Neural Markers of Responsiveness to the Environment in Human Sleep

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Sleep is characterized by a loss of behavioral responsiveness. However, recent research has shown that the sleeping brain is not completely disconnected from its environment. How neural activity constrains the ability to process sensory information while asleep is yet unclear. Here, we instructed human volunteers to classify words with lateralized hand responses while falling asleep. Using an electroencephalographic (EEG) marker of motor preparation, we show how responsiveness is modulated across sleep. These modulations are tracked using classic event-related potential analyses complemented by Lempel-Ziv complexity (LZc), a measure shown to track arousal in sleep and anesthesia. Neural activity related to the semantic content of stimuli was conserved in light non-rapid eye movement (NREM) sleep. However, these processes were suppressed in deep NREM sleep and, importantly, also in REM sleep, despite the recovery of wake-like neural activity in the latter. In NREM sleep, sensory activations were counterbalanced by evoked down states, which, when present, blocked further processing of external information. In addition, responsiveness markers correlated positively with baseline complexity, which could be related to modulation in sleep depth. In REM sleep, however, this relationship was reversed. We therefore propose that, in REM sleep, endogenously generated processes compete with the processing of external input. Sleep can thus be seen as a self-regulated process in which external information can be processed in lighter stages but suppressed in deeper stages. Last, our results suggest drastically different gating mechanisms in NREM and REM sleep.

Key words: complexity; EEG; NREM; REM; sensory processing; sleep
hypothesis proposed that behavioral unresponsiveness was achieved at an early stage, through the blockade of sensory information at the thalamic level (McCormick and Bal, 1994). However, familiar and salient stimuli trigger an awakening more easily (Oswald et al., 1960; Formby, 1967), suggesting the preservation of basic sensory processes. Even when sleep is preserved, familiar stimuli exhibit different brain responses (Perrin et al., 1999). In recent years, several studies have shown that external information processing may extend far beyond automatic operations: from the detection of semantic incongruity (Bastuji and Garcia-Larrea, 1999; Ibáñez et al., 2006) or the violation of simple rules (Ruby et al., 2008; Strauss et al., 2015) to the formation of new associations (Arzi et al., 2012; de Lavilleon et al., 2015). The cognitive processes involved during sleep therefore appear more elaborated than previously thought.

However, although these studies have demonstrated the preservation of cognitive abilities during sleep, the neural mechanisms allowing sleepers to process or isolate from external stimulations remain unse t t l e d. T h e role of evoked slow oscillations, such as K-complexes, a hallmark of NREM sleep, is particularly unclear (Halász, 2005). K-complexes have sometimes been related to arousal systems (Siclari et al., 2014) providing windows of wakefulness to sleepers (Destexhe et al., 2007). Conversely, they have also been described as local down states (Cash et al., 2009) entailing large-scale neuronal silencing (Vyazovskiy and Harris, 2013) and protecting sleep against external stimulations (Wauquier et al., 1995; Bastien et al., 2000). This tendency neurons have to alternate between periods of activations (up states) and silencing (down states) (Sanchez-Vives and McCormick, 2000) is termed “neuronal bistability” and has been proposed as a mechanism limiting sensory processing at the cortical level (Tononi and Massimini, 2008; Figorini et al., 2015). Sleep spindles are also thought to enable sleepers’ isolation by blocking incoming input at the thalamic level (McCormick and Bal, 1994).

However, evidence supporting this hypothesis is scarce (Schabus et al., 2012), whereas cellular recordings showed remarkably preserved responses to sounds during spindles (Sela et al., 2016).

REM sleep, on the other hand, has been far less studied. Paradigmatically, even if brain activity in REM sleep resembles wakefulness and consciousness is regained (Hobson and Pace-Schott, 2002), sleepers remain largely unresponsive (Ermis et al., 2010). It has been proposed that dreams themselves would compete and block the processing of external inputs (Nir and Tononi, 2010), but direct evidence is still missing.

Here, to explore responsiveness during human sleep, we relied on a paradigm that aimed at inducing task-dependent responses in the sleeping brain during naps (Kouider et al., 2014). In practice, participants fell asleep while categorizing spoken words with lateralized responses. Using EEG recordings, we explored whether stimuli elicited brain activations corresponding to the motor preparation of the correct response (i.e., lateralized readiness potential [LRP]) (Masaki et al., 2004; Smulders et al., 2012). The presence of such LRP implies that stimuli were not only encoded at the sensory level but also processed at a high level of semantic representation and propagated up to the preparation of a motor response (see Kouider et al., 2014). The presence of an LRP highlights the maintenance of complex distributed processes during sleep. Although our previous study was limited to daytime sleep (nap study), here we investigated all sleep stages in a full-night protocol. Furthermore, the presence and magnitude of the LRP was further evaluated in light of brain activity before and after the stimulus so as to provide novel insights on how the brain manages to alternatively track its environment or rather preserve sleep.

Materials and Methods

Participants

Twenty-three right-handed French native speakers (16 females, age 21–31 years) with no history of neurological or sleep disorders participated in this study. Participants underwent an interview with a sleep specialist and filled in questionnaires determining their sleep habits and their propensity to fall asleep in noisy environments. Participants did not complain of any sleep disorder. They were monitored for 7–10 d before the recording session through actigraphy and sleep diaries to ensure regular sleep/wake rhythms. Among these 23 participants, 5 participants (4 females) were discarded from our analyses either for technical issues affecting the recordings (N = 1) or because the experiment was aborted (N = 4, participants experiencing difficulties to fall asleep with auditory stimulations). The protocol had been approved by the local ethics committee (Comité de Protection des Personnes, Ile-de-France I, Paris, France).

