Sleep Disordered Breathing in the Lung Transplant Population, a Case Report and Literature Review

Abstract

Background: Patients waiting for and having received lung transplantation commonly demonstrate known risk factors for sleep disordered breathing (SDB) including central sleep apnea (CSA). However, little is known about the prevalence or clinical implications of SDB in this patient population.

Methods: We report a 61 year old man who underwent uncomplicated bilateral lung transplantation. Approximately 5 weeks post-transplant, the patient developed fatigue, somnolence, and mild intermittent confusion. Extensive cardiologic and neurologic evaluations were unrevealing regarding cause of symptoms. Diagnostic polysomnogram revealed severe central sleep apnea with a periodic respiratory pattern.

Results: The patient was initiated on positive airway pressure therapy during sleep with an adaptive servoventilation device which successfully treated central sleep apnea and improved daytime symptoms. A repeat diagnostic polysomnogram performed 8 months post-transplant showed resolution of central sleep apnea.

Conclusion: SDB including CSA is a poorly understood occurrence in the lung transplantation patient population that warrants clinical attention and further investigation.

Case Report

A 61 year old man underwent bilateral lung transplantation for respiratory failure due to severe chronic obstructive pulmonary disease (COPD). The surgical course was without significant complication and the patient was extubated without difficulty. The immediate post-operative period was significant for an isolated episode of atrial fibrillation with rapid ventricular response requiring cardioversion. Standard immunosuppression and rehabilitation were initiated per protocol, and the patient was discharged to home approximately 3 weeks post-transplant.

5 weeks post-transplant, the patient noted progressive fatigue and daytime sleepiness as well as intermittent mild confusion and a single near-syncopeal event. The patient denied anginal chest discomfort, signs of heart failure, or focal neurologic findings. On further questioning, the patient reported frequent nocturnal awakenings, non-restorative sleep, daytime somnolence and his care-taker noted the patient to have respiratory pauses in his sleep which occurred in a cyclical pattern.

On physical examination the patient was afebrile, normotensive, had a room-air saturation of 97%, was neurologically intact, and demonstrated no findings of heart failure. Serologic laboratory evaluation was without evidence of cardiac ischemia, renal or other organ failure. Electrocardiogram and telemetry demonstrated sinus rhythm without ischemia, ectopy or arrhythmia. Transthoracic echocardiogram revealed normal bilateral ventricular size and function with estimated pulmonary arterial systolic pressure of 30 mmHg. CT scan of the chest showed normal post-surgical changes with no evidence of pulmonary emboli or parenchymal lung abnormalities. Neurologic evaluation including EEG as well as brain CT and MRI were without significant finding.

A laboratory based diagnostic polysomnogram (PSG) was performed approximately six weeks post-transplant, and showed an apnea hypopnea index (AHI) of 56 consisting of predominantly central apnea events. The central apneas occurred exclusively in supine non-REM sleep and were characterized by a periodic occurrence with a respiratory pattern which was at times crescendo-decrescendo in waveform and had a cycle time of 35-40 seconds (Figure 1). The study was performed on room air and 100% of the patient’s sleep time was spent with oxygen saturations greater than 90%.
A laboratory based titration PSG was subsequently performed with persistence of central apnea events and periodic breathing with continuous positive airway pressure (CPAP) titration. Bi-level with adaptive servoventilation technology was then initiated, with an immediate resolution of central events (Figure 2). The patient was subsequently initiated on adaptive servo ventilation therapy, and noted a significant improvement in his fatigue, somnolence and confusion.

8 months post-transplant, a repeat laboratory based diagnostic PSG revealed no evidence of central sleep apnea or periodic breathing and only mild obstructive SDB with AHI of 7 and no significant hypoxia. The patient was subsequently switched to standard CPAP therapy with good clinical response.

Discussion

Lung transplantation is a validated treatment for advanced COPD. Commonly recognized post-transplant occurrences include acute rejection, infection, bronchiolitis obliterans syndrome (BOS), gastro esophageal reflux disease (GERD), hypertension, and depression [1,2]. Less understood is the prevalence, clinical presentation of, risk factors for, and optimal treatment strategies of SDB in this population. SDB encompasses a group of diagnoses including CSA, obstructive sleep apnea (OSA), sleep related hypoventilation, and sleep related hypoxemia [3]. Risk factors for SDB in the general population include obesity, large neck circumference, male gender, age, cardiologic disorders, and underlying lung disease [4,5]. Additional incurred risk factors for SDB in lung transplantation patients include acute weight changes, thoracic volume and pressure changes, altered chemoresponsiveness, lung denervation, and use of narcotic as well as mineralocorticoid altering medications.

