The fate of incoming stimuli during NREM sleep is determined by spindles and the phase of the slow oscillation

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INTRODUCTION

Classically, brain processes are considered as essentially reflexive and mainly driven by external stimuli. In this perspective, brain function is predominantly geared interpreting incoming stimuli and programming motor output. Another view posits that the bulk of brain activity is intrinsic, spontaneous (i.e., it emerges in the absence of any identified external stimulus), and continuously maintains and processes information (Raichle, 2006). Consistent with this view, the energy required for the brain to respond to external stimuli is extremely small compared to the ongoing amount of energy that the brain normally and continuously expends (Raichle and Mintun, 2006). During wakefulness, spontaneous fluctuations of brain activity profoundly modify brain responses to external information. For instance, conscious perception of the external world is dependent upon pre-stimulus activity in the somatosensory (Palva et al., 2005; Boly et al., 2007) as well as visual domain (Hanslmayr et al., 2007; Hesselmann et al., 2008a,b). Likewise, cortical responses to external stimuli should be modulated by the spontaneous non-rapid eye movement (NREM) sleep background activity materialized by sleep spindles and slow-waves (SW), which are associated with specific activity patterns in thalamic and cortical neurons during sleep. Accordingly, animal (Steriade, 1991) and human (Elton et al., 1997; Cote et al., 2000) studies clearly suggested that sensory transmission is blocked at the thalamic level during sleep spindles due to the recurrent inhibition of thalamocortical neurons by reticular thalamic cells and thereby might even herald individual resilience to disruptive stimuli such as environmental noise in human subjects (Dang-Vu et al., 2010). Likewise, meaningful events can be detected. Altogether, our results emphasize the notion that spontaneous fluctuations of brain activity profoundly modify brain responses to external information across all behavioral states, including deep NREM sleep.

Keywords: AEP, sleep, spindles, spontaneous activity, slow-wave phase, EEG/fMRI, fMRI, EEG

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processing of somatosensory inputs during deep NREM sleep was shown to be strongly influenced by the phase of the slow oscillation at which stimuli were delivered (Massimini et al., 2003). The amplitude of early evoked potentials increased during the downswing of the electroencephalography (EEG) slow fluctuation, potentially in relation to a progressive decrease in input resistance of cortical neurons (Contreras et al., 1996) and heightened probability of synaptic release (Massimini and Amzica, 2001) which is highest during the transition from the cellular down to up state (i.e., negative-to-positive EEG slope transition; Massimini et al., 2003).

In humans, EEG (Bastuji and Garcia-Larrea, 1999) and neuroimaging (Portas et al., 2000) studies primarily insisted on the persistence of brain responses to auditory stimuli. With regards to EEG the extent of residual auditory information processing during sleep has been extensively studied using event-related potentials (ERPs; for review see Campbell and Colrain, 2002). Therein it has been consistently reported that the early N1 component is gradually decreased from sleep onset to stage 2 NREM sleep whereas the P2 amplitude is increased (Campbell et al., 1992). Latter effect has also been associated with an attenuation of the classical processing negativity (de Lught et al., 1996) which is thought to reflect additional attentional processing of attenuated stimuli during waking (Näätänen et al., 1992). Consequently, absence of this overlapping negativity is regarded to reflect inhibition of information processing (Campbell et al., 1992). In addition, during the occurrence of sleep spindles the amplitude of the P2 component has been found to be even more amplified (Elton et al., 1997). This is a further inhibition of stimulus processing during sleep. Besides these modified wakening components NREM sleep (auditory) ERPs further consist of unusual large amplitude components. Specifically these are the N350 (between 250 and 400 ms), the N550 (between 500 and 800 ms), and the P900 (between 800 and 1300 ms). While the N350 possibly reflects an active inhibition of sensory processing during sleep onset, the later components (N550 and P900) form part of the very large amplitude stimulus-elicted K-complex and thus might reflect or overlap with the generation of delta oscillations for the sake of sleep protection (for review see Bastien et al., 2002; Colrain and Campbell, 2007).

