<table>
<thead>
<tr>
<th>Title</th>
<th>Classifying vulnerability to sleep deprivation using baseline measures of psychomotor vigilance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Author(s)</td>
<td>Patanaik, Amiya; Kwoh, Chee Keong; Chua, Eric C. P.; Gooley, Joshua J.; Chee, Michael W. L.</td>
</tr>
<tr>
<td>Date</td>
<td>2015</td>
</tr>
<tr>
<td>URL</td>
<td><a href="http://hdl.handle.net/10220/26277">http://hdl.handle.net/10220/26277</a></td>
</tr>
</tbody>
</table>

© 2015 Associated Professional Sleep Societies (APSS). This paper was published in Sleep and is made available as an electronic reprint (preprint) with permission of Associated Professional Sleep Societies (APSS). The published version is available at: [http://dx.doi.org/10.5665/sleep.4664]. One print or electronic copy may be made for personal use only. Systematic or multiple reproduction, distribution to multiple locations via electronic or other means, duplication of any material in this paper for a fee or for commercial purposes, or modification of the content of the paper is prohibited and is subject to penalties under law.
CLASSIFYING VULNERABILITY TO SLEEP DEPRIVATION USING BASELINE PVT MEASURES

Classifying Vulnerability to Sleep Deprivation Using Baseline Measures of Psychomotor Vigilance

Amiya Patanaik, BTech1; Chee Keong Kwoh, PhD2; Eric C.P. Chua, PhD2; Joshua J. Gooley, PhD2; Michael W.L. Chee, MBBS2

1School of Computer Engineering, Nanyang Technological University, Singapore; 2Centre for Cognitive Neuroscience, Neuroscience and Behavioral Disorders Program, Duke-NUS Graduate Medical School, Singapore

Objective: To identify measures derived from baseline psychomotor vigilance task (PVT) performance that can reliably predict vulnerability to sleep deprivation.

Design: Subjects underwent total sleep deprivation and completed a 10-min PVT every 1–2 h in a controlled laboratory setting. Participants were categorized as vulnerable or resistant to sleep deprivation, based on a median split of lapses that occurred following sleep deprivation. Standard reaction time, drift diffusion model (DDM), and wavelet metrics were derived from PVT response times collected at baseline. A support vector machine model that incorporated maximum relevance and minimum redundancy feature selection and wrapper-based heuristics was used to classify subjects as vulnerable or resistant using rested data.

Setting: Two academic sleep laboratories.

Participants: Independent samples of 135 (69 women, age 18 to 25 y), and 45 (3 women, age 22 to 32 y) healthy adults.

Measurements and Results: In both datasets, DDM measures, number of consecutive reaction times that differ by more than 250 ms, and two wavelet features were selected by the model as features predictive of vulnerability to sleep deprivation. Using the best set of features selected in each dataset, classification accuracy was 77% and 82% using fivefold stratified cross-validation, respectively.

Conclusions: Despite differences in experimental conditions across studies, drift diffusion model parameters associated reliably with individual differences in performance during total sleep deprivation. These results demonstrate the utility of drift diffusion modeling of baseline performance in estimating vulnerability to psychomotor vigilance decline following sleep deprivation.

Keywords: classification, differential vulnerability, diffusion model, psychomotor vigilance task, sleep deprivation

Citation: Patanaik A, Kwoh CK, Chua EC, Gooley JJ, Chee MW. Classifying vulnerability to sleep deprivation using baseline measures of psychomotor vigilance. SLEEP 2015;38(5):XXX–XXX.

INTRODUCTION

The negative consequences of sleep deprivation on cognitive performance are well documented.1,2 Lapses in attention resulting from sleep deprivation contribute to industrial catastrophes, medical errors, transportation accidents, and security breaches.4–8 Sleep deprivation is thought to impair neurobehavioral functioning by destabilizing performance, as evidenced by the capacity to perform well for short periods of time, interrupted by occasional attention failures.9 The extent to which sleep loss affects performance varies widely across individuals, however, with some subjects remaining relatively unaffected while others show severe cognitive impairment.9 Notably, between-subject differences in performance are trait like and stable across repeated exposures to sleep deprivation, irrespective of sleep history.10–12 Over the past decade, several studies have shown that subjects who are vulnerable versus resistant to the effects of sleep loss differ in their brain activation and behavioral performance when they are well rested.13–15 Such findings suggest that baseline measures can be used to predict how well a person will perform when he/she is deprived of sleep.