Experimental procedure

Task. We adapted a procedure previously used during daytime naps (Kouider et al., 2014) to a full-night protocol to explore all sleep stages. On the night of the recordings, participants were equipped for polysomnographic recordings. They went to bed, and spoken words in French were then played in isolation one after the other. These words referred either to an animal or to an object. Participants were instructed to perform a semantic-decision task by indicating the category of each word through right- and left-hand responses. They were asked to categorize words as long as they were awake and to resume responding in case of an awakening. Subjects were reminded to do so whenever they awoke during the night without resuming to respond. However, participants were explicitly authorized to fall asleep while performing the task. Crucially, three different lists of words were played to participants according to their vigilance state, as assessed through an online assessment of sleep stages (see below). A list was played whenever participants were in NREM sleep, another list was played during REM sleep, and a wake list was played otherwise (i.e., mainly during wakefulness). Thus, unpracticed words (i.e., novel words not previously categorized during wakefulness) were played in sleep to ensure that participants had to access the meaning of the word to prepare for the appropriate response, without relying on stimulus-response associations learned while awake.

Stimuli. Stimuli were French spoken words uttered by a female voice. Lists of 72 words were created each containing 36 words referring to animals and 36 words referring to objects. Animal and object words were matched infrequency and number of syllables using the Lexique database (New et al., 2004). The attribution of the different lists to a given vigilance state was counterbalanced across participants. Participants received on average 21.7 ± 1.0 (mean ± SEM across participants), 25.2 ± 0.7, and 6.9 ± 0.3 times the words from the wake, NREM, and REM lists, respectively.

Apparatus. Response-handles were attached to participants’ hands. The mapping between semantic categories and response-hands was counterbalanced across participants. Stimuli were played through loud speakers at ~55 dB (depending on participants’ preferences) using the Psycho-toolbox extension (Brainard, 1997) for MATLAB (The MathWorks). Stimuli were played every 6–9 s (random uniform jitter).

Data analysis

Electrophysiological recordings. EEG (N = 19 derivations, 10–20 montage), electro-oculographic (EOG, N = 2 derivations, positioned above the right canthus and under the left canthus), electromyographic (EMG, 1 derivation placed on the chin measuring muscle tone and 2 derivations on the right and left abductor pollicis brevis [thumb flexor muscle] recording muscle activity accompanying participants’ responses) and electrocardiographic (ECG, N = 1 derivation) data were continuously recorded. Video monitoring was also available. EEG, EOG, ECG, and EMG data were recorded with AgCl electrodes attached to participants’ skin and hair with an adhesive paste (EC2, Natus Neurology). Signals
EEG channels were first rereferenced to the average mastoids. The EEG in alpha oscillations or oscillations and REM sleep. Importantly, epochs showing signs of arousal (increase in theta oscillations or sleep spindles, absence of alpha oscillations, presence of saw-tooth waves, rapid eye movements, increase in theta oscillations, reduced or absent muscle tone). The experimenter typically waited for a few minutes after the transition to the new sleep stage before switching the list, so as to ensure sleep stage’s stability. The wake list was presented to participants whenever they showed signs of arousal (body movements, increase in low-amplitude desynchronized rhythms, EMG activation).

This online scoring was confirmed and refined offline (see below).

Offline sleep scoring. Wakefulness and sleep stages were scored by 2 scorers blind to experimental conditions and following established guidelines (Iber et al., 2007). Only Fz, C3, C4, and Pz EEG derivations (within the 10–20 montage) were used along EOG and EMG derivations. EEG channels were first referenced to the average mastoids. The EEG and EOG signals were high-pass filtered >0.1 Hz and then low-pass filtered <30 Hz (two-pass Butterworth filters at the fifth order). Using the same filter types, EMG data were bandpass filtered between 60 and 80 Hz. A notch-filter (second order Infinite Impulse Response filter) ~50 Hz was used on all channels to reduce line-noise. Vigilance stages were continuously scored on 20 s-long windows as wakefulness, NREM sleep Stages 1, 2, and 3 (NREM1, NREM2, and NREM3 stages, respectively), and REM sleep. Importantly, epochs showing signs of arousal (increase in alpha oscillations or oscillations >16 Hz lasting >3 s) or microarousal (<3 s) in association with trial onsets were marked. The corresponding trials were not included in further analyses to avoid potential confounds. Table 1 summarizes sleep scoring across participants.

Slow wave detection. Slow oscillations (slow waves and K-complexes) were detected in NREM sleep using an algorithm that has been presented in details previously (Riedner et al., 2007; Nir et al., 2011). Briefly, slow oscillations were detected for each EEG channel by bandpass filtering the EEG signal between 0.2 and 3 Hz (two-pass Butterworth filter at the third order). The first-order derivative was used to detect local extrema and identify single waves. Only slow oscillations with a peak-to-peak amplitude exceeding 75 μV and a duration of >0.5 s and <2 s were considered as slow waves. These slow oscillations were used to define light and deep NREM trials (see below).

EEG preprocessing. EEG data were analyzed using a combination of SPM (Functional Imaging Laboratory, University College London, London), FieldTrip (Oostenveld et al., 2011), and EEGLab (Delorme and Makeig, 2004) toolboxes running on MATLAB (The MathWorks). We examined the brain activity in response to sounds by computing event-related potentials (ERPs) and LRPs. To do so, the continuous EEG signal referenced to the average mastoids was high-pass filtered >0.1 Hz (two-pass Butterworth filter at the fifth order) and then segmented on large temporal windows around stimulus onsets ([-14, 14] s). The EEG signal was then low-pass filtered <40 Hz and notch-filtered ~50 Hz (same filter types as for the slow scoring) to reduce line noise. Epochs were resliced to focus on the activity around stimulus onsets ([-2, 7] s). Trials with absolute amplitude over central electrodes (Cz, C3, C4) exceeding a given threshold were discarded (150 μV in wake and REM sleep, and 200 μV in NREM sleep because NREM sleep contains high-amplitude physiological slow waves).

