in both groups than that observed in healthy subjects (10%), p=0.05. Change in V̇E was almost proportional to change in V̇T in patients with COPD, patients with overlap syndrome and patients with OSA (table 2, figures 1 and 2).

Neural respiratory drive decreased significantly in patients with COPD alone (25.2±3.3% vs 23.0±8.9% of maximal EMGdi, p<0.01), but increased significantly in patients with OSA (14.2±7.9% vs 23.4±10.8% of maximal EMGdi) from wakefulness to NREM sleep. However, neural respiratory drive changed little from wakefulness to stage 2 sleep in patients with overlap syndrome (29.3±15.8% vs 27.3±14.7% of maximal EMGdi, p>0.05) and in healthy subjects (12.6±3.7% vs 12.2±3.3% of maximal EMGdi, p>0.05). Respiratory rate also changed little between wakefulness and NREM sleep in all four groups (p>0.05). V̇E/EMGdi was similar between wakefulness and NREM sleep in both healthy subjects (0.69±0.18 vs 0.69±0.20) and patients with COPD alone (0.39±0.23 vs 0.37±0.21) but it decreased significantly from wakefulness to sleep in patients with overlap syndrome (0.35±0.21 vs 0.28±0.19, p<0.05) and patients with OSA (0.56±0.24 vs 0.28±0.13, p<0.01) (see table 2 and online supplementary tables E-3 and E-4, and supplementary figure E-1).

In patients with overlap syndrome the V̇E/EMGdi decreased from the state of wakefulness, to sleep in which snoring was recorded (0.35±0.21 vs 0.24±0.14, p<0.001) and further decreased during stage 2 hypopnoea events (0.15±0.12, p<0.001) (table 3; data for stage 2 sleep with or without snoring are shown in online supplementary table E-4).

### DISCUSSION

In the present study we show that unlike healthy subjects who exhibited a small (approximately 10%) difference between NREM sleep and wakefulness, V̇E was lower in stage 2 sleep than wakefulness in patients with COPD alone and those with overlap syndrome or OSA. In patients with overlap syndrome neural respiratory drive was similar between NREM sleep and wakefulness. However, neural respiratory drive from wakefulness to stage 2 sleep increased in patients with OSA but decreased in those with COPD alone, suggesting that mild or moderate OSA can partially compensate for reduction of neural respiratory drive inherent to COPD and certainly does not seem to worsen hypoventilation associated with COPD.

### Methodological issues

This is the first study to simultaneously record ventilation and neural respiratory drive from patients with COPD alone and patients with an overlap syndrome. Some studies have previously assessed ventilation during sleep using respiratory inductance plethysmography,18–19 which can be inaccurate particularly in patients with COPD.18–19 Becker et al18 used a pneumotachograph to measure ventilation during sleep in patients with COPD, but the pneumotachograph in their study was connected with a nasal mask rather than a full-face mask, leading to a potential underestimate of ventilation if patients breathed through the mouth. Using a full face mask to quantify ventilation we found a larger reduction of ventilation in NREM than that reported by Becker et al18 using a nasal mask.

The FEVi in patients with overlap syndrome was numerically, although not statistically, higher than that in patients with COPD alone. Because obesity is an important factor contributing to development of OSA and because COPD is a chronic wasting disease sometimes associated with weight loss as disease progresses, it could be argued that patients with severe COPD

### Table 1 Basic information for all subjects

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>COPD (n=19)</th>
<th>Overlap (n=16)</th>
<th>Normal (n=12)</th>
<th>OSA (n=14)</th>
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<tr>
<td>Age (years)</td>
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<td>61.5±10.2</td>
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<td>BMI (kg/m²)</td>
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<td>23.5±3.8</td>
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<td>FVC (%pred)</td>
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<td>68.5±17.4</td>
<td>98.6±9.6</td>
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<tr>
<td>FEV₁ (%pred)</td>
<td>38.5±16.3</td>
<td>47.5±15.2</td>
<td>98.1±9.2</td>
<td>99.1±14.6</td>
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<tr>
<td>FEV₁/FVC%</td>
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<td>54.6±11.4</td>
<td>79.6±3.4</td>
<td>80.4±6.3</td>
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<td>mMRC</td>
<td>1.8±0.9</td>
<td>1.6±0.5</td>
<td>0±0</td>
<td>0±0</td>
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<tr>
<td>EMGdi, max (μV)</td>
<td>176.9±72.1</td>
<td>192.3±56.6</td>
<td>156.1±50.3</td>
<td>166.2±40.8</td>
</tr>
<tr>
<td>EMGdi, min (μV)</td>
<td>29.5±13.3</td>
<td>29.3±15.8</td>
<td>12.6±3.7</td>
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<td>AHI (events/hour)</td>
<td>1.9±1.6</td>
<td>20.5±14.1</td>
<td>1.6±1.4</td>
<td>25.6±18</td>
</tr>
</tbody>
</table>

AHI, Apnoea Hypopnoea Index; BMI, body mass index; EMGdi,max, the maximum of the diaphragm electromyogram; mean SaO2, mean nocturnal oxygen saturation; mini SaO2, minimum nocturnal oxygen saturation; mMRC, modified British Medical Research Council; TST90%, time spent with saturation below 90%; TST, total sleep time.