One of the most sensitive measures of performance impairment by sleep deprivation is vigilance,20 commonly assessed using the psychomotor vigilance task (PVT).21 During the PVT, participants maintain their fastest possible reaction time (RT) to a simple visual stimulus presented at random interstimulus intervals. Recently, it was shown that subjects who are vulnerable to the effects of total sleep deprivation on PVT performance show slower and more variable response times when they are well rested.17 In separate work, subjects categorized as vulnerable or resistant to sleep deprivation differed in their diffusion drift parameters derived from PVT RT data sampled at baseline.19 These studies demonstrate that baseline PVT performance carries information about vulnerability to subsequent sleep deprivation, but it remains unclear whether features of rested PVT performance can be used to classify a person’s relative performance in the sleep deprived state.

The current study sought to assess the reliability of classifying subjects as vulnerable or resistant to sleep deprivation, using baseline features of PVT performance. We used two independent datasets to carry out our analysis involving PVT data collected from different laboratories under different experimental conditions. We extracted standard RT metrics, diffusion model parameters, and features derived from spectral analysis of RTs, and used a support vector machine (SVM) classifier with stratified fivefold cross-validation (CV5) to estimate generalization error. The objectives of this work were fourfold:

1. Identify and rank candidate features derived from baseline PVT response times that predict vulnerability to sleep deprivation.
2. Select a compact feature set from the candidate features resulting in maximum prediction accuracy, and measure
the performance of this classification model for each dataset.

3. Evaluate the reliability of classification by training the model on one dataset and testing it on another one.

4. Evaluate test-retest performance of classification by training the model using baseline data collected during one study visit, and then testing it using baseline data collected more than 5 mo later from the same set of individuals.

MATERIAL AND METHODS

Subjects
In the current investigation, we analyzed PVT data collected from two laboratories. For Dataset 1, a total of 135 subjects (69 females, age 18–25 y) who participated in five different functional imaging studies were evaluated. All five experiments in this dataset were conducted at the Cognitive Neuroscience Laboratory, under similar experimental conditions. For Dataset 2, 45 healthy ethnic-Chinese subjects (3 females, age 22–32 y) were enrolled in a laboratory study at the Chronobiology and Sleep Laboratory (CSL) as part of a previous study. Both datasets shared some common recruitment criteria. For example, health was assessed using screening questionnaires and self-reported medical history. Participants who took medications or consumed nicotine products were excluded. In the week before the laboratory study, participants were required to maintain a consistent sleep-wake schedule (6.5–9 h of sleep every day in Dataset 1, and 8 h in bed for sleep in Dataset 2), which was verified by actigraphy monitoring (Actiwatch-L or Actiwatch 2, MiniMitter, Inc., Bend, OR). In the week prior to the study, subjects were asked to avoid caffeine, alcohol, and over-the-counter medications. Informed consent was obtained from all participants, and research procedures were approved by the National University of Singapore Institutional Review Board (IRB) and the SingHealth Centralized IRB for Dataset 1 and Dataset 2, respectively.

Sleep Deprivation Procedures
Dataset 1: Subjects arrived at the laboratory at 19:30 and were kept awake continuously overnight under supervision of a research assistant. A handheld 10-min PVT was administered every hour from 20:00 to 05:00 (10 test periods). Subjects were seated upright during testing and were exposed to ordinary room light. Participants’ movements were not restricted between PVT tests.

Dataset 2: Subjects underwent total sleep deprivation in a laboratory suite that was shielded from external time cues. Participants arrived in the evening and went to bed at their regular bedtime. After 8 h in bed for sleep, subjects were kept awake for at least 26 h using constant routine (CR) procedures, as previously described. During the CR procedure, subjects remained in bed in a semirecumbent position, with exposure to dim ambient lighting (< 5 lux). The PVT was administered every 2 h (starting 2.5 h or 4.5 h after wake time) by computer using E-Prime 2 Professional software (Psychology Software Tools, Inc., Sharpsburg, PA). Visual stimuli were presented on a liquid crystal display monitor placed on an over-bed table, which allowed subjects to take the PVT while remaining in bed. After undergoing sleep deprivation, participants were invited to return to the laboratory at least 5 mo later to complete additional testing. A subset of subjects (n = 34) took part in the follow-up study. Subjects reported to the laboratory in the mid-afternoon (between 14:00–18:00) and completed two 10-min PVTs taken 2 h apart from one another under conditions that were similar to the first study visit. The research protocols for both datasets are summarized in Figure 1.