Conditions of interest. In NREM sleep, brain activity varies immensely: from NREM1 in which slow hallmarks (slow oscillations and sleep spindles) are not yet visible to NREM3 where they predominate (Iber et al., 2007). First, to ensure that observed sensory processes are not due to (even brief) awakenings, NREM1 stage was discarded as well as trials in which any sign of arousal could be observed in the EEG signal. Second, NREM2 and NREM3 trials were separated according to the presence of slow waves associated with stimulus onset. We thus divided trials in four different conditions of interest: (1) wake: correct trials during which participants were scored as awake and the wake list was played; (2) light NREM sleep: trials scored as NREM2 or NREM3, during which participants were unresponsive and in which the NREM list was played but no slow-wave were detected (on a [-2, 3] s window time-locked to stimulus onset); (3) deep NREM sleep: trials scored as NREM2 or NREM3, during which participants were unresponsive and in which the NREM list was played and slow waves were detected (on a [-2, 3] s window time-locked...
to stimulus onset); (4) REM sleep: trials scored as REM sleep, during which participants were unresponsive and in which the REM list was played. Thus, light NREM, deep NREM, and REM trials comprised only words that had not been practiced before, when participants were awake and responsive (no wake list word). For each condition, only participants with at least 70 trials were included. On average, there were 174 ± 20 trials in the wake condition (mean ± SEM across N = 17 participants), 232 ± 28 in light NREM sleep (N = 15), 400 ± 25 in deep NREM sleep (N = 18) and 186 ± 9 in REM sleep (N = 18). Light NREM trials occurred almost exclusively in NREM2 (93 ± 1.2%, N = 15 participants), whereas deep NREM trials correspond to trials occurring either in NREM2 or NREM3 stages (56% and 44%, respectively, ± 3.5% in 18 participants). Thus, light NREM sleep trials corresponded to a state in which slow oscillations could be observed (NREM2) but not in the vicinity of stimulus onset, whereas deep NREM sleep trials corresponded to a state in which slow oscillations were observed in association with stimuli regardless of the presence of continuous trains of slow waves (e.g., in NREM3). Nonetheless, we obtained qualitatively similar results (see Results) when defining light and deep NREM sleep as NREM2 and NREM3 stages, respectively. We also examined trials corresponding to the definition of light, deep NREM, and REM sleep, but during which the wake list was presented to sleepers instead of the NREM or REM lists. Indeed, participants were switched to the wake list whenever the experimenter was unsure of the state of the sleeper (later verified off-line), at transitions between states or when the experimenter had to attend to another participant (two participants participated in the experiment on each recording night). Although these trials containing practiced words were not randomly intermixed with the unpracticed words, they were scored offline as either REM or NREM sleep by two scorers blind to the list used and revealed ERPs and power spectra nearly identical to the ones obtained in trials in which the sleep lists were played (see Fig. 3). Thus, these trials allowed us to investigate how practiced (i.e., overtly categorized) words were processed during sleep, although precautions must be taken when interpreting such results (see Discussion). Participants with at least 30 trials per condition were included for this analysis.

ERPs. ERPs were computed by averaging the EEG signal across trials for a given experimental condition after baseline correction ([−0.2, 0] s).

LRPs. LRPCs allow the monitoring of action selection and preparation (Smulders et al., 2012). LRPCs are usually computed with EEG data time-locked to motor responses but can also be computed time-locked to stimuli (Leuthold, 2003; Töllner et al., 2012). In our case, due to the absence of responses during sleep, we computed LRPCs on stimulus-locked data. LRPCs were computed using ERPs recorded from the right (C4) and left (C3) electrodes placed over motor cortices as follows:

$$LRP = \frac{(C_{\text{right-hand}} - C_{\text{left-hand}}) + (C_{\text{left-hand}} - C_{\text{right-hand}})}{2}$$

ERPs over C3 and C4 electrodes were computed similarly as described above, except for the baseline correction (−2, 0] s). Using this formula, LRPCs are characterized by a negative deflection starting before participants’ response. In subsequent analyses, we extracted the LRP magnitude over the temporal windows in which significant negative clusters were observed (see Figs. 2, 3). To examine how this LRP magnitude was dynamically related to other markers of responsiveness, we computed LRPCs on windows of 60 consecutive left- and right-response trials slid every trial (see Figs. 4, 6, 7). The LRP-negative potential was extracted as a positive value (LRP magnitude) by multiplying the LRP formula by −1. A similar operation was performed on the N500 and N550 potentials amplitude (see Figs. 6, 7) to extract their magnitude. Other variables of the EEG signal were estimated on the same windows such as the Lempel-Ziv complexity (LZc; see Fig. 4) or ERP components (see Figs. 6, 7). These variables were z-score normalized across trials for each participant and then aggregated across participants to examine their correlation.

Time-frequency analyses. To better understand how sleep rhythms impact sensory processing, we computed the time-frequency decomposition of the EEG signal in response to stimuli. To do so, we applied FFT on 1.28-s-long windows (padding ratio of 2) on the preprocessed EEG signal (see above). The resulting power for each frequency and time was expressed as the ratio of the power at the corresponding frequency and time over the baseline activity ([−1.5, 0] s) at the same frequency. LRP magnitude and the activity within the slow waves ([11, 6] Hz) and spindle ([11, 16] Hz) bands were extracted on windows of 60 consecutive right-hand and left-hand responses.

