REVIEW

New Metrics from Polysomnography: Precision Medicine for OSA Interventions

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Abstract: Obstructive sleep apnea (OSA) is a highly preventable disease accompanied by multiple comorbid conditions. Despite the well-established cardiovascular and neurocognitive sequelae with OSA, the optimal metric for assessing the OSA severity and response to therapy remains controversial. Although overnight polysomnography (PSG) is the golden standard for OSA diagnosis, the abundant information is not fully exploited. With the development of deep learning and the era of big data, new metrics derived from PSG have been validated in some OSA consequences and personalized treatment. In this review, these metrics are introduced based on the pathophysiological mechanisms of OSA and new technologies. Emphasis is laid on the advantages and the prognostic value against apnea-hypopnea index. New classification criteria should be established based on these metrics and other clinical characters for precision medicine.

Keywords: obstructive sleep apnea, precision medicine, polysomnography, metrics

Introduction

Obstructive sleep apnea (OSA) is a chronic disease affecting almost one billion people around the world.¹ It is characterized by partial or complete upper airway collapse during sleep, leading to sleep fragmentation and intermittent hypoxemia. The consequences brought by OSA include disturbance of sleep structure, daytime sleepiness, various complications and decreased quality of life. Studies have revealed that the economic burden of OSA is substantial.²

The metric used for evaluating OSA severity is the apnea-hypopnea index (AHI), which mainly describes the frequency of respiratory events. Despite its convenience and wide acceptance, several defects remain which oversimplify the complexity of OSA and lead to the poor correlation with OSA consequences. First, the definition of AHI treats apnea and hypopnea equivalently. The severity of airway obstruction is associated with degree of desaturation and arousals. Studies have demonstrated that OSA patients with different respiratory patterns presented different disease severity.³ The predominance of apnea or hypopnea could be a potential subtype correlated with certain OSA comorbidity. Second, the different rules for hypopnea definition in clinical practice may bring inconsistencies to the diagnosis and severity evaluation of OSA, leaving some asymptomatic OSA patients untreated.⁴ The association of OSA with comorbidities using different criteria could be discrepant.⁵ Another nonnegligible issue is that AHI does not directly reflect the pathogenesis of OSA. Other auxiliary metrics like lowest pulse oxygen saturation (LSpO₂), percentage time below 90% oxygen saturation (T90) and oxygen desaturation index (ODI) evaluate event-related desaturation from a single aspect but have not been recommended as the diagnostic criteria. Degree, duration of desaturation, respiratory event duration, and intensity of arousals could all represent certain aspects of OSA pathophysiology, yet none of them are integrated in AHI. These defects could be the reasons for poor relationship between OSA and relevant comorbidities.^{6,7} Furthermore, the adherence of continuous positive airway pressure (CPAP), the primary therapy, is relatively low. In some cases, CPAP showed an insignificant effect on OSA comorbidities while CPAP tends to be the one-size-fits-all therapy for OSA.^{8,9}

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Figure I Summary of PSG physiological signals, new metrics and utilization of these metrics towards precision medicine. Abbreviations: ECG, electrocardiogram; EEG, electroencephalogram; OSA, obstructive sleep apnea; PSG, polysomnography.

Since the term of polysomnography (PSG) was first created in 1974,¹⁰ it has been applied as the golden standard diagnostic examination for OSA. A typical overnight PSG examination contains plenty of biological signals such as electroencephalogram (EEG), electrooculogram, nasal airflow, electromyogram, electrocardiogram (ECG), respiratory effort and blood oxygen. These simultaneously recorded signals can provide a great deal of information on patients' sleeping, respiratory and cardiovascular status. Apart from the original signals, data generated by advanced technologies like machine learning can further identify certain subgroups of patients who may benefit from personalized treatment. Moreover, efforts have been made to establish large data sets for researchers. Online resources like the National Sleep Research Resource¹¹ and PhysioNet¹² have thousands of PSG recordings which offer the opportunity for data training and algorithm validation. While automated signal processing algorithms have been proposed for decades and some were verified by large samples, these algorithms have not been widely accepted or commercialized for clinical practice. The reality urges reformation in clinical evaluation and treatment of OSA.