Neuroimaging studies suggested that the brain can even detect meaningful auditory events (like a subjects own name) during NREM sleep in which it elicits significant responses in the amygdala and prefrontal cortex in addition to the bilateral activation of auditory cortex, thalamus, and caudate nuclei seen in response to simple auditory stimuli (Portas et al., 2000). In contrast, Czisch et al. (2002) reported that the brain response to auditory stimulation was decreased during NREM sleep as compared to wakefulness, interpreting their results in terms of a sleep-protective deactivation of primary sensory areas. In a follow up study the same authors demonstrated that the stimulus-induced negative BOLD effects – again primarily found in light NREM sleep – correlated positively with EEG signs of hyperpolarization (i.e., K-complexes and delta power) suggesting “true cortical deactivation upon stimulus presentation” (Czisch et al., 2004). However, all these studies considered NREM sleep as a homogeneous and steady state and did not account for the potential influence of spontaneous ongoing brain activity. The data reported in this paper makes use of the more advanced event-related paradigm focusing directly on evoked brain activity.

Here we now use simultaneous EEG and fMRI in order to characterize brain responses to tones during light and deep NREM sleep in non-sleep-deprived healthy individuals. Specifically, we examined brain responses to auditory stimuli during NREM stage 2–4 sleep in relation to the presence of sleep spindles (fMRI data published previously in Dang-Vu et al., 2011) and the phase of the slow oscillation.

MATERIALS AND METHODS

Thirteen – out of 19 recorded (9 females; age range = 18–25; mean age = 21.3) – healthy subjects were successfully scanned during the first half of the night in a Siemens Allegra 3 T scanner with a full 64 channel montage of MR-compatible EEG electrodes.

Participants were right-handed and gave their written informed consent. After the experiment participants received a financial compensation for their participation in this study, which was approved by the Ethics Committee of the Faculty of Medicine of the University of Liège. Participants were free of any medical, traumatic, psychiatric, or sleep disorder history, as assessed by a semi-structured interview. No participant complained of excessive daytime sleepiness (Epworth Sleepiness Scale; Johns, 1991) or sleep disturbances (Pittsburgh Sleep Quality Index Questionnaire; Buysse et al., 1989). All participants had normal scores at the Beck Anxiety and Beck Depression Inventory, and were non-smokers, moderate caffeine and alcohol consumers, as well as refraining from medication. None had worked on night shifts during the last year or traveled through more than one time zone during the last 2 months. Extreme morning and evening types, as assessed by the Horne–Ostberg Questionnaire (Horne and Ostberg, 1976), were not included.

Subjects were not sleep-deprived and followed a 4-day constant sleep schedule, as controlled by wrist actigraphy (Actiwatch, Cambridge Neuroscience, UK) and sleep diaries. Volunteers were requested to refrain from all caffeine and alcohol-containing beverages and intense physical activity for 3 days before participating in the study. Subjects reported to the laboratory at 9 pm.

EEG AND fMRI METHODOLOGY

Electroencephalography was recorded simultaneously to fMRI acquisitions, during the first half of the night, utilizing two MR-compatible 32-channel amplifiers (BrainAmp MR plus, Brain Products GmbH, Gilching, Germany) and a MR-compatible EEG cap (BrainCap MR, Falk Minow Services, Herrsching-Breitbrunn, Germany) with 64 ring-type electrodes. EEG caps included 62 scalp electrodes which were online referenced to FCz, as well as 1 electrooculogram (EOG) and 1 electrocardiogram (ECG) channel. Using abrasive electrode paste (ABRALYT 2000;
Functional volumes were analyzed by using Statistical Parametric Mapping 8 (SPM8; http://www.fil.ion.ucl.ac.uk/spm/) implemented in MATLAB (MathWorks). The series of consecutive fMRI volumes corresponding to selected stage 2–3, stage 3–4, or wake periods were selected from the complete fMRI time series and volumes corresponding to selected stage 2–3, stage 3–4, or wake epochs lasting more than 2 min were considered for the analysis of sleep spindles. Only stable stage 2–3 or stage 3–4 epochs lasting more than 2 min were selected for the analysis of sleep spindles (for details see Dang-Vu et al., 2011) and SW activity, respectively. For EEG spindle and SW analysis we used Fz, Cz, and Pz. For display and statistics we selected the Cz lead as this electrode allows to depict spindle as well as SW effects.