LZc. The LZc measures the complexity of a given signal by estimating its compressibility (Ziv and Lempel, 1977). A temporally unpredictable signal will have a low compressibility and therefore a high LZc value. Such an unpredictable signal will be considered as a highly complex signal. Previous studies have shown that LZc accurately tracks the level of consciousness in patients, healthy subjects under anesthesia and during sleep (Casali et al., 2013; Abásolo et al., 2015; Schartner et al., 2015). For example, when the EEG signal is populated by high-amplitude synchronous slow waves as in deep NREM sleep, the signal becomes more predictable and the associated LZc value decreases compared with wakefulness or REM sleep (see Fig. 4d). To calculate the LZc on the continuous EEG, we implemented the approach developed by Schartner et al. (2015). Thus, the complexity was computed at the sensor level. To filter the raw EEG, the MATLAB FDAtool was used to create equiripple Finite Impulse Response FIR filters with linear phase to avoid phase distortion in the EEG. The EEG was first low-pass filtered (<85 Hz (11th order) and then notch-filtered (~50 Hz to reduce line noise (296th order). A surface Laplacian was then applied to the EEG data (BCILAB plug-in for the EEGLAB toolbox) (Delorme and Makeig, 2004; Kober and Makeig, 2013), with a neighbor count of four, to reduce the influence of volume conduction. The data were then epoched around stimulus onset ([−2, 6] s), and the LZc was extracted for each trial by integrating the EEG data over all the sensors on 500-ms-long windows slid every 50 ms. Details of the LZc algorithm can be found previously (Schartner et al., 2015). Briefly, for each window, the mean of the EEG signal was subtracted for each sensor and linear trends over the entire epochs were removed. A Hilbert transform was then applied to the signal to extract its envelope. Each channel was then binarized: values above the mean value of the envelope for the corresponding epoch and sensors were coded as 1, and 0 otherwise. The resulting binary matrix was then reshaped sensor-wise into a vector containing the time-points for all channels. The complexity was computed on such vectors and normalized by the complexity calculated on a randomly shuffled version of the same vector. After normalization, LZc takes values between 1 (minimally compressible, i.e., as predictable as the shuffle data) and 0 (fully compressible, i.e., predictable signal). The normalized complexity was thus computed for each trial and averaged across vigilance states (as scored offline) over the prestimulus baseline ([−1.5, 0] s) to compare the overall level of complexity between these different states (see Fig. 4a). We also examined how the complexity was affected by stimulation by averaging the LZc on a [−1.5, 6] s window for each vigilance state separately in an approach similar to when calculating ERPs (see Fig. 4b). To better compare the dynamics, LZc was normalized (ratio) by the level of complexity in the prestimulus window

### Table 1. Sleep scoring and conditions of interest

<table>
<thead>
<tr>
<th></th>
<th>Wake</th>
<th>NREM1</th>
<th>NREM2</th>
<th>NREM3</th>
<th>REM</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duration (min)</td>
<td>88.5 (±9.6)</td>
<td>45.3 (±5.6)</td>
<td>186.3 (±13.4)</td>
<td>79.7 (±7.3)</td>
<td>76.9 (±6.5)</td>
<td>499.6 (±6.6)</td>
</tr>
<tr>
<td>% of trials</td>
<td>17.8 (±1.6)</td>
<td>9.7 (±1.0)</td>
<td>39.6 (±2.0)</td>
<td>16.4 (±1.4)</td>
<td>16.5 (±0.8)</td>
<td>100</td>
</tr>
<tr>
<td>Marked arousals</td>
<td>0</td>
<td>12.3 (±1.6)</td>
<td>13.4 (±1.8)</td>
<td>2.6 (±0.5)</td>
<td>7.0 (±1.5)</td>
<td>35.8 (±3.8)</td>
</tr>
</tbody>
</table>

*Data mean (SEM), N = 18. Participants’ vigilance was scored according to established guidelines as wakefulness, NREM1, NREM2, NREM3, and REM stages on 20-s-long windows (see Materials and Methods). Duration indicates the average time participants spend in each stage. Each trial was attributed a sleep stage defined as the most conservative scoring obtained on a [−2, 8] s window around stimulus onset (% of trials). Marked arousals indicates the mean number of trials that were rescored as wakefulness due to an arousal.*
Statistics. To correct for multiple comparisons when examining statistical differences between two time series (see Figs. 2–5), we used a cluster-permutation approach (Maris and Oostenveld, 2007). Each cluster was constituted by the samples that consecutively passed a specific threshold (here, p < 0.1 for LRPs and p < 0.05 otherwise). As demonstrated previously (Maris and Oostenveld, 2007), this method controls for Type 1 errors independently of this threshold. For each cluster, we computed the sum of the t values of all the samples within the cluster. Then, we compared the cluster statistics of each cluster with the maximum cluster statistics of 1000 random permutations and obtained a nonparametric p value (Monte-Carlo p value; $p_{\text{cluster}}$). Significant clusters are displayed as horizontal bars on plots, and $p_{\text{cluster}}$ values are reported in the main text and figures’ legends. For scalp topographies of correlation analyses (Figs. 6, 7), the False Detection Rate (FDR) method was used to correct individual p values (Benjamini and Yekutieli, 2011).

Results
Sleepers can classify words during sleep
A classical LRP was observed when participants were awake and responsive (Fig. 2a; significant cluster: [0.490, 2.025] s, $p_{\text{cluster}} = 0.002, N = 17$ participants), characterized by a large negative deflection starting before the average response time (black arrow) over motor electrodes (Fig. 2a, inset). When focusing on light NREM sleep, we observed a similar significant negative deflection for the LRP (Fig. 2c; significant cluster: [3.035, 3.775] s, $p_{\text{cluster}} = 0.01, N = 15$ participants), although the corresponding words had not been previously categorized during wakefulness. We obtained a similar LRP even when discarding all words that had been previously presented during an even brief ($<3$ s) arousal (cluster: [2.830, 3.310] s, $p_{\text{cluster}} = 0.043, N = 15$ participants), thus when focusing on words that had been presented exclusively during sleep throughout the entire recording session. As in wakefulness, this negative deflection was maximal over motor cortices (central electrodes; see Fig. 2c, inset) but was delayed compared with wake trials. This LRP is strikingly similar to the one observed in previous nap studies (Kouider et al., 2014).

However, no significant deviation in the LRP was observed in deep NREM sleep or REM sleep (Fig. 2b,d), suggesting, at first glance, that processing of sensory information up to the decision level is abolished in deeper stages of sleep. Interestingly, light and deep NREM sleep also differed regarding the evoked responses to stimuli (ERPs). Light NREM sleep was characterized by a reduction of the large negativity evoked by stimuli in NREM sleep (N550) (Picton, 2010). This N550 is often attributed to evoked
slow waves (such as K-complexes) (Bastien et al., 2000), which entails large-scale neuronal silencing (Vyazovskiy and Harris, 2013) and could explain the absence of an LRP when the N550 is larger. Importantly, defining light and deep NREM sleep more classically as NREM2 and NREM3 stages, respectively, led to similar results (i.e., presence of an LRP in light NREM ([4.1, 4.6] s, $p_{\text{cluster}} = 0.038$) and absence in deep NREM).