Assessment of Vulnerability to Sleep Deprivation
During total sleep deprivation, cognitive performance usually reaches its nadir in the early morning hours, typically between 04:00 and 08:00, when the sleep homeostat and circadian clock interact to promote high levels of sleepiness. This is also the period when sleepiness-related motor vehicle accidents are most likely to occur. We therefore analyzed PVT performance during this time window as a measure of susceptibility to sleep deprivation. Subjects were categorized as vulnerable or resistant based on a median split on the number of lapses, defined as RTs that exceed 500 msec (Figure 1). PVTs marked in red). For Dataset 1, 70 subjects were categorized as vulnerable (≥ 5 lapses). For Dataset 2, 25 subjects were categorized as vulnerable (≥ 23 lapses), of whom 19 completed the follow-up study. The large difference in the median lapse value is likely a result of different experimental conditions between the two studies (see Discussion). The divergence in performance between vulnerable and resistant groups after their usual bedtime is shown in Figure 2. For Dataset 1, the first two PVT tests administered at 20:00 and 21:00 were used as the baseline. For Dataset 2, the third and fourth PVT sessions, which were taken during the mid-afternoon, were used as the rested baseline, as PVT measurements for the follow up session were available for the same time period.

RT-Derived Features
As detailed in the next paragraphs, we used a combination of standard RT metrics, features derived from the drift diffusion model (DDM), and spectral analysis of RTs.

Standard RT Metrics
The most widely used PVT outcome metric is the number of lapses followed by mean RT, mean 1/RT, fastest 10% RT, median RT, slowest 10% RT, and the slowest 10% 1/RT. The reciprocal RT is also referred to as response speed (response speed = 1/RT). In an analysis of various PVT outcome metrics during partial or total sleep deprivation, Basner and Dinges found that metrics involving response speed (RS) and lapses were the most sensitive to sleep loss. It was therefore recommended that these measures be used as the primary outcomes measures of the 10-min version of the PVT. In our analysis we considered lapses, mean RT, mean RS, lowest 10% RS, fastest 10% RT, median RT, and standard deviation of RT. We also included mean absolute deviation (MAD) from the mean, and ΔRT > 250, which is the number of consecutive RTs that differ by more than 250 msec. The latter was included based on a previous finding that, under baseline conditions, subjects who were categorized as vulnerable to sleep deprivation showed a greater number of consecutive RTs that differed more than 250 msec compared to resistant individuals.
Predicting Vulnerability to Sleep Deprivation—Patanaik et al.

Metrics Derived From DDM

The Ratcliff DDM is a powerful model of perceptual decision making for single-choice RT experiments such as the PVT. The model decomposes the decision process into decision and nondecision components (Figure 3A). The nondecision component refers to time spent encoding the sensory input (predecision time) as well as time spent in executing the decision (post-decision time). Decision-making itself is conceived to be a noisy process involving the accumulation of information over time that can be modeled mathematically as a diffusion process. An attractive feature of diffusion modeling is that it can predict the response time distribution under different contexts and varying levels of noise. The model has been tested by manipulating various facets of the decision process and then observing the corresponding change in diffusion parameters. More importantly, DDM parameters estimated prior to sleep deprivation have been shown to differ in groups of subjects categorized as vulnerable or resistant to total sleep deprivation, even when baseline standard RT metrics were similar.

In the Ratcliff DDM used here, nondecision time was assumed to vary from trial to trial according to a uniform distribution with mean $T_n$ and width $S_n$. Decision time was modeled using a single boundary diffusion process with a drift parameter. Evidence was assumed to accumulate from the starting point (at 0) until the boundary $a$ was reached. The drift parameter was also allowed to vary across trials according to a normal distribution with mean $\xi$ and standard deviation $\eta$ (Figure 3B). This results in a model with five parameters $\Theta = [a, T_n, S_n, \xi, \eta]$; however, not all parameters of the model are uniquely identifiable. This is because the boundary parameter $a$ can be scaled by equally scaling the drift parameters $\xi/a$ and $\eta/a$ are used. For the sake of simplicity, from this point onward when we refer to the drift parameters it is implicitly assumed that they are normalized by the boundary parameter. We also included the parameter ratio $\xi/\eta$, which is the diffusion drift signal-to-noise ratio. This parameter is known to closely track alertness. While estimating the parameters of the model, we combined RT data from two consecutive PVT sessions taken during the baseline rested state. This was necessary to get reliable estimates of the parameters. The parameter estimation process is described in detail elsewhere.