This paper aims to review recent advances in PSG analysis. In particular, the prognostic value of novel metrics towards OSA comorbidities and personalized treatment is discussed (Figure 1). The review will be organized in the following way: (1) the pathophysiological association of OSA and each comorbidity; (2) the association of these comorbidities with new metrics that reflect certain pathophysiological mechanisms; (3) new metrics validated for personalized treatment; (4) limitations and future directions of these metrics for precision medicine. Importance is laid on the role of OSA pathophysiological mechanisms for discovering clinically oriented PSG metrics.

Pathogenesis of OSA with Related Comorbidities

Cardiovascular Disease

Many clinic-based case control and longitudinal studies have demonstrated that OSA patients have a two- to fivefold increase in cardiovascular disease (CVD) risk after multivariate adjustment including body mass index.¹³ Although exact causal association is still unclear, both animal and human studies suggest that sympathetic neurohormonal activation, oxidative stress and systemic inflammation induced by sleep fragmentation and intermittent hypoxemia, which is similar to reperfusion injury, may play an important pathophysiological role.¹⁴ These molecular level changes lead to dysfunction of vascular endothelium, atherosclerosis and hypercoagulability, which

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are the fundamental reasons for hypertension, coronary artery disease and cerebrovascular disease. Apart from these common mechanisms, alteration in intrathoracic pressure also brings adverse outcomes to certain disease. For example, in atrial fibrillation, recurrent and abrupt negative changes in intrathoracic pressure during apnea may lead to structural and functional atrial remodeling and electrophysiological alterations. The same changes in intrathoracic pressure may also increase left ventricular transmural pressure and related afterload, causing reduced stroke volume and coronary blood supply. These mechanisms may precipitate myocardial ischemia and remodeling and lead to heart failure.

Hyperglycemia and Type 2 Diabetes

Cross-sectional and retrospective studies proved bilateral relationship between OSA and type 2 diabetes.¹⁵ Like cardiovascular diseases, intermittent hypoxemia and sleep fragmentation are also the cardinal pathways towards dysregulation of glucose metabolism and type 2 diabetes. Besides sympathetic activation, oxidative stress and systemic inflammation, these pathophysiological mechanisms also result in activation of hypothalamic–pituitary–adrenal axis and induction of adipocytokines. Such changes further precipitate attenuation of glucose-induced insulin secretion from pancreatic β cells and insulin resistance in peripheral tissues and organs.^{16,17}

Cognitive Impairment and Alzheimer's Disease

Considering the importance of sleep continuity, slow wave sleep and sleep spindles in neurogenesis, alertness, memory formation and consolidation, sleep fragmentation caused by repeated micro-arousals may continuously influence the sleep structure. Studies have found that long term OSA alters white matter integrity and aggregation of β -amyloid and tau pathology. Related pathophysiological changes include inflammation, oxidative stress, cerebral edema and endothelial dysfunction.¹⁸ Moreover, negative intrathoracic pressure caused by OSA could influence hemodynamics and glymphatic system. The latter has been recognized as the key pathway for brain fluid clearance and waste removal via glia-supported perivascular channels which is more active during sleep.¹⁹

Taken together, the primary mechanisms of OSA towards its related comorbidities include intermittent hypoxemia, sleep fragmentation and intrathoracic pressure oscillation. The corresponding changes such as oxidative stress, inflammation and sympathetic neurohormonal activation further cause alterations of the micro-environment and damage of target organs. Metrics representing these pathophysiological mechanisms could, in theory, have more prognostic value than AHI. Given the advantages of PSG mentioned above, it is time for clinicians and manufacturers to evaluate and promote proper PSG metrics for better clinical practice. New metrics that have been validated in clinical utility of OSA comorbidity and treatment are introduced in the next section.