In REM sleep, only words previously heard in wake are classified

Does the absence of an LRP in REM sleep mean that stimuli were not processed at all, or is this absence due to the complexity of the task at hand? To investigate this issue, we took advantage of the fact that words from the wake list were sometimes presented in REM sleep (see Materials and Methods), albeit not intermixed with unpracticed words, which would have allowed a stronger comparison (see Discussion).

Importantly, the ERPs and power spectra were similar when the practiced and unpracticed words were played (Fig. 3), but a clear LRP was observed for the wake list only ([2.750, 3.200] s, $p_{\text{cluster}} < 0.05$) but not for unpracticed words (black bar represents cluster for the comparison between the LRPs for the wake and REM lists, $p_{\text{cluster}} < 0.05$). Yet, the power spectrum and ERPs for both practiced and unpracticed words are highly similar and different from wake trials (red curves). Shaded areas represent the SEM computed across participants.
The processing of information depends on the degree of neural complexity

We then set out to examine whether the overall complexity of the pre-stimulus EEG signal was predictive of sleepers’ ability to process information up to the decision level. To do so, we used a recently developed approach (Casali et al., 2013) consisting in measuring the temporal predictability of the EEG signal by reducing it to a single value: the LZc (Ziv and Lempel, 1977). Applying this methodology to our data (Fig. 4a) revealed a strong modulation of LZc according to the vigilance state (ANOVA: $F(3) = 9.67, p = 2 \times 10^{-3}$). Such modulation indicates that, in wakefulness, the EEG signal is maximally complex (i.e., unpredictable), whereas, in deep NREM sleep, the EEG signal becomes the most predictable and therefore less complex according to the Lempel–Ziv algorithm. We could separate sleep and wake stages along a gradient matching the phenomenology associated with these states (Nir et al., 2013), confirming initial results in humans (Casali et al., 2013) and animals (Abásolo et al., 2015). Precisely, the baseline LZc was maximal in wakefulness (paired t tests across participants with other sleep stages: all $p < 0.005$) and minimal in deep NREM sleep. Light NREM and REM sleep had intermediary values, with REM sleep being the closest to wakefulness (REM vs wake: $t_{(17)} = -3.3$, $p = 0.004$; REM vs light NREM: $t_{(17)} = 5.4$, $p = 5 \times 10^{-3}$; light vs deep NREM: $t_{(17)} = 9, p = 7 \times 10^{-8}$).

We then investigated whether this baseline LZc was predictive of LRP magnitude. LRP magnitude was extracted on the temporal windows in which a LRP was observed in wakefulness (Fig. 2a), light NREM sleep (Fig. 2c), and REM sleep (when practiced words were played; Fig. 3). Importantly, the baseline LZc was positively correlated with LRP magnitude in wakefulness and light NREM sleep (Perron’s coefficient: $r = 0.08$ and 0.13, $p < 0.005$ across participants, respectively), but this correlation was reversed in REM sleep ($r = -0.23$, $p = 5 \times 10^{-24}$; $N = 1937$ samples in 17 and 15 participants, respectively). Comparing LZc in REM sleep when practiced words were presented (LRP present) and when unpracticed words were presented (LRP absent) revealed again that the presence of the LRP was associated with a lower complexity (paired t test: $t_{(17)} = 3.5$, $p = 0.004$). This result contrasts with the correlation observed in NREM sleep and wakefulness, suggesting an inverse relationship between levels of consciousness and the degree of connectedness to the environment (Sanders et al., 2012) in NREM and REM sleep.

We also examined how the LZc was modulated in response to stimuli above and beyond its baseline level (Fig. 4b). Stimulus onset was followed by a decrease in LZc (i.e., increase in signal predictability), potentially due to the stereotypical ERPs. And indeed, this decrease in LZc was more pronounced for the sleep stages in which high-amplitude ERPs were observed, such as in NREM sleep (Fig. 2). Accordingly, this decrease in LZc did not reach significance in REM sleep, a state in which auditory stimulation has less impact on brain activity (Bastuji and Garcia-Larrea, 1999; Picton, 2010). Interestingly, in NREM and REM sleep, this initial decrease was followed by an increase in LZc.
across sensors to emphasize regional differences. The decrease associated with the LRP is centrally distributed for slow waves and sleep spindles despite their originally frontal distribution, an increase within the slow-wave (deep NREM) for electrode Cz. Stimulus onset was accompanied by the EEG signal in response to sounds in NREM sleep (both light and bands associated with wakefulness (alpha: [9, 11] Hz, beta: [11, 16] Hz: spindle range, gray curve). Horizontal bars represent the significant clusters determined across participants (pCluster < 0.05). Shaded areas represent the SEM computed across participants. Insets, Scalp topographies of the power within the slow-wave and spindle ranges at trial onset ([0, 2] s) and during the LRP window ([2.9, 3.8] s). Power was z-scored across sensors to emphasize regional differences. The decrease associated with the LRP is centrally distributed for slow waves and sleep spindles despite their originally frontal distribution, suggesting a local suppression of sleep rhythms. Sensory processes locally disrupt sleep rhythms

So, are the LRP s observed in sleep and the associated increase in Lz reflecting the consequence of a partial awakening? To answer these questions, we examined the consequences of auditory stimulation on sleep itself. Figure 5a shows the time-frequency decomposition of the EEG signal in response to sounds in NREM sleep (both light and deep NREM) for electrode Cz. Stimulus onset was accompanied by an increase within the slow-wave (< 6 Hz; pCluster = 0.001, N = 18 participants) and spindle ([11, 16] Hz; pCluster = 0.002) ranges (Fig. 5b). This increase was followed by a decrease within these two ranges (< 6 Hz: [2.1, 6.3] s; [11, 16] Hz: [2.4, 5.0] s, both pCluster < 0.001), interestingly, at a time corresponding to the appearance of an LRP in light NREM sleep. This result suggests that the preparation of task-related responses translates into a local-in-time modulation of sleep rhythms and hence of sleep depth. Such modulation was also local-in-space. Indeed, while the increase within the slow-wave and spindle range after stimulus onset showed a typical frontal topography (Fig. 5b), the following decrease had a different topography and was prominent over central electrodes (i.e., over the sensors showing the LRP). It is important to note that this decrease within the slow-wave and spindle ranges was not accompanied by an increase in frequency bands associated with wakefulness (alpha: [9, 11] Hz, beta: >16 Hz), which is concordant with the fact that trials with signs of arousal were discarded from our analyses. Thus, in NREM sleep, the processing of auditory information resulted in a local (in time and space) modulation of NREM oscillations in the absence of any observable trace of awakening on the scalp.