Functional MRI time series were acquired using a 3T MR scanner (Allegra, Siemens, Germany). Multislice T2*-weighted fMRI images were obtained with a gradient echo-planar sequence using axial slice orientation (32 slices; voxel size: 3.4 mm × 3.4 mm × 3 mm; matrix size = 64 × 64 × 32; TR = 2460 ms; TE = 40 ms; flip angle = 90°; delay = 0). Subjects were scanned during the first half of the night, starting at around midnight. They were asked to relax, try to sleep in the scanner and not pay attention to occasional tones, while fMRI and EEG data were acquired continuously. They stayed until they indicated by button press that they would like to go out, or for a maximum of 4000 volumes (about 164 min). The number of acquired volumes varied between 1195 and 4000 [3401 ± 965 volumes or 139 ± 40 min (mean ± SD)]. A structural T1-weighted 3D MP-RAGE sequence (TR = 1960 ms, TE = 4.43 ms, inversion time = 1100 ms, FOV = 230 mm × 173 mm, matrix size = 256 × 192 × 176, voxel size = 0.9 mm × 0.9 mm × 0.9 mm) was also acquired in all subjects.

In addition, waking sessions before and after sleep were selected to identify default brain activations associated with the occurrence of tones during wakefulness (TW). The analysis of waking fMRI data characterized the brain responses to TW compared to waking baseline activity.

Functional volumes were analyzed by using Statistical Parametric Mapping 8 (SPM8; http://www.fil.ion.ucl.ac.uk/spm/) implemented in MATLAB (MathWorks). The series of consecutive fMRI volumes corresponding to selected stage 2–3, stage 3–4, or wake periods were selected from the complete fMRI time series and constituted a “session.” fMRI time series were corrected for head motion, spatially normalized (voxel size = 2 mm × 2 mm × 2 mm; resampled using spline interpolation) to an echo-planar imaging template conforming to the Montreal Neurological Institute space, and spatially smoothed with a Gaussian kernel of 8-mm FWHM.

The analysis of fMRI data, based on a mixed effects model, was then conducted in two serial steps, accounting respectively for intraindividual (fixed) and interindividual (random effects) variance in SPM8.

In a first analysis (as previously published in Dang-Vu et al., 2011), spindles were identified on band pass filtered EEG data between 11 and 15 Hz, using an automatic detection algorithm by thresholding the spindle root mean square signal at its 95th percentile and post hoc visually checking for correct classification (Mölle et al., 2002). Brain responses related to spontaneously occurring spindles and SW have already been described elsewhere (Schabus et al., 2007; Dang-Vu et al., 2008) and are not reported in the following. Responses to sounds corresponded to systematic deviations of BOLD signal over and above the baseline activity during NREM sleep having taken into account the activity related to characteristic oscillations of NREM sleep.