Metrics Derived From Spectral Analysis of RTs

Given that spatiotemporal features (i.e., structures in the ordering and positioning of the RTs) might not be captured by standard PVT metrics or the DDM, we used discrete wavelet transform (DWT) to extract multiresolution features from RTs. The DWT effectively addresses the tradeoff between time and frequency resolution.
frequency resolution in signal analysis and can handle non-stationary signals as well. It decomposes the signal into a hierarchical set of low frequency and high frequency components called approximations and details, respectively. The DWT was computed by applying successive low-pass and high-pass filtering in the time domain, resulting in a multilevel decomposition of the RTs at different scales. Intuitively, the DWT can be thought of as a mathematical microscope optimized to capture temporal structures on finer and finer scales. Because of the way DWT operates, the length of the signal has to be a power of 2. Because the number of RTs collected per PVT can vary widely across individuals, it is necessary to either truncate or to artificially extend the number of trials associated with each participant. To avoid introducing artifacts because of signal extension, and to maintain consistency across subjects and datasets, we considered the last $2^6 = 64$ PVT trials from each test.

Samples from two consecutive PVTs taken during baseline were combined, resulting in uniform sample size of 128, and a six-level DWT was applied. For each level $l$, the mean absolute value ($MAV_l$) was computed from the detail coefficients at that level. Details of the DWT analysis are presented in the supplemental material. The selected features were normalized to a unit sphere.

**Feature Selection**

As summarized in the previous section, we examined a total of 20 PVT features of the baseline data, including nine standard RT features, five DDM features, and six wavelet features. The task of predicting the class label (vulnerable or resistant) from baseline RT data is essentially a supervised classification problem. It is supervised in the sense that a pattern recognition algorithm (support vector machines [SVMs] in this case, discussed in the next section) was trained on a set of labeled data (i.e., it is known whether the subject is vulnerable or resistant). The trained algorithm was then applied to new test data. The term ‘generalization error’ refers to the prediction error with respect to the new data and measures how well a learning algorithm generalizes to unseen data. The stratified $k$-fold cross validation method with $k = 5$ or 10 has been shown to be superior to other error estimation methods. We chose the fivefold stratified cross-validation (CV5) method in our analysis as tenfold cross-validation would result in very small sample per fold because of the smaller sample size of Dataset 2. To carry out the cross-validation, the original data was randomly partitioned into five equally sized subsamples. Each subsample had the same proportion of vulnerable and resistant subjects. Of the five subsamples, one was retained as the test set and the remaining four were used to train the model. This was repeated five times, with each subsample used exactly once for testing. The accuracy was then aggregated over all the subsamples that constitute the CV5 accuracy.

In any supervised classification problem, identifying the essential features is critical to the performance of the classifier. Some of the features might be irrelevant in the sense that they provide no additional information from the point of view of class prediction. Features could also be redundant, i.e., in the presence of other relevant features they provide no additional information. Moreover, irrelevant and redundant features increase computational complexity and might introduce noise into the system, reducing the performance of the classifier. Therefore, a subset of features must be selected to optimally predict vulnerability. A naïve way of achieving this is to consider all possible combinations of features and select the one that minimizes the generalization error. Unfortunately, this
is computationally inefficient even for a moderately large number of features. Additionally, because the generalization error has to be estimated from data, a large number of searches increases the likelihood of overfitting, especially in the case of a small dataset.

For our analyses, we first eliminated highly redundant or irrelevant features to shortlist a critical set of candidate features based on minimal redundancy-maximal relevance (mRMR). This was conducted independently for each dataset. The mRMR is a multivariate feature selection method that is superior to univariate methods such as the t test and F-test based methods (refer to Saey et al. for a review) and is widely used within machine learning and clinical communities. For example, mRMR has been used for feature selection in predicting drug-target interaction, microarray gene expression data, and classification using electroencephalographic signals, among others. The mRMR criterion subtracts the minimal redundancy from maximum relevance, thereby combining both relevance and redundancy information into a single score for feature ranking (see supplemental material for details). It was observed that the classification accuracy stabilized by the eighth ranked feature for both datasets, when each feature was considered incrementally. Therefore, only the top eight features were selected as the candidate feature set for further analysis.