In REM sleep, brain responses were qualitatively different (Fig. 5c,d): auditory inputs clearly disturbed REM sleep with an initial increase in higher frequencies (> 12 Hz, pCluster = 0.01). This increase was followed by a sustained decrease within the theta range ([4, 8] Hz, pCluster < 0.001), reflecting a stimulus-induced perturbation of theta oscillations, which are a hallmark of REM sleep in animals (Buzsáki, 2006). Interestingly, these increases were both maximal over central electrodes where auditory and motor components are observed in wakefulness (Picton, 2010; Smulders et al., 2012). However, these transient markers of arousal did not lead to the appearance of a clear LRP (Fig. 2) unless practiced words were presented (Fig. 3).

Neuronal bistability gates sensory processing in NREM sleep

In NREM sleep, brain responses to stimuli were characterized by strong evoked potentials, such as the P200 and the N550 (Fig. 6a). These two potentials are of high interest because the P200 is thought to reflect an activation within the primary sensory cortices corresponding to the eliciting event (i.e., dependent of the sensory modality), whereas the N550 is thought to reflect a broad modality-independent neuronal silencing, maximal in frontal areas (Laurino et al., 2014; Halász, 2015). The respective scalp topographies of these potentials corroborated such view (Fig. 6a, right). According to this interpretation, the P200 could represent a marker of the brain’s responsiveness to external events, whereas the N550 could be associated with sleep protection against external perturbations.

We thus examined whether the auditory responses (evoked-potentials associated with stimulus onset) were predictive of the appearance of an LRP. Interestingly, in light NREM sleep, the...
Neural bistability gates sensory processing in NREM sleep. Two distinct potentials are clearly visible: a positivity ~200 ms (P200) maximal at centroparietal electrodes (see scalp topography on the top right); and a negativity ~550 ms (N550) predominant in deep NREM sleep (trials associated with slow waves) and maximal at frontal electrodes (see scalp topography on the bottom right). Shaded areas represent the SEM computed across participants (N = 18). Scalp topographies were established by averaging the voltage over windows around the two potentials of interest (see gray areas on ERP plot). These values were averaged across participants and z-scored across channels to emphasize regional differences. Correlations between the LRP magnitude and the P200 magnitude (left) or the N550 magnitude (right, opposite of amplitude) for trials in light NREM sleep (Fig. 6a). This correlation was maximal over centroparietal electrodes. Thus, the larger the P200, the larger the LRP in light NREM sleep. A much less robust correlation was observed with the magnitude of the N550 (r = −0.04, p = 0.02), which could be explained by the fact that, in light NREM, the N550 was almost abolished.

On the other hand, in deep NREM sleep, there was a clear negative correlation between the N550 and the LRP magnitude (Fig. 6c; r = −0.10, p = 10−4; N = 6037 samples in 18 participants). Thus, the more pronounced the N550, the smaller the LRP in deep NREM sleep. Interestingly, the relationship between LRP magnitude and the P200 was reversed between light and deep NREM sleep. Contrary to light NREM sleep, the LRP magnitude was negatively correlated with the P200 magnitude (Pearson coefficient at Cz: r = −0.12, p = 10−20) despite the fact that the P200 increases in amplitude from light to deep NREM sleep (Fig. 6a), which should favor sensory processing according to the positive relationship observed in light NREM sleep. Such reversal could be due to the fact that the P200 activation can trigger a down state (N550) in deeper stages of NREM sleep, which would ultimately inhibit information processing and reverse the relationship between the P200 and the LRP (see Discussion).

Auditory evoked potentials are predictive of motor preparation in REM sleep

We applied the same approach to REM sleep. Auditory stimuli in REM sleep also evoked archetypal potentials: a positivity ~200 ms followed by a negativity ~500 ms (Fig. 7a). However, the topography and temporal profile of these potentials are quite different from NREM potentials. In REM sleep, the P200 is usually associated with the wake auditory P200, generated over the primary and secondary cortices (Picton, 2010). The scalp topography of the P200 with a maximum over central electrodes is concordant with this view. Contrary to light NREM sleep, the P200 was negatively correlated with the
LRP magnitude in REM sleep (Pearson coefficient at Pz: \(r = -0.14, p = 4 \times 10^{-11}, N = 2622\) samples in 18 participants; Fig. 7b). This negative correlation was maximal over occipital electrodes.

The N500, a potential that is prominent over occipital electrodes in REM sleep, was positively correlated with LRP magnitude (Pearson coefficient at Pz: \(r = 0.15, p = 3 \times 10^{-14}\); Fig. 7c), an opposite relationship compared with the one found for the N550 potential in NREM sleep. Similar correlations were observed when considering the REM trials in which words practiced in wake were played (P200: \(r = -0.12, p = 8 \times 10^{-4}\); N500: \(r = 0.15, p = 7 \times 10^{-15}\)). These results indicate that the evoked potentials associated with auditory stimuli are predictive of later and more complex stages of processing (here motor preparation) and suggest that these auditory potentials are not an unsppecific reaction to external stimuli but rather reflect cortex’s responsiveness to these inputs.

