Endotyping Sleep Apnea One Breath at a Time
An Automated Approach for Separating Obstructive from Central Sleep-disordered Breathing

Ankit Parekh1, Thomas M. Tolbert1, Anne M. Mooney1, Jaime Ramos-Cejudo2, Ricardo S. Osorio2, Marcel Treml3, Simon-Dominik Herkenrath3, Winfried J. Randerath3, Indu Ayappa1, and David M. Rapoport1

1Division of Pulmonary, Critical Care, and Sleep Medicine, Icahn School of Medicine at Mount Sinai, New York, New York; 2Center for Sleep and Brain Health, Department of Psychiatry, New York University Grossman School of Medicine, New York, New York; and 3Institute of Pneumology at the University of Cologne, Department of Pneumology and Allergology, Center of Sleep Medicine and Respiratory Care, Bethanien Hospital, Solingen, Germany

Abstract

Rationale: Determining whether an individual has obstructive or central sleep apnea is fundamental to selecting the appropriate treatment.

Objectives: Here we derive an automated breath-by-breath probability of obstruction, as a surrogate of gold-standard upper airway resistance, using hallmarks of upper airway obstruction visible on clinical sleep studies.

Methods: From five nocturnal polysomnography signals (airflow, thoracic and abdominal effort, oxygen saturation, and snore), nine features were extracted and weighted to derive the breath-by-breath probability of obstruction (Pobs). A development and initial test set of 29 subjects (development = 6, test = 23) (New York, NY) and a second test set of 39 subjects (Solingen, Germany), both with esophageal manometry, were used to develop Pobs and validate it against gold-standard upper airway resistance. A separate dataset of 114 subjects with 2 consecutive nocturnal polysomnographies (New York, NY) without esophageal manometry, were used to develop Pobs and validate it against gold-standard upper airway resistance. A separate dataset of 114 subjects with 2 consecutive nocturnal polysomnographies (New York, NY) without esophageal manometry, was used to derive the breath-by-breath probability of obstruction (Pobs).

Measurements and Main Results: A total of 1,962,229 breaths were analyzed. On a breath-by-breath level, Pobs was strongly correlated with normalized upper airway resistance in both test sets (set 1: cubic adjusted [adj.] $R^2 = 0.87$, $P < 0.001$, area under the receiver operating characteristic curve = 0.74; set 2: cubic adj. $R^2 = 0.83$, $P < 0.001$, area under the receiver operating characteristic curve = 0.7). On a subject level, median Pobs was associated with the median normalized upper airway resistance (set 1: linear adj. $R^2 = 0.59$, $P < 0.001$; set 2: linear adj. $R^2 = 0.45$, $P < 0.001$). Median Pobs exhibited low night-to-night variability [intraclass correlation(2, 1) = 0.93].

Conclusions: Using nearly 2 million breaths from 182 subjects, we show that breath-by-breath probability of obstruction can reliably predict the overall burden of obstructed breaths in individual subjects and can aid in determining the type of sleep apnea.

Keywords: sleep apnea; esophageal pressure swings; airflow limitation; upper airway resistance; machine learning

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Correspondence and requests for reprints should be addressed to Ankit Parekh, Ph.D., Division of Pulmonary, Critical Care, and Sleep Medicine, Icahn School of Medicine at Mount Sinai, New York, NY 10029. E-mail: ankit.parekh@msm.edu.

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At a Glance Commentary

Scientific Knowledge on the Subject: Differentiating central from obstructive sleep apnea is critical in guiding treatment. This differentiation is largely dependent on classifying apneas and hypopneas using an assessment of inspiratory effort. Together with flow, effort determines upper airway resistance. Noninvasive signals that are surrogates of inspiratory effort are sufficient to classify apneas. However, for hypopneas, the gold standard for quantifying upper airway resistance is invasive esophageal manometry, which is not well tolerated and results in sleep disruption. As such, noninvasive surrogates of upper airway resistance are imperative to classify hypopneas, and thus, separate central from obstructive sleep apnea.

What This Study Adds to the Field: Our study shows that a probability of obstruction derived using a feature-engineered machine learning approach is a reliable and noninvasive surrogate of upper airway resistance and can successfully distinguish central from obstructive sleep apnea both on a breath-by-breath level and on a subject level. Our probability of obstruction, which is derived within a matter of minutes, can determine the primary type of a subject’s sleep apnea and aid in determining risks associated with untreated disorder and informing treatment approaches.