In a second analysis, two tone categories were considered, depending on whether the stimulus appeared before [tone 0–300 ms pre-SW-peak (TPre)] or (max. 300 ms) after the peak negativity of the (stage 3–4) slow oscillation [post-SW-peak (TPost)]. The SW-peak negativity was defined as the highest negative component (a negative peak between two zero crossings with voltage \(-35 \mu V\)) in a frontal EEG array (Fp1, Fp2, F3, F4, F7, F8, Fz, F1, F2, AF3, AF4, F5, F6, AF7, AF8, AFz). We assessed brain responses as simple main effects of tones relative to baseline activity (independent of the phase of the slow oscillation) as well as differential main effects between the two tone categories (pre to post SW-peak differences). In order to find the instant at which the sound volume changed at its fastest rate, we identified the downward zero crossing of the second derivative of the Gaussian sound envelope. This timepoint (88 ms after sound onset) was taken as the moment at which the sound became detectable by the volunteer which we consequently used as onset for EEG and fMRI analysis.

In order to take into account the effects of all identifiable neural events on the BOLD signal during sleep, sleep spindles and SW power were also modeled in the analysis. To take into account artifacts related to cardiac cycle, an estimation of R–R intervals derived from ECG was included as regressor of no interest in all individual design matrices. Movement parameters estimated during realignment (translations in x, y, and z directions and rotations around x, y, and z axes) and a constant vector were also included in the matrix as variables of no interest. High-pass filtering was implemented in the matrix design using a cut-off period of 128 s to remove low frequency drifts from the time series. Serial correlations in fMRI signal were estimated using an autoregressive (order 1) plus white noise model and a restricted maximum likelihood (ReML) algorithm.

Statistical inferences were conducted after correction for multiple comparisons either on the whole brain volume (\(p_{FWE} < 0.05\)) or for regions of interest previously identified in the literature using small volume correction (\(p_{SVC} < 0.05\)). The Supplementary Material (Table S1) is not corrected for multiple comparisons, and...
consequently no statistical inferences were conducted for those areas.

AUDITORY STIMULATION DURING SLEEP
Throughout sleep pure tones were presented binaurally using headphones. Tones had a frequency of 400 Hz, a duration of 300 ms and were presented at each TR (2460 ms) with a probability of 70%. Within a given volume the sound could occur anywhere with the 2460 ms scan frame. Yet in 30% of the cases the sound was not presented within that volume. This resulted in a median ISI of 2910 ms and SD of 10706 ms.

The intensity of tones was held constant throughout the night and adjusted individually during a test scanning session reproducing the same background noise than the experimental one. Subjects were requested to adjust the tone loudness to a level which was discernible but not disturbing. For definition of the early auditory ERP components we used latency ranges similar to Crowley and Colrain (2004) but shifted them 40 ms in time (N1: 115–190 and P2: 190–290 ms) after stimulus onset in order to account for delayed responses in fMRI scanner environments (cf. Novitski et al., 2001).

TPre and TPost epochs were classified on 0.5–20 Hz bandpass filtered EEG data if a frontal SW peak “exceeding” $-35 \mu V$ was identified within 300 ms after or before the tone, respectively (cf. Figure 1). Each analyzed epoch then ranged from $-1500$ ms pre to 1500 ms post tone stimulation. One subject had to be excluded for SW-EEG averaging due to bad signal quality. Statistical analysis between TPre and TPost were conducted on SW phase-sorted averages (i.e., after identifying all TPre and TPost SW-peaks, we re-aligned the single trials to the SW-peak, and then averaged across

![Figure 1](https://example.com/figure1.png)

Figure 1 | Grouping of brain activity to tones according to the spontaneous background activity characterized by NREM (A) sleep spindles and (B) slow-wave activity. Note that during sleep sounds were randomly presented (at 70% of every TR) and post hoc categorized according to (i) the absence or presence of spindles (stage 2–3) or (ii) the phase of the slow oscillation (stage 3–4) when the tone occurred.
trials and subjects per tone category) which were previously low-passed filtered at 4 Hz. In order to be able to interpret the effect of tones on the slow oscillation we also calculated spontaneous SW’s (in a time window of −1500 to 0 ms before tone stimulation) for comparison. The peak negativity (“exceeding” −35 μV) of these spontaneous SW’s had to occur 400 ms after epoch start (−1100 ms pre-stimulus) as well as at least 800 ms before tone onset in order to only identify spontaneous SW’s without any influence of surrounding tones on the spontaneous waveform. In addition, we classified SWs depending on the amplitude of the peak negativity (small SW: max. −50 mV; big SW: more negative than −50 mV) for further analyses. For stage 2–3 sleep analysis of spindles, tones were categorized according to their occurrence outside (TN) or within detected spindles (TS; cf. Figure 1, Dang-Vu et al., 2011).

