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► **To cite this version:**

David Angulo-Garcia, Maëva Ferraris, Antoine Ghestem, Lauriane Nallet-Khosrofian, Christophe Bernard, et al.. Cell Assemblies in the Cortico-Hippocampal-Reuniens Network during Slow Oscillations. *Journal of Neuroscience*, 2020, 40, pp.8343 - 8354. 10.1523/jneurosci.0571-20.2020 . hal-03582770

HAL Id: hal-03582770

<https://amu.hal.science/hal-03582770>

Submitted on 21 Feb 2022

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Cell Assemblies in the Cortico-Hippocampal-Reuniens Network during Slow Oscillations

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The nucleus reuniens (NR) is an important anatomic and functional relay between the medial prefrontal cortex (mPFC) and the hippocampus (HPC). Whether the NR controls neuronal assemblies, a hallmark of information exchange between the HPC and mPFC for memory transfer/consolidation, is not known. Using simultaneous local field potential and unit recordings in NR, HPC, and mPFC in male rats during slow oscillations under anesthesia, we identified a reliable sequential activation of NR neurons at the beginning of UP states, which preceded mPFC ones. NR sequences were spatially organized, from dorsal to ventral NR. Chemical inactivation of the NR disrupted mPFC sequences at the onset of UP states as well as HPC sequences present during sharp-wave ripples. We conclude that the NR contributes to the coordination and stabilization of mPFC and HPC neuronal sequences during slow oscillations, possibly via the early activation of its own sequences.

Key words: cell assemblies; hippocampus; network; nucleus reuniens; medial prefrontal cortex; slow oscillations

Significance Statement

Neuronal assemblies are believed to be instrumental to code/encode/store information. They can be recorded in different brain regions, suggesting that widely distributed networks of networks are involved in such information processing. The medial prefrontal cortex, the hippocampus, and the thalamic nucleus reuniens constitute a typical example of a complex network involved in memory consolidation. In this study, we show that spatially organized cells assemblies are recruited in the nucleus reuniens at the UP state onset during slow oscillations. Nucleus reuniens activity appears to be necessary to the stability of medial prefrontal cortex and hippocampal cell assembly formation during slow oscillations. This result further highlights the role of the nucleus reuniens as a functional hub for exchanging and processing memories.

Introduction

Information exchange between the hippocampus (HPC) and the medial prefrontal cortex (mPFC) is essential for different memory processes, including consolidation (Siapas and Wilson, 1998; Frankland and Bontempi, 2005; Preston and Eichenbaum, 2013; Maingret et al., 2016; Eichenbaum, 2017; Kitamura et al., 2017; Latchoumane et al., 2017; Ferraris et al., 2018). The representation of information is supported by the recruitment of neuronal cell assemblies, which refer to a group of neurons that are

coactivated repeatedly for a given brain operation and thus represent a distinct entity embedded within neuronal networks (Hebb, 1949; Buzsáki, 2010; Buzsáki, 2019). In both HPC and mPFC, cell assemblies fire in a fine time-resolved manner (Skaggs and McNaughton, 1996; Nadasdy et al., 1999; Lee and Wilson, 2002; Euston et al., 2007; Luczak et al., 2007; Pastalkova et al., 2008; Peyrache et al., 2009; Battaglia et al., 2011). These neuronal assemblies display a precise temporal sequential activation, which reflects the encoded information (Nádasdy, 2000; Foster and Wilson, 2006; Ji and Wilson, 2007; Davidson et al., 2009; Marre et al., 2009; Pfeiffer, 2020). Such activity is prevalent during sleep (Pavlidis and Winson, 1989; Wilson and McNaughton, 1994; Nadasdy et al., 1999; Atherton et al., 2015), particularly during NREM sleep, and is organized in space and time by a set of oscillations, such as hippocampal sharp-wave ripples (SPW-Rs), cortical slow oscillations (SOs), and spindles (Sirota et al., 2003; Sirota and Buzsáki, 2005; Staresina et al., 2015; Maingret et al., 2016). How these cell assemblies are finely coordinated between the HPC and the mPFC to support memory consolidation is not well understood. The thalamic nucleus reuniens (NR), which bidirectionally connects the HPC and the

Received Mar. 9, 2020; revised July 22, 2020; accepted Sep. 1, 2020.

Author contributions: D.A.-G., M.F., and P.P.Q. analyzed data; D.A.-G. and P.P.Q. wrote the first draft of the paper; D.A.-G., M.F., C.B., and P.P.Q. edited the paper; D.A.-G. and P.P.Q. wrote the paper; M.F. and P.P.Q. designed research; M.F., A.G., L.N.-K., and P.P.Q. performed research.

*D.A.-G. and M.F. contributed equally to this work.

The authors declare no competing financial interests.