Methods

Subjects
A total of three datasets, comprising data from a total of 182 subjects, were used in this study (Figure 1). Initial data (used for development and initial testing) consisted of 29 patients (Table 1) seen at the New York University Sleep Disorders Center who previously underwent routine NPSGs with esophageal manometry as part of other research protocols (see Reference 12 for inclusion and exclusion criteria). The study was approved by the Institutional Review Board at the New York University School of Medicine, and all subjects provided written informed consent. A second set of data, consisting of 39 subjects (Table 1) who underwent routine NPSGs and a total of 182 subjects, were used in the present study (Figure 1). Initial data (used for development and initial testing) consisted of 114 patients who underwent 2 consecutive NPSGs without esophageal manometry was used to assess night-to-night variability of the derived probability of obstruction. Details on this data are provided in the online supplement.

NPSG Protocol
All NPSGs were collected using standard clinical equipment (Sandman; Embla Systems Inc. at New York University and SOMNOlab in Solingen, Germany). Each NPSG included signals for electroencephalography, electromyography, electrooculography, airflow using a nasal cannula/pressure transducer system, thoracic and abdominal effort, snoring, and oxygen saturation. In NPSGs with esophageal manometry, pressure (Pes) swings were measured with an esophageal catheter.
consisting of a thin catheter ending in a 10-cm latex balloon (Ackrad Labs) that was placed transnasally following lidocaine anesthesia and positioned in the lower third of the esophagus. Esophageal pressure measurements were made with a 100 cm H₂O pressure transducer (Validyne in data from New York University and UniTip catheters, UNISENSOR AG, in data from Solingen, Germany) (17).

**Novel Breath-by-Breath Method for Endotyping Obstruction**

Figure 2 depicts our overall approach of constructing the breath-by-breath probability of obstruction (P_{obs}). We adopted a feature-engineered, simplified machine learning approach to transform distinct patterns on routine NPSG to a breath-by-breath probability indicative of upper airway obstruction. Although the patterns were initially conceived on the basis of visual inspection, their identification in our algorithm was automated. Our primary assumptions driving the transformation were that 1) increased effort associated with a reduction in flow (i.e., elevated upper airway resistance) is indicative of obstruction of the upper airway (high P_{obs}); 2) sufficiently reduced effort associated with a reduction in flow (i.e., low upper airway resistance) is indicative of preserved airway patency or decreased inspiratory effort in harmony with reduced airway patency (low P_{obs}); and 3) distinctive patterns on routine NPSG are representative of immediate physiologic consequences of either increased or reduced effort. All analyses were automated and written in MATLAB (MathWorks) and C++ and are publicly available (github.com/aparek/pobs).

**Signal and feature selection.** A total of five signals (airflow, thoracic and abdominal effort, oxygen saturation, and snoring) collected during routine NPSG were selected as signals of interest. All signals other than O2 saturation were uncalibrated. Periods of NPSG where signals were invalid (e.g., disconnect) were excluded in an automated fashion. On the basis of the experience of the authors in research and clinical scoring, we extracted a total of nine features from the five signals, some of which are reported in previous studies (3, 18–20). A description of the selected features is given in Table 2 (see online supplement for detailed derivation of each feature and signal specific automated preprocessing). Scored sleep stages and marked respiratory events (e.g., apneas and hypopneas) were not used for the proposed approach.

**Development of breath-by-breath P_{obs}**

The six development studies from the 29 studies acquired at NYU were visually assessed by four experts and were unanimously agreed to have predominantly either OSA (three studies) or CSA (three studies) pathophysiology and hence considered to be extremes in this set of data. Sleep apnea manifests as periodic reductions in airflow observed on a nasal cannula/pressure transducer system. As such, the first step in the diagnosis of sleep apnea is the identification of relatively small breaths. To this end, breaths within a study were identified, segmented, and labeled as either small (normalized amplitude < 85%) or normal (normalized amplitude between 85 and 200%); see detailed methodology in online supplement. Each small breath in the six development set studies was assigned two weights based on the presence/absence of a given feature listed in Table 2: a weight suggesting the breath was part of a sequence of breaths during an ongoing upper airway obstruction event (likely OSA), and a weight suggesting the breath was part of a sequence of breaths during an ongoing event with preserved airway patency (likely CSA). For