Even though a selected candidate feature is deemed relevant, including all such features does not necessarily result in a better classification rate as compared to using a smaller feature set. Hence, relevance does not imply optimality. As a result “the m best features are not the same as best m features.” To maximize the accuracy of the classifier, a further compact subset of features was selected from the candidate features using a wrapper-based approach. A wrapper is a feature selector that is “wrapped” with the classifier to select features that result in the lowest generalization error. This is very similar to the naïve method described earlier, but used on a much smaller candidate feature set employing some heuristic schemes instead of trying out all possible combinations of features. We employed both the incremental forward and backward selection wrapper schemes detailed previously. In the incremental forward selection scheme, we started with an empty set. Features from the candidate set that resulted in the highest accuracy were added one by one until further addition did not improve classification accuracy. Similarly, in the incremental backward selection scheme, we started with all features in the candidate set and the least significant feature (in terms of accuracy obtained on remaining features) was incrementally removed until further removal of a feature reduced classification accuracy.

Classifier

For classification we used SVMs with a radial basis function kernel. SVMs are powerful supervised classification methods with strong theoretical foundations in statistical learning theory and structural risk minimization. In addition to being simple to implement, SVMs provide very good classification accuracy and have a high tolerance of noise (see Kotsiantis for a review). A SVM is similar to logistic regression in the sense that both search for a linear separation between the classes. The key difference between the two methods is that a standard SVM tries to maximize the margin between the two classes instead of minimizing the logistic cost function. For data that are not linearly separable, a kernel function is used to transform the original data to a high-dimensional space, where the data is linearly separable. The exact details of the implementation are discussed in the supplemental material. All analyses were implemented in Matlab 2013b, The MathWorks, Inc., Natick, MA, United States. The SVM was implemented in Matlab using LIBSVM. The complete feature selection process is summarized in Figure 4.

Statistical Analyses

Standard RT metrics are known to be stable and reproducible across studies. To test the reproducibility of the DDM parameters, parameters estimated from baseline data of Dataset 1 collected in the first visit were compared with parameters estimated from the follow up study using two-way repeated-measures analysis of variance (ANOVA) with group (vulnerable versus resistant) as the between-subject factor and schedule (first visit versus follow-up) as the within-subject factor.

The variation in DDM parameters for vulnerable and resistant subjects on the evening before and the morning after sleep deprivation has been reported previously. As PVT measurements for Dataset 2 were available across the day, DDM parameters were also estimated for the vulnerable and resistant
groups at different time of the day. This was also done using repeated-measures ANOVA (group as between-subject and time of day as within-subject factor, with data binned for two sessions of the PVT). For ANOVAs with statistically significant interaction, post hoc $t$ tests were used to examine simple effects of group and time. The classifiers were compared using McNemar test with Yates correction. Statistical analyses were performed using SPSS (IBM Corp., New York, NY). Statistical significance was set at $a = 0.05$.

RESULTS

In each dataset, the top eight features ranked according to their mRMR score are shown in Figure 5. As described in Methods, a wrapper-based approach was used to select a subset of features that minimized the generalization error. In the next section we report classifier accuracy as assessed by cross validation (CV5 accuracy) of features using the optimal feature set.

Model Performance in Dataset 1

A CV5 accuracy of 77% was obtained with sensitivity of 85.7% and specificity of 67.7% by using five features: diffusion drift $\xi$, variability in diffusion drift $\eta$, range of nondecision time $S_t$, the number of consecutive RTs that differed by more than 250 msec ($\Delta RT > 250$), and wavelet feature $\text{MAV}_6$ (Figure 6A). The receiver operating characteristic (ROC) curve and associated confusion matrix are presented in Figure 6A. The area under the ROC curve (AUC) is often used as a single-value representation of overall classifier performance and is equivalent to the probability that the classifier will rank a randomly chosen positive instance higher than a randomly chosen negative instance. For Dataset 1, the AUC for the classifier was 0.75.

Model Performance following Training on Dataset 1 and Testing on Dataset 2

We trained the model on Dataset 1, using the corresponding five features deemed most optimal and applied this feature set to Dataset 2. An accuracy of 71.1% was obtained with sensitivity of 80% and specificity 60%. On varying the threshold for the classifier, a more balanced result was obtained with sensitivity of 72% and specificity of 70% while the accuracy remained unchanged. The ROC and confusion matrix at the default threshold as well as the balanced threshold are presented in Figure 7. AUC for the classifier was 0.73.