CSA is defined by the cessation of airflow without concurrent respiratory effort, and may clinically present in a broad range of respiratory patterns [6]. This diversity is reflective of the complex physiologic regulation, which includes contributions from chemo-receptor responsiveness, circulatory response times, and ventilatory capacity. An established mode of classification for central apnea is based on the concept of loop gain, defined as the response to a disturbance (ventilatory response) in ratio to the disturbance itself (apnea). Components of the overall loop gain of the system include; controller gain (chemo-responsiveness),
plant gain (ventilatory efficiency), and mixing gain (including 
circulatory times and gas delivery) [7]. High loop gain models such
as Cheyne-Stokes and other periodic respiratory patterns have a
ratio of ≥ 1, resulting in an easily destabilized ventilatory system.

High loop gain CSA models are common in patients with congestive
heart failure, atrial fibrillation, cerebral vascular accident, and
renal failure [8], a specific cause for the occurrence of central
apnea in our patient is not clearly established. Our patient had
no history of SDB prior to lung transplantation, and at the time of
presentation demonstrated no clinical evidence of a neurologic
or cardiogenic process. Additionally, while daytime fatigue and
sleepiness can be associated with central apnea, an association
with our patient’s confusion and near-syncopal episode is less
supported. That said, it is feasible that in the setting of recent
transplantation, our patient developed a high-gain system due to
a large increase in plant gain (transplanted lungs) with additional
increases in controller and mixing gain which subsequently
equilibrated over time.

Although it is not unexpected that patients with respiratory
failure awaiting lung transplantation would have a high
prevalence of SDB, there is surprisingly little data from quality
studies to guide medical decision-making. In a prospective study,
Malouf et al. performed room air PSG on 80 patients awaiting
lung transplantation. 32 of 80 (40%) patients met SDB criteria
defined as a respiratory disturbance index (RDI) ≥ 10 (central and
obstructive events were not distinguished), or ≥ 10% of sleep
time with oxygen saturations ≤ 90% [9]. A retrospective review
of 60 patients with end-stage lung disease undergoing evaluation
for lung transplantation demonstrated 40 of 60 (67%) to have
OSA (defined by RDI ≥ 5, with no data on central apnea events
provided) with the majority having moderate-to-severe OSA.
While there was no association between underlying diagnosis
and the presence of OSA, associations were found between AHI
severity and higher DLCO% as well as lower nocturnal oxygen
saturations [10].

There is a similar paucity of data regarding the occurrence of SDB
after lung transplantation. In the previously mentioned study,
Malouf et al. completed post-transplant PSG on 25 patients
including 11 who had met SDB criteria prior to transplantation.
SDB resolved in 6 of the 11 (55%) patients with prior SDB, but
new SDB developed in 4 of 14 (29%) patients without prior SDB
findings. The presence of SDB did not impact pre or post-
transplant survival. A single center cross sectional study of 24
patients at least one year post-transplant showed 15 of 24 (63%)
to have SDB (defined as AHI ≥ 10), with 25% demonstrating
predominant CSA. Patients who developed SDB were shown
to have elevated systolic blood pressure, greater age, greater
change in BMI, more recent date of transplantation, diagnosis
of emphysema, observed snoring, and an association between
Cyclosporine use and central apnea was suggested (P=0.006)
[11]. Our patient was not taking Cyclosporine, and further, its
use has not been associated with central apnea in other solid-
organ transplant populations [12,13]. Lastly, a prospective
study evaluating risk factors for BOS performed PSG in 14 post-
transplant patients and found 100% to meet SDB criteria (defined
as AHI>5) with 3 patients demonstrating mild central apnea
activity. No associations between SDB and esophageal reflux or
SDB and BOS were found.

Conclusion
Our case report and literature review highlight the uncertainties
to the frequency as well as spectrum of clinical presentations by
which SDB and in particular CSA may present in patients prior
to and after lung transplantation. Despite growing data showing
that SDB is common in this patient population, there are no
established screening guidelines or treatment recommendations.
Thus, data from additional longitudinal studies are needed to
identify patient risk factors as well as to clarify the impact of
SDB on unique clinical consequences in these patients such as
health related quality of life, depression, GERD, hypertension
and bronchiolitis obliterans syndrome. In the interim, providers
treating this patient population should have a low threshold to
screen for SDB and should treat accordingly.
References

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