**Figure 1.** Datasets used in the study. Demographics and characteristics of the subjects are detailed in Table 1. Demographics for dataset 3 are detailed in the online supplement. For the leave-one-subject-out cross-validation, at each instance, n = 22 subjects are used as a training set, whereas n = 1 subject is used as a test set. The performance metrics from each instance are then tabulated and used to determine the performance of the model.
Table 1. Subject Characteristics

| Comparison with Gold-Standard Resistance | Development (n = 6) | Test Set 1 (n = 23) | Test Set 2 (n = 39) |
|------------------------------------------|--------------------|---------------------|---------------------|
| Demographics                             |                    |                     |                     |
| Age, yr                                  | 56 ± 19            | 50 ± 17             | 51 ± 16             |
| Sex, M/F                                 | 5/1                | 21/2                | 31/8                |
| BMI, kg/m²                               | 35.9 ± 10.5        | 34.3 ± 8.7          | 28.7 ± 5.0          |
| Sleep-disordered breathing               |                    |                     |                     |
| OAI, h⁻¹                                 | 10.6 ± 9.8         | 7.8 (23.4)          | 3.0 ± 8.0           |
| CAI, h⁻¹                                 | 0.8 (11)           | 0.6 (3.1)           | 0.5 ± 1.6           |
| AHI₃A, h⁻¹                               | 68.2 ± 34.4        | 55.2 ± 26.4         | 13.2 ± 12.0         |
| Breath size distribution*                |                    |                     |                     |
| Imputed breaths                          | 5,161 (14.6%)      | 15,689 (11.5%)      | 5,573 (0.1%)        |
| Small breaths                            | 8,932 (25.2%)      | 40,317 (29.5%)      | 59,836 (21.5%)      |
| Normal breaths                           | 21,305 (60.2%)     | 80,907 (59.0%)      | 211,594 (76.4%)     |
| Total number of breaths                  | 35,398             | 136,913             | 276,803             |

Values are presented as mean ± SD. In cases of non-normally distributed data, values are represented as median (interquartile range). Test set 1: New York, NY. Test set 2: Solingen, Germany.

*For the breath size distribution, values in parentheses represent percent of total number of breaths.

Figure 2. The proposed approach for estimating breath-by-breath probability of obstruction using patterns from routine nocturnal polysomnograms. (A) Input and feature engineering: raw and uncalibrated signals from routine nocturnal polysomnograms are preprocessed, and each breath is given weights according to the presence or absence of selected features. The color for each feature indicates our perceived importance of the feature used in assigning the weights (see Table 2). (B) Scores to probabilities: obstructive and central scores for each breath are determined using the sum of all weights. A logistic model is then learned to transform the raw scores to breath-by-breath probability of obstruction. Note that pressure is used only for comparison of our estimated breath-by-breath probabilities with gold-standard resistance (ΔPes/flow) and is not used developing the obstructive/central scores. **Absence of effort** feature is used only for imputed breaths during apneas. Abd = abdomen; Pes = esophageal pressure; Rib = ribcage; SpO₂ = oxygen saturation.
The two breath-by-breath scores in the six studies were used as predictors, and the corresponding breath-by-breath normalized resistance was used as the response to train a logistic regression model. The aim of the logistic regression model was to provide breath-by-breath probabilities that could indicate likelihood of obstruction. The learned logistic regression model transforms breath-by-breath scores to a breath-by-breath probability of obstruction, i.e., P_{obs} (0 = most likely not obstructive; 1 = most likely obstructive). A cutoff of 200% for normalized resistance was used to define low versus high normalized resistance, on the basis of our previous study as well as on the observation by our group and others that airway resistance in normal subjects can double during sleep onset (12, 21).

*Internal and external validation of breath-by-breath P_{obs}.* We applied the learned logistic regression model on the 23 remaining studies in the initial test set from New York and the 39 studies in the second test set from Solingen, Germany. The learned logistic regression model was used to transform breath-by-breath scores to P_{obs} on each small breath across the test sets and compared against the corresponding normalized resistance. Logistic regression model estimated probabilities (e.g., P_{obs}) can either be used as continuous values (i.e., 0 to 1) or as classes by dichotomizing (e.g., low vs. high) them using a predefined threshold (usually 0.5). Here, we assessed the predictive value of the learned logistic model using both approaches: using continuous values for normalized resistance and P_{obs} as well as using dichotomized P_{obs} and normalized resistance, that is, low (P_{obs} < 0.5; normalized resistance < 200%) and high (P_{obs} ≥ 0.5; normalized resistance > 200%).