Metrics for OSA Comorbidities

Arousal Intensity

Arousal index is routinely used to evaluate arousals in sleep. However, like AHI, the arousal index only captures frequency of arousals during sleep. The extent of each arousal varies a lot during a respiratory event which could lead to different degrees of autonomic activation. Azarbarzin et al created an automatic algorithm to assess the extent of arousal. Wavelet transform analysis was used to capture EEG signals' time and frequency characteristics. A scale from zero (barely perceptible) to nine (most intense), named arousal intensity, was applied during the evaluation.²⁰ Despite its variability among subjects, arousal intensity has been proved to be reproducible²¹ and heritable.²² In a sample of 20 PSGs from patients receiving sleep monitoring, they found a strong relationship between arousal intensity and the heart rate change associated with respiratory events (average r: 0.95 ± 0.04). The latter was proved to be associated with an increased risk of both fatal (HR 1.68, 95% CI 1.22–2.30) and non-fatal (HR 1.60, 95% CI 1.28–2.00) cardiovascular disease events in the Sleep Heart Health Study (SHHS).²³ However, arousal intensity may not be the only factor which affects events related heart rate change.²⁴ No studies have been performed to evaluate the association of arousal intensity directly with OSA comorbid conditions.

Odds Ratio Product

Conventionally, sleep stages are scored based on recommended criteria in each 30s epoch. This criterion has not been changed in the digital era. In fact, sleep state changes continuously rather than an intermittent 30s epoch. Details about variation of EEG signals indicating sleep depth or sleep quality could be neglected within sleep stages. An automated algorithm for estimating sleep depth was developed by Younes et al.²⁵ Fast Fourier transform was used for power spectrum analysis in each 3s EEG consecutive epochs. Odds ratio product (ORP) was calculated with a range from 0 to 2.5. In the validation test, an ORP less than 1.0 predicted sleep and an ORP larger than 2.0 predicted wakefulness in more than 95% of 30s epochs. This high consistency indicated an adequate measure of sleep depth. A recent study has found that the correlation between the right and left hemisphere ORP measures (interhemispheric sleep depth coherence) predicted occurrence of car accidents 2 years after the sleep study in patients with OSA. Higher sleep depth coherence was associated with lower risk of accidents after multivariate adjustments.²⁶ ORP could be a measure of susceptibility to adverse neurocognitive outcomes. However, no significance was observed between ORP and hypertension in the Multi-Ethnic Study of Atherosclerosis.²⁷ Associations with other metabolic outcomes remain untested.

The same EEG processing method was used by Mullins et al, EEG power spectra was obtained using fast Fourier transform. EEG slowing (a ratio of slow frequencies to fast frequencies) and sleep spindle density (spindle events per minute) were calculated. They found greater EEG slowing during REM and lower sleep spindle density during NREM were associated with worse psychomotor vigilance and simulated driving performance in OSA patients.²⁸ However, the results need to be interpreted cautiously due to the small sample of 8 patients and limited statistical power. Confirmation in a larger sample is required.

Hypoxic Burden and Related Metrics

Since AHI cannot fully represent event related desaturation and other traditional metrics only capture certain aspects of hypoxemia, metrics with combined characteristics have been proposed in recent years. The most validated metric is hypoxic burden, which encapsulates the frequency, duration, and depth of the respiratory event related desaturation. The total hypoxic burden is defined as the sum of individual burdens divided by total sleep time. The prognostic value of hypoxic burden has been proved in many studies. Azarbarzin et al reported significant association between hypoxic burden and cardiovascular disease related mortality in the Osteoporotic Fractures in Men Study (MrOS) and the SHHS.²⁹ Patients with the highest hypoxic burden had increased risk of mortality after adjusting multiple factors including AHI, T90 and LSpO₂ (HR 2.73, 95% CI 1.71–4.36 in MrOS and HR 1.96, 95% CI 1.11–3.43 in SHHS). In contrast, AHI could not predict mortality risk in the MrOS. Similar results were found when considering the risk of incident heart failure. Increased hypoxic burden per 1 SD were associated with incident heart failure (HR 1.22, 95% CI 1.02–1.45 in MrOS and HR 1.18, 95% CI 1.02–1.37 in SHHS), while AHI was insignificant.³⁰ Other positive findings including hypertension,²⁷ stroke³¹ and chronic kidney disease.³² These significant results suggest that hypoxic burden is an important metric for OSA evaluation.