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are less likely to develop OSA because of weight loss. However, the current dataset (by showing patients with COPD with polysonographically proven OSA) shows that this conjecture is insufficient to prevent OSA in patients with COPD.  

However, we note that while there was no statistically significant difference in FEV1 between patients with COPD alone and those with overlap syndrome, type II error cannot be absolutely excluded given the small numbers dictated by a physiological study of this nature. We would also point out that the similarity in mMRC score and EMGdi at rest between patients with COPD alone and those with overlap syndrome also suggests the severity of airway obstruction between the two groups is similar. In the present study, the prevalence of overlap syndrome (46%) seems to be high for a COPD cohort study, in particular for those with low BMI. However, a high prevalence of overlap syndrome (46%) in the COPD cohort study, despite the low BMI, may thus be attributable to Asiatic craniofacial morphology.

Oesophageal pressure combined with airflow recordings has recently been used to quantify upper airway resistance. However, oesophageal pressure is influenced by lung volume and airflow and thus has a potential limitation in the assessment of upper airway resistance in patients with OSA which is characterised by changes in lung volume and airflow. Classically a catheter positioned in the pharynx has been used to measure upper airway resistance by measurement of pharyngeal pressure. However, upper airway resistance derived from measurement of pharyngeal pressure is a difficult technique and may be variable during both wakefulness and sleep. Besides providing data regarding neural respiratory drive, EMGdi has an advantage over oesophageal pressure in the assessment of upper airway resistance because it is independent of change in airflow and lung volume. Normally, VT is achieved in response to

![Table 2 Diaphragm EMG, ventilation during wakefulness and NREM sleep](image)

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>COPD (n=19)</th>
<th>Overlap (n=16)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Wakefulness</td>
<td>NREM</td>
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<tr>
<td>EMGδmax%max</td>
<td>29.5±13.3</td>
<td>23.0±8.9</td>
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<tr>
<td>Vt (L)</td>
<td>8.6±2.0</td>
<td>6.5±1.5</td>
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<tr>
<td>Vt/EMGδmax</td>
<td>0.47±0.11</td>
<td>0.37±0.09</td>
</tr>
<tr>
<td>ETCO2 (%)</td>
<td>4.2±0.5</td>
<td>4.5±0.6</td>
</tr>
<tr>
<td>RR</td>
<td>18.4±4.1</td>
<td>17.8±2.1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Normal (n=12)</th>
<th>OSA (n=14)</th>
</tr>
</thead>
<tbody>
<tr>
<td>EMGδmax%max</td>
<td>12.6±3.7</td>
<td>12.2±3.3</td>
</tr>
<tr>
<td>Vt (L)</td>
<td>8.3±1.6</td>
<td>7.5±1.3</td>
</tr>
<tr>
<td>Vt/EMGδmax</td>
<td>0.50±0.09</td>
<td>0.47±0.10</td>
</tr>
<tr>
<td>ETCO2 (%)</td>
<td>4.8±0.3</td>
<td>5.0±0.3</td>
</tr>
<tr>
<td>RR</td>
<td>16.7±2.3</td>
<td>15.6±2.3</td>
</tr>
</tbody>
</table>

Δ%, percentage change compared with wakefulness; EMGδmax%, percentage of maximal EMGδ. ETCO2, end-tidal CO2; NREM, non-rapid eye movement; p value, comparison of wakefulness and NREM; RR, respiratory rate; VE, minute ventilation; VT, tidal volume; VT/EMGδ, the ratio of tidal volume to peak root mean square of EMGδ.

![Figure 1 Diaphragm electromyogram (EMG) recording from five pairs of oesophageal electrodes and airflow from pneumotachography during polysomnography in patients with COPD alone, patients with overlap syndrome, normal subjects and patients with obstructive sleep apnoea (OSA).](image)
neural respiratory drive reflected by EMGdi after overcoming total respiratory resistance, including elastic recoil of the lung, lower airway resistance and upper airway resistance. VT/EMGdi reflects changes in upper airway resistance if lung mechanics and lower airway resistance remain the same; one limitation of our study which should therefore be acknowledged is that lung volumes were not measured during sleep. OSA is characterised by an increase in upper airway resistance from wakefulness to snoring and a further increase to hypopnoea. In line with this concept, VT/EMGdi decreases significantly from wakefulness to snoring and further decreases to hypopnoea in patients with overlap syndrome, which supports using change in VT/EMGdi to assess changes in upper airway resistance.