**BOLD SIGNAL MODULATION BY SPINDLES (fMRI DATA REPRODUCED FROM DANG-VU ET AL., 2011)**

The mean number of sounds delivered without (TN) or with ongoing spontaneous spindles (TS) per subject was 534.3 (SD = 198.8) and 30 (SD = 11.2), respectively. As reported previously (Dang-Vu et al., 2011), tones delivered in the absence of sleep spindles (TN) were also – like TW (Figure 2A) – associated with responses in thalamus and primary auditory cortex (Heschl’s transverse gyrus) (Figure 2B), confirming that sounds can be processed in stage 2–3 NREM sleep. Significant additional TN responses were found in a set of cortical and subcortical areas, in the pons, cerebellum, middle frontal gyrus, precuneus, and posterior cingulate gyrus, all areas known to respond to auditory stimuli (Holcomb et al., 1998; Portas et al., 2000; Gaab et al., 2003). In contrast, at the same statistical threshold (p < 0.05

**FIGURE 2 | Brain regions activated in relation to tones during waking, light NREM sleep but outside sleep spindles and light sleep during spindles (data reproduced from Dang-Vu et al., 2011).** (A) Significant responses associated with tones presented during waking (TW). Note that tones during wakefulness (TW) induced responses in the primary auditory cortex (Heschl’s transverse gyrus) and the thalamus, in an area compatible with the medial geniculate nucleus. (B) Significant responses associated with tones presented during stage 2–3 sleep, in the absence of ongoing spindles (TN). These responses are located in the thalamus, primary auditory cortex, brainstem, cerebellum, middle frontal gyrus, precuneus, and posterior cingulate gyrus. The brainstem response encompasses areas compatible with the cochlear nuclear groups the trapezoid bodies and the superior olivary complex. (C) Significant responses associated with tones presented within spindles in stage 2–3 sleep (TS). Here neural populations that process sound are found in the nuclei of the lateral lemniscus of the brainstem (insert, marked by arrow). Lower panels depict the fitted responses in the thalamus [x = −12, y = −22, z = −6; left panel] and the auditory cortex [x = 58, y = −14, z = 6; right panel] associated with sounds delivered with (red) or without (blue) ongoing spontaneous spindles. The curves correspond to the mean and the shaded areas to the SEM. Functional results are displayed on an individual structural image (displayed at p < 0.001, uncorrected). (Modified from Dang-Vu et al., 2011. Copyright by the National Academy of Sciences.)
Table 1 | Brain responses to tones during deep NREM sleep.

| Region                        | SW phase independent effect | SW phase dependent differences (TPost > TPre) |
|-------------------------------|-----------------------------|---------------------------------------------|
|                               | x   | y   | z   | Cluster size | Z score | pSVC     | x   | y   | z   | Cluster size | Z score | pSVC     |
| Thalamus (Portas et al., 2000) | −14 | −14 | 2   | 320        | 4.96    | <0.001*  |     |     |     |             |         |          |
| Heschl gyrus (Lockwood et al., 1999) | 44  | −24 | 14  | 84         | 4.17    | 0.002    |     |     |     |             |         |          |
| Superior temporal gyrus (STG; Czisch et al., 2009) |     |     |     |             |         |          | 68  | −44 | 18  | 49         | 3.45    | 0.017     |