Similar to hypoxic burden, other studies combined information of event related desaturation in different ways and also found significant associations with OSA comorbidities. Cao et al invented a new metric named sleep breathing impairment index (SBII), which was the product of area under desaturation curve and respiratory event duration. In a cross-sectional study with 140 male OSA patients, higher SBII was associated with moderate-to-high Framingham CVD risk.³³ The incorporation of respiratory event duration may have advantages over hypoxic burden. Wang et al evaluated prognostic value of the change in the percentage of SpO₂ per second after obstructive apnea, expressed as oxygen desaturation rate (ODR). Participants with faster ODR demonstrated a stronger association with the elevation of both awake and sleep blood pressure (BP) levels and short-term BP variability.³⁴ ODR might be a marker for sympathetic activation.

Photoplethysmography and Pulse Wave Amplitude Drops

Photoplethysmography (PPG) is another signal derived from pulse oximetry, which assesses the blood volume changes in the microcirculation of the fingertip. Various conditions like cardiovascular disease, hypertension, diabetes and mental

health were proved to be effectively assessed by PPG in previous studies.³⁵ However, the relatively low accuracy with raw signals and inadequate validation limited its clinical application.

Pulse wave amplitude (PWA) drops are derived from the variance of PPG signal amplitude. PWA is calculated as the difference between the peak and nadir values of PPG waveform. PWA drops with an amplitude >30% compared to baseline and a duration >4 heartbeats are considered clinically significant. A former study showed that PWA drops reflect peripheral vasoconstriction resulting from sympathetic activation.³⁶ Respiratory events related PWA drops were associated with cortical activity and could serve as a marker of cerebral response to respiratory events.³⁷ Hirotsu et al found that certain PWA drop features (lower PWA-drop index, longer duration and greater area under the curve) increased odds of hypertension, diabetes, or CV event after multivariate adjustment. An independent association of PWA drops index with AHI and total arousal index was also found in this study.³⁸

Pulse Arrival Time

Apart from creating new metrics with single signals, metrics generated from multi-signals may contain more pathophysiological information. Pulse arrival time (PAT) is such a composite metric acquired from ECG and oximeter. It is a widely accepted surrogate of pulse transit time, which correlates well with systolic BP and has been used in monitoring BP during sleep.³⁹ PAT is defined as the time interval between the EEG R wave and pulse arrival by PPG. Kwon et al evaluated prognostic value of PAT response to obstructive respiratory events, which was defined as the area under the PAT waveform following respiratory events. They found that PAT response was associated with key subclinical CVD markers (left ventricle mass, carotid plaque burden and coronary artery calcium) as well as with incident CVD (adjusted HR 1.18, 95% CI 0.99–1.40).⁴⁰ These findings suggested average PAT response to respiratory events, a reflection of dynamic sympathetic activity, could be valuable for risk stratification of patients with sleep apnea.