This work was supported by Fondation Recherche Médicale Grant FDT201805005246 to M.F. D.A.-G. was supported by A*MIDEX Grant ANR-11-IDEX-0001-02 and Vicerrectoria de Investigaciones-Universidad de Cartagena Project 038-2019. We thank Wesley Clawson for proofreading the manuscript.

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<https://doi.org/10.1523/JNEUROSCI.0571-20.2020>

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mPFC (Herkenham, 1978; Van der Werf et al., 2002; Vertes, 2006; Varela et al., 2014), plays a key role in memory consolidation (Loureiro et al., 2012; Cassel et al., 2013; Pereira de Vasconcelos and Cassel, 2015; Dolleman-van der Weel et al., 2019; Klein et al., 2019). In particular, the NR synchronizes γ bursts, which can open temporal windows for information exchange (Buzsáki and Draguhn, 2004; but see Buzsáki and Schomburg, 2015), between the HPC and the mPFC during SOs (Ferraris et al., 2018). The NR is therefore ideally poised to orchestrate the dynamics of cell assemblies in both regions.

We have recorded and manipulated the NR-mPFC-HPC circuit during SOs under anesthesia, which resembles sleep patterns and provides long periods for analysis. We find that cell assemblies are recruited in the NR at the onset of the UP state of SOs and that NR activity is needed for the stability of HPC and mPFC cell assemblies.

Materials and Methods

Experimental model and subject details. All experiments were performed in accordance with experimental guidelines approved by Aix-Marseille University Animal Care and Use Committee under the authorization #01451-02. All rats were group housed to avoid social isolation-induced stress (Manouze et al., 2019). A total of 18 rats were used in this study. Part of these data (14 Wistar Han rat data) was used in a previously published study (Ferraris et al., 2018), and 4 Wistar Han rats are original data. They include local field potentials (LFPs) and single-unit recordings made in the mPFC, HPC, and NR of anesthetized rats.

Animal surgery. Wistar Han IGS male rats (250–400 g; Charles River; RRID:RGD_2308816) were anesthetized with urethane (1.5 g/kg, i.p., catalog #U2500, Sigma Millipore) and ketamine/xylazine (20 and 2 mg/kg, i.m., Renaudin Cip; 3400957854195 and Centravet, catalog #ROM001, respectively), additional doses of ketamine/xylazine (2 and 0.2 mg/kg) being supplemented during the electrophysiological recordings. The heart rate, breathing rate, pulse distension, and the arterial oxygen saturation were also monitored with an oximeter (MouseOx, Starr Life Science) during the entire duration of the experiment to ensure the stability of the anesthesia and monitor the vital constants. The head was secured in a stereotaxic frame (Kopf #962, Phymep), and the skull was exposed and cleaned. Two miniature stainless-steel screws, driven into the skull above the cerebellum, served as ground and reference electrodes. Up to three craniotomies were performed to target, from bregma: the prelimbic area of the mPFC at 3 mm AP and 0.8 mm ML; the CA1 field of the intermediate HPC at –5.6 mm AP and 4.3 mm ML; and the NR at –1.8 mm AP and –2 mm ML. Silicon probes (NeuroNexus) were used to record from these structures: a A1x32-Edge-5 mm-20-177-H32-15 probe placed at [–2.5 –3.1] mm from brain surface to reach mPFC layer 5; a A1x32-Edge-10 mm-20-177-H32-50 32-site probes placed at –7.2 mm from brain surface to reach the NR; and a HPC A1x32-6 mm-50-177-H32-15 probe placed at [–2.8 –3.0] mm perpendicularly to the CA1 field from stratum oriens to stratum lacunosum moleculare in the HPC. All the probes were lowered inside the brain with a motorized manipulator (IVM single, Scientifica).

For NR inactivation experiments ($n=5$ rats), a local injection of a fluorophore-conjugated muscimol (BODIPY-MSCI TMR-X Conjugate, catalog #M23400, Invitrogen) was performed in the NR, and data from the mPFC and the HPC (CA1) were simultaneously acquired. The injection needle (33 gauge, catalog #87930, catalog #7803-05, Hamilton) was inserted in the NR (using the same depth coordinates as the probes and mounted on the same micromanipulator) and 0.70 nmol of muscimol in 0.3 μ l of PBS (Ferraris et al., 2018) was delivered over 60 s through a micropump (UltraMicroPump UMP3-1, WPI). The needle was left in place for 3 additional minutes to allow for adequate diffusion of the drug, then carefully removed.

At the end of the recording, the animals were injected with a lethal dose of pentobarbital Na (150 mg/kg, i.p.) and perfused intracardially with 4% PFA solution in PB (0.12 M). The position of the electrodes (DiIC18(3), catalog #46804A, InterChim) was applied on the back of the

probe before insertion and was confirmed histologically on Nissl-stained 60 μ m sections (NeuroTrace 500/5225 Green Fluorescent Nissl Stain, catalog #N21480, Invitrogen). Only experiments with appropriate position of the probe were used for analysis (see Fig. 3A).