**Markers of responsiveness are modulated within sleep cycles**

The presence of LRP in light NREM but not deep NREM sleep or REM sleep (when unpracticed words are presented) could be due to the fact that light NREM sleep is more pervasive to external information. Another interpretation would be that the ability to prepare for the adequate motor response slowly decays with the time spent asleep. With light NREM sleep occurring first, the LRP would be prominent in this state. To test this possibility, we examined how the LRP magnitude was modulated within sleep cycles. Sleep cycles were detected using participant’s hypnograms (see Materials and Methods). For each cycle, we retrieved the dynamics of the LZc, the slow-wave power (i.e., power, a proxy for slow-wave density), and the LRP magnitude in the NREM part of the cycle (Fig. 8). In accordance with the archetypal profile of sleep cycles, slow-wave power density gradually increased at the beginning of the sleep cycle (descending slope) but decreased toward the end of the cycle (ascending slope and transition to REM sleep) (Halász, 2015). The LZc mirrored this pattern with a gradual decrease in overall complexity followed by a steep increase. Interestingly, the LRP magnitude followed the LZc dynamics and increased again toward the end of the cycle (U shape). This result suggests that the capacity to process information is dynamically related to signal complexity and neural bistability. Thus, the beginning and end of NREM episodes might represent temporal windows in which monitoring of the surrounding environment is possible.

We also examined the effect of the progression within the night on LRP magnitude. To do so, each night was divided into thirds and the LRP magnitude was computed in light NREM, deep NREM, and REM trials in each third separately. In light NREM sleep, a linear decrease was observed across the night (linear regression: \(\beta = -0.41, p = 0.002, N = 15\) participants). A post hoc comparison showed a highly significant difference between the LRP magnitude in the first and last third of the night \((t_{(14)} = 4.03, p = 0.001)\) with only the first third of the night showing a significant LRP \((t_{(14)} = 3.15, p = 0.007)\). Such within-nights effect was not observed in deep NREM or REM trials (linear regressions: \(\beta = -0.20, p = 0.44\) and \(\beta = -0.08, p = 0.58\), respectively). However, because sleep lists were repeated across the night, it is impossible to disentangle in this study a potential effect of early and late sleep from an effect of word-repetition.
The ability to process information is dynamically modulated within sleep cycles. Modulation of the LRP magnitude (colored dots), the LZc (black curve), and the δ-power (gray curve) within the NREM sleep part of sleep cycles. Colors of dots (LRP magnitude) represent the proportion of light and deep NREM trials included in the corresponding bin. A classical increase in δ-power (a proxy for slow-wave density) is observed corresponding to the transition from light to deep NREM sleep. This increase in δ-power is accompanied by a decrease in LZc and LRP magnitude. Both LZc and LRP have the tendency to increase again at the end of the NREM cycle, paralleling the transition from deep NREM sleep to REM sleep. LRP, LZc, and δ values were estimated within each sleep cycle on fixed windows (see Materials and Methods). Sleep cycles were then binned (n = 30) so as to average cycles with different durations, and values were normalized across the entire cycle to better visualize the dynamics of each variable of interest (expressed here in arbitrary units [a.u.]).

Discussion

Complex and distributed processes in the sleeping brain

Participants had to categorize spoken words based on their semantic category through lateralized hand responses while falling asleep. To assess participants’ responsiveness, we computed an LRP (Smulders et al., 2012), which indicates whether brain activity is lateralized according to the expected response side. Accordingly, a negative deflection was observed in wakefulness around response time (Fig. 2a). This negativity was conserved in light NREM sleep when unpracticed words were played despite the absence of overt responses (Fig. 2c), replicating previous findings in naps (Kouider et al., 2014). Because the stimulus-response mapping was counterbalanced across participants, the presence of an LRP for novel words in light NREM sleep reflects the maintenance of complex and distributed processes going from the encoding of the auditory information to the preparation of the appropriate motor response. The maintenance of such complex processing chain could be explained by the automation of the task-set during wakefulness (Kouider et al., 2014).

No LRP was observed in deep NREM sleep or REM sleep (for unpracticed words), suggesting that part of this chain is disrupted in these stages. However, when considering practiced words (wake list, Fig. 1), an LRP was observed in REM sleep (Fig. 3). The presence of an LRP for practiced words only could be interpreted as a failure to access the meaning of novel words during the REM period. Under such circumstances, sleepers could still rely on stimulus-response contingencies learned during wakefulness for practiced words. However, because the practiced and unpracticed words were not intermixed, we cannot rule out the possibility that the presence or absence of an LRP stems from a difference in terms of sleep (i.e., sleep depth) rather than stimulus type (practiced or not).

The complexity of brain dynamics interacts with the processing of the environment

Certain periods of sleep proved more propitious for the induction of task-dependent responses. To understand the relationship between the background neural activity and brain’s responsiveness, we computed the temporal predictability of the EEG signal using LZc (Ziv and Lempel, 1977). LZc has been used to determine the level of consciousness during anesthesia or in brain-damaged patients (Casali et al., 2012; Sarasso et al., 2015; Schartner et al., 2015). We confirmed previous findings in sleep (Casali et al., 2013; Abásolo et al., 2015) and LZc matched sleep’s phenomenology (Nir et al., 2013), dreams (i.e., conscious contents) being more frequent and complex in REM compared with light NREM but quite rare in deep NREM sleep.

LZc also predicted the LRP magnitude (Fig. 4c). In light NREM sleep, an increased propensity to respond to external input was associated with higher levels of complexity before stimulus onset. In REM sleep, however, this relationship was reversed, which could be related to the peculiarity of REM sleep: a state of consciousness disconnected from the environment (Hobson, 2009). This result emphasizes the difference between consciousness and responsiveness as observed in anesthesia (Sanders et al., 2012).

Neuronal bistability and the gating of sensory processing

In contrast with light sleep, no LRP was observed in deep NREM sleep. Light and deep NREM sleep were defined based on the presence or absence of slow oscillations associated with stimuli. Accordingly, light NREM showed a drastically reduced N550 potential (Figs. 2, 6), a potential usually linked to the down states of stimuli-evoked K-complexes (Bastien et al., 2002). Down states represent episodes of neuronal silencing (Steriade, 2003) and perturb the encoding and integration of information (Schabus et al., 2012; Pigorini et al., 2015). The presence of a N550 could therefore prevent the further processing of information and abolish the LRP in deep NREM sleep. This hypothesis is buttressed by the positive correlation between the N550 and LRP magnitudes (Fig. 6c).