Conventional Metrics with New Clinical Significance

Apart from the new metrics mentioned above, other traditional metrics, once neglected, were also evaluated in recent years. The prognostic value of fast sleep spindles (13–15 Hz) on Alzheimer's disease (AD) and Mild Cognitive Impairment (MCI) was evaluated by Gorgoni et al. A significant parietal fast spindle density decrease was found in AD and MCI patients, positively correlated with Mini-mental State Examination scores (r = 0.33, p = 0.03).⁴¹ The results showed that spindle density changes are specific for frequency and location in MCI and AD. Certain spindle changes are related to the severity of cognitive impairment and early onset of MCI. Butler et al tested whether apnea-hypopnea event duration predicted all-cause mortality in SHHS. They found that shorter respiratory event duration was associated with all-cause mortality (adjusted HR 1.31, 95% CI 1.11–1.54).⁴² While it seems contradictory with the idea that longer event duration associates with heavier hypoxemia, shorter event duration may be a marker for low arousal threshold and autonomic nervous augmentation. Baumert et al measured mean nocturnal respiratory rate during sleep in the MrOS sleep study and the Study of Osteoporotic Fractures (SOF). After adjusting multiple covariates, patients with respiratory rate \geq 16 breaths per minute were independently associated with CVD mortality (HR 1.57, 95% CI 1.14–2.15 in MrOS and HR 2.58, 95% CI 1.41–4.76 in SOF) and all-cause mortality (HR 1.18, 95% CI 1.04–1.32 in MrOS and HR 1.50, 95% CI 1.02–2.20 in SOF).⁴³ Elevated respiratory rate may reflect dysfunctional central respiratory control and lead to hypocapnia related electrolyte abnormalities.

Cluster Analysis

Since OSA is recognized as a heterogeneous disease, efforts have been made to discriminate different subtypes for clinical practice. A widely applied method for OSA is cluster analysis. It is an unsupervised machine learning method with potential to find undiscovered patterns of diseases. Zinchuk et al used principal components-based cluster analysis to identify polysomnographic features. Based on OSA pathophysiological domains, seven clusters were identified in the US Veteran cohort. They further evaluated prognostic value of these PSG based subtypes. Patients labeled as certain OSA subtypes had significant risk of CVD outcomes and incident diabetes. The addition of these subtypes further improved diabetes risk prediction. Meanwhile, categorized AHI was not associated with increased CVD risk.^{44,45} These subtypes may represent certain pathophysiological characteristics including hypoxemia and sympathetic activation and were

superior in predicting OSA comorbidities comparing to AHI. The utility of unsupervised learning method with abundant PSG information could serve as an important clinical guidance towards OSA evaluation.

Endotypes for Personalized OSA Treatment

Despite the first line treatment recommended by the American Academy of Sleep Medicine, the effect and adherence of CPAP vary a lot in OSA patients. Other treatment options have been proved to have a modest improvement. This situation urges the need for personalized treatment. Endotypes based on OSA pathophysiology (PALM scale) were proposed, aiming at identifying subgroups of patients responding to specific OSA treatments.⁴⁶ However, this classification required invasive examinations like epiglottic pressure catheter which limited the clinical application. Novel approaches using PSG signals to determine OSA pathophysiology have been developed. Terrill et al quantified the loop gain, a part of ventilatory control, using PSG signals in 28 OSA patients. The method was compared with the standard procedure (CPAP drop method) and showed good agreement (r = 0.63, p < 0.001). It further predicted reduction in loop gain with oxygen and acetazolamide therapy.⁴⁷ Similar to loop gain, other OSA endotypes including low arousal threshold, poor pharyngeal dilator muscle effectiveness was also identified by PSG monitoring.^{48,49} These nonanatomical endotypes have also demonstrated the clinical value of guidance for non-CPAP treatment. Patients with less upper airway collapsibility and low loop gain could benefit from an oral appliance and upper airway surgery.⁵⁰ OSA due to poor muscle compensation could be improved from the upper airway muscle stimuli like desipramine.⁵¹ Hypnotics may be effective in patients with low arousal threshold.^{52,53} Details can be obtained from this former excellent review.⁵⁴ These brand-new algorithms provided insights into personalized treatment. Due to the limited study samples, single night PSG data and multiple potential confounders, these endotypes need more evidence for clinical utility.