Data collection and initial analysis. Extracellular signal recorded from the silicon probes was amplified (1000 \times), bandpass filtered (1 Hz to 5 kHz), and acquired continuously at 32 kHz (64-channel DigitalLynx; NeuroLynx) at 16-bit resolution. Raw data were preprocessed using a NEUROSUITE (Hazan et al., 2006; <http://neurosuite.sourceforge.net/>; RRID:SCR_008020). After recording, the signals were downsampled to 1250 Hz for the LFP. Spike sorting was performed automatically using KLUSTAKWIK (Harris et al., 2000) (<http://klustakwik.sourceforge.net/>; RRID:SCR_008020; RRID:SCR_014480), followed by manual adjustment of the clusters, with the help of autocorrelogram, cross-correlogram, and spike waveform similarity matrix (KLISTERS software, RRID:SCR_008020) (Hazan et al., 2006). After spike sorting, the spike features of units were plotted as a function of time, and the units with signs of substantial drift over the period of recording were discarded. Moreover, only units with clear refractory periods and well-defined cluster were included in the analyses (Harris et al., 2000). Recording sessions were divided into brain states of theta and SO periods. The epochs of stable SO periods were visually selected respectively from the ratios of the whitened power in the SO band ([0.5 2] Hz) and the power of the neighboring band ([20 30] Hz) of mPFC or NR LFP and assisted by visual inspection of the raw traces (Quilichini et al., 2010).

Neurons were assigned as “NR neurons” by determining the approximate location of their somata relative to the recording sites, the known distances between the recording sites, the histologic reconstruction of the recording electrode tracks, and subsequent estimation of the recording sites. All the neurons recorded from sites located near the close contour of the NR were discarded. Neurons located at a minimal distance of 200 μ m of NR border and located within contours of the ventro-median, submedian or antero-median thalamic nuclei were classified as “other thalamic neurons” and used in the analysis (Ferraris et al., 2018).

Data postprocessing. All the analysis was performed using custom-written MATLAB (RRID:SCR_001622; RRID:SCR_008020) scripts. From the spike times, the instantaneous firing rates of each cell were calculated by counting the number of spikes inside a window of 50 ms, in overlapping intervals of 10 ms. The population firing rate was estimated by averaging the single-cell firing rates at each 10 ms interval. Both the population rate and the single-cell firing rate were smoothed with a Gaussian kernel of 5 ms width. Peaks of population activity (AP) were identified as the points where the population rate amplitude was larger than the average rate plus 1 SD. A separation between two consecutive peaks of at least 600 ms was imposed on the peak detection algorithm to avoid multiple peaks of activity within one UP state. For visualization purposes, all firing rate’s heat maps were normalized with the peak firing rate for each cell to guarantee a variation between [0:1].

Phase analysis and sequence identification. LFP signals (from mPFC or NR) during SO phase were bandpass filtered between 0.5 and 2 Hz with a second-order Butterworth filter to extract only the UP/DOWN transitions. The time evolution of the phase during the UP/DOWN cycle was extracted performing the Hilbert transform of the filtered LFP (from the mPFC or the NR) and was visually checked. The [0.5 2] Hz LFP oscillation is most likely volume-conducted from the cortex in the NR (for review, see Buzsáki et al., 2012; Herreras, 2016), and this was checked by computing the current source density analysis as the second spatial derivative of the NR LFPs and smoothed spatially with a triangular kernel (Quilichini et al., 2010; Ferraris et al., 2018), which resulted in a flat signal on all the channels (data not shown).

Similar results regarding the detection of UP/DOWN transitions were obtained by constructing the phase as a linear interpolation between peaks and troughs of the filtered LFP signals (Belluscio et al., 2012). We also obtained similar results by extracting UP states from the mPFC multiunit activity. UP states were identified as periods of neuronal firing between periods of at least 100 ms of global silence (without spike). The time of the first spike marking a transition between the silence period and the active one is defined as the onset of the UP state. UP states offset corresponds to the time of the last spike marking a