Model Performance in Dataset 2

For Dataset 2, diffusion drift $\xi$, range of nondecision time $S_t$, and the number of consecutive RTs that differed by more than 250 msec ($\Delta RT > 250$) were deemed most important. A CV5 accuracy of 82.2% was obtained with sensitivity of 84% and specificity of 80% using three features: diffusion drift $\xi$, range of nondecision time $S_t$, and the number of consecutive RTs that differed by more than 250 msec ($\Delta RT > 250$). The ROC and associated confusion matrix are presented in Figure 6B. For Dataset 2, the AUC for the classifier was 0.74.
Reproducibility of Classification across Testing Episodes in the Same Participants

Using the optimal feature set selected in Dataset 2, when the model was trained on baseline PVT data collected during the first visit and then tested on PVT data from the follow-up visit, we obtained an accuracy of 79.4%, with sensitivity of 73.7% and specificity of 86.7%.

Classification Using the Most Sensitive PVT Measures

Although standard PVT metrics were not selected by the model, we considered the possibility that such measures might nonetheless carry information about vulnerability to sleep deprivation. Using lapses, mean RS, and slowest 10% RS measured at baseline, three measures previously found to be sensitive to total sleep deprivation, we obtained a CV5 accuracy of 64.4% on Dataset 1 and CV5 accuracy of 68.9% on Dataset 2. Compared to the best performing features for each dataset, classification accuracy using these PVT measures was significantly poorer in the case of Dataset 1 ($\chi^2 = 10.2, P < 0.005$) but not significantly different for the smaller Dataset 2, ($\chi^2 = 1.8, ns$).

DDM Variability across Visits and with Time; Computational Performance

Except for across-trial variability in drift $\eta$ ($F_{1,32} = 4.65, P < 0.05$), DDM parameters were stable across study visits (Figure 8). Vulnerable subjects had lower mean drift ($F_{1,32} = 11.6, P < 0.005$) and higher variability in nondecision time ($F_{1,32} = 13.6, P < 0.001$) across both visits compared to resistant subjects. Although DDM parameters were significantly affected by time of the day (Figure S1, supplemental material), vulnerable subjects had lower mean diffusion drift ($F_{1,43} = 4.03, P < 0.001$) and higher variability in non-decision time ($F_{1,43} = 27.5, P < 0.001$) irrespective of time elapsed since wake.

The feature extraction process was reasonably quick (~10 min/subject) when run on a contemporary workstation (six core Intel® Xeon® 3.2 GHz with 16 GB of RAM), with the majority of time spent estimating the DDM parameters. Overall, the $\chi^2$ DDM fit (at baseline) for Dataset 1 was 19.2 ± 6.1 and for Dataset 2 it was 16.8 ± 7.2. The critical value for $\chi^2$, df = 14 was 26.1.

DISCUSSION

We observed large between-subject differences in PVT performance during sleep deprivation, such that persons vulnerable to sleep deprivation had on an average three to eight times more lapses than members of the resistant group. Although prior studies have identified baseline differences between individuals who are either vulnerable or resistant to sleep deprivation, the ability to predict performance vulnerability using baseline data has not been systematically examined. Here, we examined features beyond summary statistics conventionally used in assessing RTs, including measures derived from the DDM and spectral analysis. Using two independent datasets, we identified a subset of PVT features that can be used to...
predict relative vulnerability to total sleep deprivation with about 77–82% accuracy.

SLEEP, Vol. 38, No. 5, 2015

classify relative vulnerability to total sleep deprivation with about 77–82% accuracy.

SLEEP, Vol. 38, No. 5, 2015

Features Most Useful for Discriminating Vulnerable and Resistant Participants

Despite substantial differences in the way that PVT data were collected across the two studies considered here, DDM parameters and wavelet MAV parameters were among the top baseline features associated with relative vulnerability to sleep deprivation. In terms of best performing features, three features were selected in both datasets: diffusion drift $\xi$, range of non-decision time $S$, and the number of consecutive RTs that differ by more than 250 msec ($\Delta RT > 250$). Interestingly, none of the standard PVT performance metrics (e.g., mean RT and lapses) were selected when the mRMR criterion was used. The empirical data collected suggest that relative to standard RT metrics, the DDM better captures useful information embedded in RT data that can distinguish persons vulnerable to vigilance decline following sleep deprivation by decoupling RT into distinct components. This is consistent with a previous study where diffusion parameters measured at baseline predicted vulnerability despite the absence of significant differences in baseline standard RT metrics. This might not be surprising upon inspecting the variation of the DDM parameters for the two groups across the day for Dataset 2 (Figure S1). Although the mean diffusion drift and range of nondecision time showed statistically significant differences across groups irrespective of time of day, the mean non-decision time and across-trial variability parameters showed interesting variations throughout the day. Depending on the time of the day, DDM parameters moved in opposite directions; i.e., some DDM parameters had a tendency of increasing the RT while others had a tendency to decrease it. In other words, the decision and nondecision components can trade off with each other without affecting overall observed performance. This has also been demonstrated using simulations.