*Subject-level validation of breath-by-breath P_{obs}.* We further tested the utility of the P_{obs} in classifying subjects as either those with predominantly low or high resistance, indicating a subject likely to have either CSA or OSA pathophysiology.
and Bland-Altman (22) plots were used to assess the absolute agreement, single measurement correlations (ICC, one-way random effects, feature-engineered model. Intraclass correlation was used to assess correlations. Parekh, Tolbert, Mooney, et al. (23) assessed the subject level relationship between median \( P_{\text{obs}} \) and normalized resistance using median \( P_{\text{obs}} \) as a continuous value and as dichotomized into two classes (i.e., “likely central” as those with median \( P_{\text{obs}} < 0.5 \), and “likely obstructive” as those with median \( P_{\text{obs}} \geq 0.5 \)).

### Statistical Analyses

All statistical analyses were performed using IBM SPSS 24 and MATLAB R2020a (MathWorks). Normality of data was tested using the Shapiro-Wilk test. Spearman’s rank correlation was used to assess correlations. Binomial logistic regression with tenfold cross validation was used to learn the logistic model from the development set (\( n = 6 \)). Normalized resistance (%) was log transformed. We assessed the relationship between \( P_{\text{obs}} \) and normalized resistance, both treated as continuous values, using median-fit lines. For these median–median plots, \( P_{\text{obs}} \) values were divided into 100 equal-width bins ranging from 0 to 1 (i.e., bin width of 0.01). Bins with fewer than 10 breaths were discarded. Strength of relationships on the median–median plots was determined using the coefficient of determination (\( R^2 \)) from linear and cubic fit lines obtained using linear and polynomial regression, respectively. When both \( P_{\text{obs}} \) and normalized resistance were dichotomized into classes, the primary metrics for assessing classification performance of the learned logistic regression model were area under the receiver operating characteristic curve (AUC-ROC), accuracy (\( \text{ACC} = \% \) of breaths correctly classified), and Cohen’s kappa scores. We used paired \( t \) tests to assess significant differences in classification metrics between the random forest classification models and the simplified feature-engineered model. Intraclass correlations (ICC, one-way random effects, absolute agreement, single measurement) and Bland-Altman (22) plots were used to assess the night-to-night variability in median \( P_{\text{obs}} \). A two-tailed \( P \) value of less than 0.05 was considered indicative of statistical significance for all tests.

### Results

A combined total of 1,962,229 breaths from data in 182 subjects were analyzed. The computational runtime of deriving \( c \) was 3.1 ± 1.3 min (mean ± SD) for a single study with between 5,000 and 7,000 breaths.

#### Efficacy of Learned Logistic Regression Model

The six NPSGs in the development set had a roughly equal distribution of breaths with low (\( n = 3,360 \) breaths) versus high (\( n = 3,502 \) breaths) resistance. The logistic model fit for the development set was statistically significant (Nagelkerke \( R^2 = 21.4 \); see Table E1 in the online supplement). Both obstructive and central scores were significant predictors of normalized resistance. However, compared with the model with only the obstructive score, the model with both obstructive and central scores reduced the model deviance by less than 1% (data not shown).

#### Internal and External Validation of Breath-by-Breath \( P_{\text{obs}} \)

Example tracings and the corresponding \( P_{\text{obs}} \) values for three representative subjects from the initial test set are shown in Figure 3. Tracings for three subjects who had a predominance of apneas over hypopneas are shown in Figure E1. An example tracing for a representative subject from the second test set is shown in Figure E2.

On a breath-by-breath level, \( P_{\text{obs}} \) was significantly correlated with normalized resistance (Figure 4). Visually, the median–median data in Figure 4 appeared to be polynomial in nature, which was confirmed by the higher \( R^2 \) with the cubic fit as opposed to the linear fit (cubic adjusted \( \text{adj.}R^2 = 0.87 \) vs. linear \( \text{adj.}R^2 = 0.79 \)). Similarly, across the 39 subjects in the second test set, the \( P_{\text{obs}} \) was significantly correlated with normalized resistance (Figure 4B). In addition, as in the first test set, the relationship between \( P_{\text{obs}} \) and normalized resistance in the second test set appeared to be polynomial (cubic \( \text{adj.}R^2 = 0.83 \) vs. linear \( \text{adj.}R^2 = 0.78 \)). It should be noted that although we excluded apneas when assessing the relationship between the estimated probabilities and gold-standard resistance to avoid a 0/0 calculation, \( P_{\text{obs}} \) appeared to separate obstructive from central apneas (Figure E1). Further, \( P_{\text{obs}} \) appeared to correctly identify the likely obstructive versus likely central apneic components within mixed apneas (details in the online supplement).