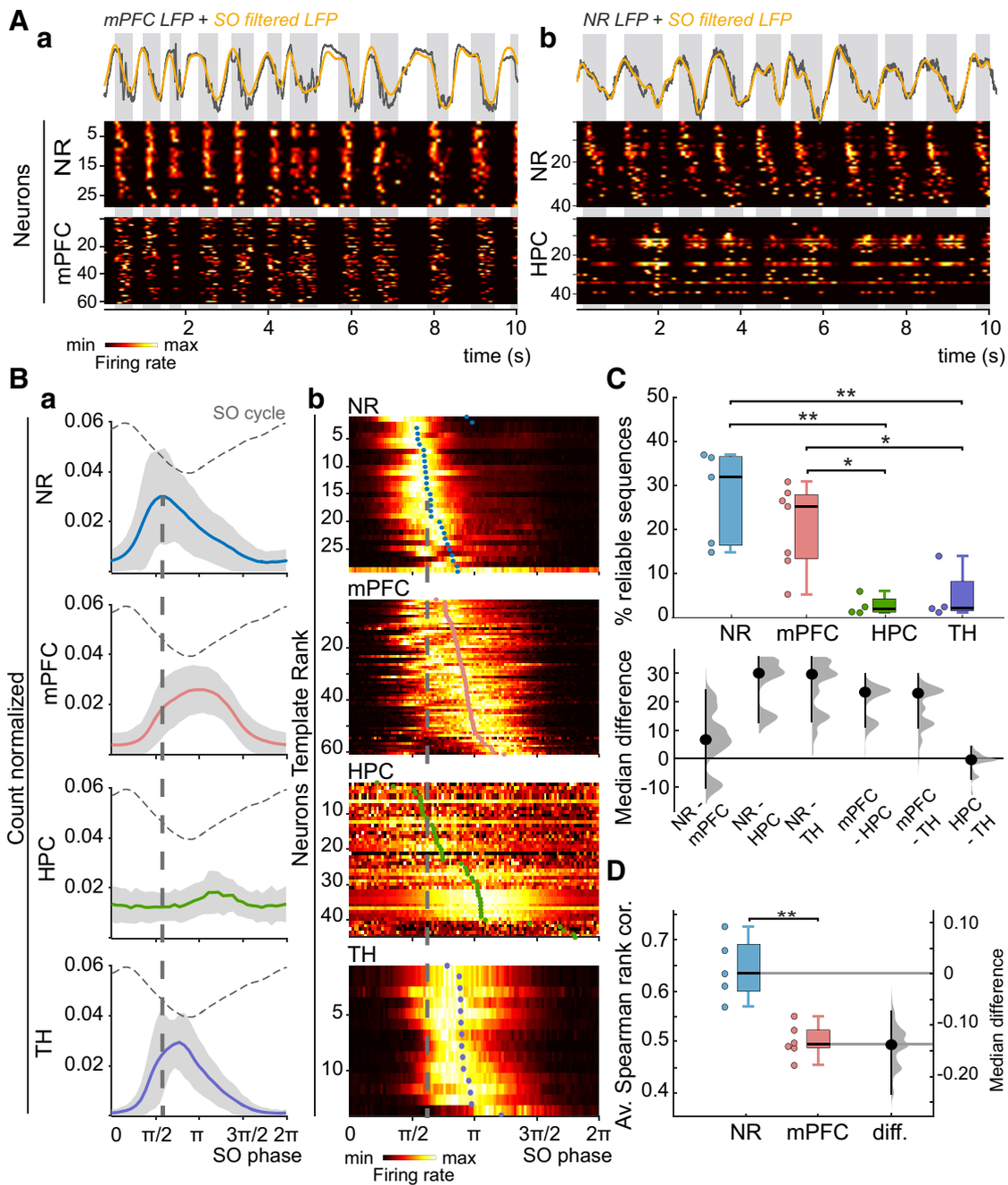


Figure 1. Sequential dynamics of cell assemblies. **A**, Top, LFPs from (**Aa**) mPFC and (**Ab**) NR (mainly volume-conducted from the cortex) in gray, with superimposed [0.5 2] Hz filtered LFPs (yellow), where the UP states are marked by the gray boxes, from two typical recordings. Bottom, The corresponding template heat maps of normalized NR unit activities with (**Aa**) mPFC or (**Ab**) HPC normalized unit activities show the repetition of neuronal assemblies, particularly in the NR and the mPFC. The identification number of the recorded neurons is indicated on the left axis of the heat maps. **Ba**, Phase distribution of the NR, mPFC, HPC, and TH population firing (grouped data, all recordings) with reference to the SO phase (indicated by the dashed curve). Dashed line indicates the most preferred phase of NR neurons, showing that their firing precedes other cells located in the mPFC, HPC, and TH at the onset of the UP state. **Bb**, Heat maps of normalized distribution of neurons preferred SO phase for sample representative firing patterns of NR (same neurons as in **Aa**), mPFC (same as in **Aa**), HPC (same as in **Ab**), and TH (representative example). The activation order for each neuron is calculated as the average preferred phase. Heat maps are ordered according to the increasing value of average preferred phase (colored circles), defining a template order. The sequential activation is clearly apparent in NR and mPFC neurons, but neither in the HPC nor in the TH (the phase distribution extends over at least $\pi/2$ for each neuron). **C**, Top, The largest percentage of reliable sequences is found in the NR and mPFC compared with the HPC and the TH. Bottom, The median differences for 6 comparisons are shown in a Cumming estimation plot. Each median difference is plotted on the lower axis as a bootstrap sampling distribution. Dots represent median differences. Ends of the vertical error bars represent 95% CIs. (* $p < 0.05$, ** $p < 0.01$) **D**, The average Spearman correlation shows that sequences in NR are more reliable than those of mPFC. The estimation plot on the right side shows the median difference (dots) and the 95% CI (diff., median difference).