Coordinates (x, y, z) are expressed in millimeter in the Montreal Neurological Institute (MNI); Z scores result from the statistical parametric analysis; pSVC refers to the probability of the null hypothesis (i.e., absence of activity change associated with tones in deep NREM sleep), after correction for multiple comparisons either on the whole brain volume (x) or on small volumes of interest identified in the literature (references in brackets).
Furthermore, we tested whether the amplitude of the slow oscillation during sound delivery has a modulatory effect on the subsequent differences (TPre vs. TPost) in the positive component after tone delivery. We revealed that the positive component following tone delivery is markedly enhanced if the tone is arriving at the positive going slope of the slow oscillation (TPost) in both amplitude conditions (peak amplitude small and big SW: $t_{11} = 2.21$, $p < 0.05$ and $t_{11} = 5.91$, $p < 0.001$, respectively; or positive area under component for small and big SW: $t_{11} = 3.63$, $p = 0.004$; $t_{11} = 6.00$, $p < 0.001$, respectively). Yet, the TPre to TPost difference for the positive component was identical for tones arriving during small or big SWs.

**DISCUSSION**

Our study characterizes the modulation of brain responses to auditory stimulation by spontaneous NREM sleep oscillations (sleep spindles and slow oscillations). As expected, sounds during wakefulness elicited responses in the thalamus and primary auditory cortex. These responses – although somewhat altered in size and location – persisted during NREM sleep, except during presence of light NREM spindles. Yet interestingly, responses at a higher cortical level became less consistent or even absent during spindles and the negative going phase of the (deep NREM) slow oscillation.

At the EEG level we found that ERPs elicited during NREM sleep showed a reduction in the amplitude of the N1 and an increase in the amplitude of the P2 component compared to the ERPs during wakefulness (Figure 4A) in accordance with previous studies (Campbell et al., 1992; Elton et al., 1997; Cote et al., 2000).
FIGURE 4 | Grand-averaged EEG brain activity in response to sounds during (A) waking and NREM sleep as well as (B) in response to the occurrence of sleep spindle events. Note the decreasing signal complexity from waking (black line), to light (red) to deep (blue) NREM sleep (left panel) in response to tones. On the right, EEG responses to tones presented during TS (red line) or outside spontaneously occurring stage 2-3 sleep spindles (TN, blue line) are depicted. Note the massive late negativity evoked by tones falling into sleep spindles. Data are bandpass filtered between 0.5 and 20 Hz and reflect the grand-average of 12 subjects.

FIGURE 5 | Electroencephalography brain responses to sounds during deep NREM sleep. On the upper left (A) the evoked response to all tones during stage 3-4 is depicted. The lower left panel (B) shows the evoked response if tones are categorized according to the phase of the slow oscillation which was present during sound delivery. Note that tones occurring before the SW-peak negativity (TPre, red line) are differently processed (orange shading) than tones arriving at the upswing of the slow oscillation (TPost, blue line). The black dashed line depicts the (phase unsorted) TPre grand-average waveform which was shifted in time to the left so that their SW-peak negativity overlays with the TPost peak negativity for better comparison. Data are bandpass filtered between 0.5 and 20 Hz and reflect the grand-average of 12 subjects. (C) Single trial phase-sorted and re-aligned EEG brain responses to sounds during deep NREM sleep. Note that latter analysis accentuates the amplitude as no time jitter is present when averaging TPre and TPost trials according to SW-peak negativity (rather than to tones) on a trial-by-trial basis. In (C) Time 0 is therefore marking the SW-peak negativity for TPre and TPost trials. Spontaneous slow-waves are plotted for comparison in green. Data are (zero-phase lag) low-pass filtered at 4 Hz and error bars are overlaid at each sampling point. All potentials with negativity upwards.
when that information is occurring during the presence of a NREM sleep spindle. Likely this is due to the fact that thalamic neurons adopt a burst firing mode during NREM sleep which prevents faithful sensory transmission of external inputs to the cortex. Presumably, exactly that functional isolation of sensory input during sleep spindles is of benefit for internal neuronal interactions subserving brain plasticity.