Another possible solution to discriminate OSA endotypes is the application of supervised machine learning. Unlike unsupervised machine learning, supervised learning uses labeled data to predict patterns for unforeseen data. Edwards et al used regression models to find predictors for arousal threshold in 127 participants. An epiglottic pressure catheter was applied for the gold standard arousal threshold. Three PSG variables including AHI < 30, oxygen saturation nadir > 82.5%, and fraction of hypopneas >58.3% were used to predict low arousal threshold with a sensitivity of 82.2% and specificity of 84.0%.⁵⁵ The algorithm was further applied to predict the effect of drugs for OSA treatment and CPAP adherence.^{56,57}

Conclusions and Future Directions

Based on the pathogenesis of OSA and related comorbidities, this review discussed new metrics and methods aiming at OSA evaluation and treatment. Advantages of these metrics over AHI have been proved in some OSA comorbidities. PSG-oriented personalized treatments for specific OSA endotypes have been validated in some randomized control studies. This exciting scenario shows the great value of data-driven medicine. These new metrics excavated from abundant information in PSG will greatly enhance our ability to identify subtypes of OSA, explore disease pathogenesis towards certain comorbidities and discover better OSA treatment.

Despite these findings, more work is needed in future studies. First, metrics for OSA evaluation should be closely focused on pathophysiological mechanisms. For example, limitations exist in traditional indexes of hypoxemia. ODI with lowest oxygen saturation of more than 95% could be a physiological variation. Meanwhile, T90 does not consider the percentage oxygen saturation time between 90% to 95%. These defects might be contributors to negative results. Nasal flow or respiratory effort could represent the fluctuation of intrathoracic pressure yet few studies aiming at this important mechanism are carried out. Second, more validation tests towards clinical demands are required. With large cohort datasets available nowadays, PSG based new metrics could be validated easily across different ages, ethnicities and OSA comorbidities. However, most studies focus on the CVD and CVD related mortality. Associations of promising metrics like hypoxic burden with diabetes or cognitive impairment should be tested in future research. Other metrics related to certain pathophysiology of OSA like cardiopulmonary coupling⁵⁸ and cyclic alternating pattern⁵⁹ should also be taken into consideration. Third, due to the complex mechanisms of OSA and related comorbidities, assessing OSA severity by a single metric seems unrealistic. A scoring system combined with demographics, symptoms, genetics, molecular biomarkers and PSG signals may mostly satisfy the clinical practice. Such system could serve as the classification

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criteria for OSA subtypes and endotypes. This is particularly feasible with the development of big data and artificial intelligence. Last, new technologies like wearables could provide longitudinal data which is important for night-to-night, sleep stage or position related variability of these new metrics. Devices with new signals could also be introduced in OSA evaluation.

In conclusion, novel metrics derived from PSG could provide insights into OSA pathogenesis and instructions for disease assessment and treatment. More strict studies with high quality requirements for validation are needed in the future. These metrics, together with concepts of subtypes and endotypes, will facilitate better management of OSA and provide great help to the precision medicine.

Abbreviations

AD, Alzheimer's disease; AHI, apnea-hypopnea index; BP, blood pressure; CVD, cardiovascular disease; CPAP, continuous positive airway pressure; ECG, electrocardiogram; EEG, electrocencephalogram; LSpO₂, lowest pulse oxygen saturation; MCI, mild cognitive impairment; ODI, oxygen desaturation index; ODR, oxygen desaturation rate; ORP, odds ratio product; OSA, obstructive sleep apnea; PAT, pulse arrival time; PPG, photoplethysmography; PSG, polysomno-graphy; PWA, pulse wave amplitude; SBII, sleep breathing impairment index; T90, percentage time below 90% oxygen saturation.

Funding

This study was supported by the National Key Research and Development Projects of China [No. 2013BAI09B10].

Disclosure

The authors report no conflicts of interest in this work.

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