transition to a silence period. UP states < 200 ms were excluded (Luczak et al., 2007). We used circular statistics (Berens, 2009; Ferraris et al., 2018) to compute the mean phase at which each neuron fires (“preferred phase”) and to build their firing-phase histograms (see Fig. 1Ba). For visualization purposes, the resulting histograms depicted in the heat maps were normalized with the peak value of the distribution. For each cell,

we then calculated a resultant vector characterized by an angle describing the average preferred phase and a magnitude with values between [0 1], quantifying the coherence of the phases. The template order was obtained by organizing the average preferred phase in increasing order [0 2π]. To prove that sequential ordering was not because of purely random fluctuations, we also calculated the preferred phase using half of the

recording and compared with the preferred phase using the second half of the recording. Moreover, to test the possibility that sequences were a consequence of differences in firing rates between neurons, we normalized the firing rate of all recorded units in such a way that all neurons had an average 2 Hz firing rate by means of random spike removal. After spike removal, the average phases were recalculated and compared with the average phases obtained with the unnormalized spike train.

Sequence reliability, participation index, and sequence delay. We extracted the UP state duration as described by Ferraris et al. (2018). At each UP state, the local activation order was calculated by measuring the time lag between the first activation of each neuron relative to the AP in a ± 200 ms window. Ordering the latencies in increasing order resulted in the local activation order. The reliability of a given sequence within a population peak was quantified as the Spearman rank correlation of the template order and the local activation order in that particular UP state. Sequences were considered reliable above 99% significance level. For each reliable sequence, a robust fit between the local activation order and the activation latency was computed. Nonparticipating neurons in a given sequence were identified as neurons that meet one of the following criteria: (1) did not fire within the ± 200 ms window or (2) neurons appearing as outliers in the robust fit (those whose residual value were larger than twice the SD of the residuals in the robust fit). The participation index was then calculated as the mean fraction of neurons that participated in the sequences of a recording session. A second linear fit between the local activation latency and the site of the linear probe closest to the neuron gave a measure of the directionality of the sequence activation. A slope of the fit different from 0 implies an activation toward a given direction. Since this slope has units of time/electrode site, we multiplied the resulting value by 32 (the number of sites in the electrode) to account for the time required for a sequence (sequence delay) to travel along the electrode.

Activity peak triggered histogram (APTH). Once the APs have been identified according to the algorithm described in Data postprocessing, we store the values of activation time lag of the first spike of each neuron relative to the AP at each UP state in a time window of ± 200 ms centered at the AP. This allows us to compute the so-called activity peak triggered histogram for each cell. With the APTH, it is possible to assess the statistics of the firing latencies in the neighborhood of the UP state. Once the APTH was obtained, one can calculate the variability of the activation lags as a measure of the firing extent around the peak reported in Figure 4C. Variability of the APTH can be calculated in analogy to the variance of a probability distribution function where τ is the mean value of the APTH (Fig. 4C, inset).

SPW-Rs detection and analysis. The procedure of SPW-Rs detection in the HPC stratum pyramidale LFP was based on those described previously (Isomura et al., 2006; Ferraris et al., 2018). Briefly, the LFP was digitally bandpass filtered [80–250] Hz, and the power (root-mean-square) of the filtered signal was calculated. The mean and SD of the power signal were calculated to determine the detection threshold. Oscillatory epochs with a power of ≥ 5 SDs above the mean were detected. The beginning and the end of oscillatory epochs were marked at points where the power fell 0.5 SD. Once the SPW-Rs were detected, the SPW-R duration was calculated as the average between the start and the end of it. SPW-Rs are known to replay forward and backward sequences. With this in mind, we checked for different sequential activations in SPW-Rs by extracting the local firing order of neurons at each SPW-R (using the first spike as the activation latency for each neuron) and calculating a similarity matrix (cross-correlation) among all the obtained orders. The existence of well-defined clusters in the cross-correlation matrix could signal the existence of two or more different templates.

Therefore, we clustered the similarity matrices of the local orderings for each recording using a k -means algorithm with $k = 1, 2, 3, 4, 5$. The optimal clustering is the partition in k clusters that maximizes the modularity measure (for details, see Newman, 2010). For all the HPC recordings, modularity did not significantly change between $k = 1$ and $k = 2$ and definitely decreased for $k > 2$. This means that there is no convincing evidence of the existence of different templates, and we assumed the existence of a unique template. With this, the APTH for each cell was computed, taking each SPW-R half-time as the activity peak. To test whether

a cell robustly participates in the SPW-Rs, we compared the APTH against a uniform distribution with identical mean and SD. Cells whose APTH were different from the flat distribution above a 95% level were considered as robustly participating in the SPW-Rs. Cell participation ratio was obtained dividing the number of robustly participating neurons by the total number of recorded neurons for that session. Organizing the average time lag for robustly participating neurons in increasing order defined the template order for the ripple. For each SPW-R, the reliability of activation respect to the template was calculated via the Spearman rank correlation between the activation order of that SPW-R and the template order. To calculate this correlation, only the neurons that belong to the template order were considered. A correlation $> 95\%$ CI was considered reliable.