Contrary to Elton et al. (1997) but in line with Cote et al. (2000) we did not find an increased P2 amplitude for stimuli occurring during a spindle as compared to stimuli outside a spindle (Figure 4B). Yet, most interestingly tones presented during a spindle elicited a much stronger N550 component than tones presented outside a spindle (Figure 4B). According to Bastien and colleagues (2002) averaging responses to tones presented during sleep, which include K-complexes, produce a N550 component. In addition also Church et al. (1978) showed that auditory evoked K-complexes were larger when stimuli were presented during sleep spindles. It is thus possible that also the strong N550 effect seen in our data is due to the superposition of underlying K-complexes. In relation to the literature it is also interesting to note that we only find a marginally enhanced P2 component across light and deep sleep (Figure 4A); Yet we do find a strong N550 effect which is consistent with the idea that inhibition of the incoming auditory information is not occurring at an early (as reflected by P2) but later stage (in time and brain hierarchy) in accordance with our fMRI results (cf. Figure 3B).

Note that also the late negative swing of our BOLD response (at 6–8 s for spindles, and at 4–6 s for SWs) could be related to underlying K-complexes on some trials. Specifically it is interesting to note that Czisch et al. (2009) using an acoustic oddball paradigm reported a prominent negative BOLD response for (rare) tones, yet no wake-like activation of the auditory cortex. In their data only rare tones followed by an evoked K-complex were associated with a “wake-like activation of task-related areas in the temporal cortex” in accordance with data from Dang-Vu et al. (2011). Our current fMRI results showing auditory cortex activation (in the absence of sleep spindles and across SW phases) as well as a late negative BOLD swing might thus be interpreted as combination of these two effects. However, note that in our data a K-complex like N550 effect appears only when averaging EEG across all SW phases (cf. Figure 4A, red line) or tones in the presence of sleep spindles (cf. Figure 4B, red line), but not in response to tones just before or after the SW-peak negativity (cf. Figure 5B).

Yet, the exact anatomical stage at which the transfer of sensory information to auditory thalamo-cortical pathways is hindered remains uncertain. It appears that no clear-cut change in neural activity takes place at any of the early auditory relay structures during NREM sleep (Velluti, 2008), but prethalamic modifications in transmission were reported in the somato-sensory system during NREM sleep, although not specifically in relation to spindles (Rosanova and Timofeev, 2005). On the other hand, thalamic neurons are likely to hinder the faithful transmission of sensory input during spindles, because the burst firing mode that they adopt during NREM sleep, and especially during spindles, distorts the transmission of sensory inputs to the cortex in a non-linear fashion (McCormick and Feeser, 1990; Sherman and Guillery, 2002). Finally, also note that a strong recruitment of inhibitory interneurons was recently described during sleep spindles (Peyrache et al., 2011). This phenomenon might add to the absence of cortical responses to auditory stimuli during spindles as observed in our previous study and reiterated herein.

PHASE DEPENDENCE OF BRAIN RESPONSES WITH RESPECT TO THE SLOW OSCILLATION

While the phase of the slow oscillation does not appear to alter brain responses in primary sensory cortex (cf. Figure 3A), it does modulate responses at higher cortical levels as shown in superior temporal gyrus (cf. Figure 3B; for additional areas also see Table S1 in Supplementary Material). On the one hand, the results are consistent with the hypothesis that brain responses during deep NREM sleep vary as a function of the fluctuating state of thalamo-cortical circuits (Massimini et al., 2003; Rosanova and Timofeev, 2005). In accordance with Massimini and colleagues the brain appears most receptive to the environment at the negative-to-positive going slope of the slow oscillation. Yet, a direct comparison to our data is difficult as stimulation was done in another sensory modality (somatosensory vs. auditory) and EEG components of interest were markedly earlier. Last but not least we compared pre- vs. post-SW-peak phases (300 ms bins each) whereas Massimini and colleagues used smaller and differently grouped time windows (e.g., ±50 ms directly around the SW-peak negativity).