**Figure 2** Electromyogram (EMGdi)% (left panel), ventilation (middle panel) and the VT/EMGdi (right panel) in patients with COPD alone, overlap syndrome, normal subjects and patients with obstructive sleep apnoea (OSA). Ventilation decreases from wakefulness to sleep in all four groups but only in patients with COPD, overlap syndrome and OSA is the reduction statistically significant. EMGdi decreases in patients with COPD alone but increases in patients with OSA from wakefulness to non-rapid eye movement (NREM) sleep, whereas it remains the same in normal subjects and patients with overlap syndrome. VT/EMGdi decreases from wakefulness to sleep in patients with overlap syndrome and those with OSA but it changes little in normal subjects and patients with COPD alone. The decrease in ventilation is associated with a reduction of EMGdi in patients with COPD alone whereas ventilation reduction is associated with decreased VT/EMGdi in patients with overlap syndrome and those with OSA.

**Significance of findings**

This study shows that neural drive decreases from the state of wakefulness to sleep in patients with COPD alone, as previously reported. It may be expected that neural respiratory drive in patients with overlap syndrome would also decrease from wakefulness to sleep, but we found that this is not the case, consistent with our prior report that obstructive respiratory events are associated with increased neural respiratory drive. Several studies suggest that the increased neural respiratory drive during sleep is directly related to the presence of increased upper airway resistance since neural respiratory drive decreases when airway resistance is offset by, for example, treatment with continuous positive airway pressure or inhalation of Heliox (76%...
He(24%O$_2$)$_2$ The present study suggests that sleep-related reduction of neural respiratory drive characteristic of COPD could be offset by an increase in upper airway resistance as a consequence of coexistent OSA, in an adaptation potentially protective in nature.

It has been hypothesised that desaturation in patients with overlap syndrome would be more severe than that in patients with COPD alone because COPD and OSA can cause desaturation. However, this view is mainly derived from studies of patients who had predominately mild or moderate COPD or who were recruited from patients with OSA and obesity, and thus may not represent a clinical cohort of patients with severe COPD. However, a recent cohort study of non-obese patients with severe COPD showed that the number of patients with COPD alone who required oxygen supplementation was similar to that in those with overlap syndrome, suggesting the prevalence of oxygen desaturation is similar between patients with COPD alone and patients with overlap syndrome. In the present study we found that mean oxygen saturation and minimal oxygen saturation during overnight sleep were similar in patients with or without overlap syndrome. Moreover, although patients with coexistent OSA and severe COPD usually have brief periods of desaturation, prolonged desaturation (SaO$_2$<90 for longer than 5 min) occurred more often in patients with severe COPD alone than those with overlap syndrome (see online supplementary table E-2). This finding may be clinically significant for the management of patients with overlap syndrome. If sleep-related hypoventilation or desaturation in patients with severe COPD had been worsened by coexistent OSA, one would have to be cautious with nutritional supplements which may raise BMI in patients with COPD. In COPD, the present study suggests that when patients with severe COPD develop mild or moderate OSA, a sleep-related reduction of neural respiratory drive as a consequence of COPD would be relieved because of increased upper airway resistance, preventing ventilation from further decreasing. The observation that 16% of patients with COPD alone developed significant oxygen desaturation but none of patients with an overlap syndrome developed a significant desaturation may give a hint that if patients with severe COPD develop mild or moderate OSA, oxygen desaturation would not necessarily be worsen. This interesting finding is in line with the recent report that the clinical outcome in end-stage patients with overlap syndrome is better than those with COPD alone. Nevertheless, we note that pathophysiological change in patients with overlap syndrome of severe COPD and mild OSA might differ from those with mild COPD and severe OSA; in addition variation on this spectrum may differ between western patients in whom OSA is more driven by obesity and Asian patients (as studied here) in whom craniofacial morphometry may be more relevant.

In conclusion the mechanism underlying the reduction of ventilation at stage 2 sleep in patients with COPD alone differs from that in patients with overlap syndrome. Ventilation reduction in patients with COPD alone is mainly because of a decrease in neural respiratory drive whereas it is mainly a result of an increase in upper airway resistance in patients with overlap syndrome.

Contributors Conception and design: BH, GL, SX, JS, JM, MP, YL; analysis and interpretation: BH, GL, SX, RC; drafting the manuscript for important intellectual content: BH, GL, RC, JS, JM, MP, YL.

Funding The work was supported by National Natural Science Foundation of China (NSFC No. 81120108001 and 81270143). Professor Polkey’s contribution to this project was supported by the NIHR Respiratory Biomedical Research Unit at the Royal Brompton and Harefield NHS Foundation Trust and Imperial College, London UK, who part funded his salary.

Competing interests None declared.

Patient consent Obtained.

Ethics approval The Ethics Committee of the First Affiliated Hospital of Guangzhou Medical University.

Provenance and peer review Not commissioned; externally peer reviewed.

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Thorax published online November 2, 2016

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