Statistics. All results reported are based on a significance threshold $\alpha = 0.05$ (otherwise stated), and all groups included enough samples to enable rejection of the null hypothesis at that level. We used two-sample Kolmogorov-Smirnov test to assess differences between distributions, and Student's t test to evaluate differences in the mean of distributions unless otherwise stated. In parallel, results from small samples data ($n < 10$) were evaluated using estimation statistics providing the effect size and the 95% CI of the median difference between two groups. Raw data were processed at <https://www.estimationstats.com>, directly giving both results and graphs (Calin-Jageman and Cumming, 2019). The median difference between two groups is shown with a Gardner-Altman estimation plot. Raw data are plotted on the left axis. The median difference is plotted on a floating axis on the right as a bootstrap sampling distribution (see Figs. 1C,D, 3D, 4C,E, 5B). Five thousand bootstrap samples were taken; the CI is bias-corrected and accelerated. The mean difference is depicted as a dot; the 95% CI is indicated by the ends of the vertical error bar. If 95% CI includes 0, differences are considered as nonsignificant. The p value(s) reported are the likelihood(s) of observing the effect size(s), if the null hypothesis of zero difference is true. Correlation tests involving ranked variables (neuron indices and electrode sites) were performed via a Spearman rank correlation. We tested significant differences between percentages with a two-proportion Z test.

Results

Cell assemblies are recruited in NR at UP state onset

We first assessed the behavior of NR ($n = 166$ cells, $n = 5$ rats), mPFC ($n = 496$, $n = 7$), and HPC ($n = 163$, $n = 4$) neurons during SOs in anesthetized rats. SOs during anesthesia share similar features as NREM sleep (Hutt, 2011; Ferraris et al., 2018) and offer long-duration stable recordings necessary to identify statistically significant sequences. Neighboring thalamic neurons (anteromedian and ventro-medial nuclei) were also recorded as a control group ("TH," 89 neurons, $n = 4$).

During SOs, most of the neurons fired at the onset of the UP states, mainly in mPFC and NR (Fig. 1A) (Ferraris et al., 2018). We found a robust recruitment of neuronal activity at the UP state onset in NR (median percentage of recruited neurons per UP state 70.4%, min/max: 61.1%/85.9%) and mPFC (median: 62.9%, min/max: 42.1%/70%). Given the number of recorded neurons per session, 98% of the total number of UP states recorded in mPFC and NR recruit > 10 neurons. In comparison, HPC neurons showed less participation during UP state (median: 33.8%, min/max: 29.3%/46.4%). To identify any ordering in this firing activity, we ranked each neuron according to its mean preferred SO phase (Fig. 1Ba). In contrast, HPC neurons displayed better sequential activation during sharp-wave ripples (SPW-Rs), which tend to occur during the UP state (Sirota et al., 2003; Battaglia et al., 2004; Isomura et al., 2006; Buzsáki, 2015; Maingret et al., 2016; Khodagholy et al., 2017) (median: 45%, min/max: 35%/64%).

Neurons were organized in increasing order of average preferred phase, defining a template order (Fig. 1Bb). Sequences in

NR always preceded mPFC ones (Fig. 1B). As a negative control, we analyzed the activity of neighboring TH neurons. Although TH neurons displayed a strong entrainment by SOs (Ferraris et al., 2018), there was poor sequential activity (Fig. 1B). The sequential activity at UP state onset therefore appears to be specific of NR and mPFC neurons in the areas investigated here. As a further control, we analyzed the activity of NR neurons during epochs dominated by theta ([4–6] Hz) oscillations. We did not find stable sequential neuronal assemblies during theta (data not shown). Together, these results show that, at the beginning of UP states, sequential cell assembly formation occurs first in the NR, then in the mPFC, and marginally in the HPC and neighboring thalamic nuclei.