Classification Reliability and Reproducibility

PVT outcomes can be affected by multiple factors including experimental conditions, interventions, and time of day. Here, we showed that the same set of features appear to be highly discriminatory across datasets. For Dataset 1, five features were selected by the model:
diffusion drift $\xi$, variability in diffusion drift $\eta$, range of nondecision time $S_t$, $\Delta RT > 250$, and $MAV_r$. A CV5 accuracy of 77% was achieved using these features. Despite differences in experimental conditions and baseline data acquisition times between the two datasets, the model trained on Dataset 1 and tested on Dataset 2 showed only a small drop in classification performance (71% accuracy). Importantly, the model performed well despite the large difference in the average number of lapses between studies, i.e., the median split on Dataset 1 cannot be directly linked to the median split on Dataset 2. Allowing the threshold to change when the model trained on Dataset 1 was applied to Dataset 2 resulted in a more balanced classification. These results attest to the utility of our classification model for predicting relative vulnerability to sleep deprivation.

Because of aforementioned dataset differences, it might be expected that the best set of features selected by the model would be dissimilar across the two datasets. However, we found that of the five features assessed as most optimal for Dataset 1, three were again selected for Dataset 2 (diffusion drift $\xi$, range of non-decision times $S_t$, and $\Delta RT > 250$). Using these features, CV5 accuracy of 82.2% was achieved. Importantly, when the classifier was trained on PVT data collected during baseline of the first study visit in Dataset 2, performance of the model was similar to that for data analyzed during the follow-up visit (79.4% accuracy). Because the second visit occurred more than 5 mo after sleep deprivation, our results suggest that baseline diffusion drift $\xi$ and range of non-decision times $S_t$ show stable between-subject differences over time. Although the best set of features might change depending on the type of classifier used, the type of heuristic used, and the definition used for categorizing subject vulnerability, our findings nonetheless demonstrate that DDM parameters, wavelet parameters, and $\Delta RT > 250$ are useful for predicting vulnerability to decline in psychomotor vigilance following total sleep deprivation under laboratory conditions.

**Differences Between Datasets**

In Dataset 1, participants were free to walk around when not taking the PVT, and they were exposed to ordinary room light. In Dataset 2, PVT data were collected under conditions that were conducive to poorer performance. Specifically, subjects were restricted to bed in a semi-reclined position in constant dim light. Also, in Dataset 2 we used baseline PVT data collected in the late afternoon, corresponding to the mid-afternoon dip in performance. By comparison, baseline PVTs in Dataset 1 were taken in the late evening, corresponding to the wake maintenance zone when the circadian drive to remain awake is near its peak. The aforementioned experimental differences may explain, in part, the slightly better classification rates in Dataset 2 using a smaller number of features.

**Definition of Vulnerability to Total Sleep Deprivation**

In our analysis, group assignment was based on a median split using the number of lapses in the last session of sleep deprivation. The number of lapses (RTs $> 500$ msec) is the most commonly used PVT metric to assess the effects of sleep deprivation on sustained attention. We acknowledge, however, that vulnerability could be defined using other PVT outcome measures, e.g., response speed, or by individualizing the lapse threshold relative to each person’s baseline performance. Dividing each dataset into two groups enabled us to build a model that predicts a binary outcome (resilient or vulnerable), but it
is important to note that vulnerability to sleep deprivation is a continuous variable. Hence, the decision to split the dataset by the median was arbitrary, and the ‘most vulnerable’ resistant subjects were qualitatively similar to the ‘most resistant’ vulnerable subjects, separated only by a few PVT lapses during sleep deprivation. Despite this limitation, subjects were classified with nearly 80% accuracy. Of note, although resistant and vulnerable groups differed by a few lapses even during the baseline rested state (Figure 2), the feature selection process did not select lapses in either dataset. Instead, vulnerability was better predicted by underlying decision and nondecision components of the DDM.