In light NREM sleep, stimulus-evoked potentials showed an initial P200 correlating positively with LRP magnitude (Fig. 6b). The P200 has been interpreted as a local modality-dependent excitation (Laurino et al., 2014) and could reflect brain’s responsiveness to external inputs. However, in deep NREM sleep, the P200 was associated with LRP suppression (Fig. 6c). We interpret this reversal as the consequence of an increased neuronal bistability from light to deep NREM sleep.

Indeed, in deep NREM sleep, cortical neurons exhibit more down states (Steriade, 2003). Because a local excitation can trigger a down state (Sanchez-Vives et al., 2010; Menicucci et al., 2013), in deep NREM sleep, the P200 could ignite a down state in frontal areas that would shut down in turn any further processing. Thus, depending on sleep depth and neuronal bistability, the relationship of the P200 with sensory processing would reverse: from a potentiating activation to a suppressive mechanism. Brain responses to sounds could therefore follow a self-regulated process (Halász, 2015), whereby sensory activations can be retroactively suppressed. Thus, in NREM sleep, a reactive cortical gating (Tononi and Massimini, 2008), rather than a default thalamic one (McCormick and Bal, 1994), seems to take place.
Could dreams gate sensory processing in REM sleep?
The mechanisms underlying sensory decoupling in REM sleep are still unknown. The recovery of a wake-like brain activity and consciousness should favor the processing of external information, and yet external inputs are rarely integrated in dream scenery (Nir and Tononi, 2010). A recent study described the presence of slow waves in REM sleep restricted to the superficial layers of primary sensory cortices (Funk et al., 2016). These slow waves could act as a gating mechanism because superficial cortical layers are the main targets of thalamic sensory relays (Jones, 2007). We showed here, however, that stimuli had a clear impact on cortical activity: signs of arousals could be observed after stimulus onset (Fig. 5c,d) as well as robust auditory potentials (Fig. 7). These activations indicate that sensory information did reach the cortex as previously suggested (Sallinen et al., 1996; Nir et al., 2015). However, no LRP was observed when novel words were played. A potential explanation is that stimulus-evoked perturbations were confined in sensory areas.

But what could limit distributed cortical processes if cortico-cortical connectivity is regained in REM compared to NREM sleep (Massimini et al., 2010)? We propose that sensory information may compete with endogenous contents (i.e., dreams). Indeed, wake-like endogenous activations have been observed in REM sleep (Louie and Wilson, 2001; Andrillon et al., 2015). These endogenous activations could compete for the brain’s computational resources and interfere with the processing of external inputs. Accordingly, prestimulus complexity had a negative impact on LRP magnitude (Fig. 4). This “informational gating” could be mediated by a domination of top-down signaling at the expense of bottom-up processes (Nir and Tononi, 2010). Further investigations are needed to investigate the relationship between onepiece contents and sensory disconnection.

Consequences of external stimulations on sleep
How did stimulation affect sleep? In NREM sleep, we observed a two-step response (Fig. 5a,b). Right after stimulus onset, power increased within the slow-wave and spindle ranges in accordance with the fact that sleep rhythms can be evoked by external stimulations (Halász, 2015). These increases were maximal over frontal electrodes and could correspond to a protective mechanisms ensuring that sleep is preserved (Halász et al., 2014). Likewise, the suppressive N350 (Fig. 6) was maximal at frontal electrodes. After this initial increase, slow waves and spindle power were on the contrary suppressed. The associated spatial and temporal distributions overlap with the time and location of the LRP (Fig. 2c) and the increase in LZc following stimulus presentation (Fig. 4b).

A local dampening of sleep depth, accompanying sensory processing, thus followed the initial protective response. No sign of arousal was observed, but the absence of arousal at the scalp level does not preclude local awakening in motor or sensory areas (Nobili et al., 2011; Peter-Derex et al., 2015). The local modulation of sleep depth associated with sensory processing could therefore reflect, at the scalp level, the occurrence of local wake within sleep (Nobili et al., 2012). In REM sleep, auditory stimulations also perturbed theta oscillations (Fig. 5c,d), a hallmark of REM sleep (Buzsáki, 2006). Signs of cortical activation (increase in the high-frequency range) could also be observed after stimulus onset. Once again, these perturbations were maximal over central electrodes, potentially revealing the recruitment of auditory areas.

Responsiveness is dynamically modulated during sleep cycles
We also explored the dynamics of neural complexity, motor preparation, and θ-power (a proxy for slow-wave density) across sleep cycles. Both the LZc and θ-power dynamics reflected sleep cycles’ structure, with an initial descent toward deep NREM sleep (increase in θ-power, decrease of LZc) followed by a steep damping of NREM sleep at the transition with REM sleep. The LRP magnitude was also robustly modulated within sleep cycles and paralleled complexity’s dynamics. Crucially, LRP magnitude increased toward the end of sleep cycles. Thus, the LRP does not slowly decay after cycle onset but is tightly linked to sleep depth. This fragility of sleep at the transition with REM sleep could be instrumental in allowing the brain to “decide” whether to stay asleep or to wake up (Halász et al., 2004). This modulation of responsiveness during NREM sleep cycles is reminiscent of a model (Halász and Bódizs, 2013) in which the descending slope of the NREM cycle is gradually dominated by sleep-promoting mechanisms (such as evoked slow waves, as suggested by Fig. 6), whereas the ascending slope is characterized by the emergence of wake-promoting mechanisms, which could explain the recovery of the responsiveness marker toward the end of the NREM episode (Fig. 8). Interestingly, such a model would predict that external information would be better processed at the end of the NREM sleep cycle compared with the beginning, sleep depth being kept equal. Further research would be needed to confirm such a hypothesis.

Overall, our data suggest that sleepers can complexly and flexibly process information during certain sleep stages. This ability is self-regulated by the neural dynamics, allowing the sleeping brain to manage the challenging tradeoff between sensory decoupling and monitoring its environment.

References