On the other hand, the current data suggest that the slow oscillation does only modulate responses in higher associative cortices. In keeping with this observation, the slow oscillation was also found to be more associated with a pronounced modulation of neuronal firing in associative than in primary cortices at a cellular level (Steriade and McCarley, 2005). This response pattern also concords with the breakdown of local functional connectivity reported during NREM sleep (Massimini et al., 2005) and is reminiscent of auditory responses recorded in unconscious patients in whom primary auditory cortices still respond whereas higher association cortices do not (Laureys et al., 2006; Boly et al., 2004). The results suggest that the residual cortical processing during NREM sleep is insufficient for ignition of processes thought to be necessary for conscious awareness but might very well allow to trigger awakening responses to salient and personally important stimuli.

The analysis of EEG data simultaneously recorded during fMRI sessions further dissect the reciprocal interactions between spontaneous brain activity and responses to external stimuli. Importantly, EEG data allow us to probe how the intrinsic state of neural responsiveness (i.e., SW phase) influences responses to external stimuli. They allow us to go one step closer to the actual neural events subtending sound processing during NREM sleep although the exact phase relation between the scalp-recorded human sleep EEG and the underlying intracellular dynamics can only be extrapolated from animal experiments (Massimini et al., 2003; Vyazovskiy et al., 2011).

Tones delivered after the peak negativity (i.e., possibly during the ON state) probably contribute to further synchronize neural firing (cf. Figure 5), already ongoing due to the ON state. Tones delivered before SW-peak negativity (supposedly at the end of the OFF state) potentially initiate the cascade of events that lead to the ON state. The smaller amplitude of the second positive evoked component might suggest that the neural recruitment induced by sounds in the negative going slope (TPre) is smaller as compared to the – probably synchronized – TPost response during the positive
going phase. On the other hand one could argue that the auditory evoked potential during deep NREM sleep – presumably reflecting a K-complex related N550 around 650 ms (cf. Figure 5A) – “attenuates” the late positive component (also around 650 ms) of the tones arriving before SW-peak negativity (Figure 5B). Yet, the reconstruction of a spontaneous SW without tone delivery (cf. Figure 5C, green line) clearly favors the earlier interpretation. In either case the EEG data reveal a clear phase-dependent modulation of the auditory ERP response post-event which is not present before tone delivery (Figure 5C).

Future studies should elaborate on the reported findings and investigate phase-dependent effects of smaller temporal windows (100 ms or less) as well as auditory stimuli of varying complexity.

CONCLUSION

Present evidence extends previous findings and suggests that SWS is not a static phenomenon as mutually assumed by many earlier studies (Perrin et al., 1999; Portas et al., 2000; Bastuji et al., 2002; Campbell et al., 2005). Altogether, brain responses during NREM sleep appear to be non-stationary and highly dependent upon spontaneous brain activity such as prominent sleep spindles or the phase of the slow oscillation. The exact temporal window when a stimulus arrives might thus not only determine the fate of that very material during waking but likewise all stages of NREM sleep.

AUTHOR CONTRIBUTIONS

Manuel Schabus, Thien Thanh Dang-Vu, and Pierre Maquet contributed at all stages of the manuscript. Manuel Schabus and Pierre Maquet wrote up the manuscript. These and all of the following authors (Annabelle Darsaud, Christina Schmidt, Gilles Vandewalle, Mélanie Boly, Martin Desseilles, Steffen Gais) were involved in data recording, discussed the results or commented on the manuscript. Dominik Philip Johannes Heib, Christian Degueldre, Christophe Phillips, and Evelyne Balteau added important technical expertise.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at http://www.frontiersin.org/sleep_and_chronobiology/abstract/18937

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