#### Comparison with Data-driven Random Forest

Using feature weighting, the classification performance of dichotomized \( P_{\text{obs}} \) was significantly higher than the conventional random forest model (\( P < 0.01 \)) (Figure E5; see online supplement for methodological details).

### Discussion

Upper airway resistance, determined using invasive esophageal manometry, is the gold-standard measurement for classifying respiratory events as obstructive or nonobstructive, and therefore for distinguishing OSA from CSA. Using a
Figure 3. Example tracings and estimated probabilities from three representative subjects in our internal test set. Histograms on the right indicate the distribution of probability of obstruction ($P_{\text{obs}}$) throughout the night for each subject. (A) Pressure swings indicate a pattern of reduced effort on small breaths, which is also reflected in $P_{\text{obs}}$ values (near 0, majority $P_{\text{obs}} < 0.5$). (B) Pressure swings indicate a pattern of increased effort on small breaths, which is also reflected in $P_{\text{obs}}$ values (near 1). (C) Subject has patterns of reduced resistance; however, in certain cases the inspiratory flow shape indicates flow limitation. This ambiguity is reflected in distribution of $P_{\text{obs}}$ ($P_{\text{obs}}$ between 0.4 and 0.6). Overnight, this subject has a mix of obstructive and central physiology on small breaths. Abd = abdomen; Pes = esophageal pressure; Rib = ribcage; $\text{SpO}_2$ = oxygen saturation.
combined total of 1,996,629 breaths during sleep, assessed in a simplified machine learning and feature-engineered approach, we demonstrate that a breath-by-breath probability of obstruction obtained from routine NPSGs is a noninvasive surrogate for gold-standard upper airway resistance. We show that the proposed breath-by-breath probability of obstruction exhibits low night-to-night variability and aids in classifying a subject’s type of sleep apnea.

To our knowledge, our study uses one of the largest datasets consisting of NPSGs with concurrent esophageal manometry. We observed a strong correlation between Pobs and gold-standard upper airway resistance. We show that the proposed breath-by-breath probability of obstruction exhibits low night-to-night variability and aids in classifying a subject’s type of sleep apnea.

**Clinical Implications**

Developing validated tools to better characterize CSA, especially in patients with heart failure, is one of the research priorities of the American Thoracic Society (2). A recent international taskforce report advised that the principal step in the approach to treating CSA is establishing the presence of an underlying CSA pathophysiology in an individual; establishing the presence of CSA is fundamental to its treatment (6). As such, the primary step toward effective treatment of CSA is establishing presence of CSA, or absence thereof, which the proposed Pobs achieves reliably within a matter of minutes from a routine clinical NPSG. Further, identifying presence of a central ventilatory disturbance that is not based on the presence of central apneas alone is clinically meaningful as it could predict failure to respond to therapies that target upper airway obstruction (14).

Current AASM rules and practices do not adequately operationalize classifying respiratory events (apneas/hypopneas). Although some events may be clearly central (or obstructive), many events have a combined etiology, and a lack of consensus on their identification and reporting makes it difficult for labs to operationalize any event-based rules. As recently reported, the classification performance based on these rules is relatively low (23). Here, we have shown how our breath-by-breath probability may overcome these limitations. Breaths are the smallest division using which one can measure an individual’s primary type of sleep apnea. With each sleep study containing several thousand breaths, breath-by-breath analyses are robust to both physiological and nonphysiological noise. In addition, breath-by-breath analyses, such as Pobs, can break...
down and effectively characterize mixed events for which there is no consensus currently. Furthermore, rudimentary techniques can be put in place that can effectively combine $P_{obs}$ to generate an event-level probability, or as we have shown here, to a subject-level probability. Our results of near-perfect agreement of the probability scores on a subject level, when compared with gold-standard resistance, further indicate the clinical utility of breath-by-breath analyses and $P_{obs}$ in deducing subject-level outcomes.

Our approach of using probabilities as continuous values, rather than dichotomized into discrete categories, can provide noteworthy physiological insight while offering an intuitive clinical interpretation. As an example, a subject whose median $P_{obs}$ is around 0.5, say between 0.4 and 0.6, could indicate several possibilities including 1) the subject has underlying OSA physiology, but their upper airway resistance is not as elevated as others with more severe disease, 2) the subject has a mix of OSA and CSA physiology, or 3) presence of other comorbid conditions that alter the ventilatory stability. A dichotomized approach for $P_{obs}$ would obscure these physiological insights, and as such, we recommend future studies in the direction where the entire histogram of overnight $P_{obs}$ is used rather than in a binary form (yes vs. no, obstructive vs. central).