**Further Improvements in Classification**

The reliability of estimates for DDM parameters is affected by the size of the dataset\(^9,31\). To ensure that we had sufficient data for the DDM, we combined PVT RT data across two consecutive 10-min PVT sessions, separated by 1 h (Dataset 1) or 2 h (Dataset 2). DDM parameter estimates might be further improved by implementing longer PVT sessions (e.g., 20–30 min in duration) or by aggregating data across more PVT sessions. It must be kept in mind, however, that longer-duration PVTs are more affected by time-on-task effects\(^3\), and adding more PVT sessions might not be practical if our model is to be applied in real-world settings. The classification might also be more accurate if participants are studied under baseline conditions that are conducive to sleep. Adding other easily derived physiological measures, for instance, heart rate variability\(^52\), could also potentially improve the performance of the classifier.

**Figure 8**—Estimated drift diffusion model (DDM) parameters for vulnerable and resistant subjects estimated from baseline psychomotor vigilance task sessions measured across two study visits for Dataset 2. Of the 45 subjects who participated in the first study, 34 returned for a follow-up study after at least 5 mo following their initial visit to the laboratory. Baseline individual differences in mean diffusion drift and variability in nondecision time were reproducible across study visits. Vulnerable subjects had lower mean drift ($F_{1,32} = 11.6, P < 0.005$) and higher variability in nondecision time ($F_{1,32} = 13.6, P < 0.001$) across both visits compared to resistant subjects. The across-trial variability in diffusion drift appeared to be the only DDM parameter to be affected across the two studies ($F_{1,32} = 4.65, P < 0.05$). #, significant main effects of group on DDM parameters. Mean ± SEM are shown.

**Table 1**—Most optimal features selected for each dataset.

<table>
<thead>
<tr>
<th>Dataset 1</th>
<th>Dataset 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diffusion drift $\xi$ (51.9%)</td>
<td>$\Delta RT &gt; 250$ (44.4%)</td>
</tr>
<tr>
<td>MAV$_6$ (63.7%)</td>
<td>Diffusion drift $\xi$ (77.8%)</td>
</tr>
<tr>
<td>Range of non-decision time $S_r$ (65.2%)</td>
<td>Range of non-decision time $S_t$ (82.2%)</td>
</tr>
<tr>
<td>$\Delta RT &gt; 250$ (75.6%)</td>
<td>Variability in drift $\eta$ (77%)</td>
</tr>
</tbody>
</table>

The features are ordered according to their importance towards vulnerability prediction. Number in parenthesis show stratified 5-fold cross validation accuracy (CV5) achieved by including the corresponding feature as well as all features above it.
CONCLUSION
In this study, we built a classifier to predict vulnerability in sustained attention during sleep deprivation, using features derived from PVTs taken under rested baseline conditions. We included a range of features including several summary statistics, DDM parameters, and features derived from wavelet transform of the RT sequence. We found that DDM parameters, including decision and nondecision components, the number of consecutive RTs that differ by more than 250 msec, and wavelet features can be used to discriminate sleep deprivation vulnerable and resistant individuals with an accuracy of 77–82% across datasets.

ACKNOWLEDGMENTS
The authors thank Badrinarayanan Rangarajan and Ankit Das for technical discussions regarding best strategies for feature selection; and research staff and students in the Chronobiology and Sleep Laboratory and the Cognitive Neuroscience Laboratory for their assistance in carrying out these studies.

DISCLOSURE STATEMENT
The study was supported by the National Research Foundation/National Medical Research Council, Singapore under STaR 004/2008 (Chee); the Duke-NUS Signature Research Program funded by the Agency for Science, Technology and Research, Singapore; and the Ministry of Health, Singapore; the National Medical Research Council, Singapore under NIG/1000/2009 (Gooley); and the SingHealth Foundation, Singapore, under SHF/FG410P/2009 (Gooley). The authors have indicated no financial conflicts of interest. Work was performed at Cognitive Neuroscience Laboratory and Chronobiology and Sleep Laboratory, Duke-NUS Graduate Medical School, Singapore.

REFERENCES
52. Chua EC, Tan WQ, Yeo SC, et al. Heart rate variability can be used to estimate sleepiness-related decrements in psychomotor vigilance during total sleep deprivation. Sleep 2012;35:325–34.