We then quantified the reliability of these sequences. To do so, we computed at each UP state the activation latency of each neuron relative to the population peak activity (see Materials and Methods). The local activation order of the cell assembly was determined from the ascending sorting of the activation latencies. We then calculated the Spearman rank correlation between the local activation order and the template order. A rank correlation $r = 1$ indicates that the sequential activation follows exactly the template order determined previously by the average phase preferences. For NR neurons, the template order was significantly expressed in 32% of the total number of UP states (min/max = 15%/37%, Spearman test, $p < 0.01$), while mPFC reliable sequences were found in 25% of the UP states (min/max = 5%/31%; Fig. 1C). In contrast, reliable HPC sequences were only detected in 2% of the UP states (min/max = 1%/6%) and in 2% for sequences in TH neurons (min/max = 1%/14%). Although the proportion of reliable sequences in the NR and mPFC were not significantly different (Kolmogorov-Smirnov test, $p = 0.15$), the average Spearman correlation values of the reliable sequences were significantly larger in the NR than in the mPFC (Kolmogorov-Smirnov test, $p < 0.01$; Fig. 1D), indicating that the NR cell assembly sequential organization is the most consistent. Since sequences in NR preceded those of mPFC, and reliable sequences can be found in both, we checked whether the reliability in NR could affect that of mPFC. To do so, we compared the reliability index of NR and mPFC within the same UP state in the $n = 3$ paired recordings. We found a very small correlation between the two reliability indices (median r : 0.09 min/max: 0.06/0.12). To estimate the participation of neurons in sequence generation, we calculated a participation index, defined as the probability for a neuron to be involved in a given sequence (see Materials and Methods). The average participation index was consistently larger across experiments in the NR (median: 0.63, min/max: 0.55/0.81) than in the mPFC (median: 0.52, min/max: 0.36/0.60; Kolmogorov-Smirnov test, $p < 0.05$). We also calculated the average fraction of nonparticipating neurons (not fulfilling the condition of the robust fit, see Materials and Methods) and compared this with the fraction of nonparticipating neurons because of the nonfiring condition. A very small fraction of neurons was classified as nonparticipating from the outlier condition (median 2.5% in NR and 1.4% in mPFC) compared with the nonfiring condition (median 30.2% in NR and 46.0% in mPFC).

As a cross-validation, to discard the possibility that the sequential ordering was because of random fluctuations, we compared the average preferred phases using the first half of the spike train with the average phases using the second half (Fig. 2A). Both for NR and mPFC, the average phases obtained in the two halves were strongly correlated (median r : 0.94, min/max: 0.8/0.97).

The sequential firing of neurons could just be the result of an increased excitability of early firing neurons compared with neurons firing later after the onset of the UP state (Peyrache et al.,

2010). To address this issue, we recalculated the average phases and the subsequent template ordering by normalizing all the neurons' spike trains via random spike removal to have an average firing rate of 2 Hz. Average phases obtained with the normalized and unnormalized spike trains were virtually identical in NR and mPFC ($r \sim 1$ aggregated data; Fig. 2B). Additionally, we tested the correlation between the firing order and the firing rate for each NR and mPFC neuron. We found that 2 of 5 NR recordings were significantly correlated while only 1 of 7 mPFC recordings showed significant correlations (Spearman test, $p < 0.05$).

Together, these results show that mPFC and NR neurons display highly stable patterns of sequential neuronal activations at the onset of the UP state, with NR sequences being more reproducible. The sequential activation cannot be solely explained by increased excitability of early firing cells.

NR cell assemblies are spatially distributed

We also investigated the spatial distribution of the NR and mPFC cell assemblies. To do this, we correlated the template order of each neuron to their anatomic dorsoventral localization, since the silicon probes were sampling NR from the dorsal-to-ventral pole (Fig. 3A). The slope of a linear fit of such correlation provides the information on whether the activity is propagating in a given direction (Luczak et al., 2007) (see Materials and Methods; Fig. 3B). Furthermore, we quantified the degree of spatial distribution of the activity for each UP state, calculating a linear fit between the activation latency of each neuron relative to the peak activity and the dorsoventral localization. The slopes of the fit for each UP state, properly multiplied by the number of sites of the probe, resulted in the sequence delay (see Materials and Methods). Positive or negative delays are fingerprints of propagating activity in a given direction. Although both NR and mPFC can show some degree of propagation in a preferential direction (Fig. 3C), NR average delays were consistently larger than mPFC ones (Kolmogorov-Smirnov test, $p < 0.05$; Fig. 3D), reflecting a dorsal-to-ventral direction of the sequential activation. To assess the consistency of the preferential direction of propagation, we calculated the Spearman rank correlation between the rank of the template order and the anatomic location of the corresponding recording site. We found a highly significant correlation in NR recordings in most cases ($n = 4$ of 5, median $|r|$ value across all n : 0.65, $p < 0.001$), whereas it was virtually absent in mPFC ($n = 1$ of 7, $|r| = 0.35$, $p < 0.01$). There are thus more spatially organized sequences in NR compared with mPFC (z test, $p < 0.05$; Fig. 3E).

Together, these results demonstrate that NR displays at the UP state onset robust sequential activations, which are spatially organized in a dorsoventral pattern. This is particularly surprising as it suggests a topical spatial organization in the NR.