![Figure 5](image-url) **Figure 5.** Classification performance of the learned logistic model using dichotomized probability of obstruction (two classes: low vs. high, with a threshold of 0.5). The left plot shows the mean area under the receiver operating characteristic curve (AUC-ROC) across all subjects in the two sets. Confidence interval is represented by the shaded area. The dashed reference line indicates the AUC-ROC for a randomized classifier (i.e., classifies breaths as low vs. high resistance by chance). Box plots on the right show classification accuracy (ACC) and kappa for each individual subject in the internal and the external test sets. Dark red lines indicate the mean ACC values, with the light and dark shaded sections of the boxes indicating one and two standard deviations.

![Figure 6](image-url) **Figure 6.** Association of median probability of obstruction with median normalized resistance across all small breaths on a subject level in (A) test set 1 (New York, NY) and the (B) test set 2 (Solingen, Germany). Each dot represents an individual subject. Solid lines represent the fit lines, and the dashed lines represent the 95% confidence intervals. The dashed horizontal line indicates a normalized resistance of 200%, which was used as a cutoff to determine low versus high resistance. Adj. = adjusted.
Furthermore, continuous values can also be used as a decision aid to human scorers, should such a need arise.

**Limitations**

We used 6 of 29 studies for the development and training of the binary logistic model that could transform raw obstructive/central scores to probabilities. Although we had a larger set of studies for training at our disposal, traditional machine learning theory and empirical evidence show that performance saturates as more training data are added (24). Accordingly, we chose the six studies that, on a clinical reading, had unambiguously OSA or CSA physiology overall to optimize the feature weights and features. Deep learning models that could be developed in the future, however, could benefit from using our large dataset for training if additional studies can be used to validate the results. It should be noted that the proportion of female subjects in the development and the test sets (internal and external) was low. Since it is observed in routine clinical practice that sleep apnea prevalence is greater among men than women, such a skewed distribution is not surprising. However, balanced datasets may be used in a future study to assess the role of sex in the performance of our automated approach.

We validated and tested the utility of our estimated breath-by-breath probability of obstruction in two ways: 1) “as is”, i.e., using probabilities as continuous values and 2) dichotomizing them into two classes (low vs. high with a cutoff of 0.5). For both the approaches we observed a performance that is greater than the performance by a previously published manual visual algorithm (17). Logistic models provide probabilities as an outcome, and dichotomizing them is a forced choice and diminishes their overall utility (25). Thus, it is not surprising that the strength of the classification performance using the leave-one-subject-out cross-validated approach, which dichotomized the probabilities, was slightly less than when using the “as is” continuous value probabilities.

We compared our approach, which is based on a weighting scheme influenced by our perceived importance of features (feature engineered), with conventional machine learning (random forest), wherein feature importance is data driven. Although using custom weighted features may be thought of as subjective in nature, they offer translatability and are built upon years of clinical knowledge. We observe that the feature-engineered approach outperformed the conventional random forest model, which is not surprising as it has been long known that machine learning performance can be improved using as much domain knowledge as possible. We observed that both the approaches resulted in similar relative importance of the features. For example, the presence of an inspiratory shape suggesting flow limitation was considered one of the most important features. The relative importance of features that we observe here (see online supplement for details) was also echoed in the study by Randerath and colleagues (17). Further, as pointed out by Iber in his corresponding commentary (26), snoring did appear to be a relatively important feature.

**Conclusions**

Upper airway resistance quantified using invasive esophageal manometry is the gold standard for deducing the type of sleep apnea as obstructive or central and is key in determining the appropriate course of treatment. However, esophageal manometry is not well tolerated during sleep, and a lack of consensus on events with combined obstructive and central components can make characterizing the type of sleep apnea difficult. It has been long known that respiratory disturbances are associated with distinctive patterns observed during routine nocturnal polysomnograms. In most clinical scoring schemes these patterns influence, either directly or subliminally, the visual characterization of respiratory events. In this study, we develop, validate, and test a probabilistic approach that amalgamates known physiologic features of respiratory disturbances into a breath-by-breath probability of obstruction. We show that the probability of obstruction is a reliable noninvasive surrogate for the invasive gold-standard measurement of upper airway resistance and can aid in identifying a subject’s type of sleep apnea within a matter of minutes from a standard clinical sleep study.

**Author disclosures** are available with the text of this article at www.atsjournals.org.

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