NR firing is necessary to the mPFC and HPC sequence stability

The fact that NR cell assemblies occur earlier than mPFC ones raises the possibility that the NR may be involved in the sequential activation of mPFC neurons. To test this hypothesis, we used muscimol to inactivate chemically the NR (see Materials and Methods) and recorded the activity of mPFC neurons ($n = 190$ neurons, $n = 5$ experiments). Following NR inactivation, mPFC neurons showed a reduced firing activity during the UP state compared with noninactivation, control recordings (Fig. 4A) (Ferraris et al., 2018). During SOs, the distribution of interspike intervals is characterized by the presence of two peaks: one at ~ 1 Hz corresponding to the frequency of the SO, and one at

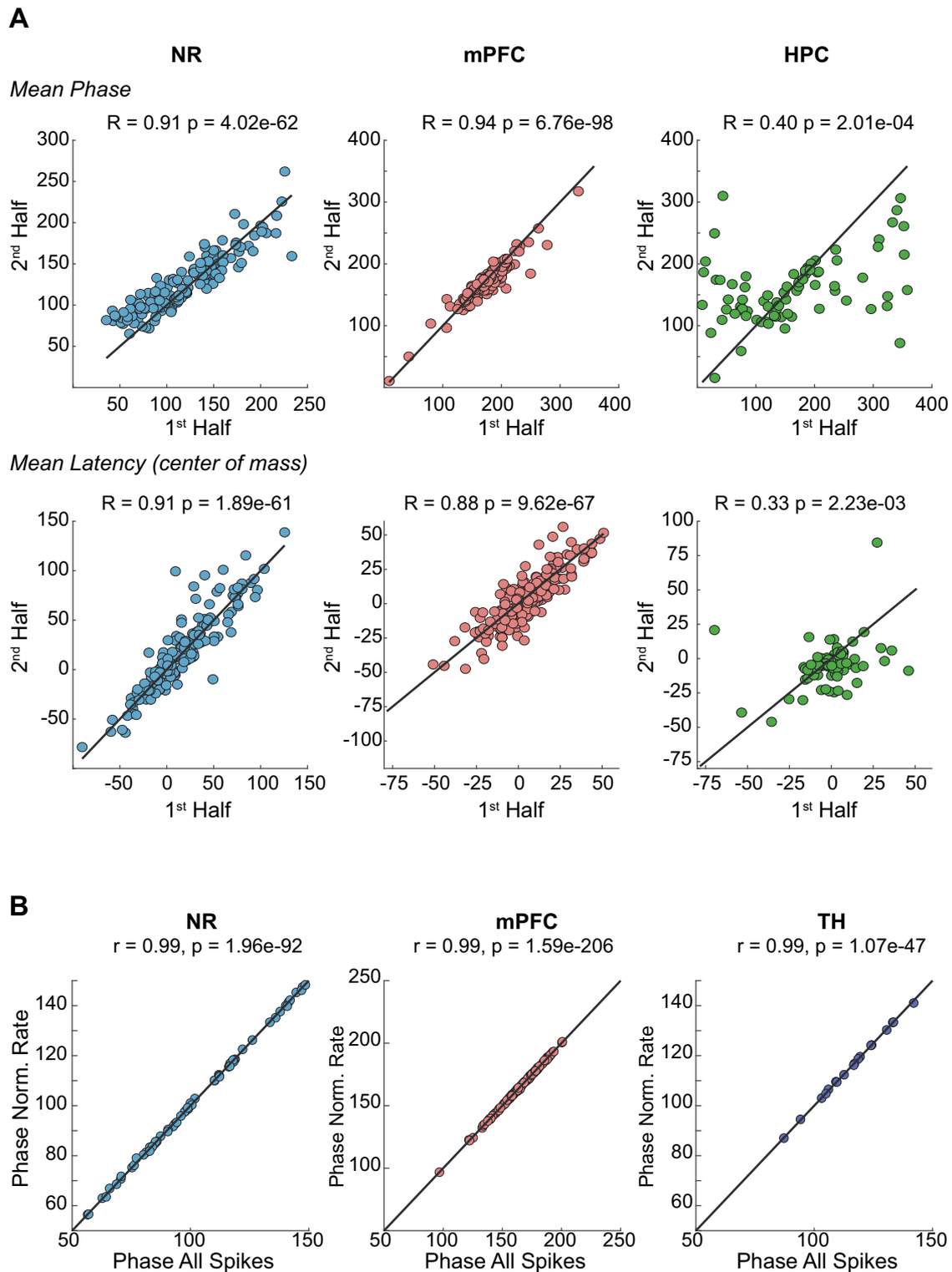


Figure 2. Cross-validations of the sequential dynamics of cell assemblies. **A**, Top, Average phases calculated separately during the first half and second half of the recordings for each region. Bottom, Same as top panels using the average latency method. Both methods lead to similar results. **B**, Average firing phases of NR, mPFC, and TH neurons with reference to the S0 calculated using the whole spike train versus the normalized firing rate (2 Hz). The lack of difference indicates that the sequential activity does not appear to be caused by a mere increase of excitability.

~100 ms, which indicates that cells fire bursts of action potentials (Fig. 4B). The distribution of interspike intervals was modified in inactivation conditions with a smaller peak at ~100 ms, reflecting reduced bursting activity during UP state (Fig. 4B). We then calculated the distribution of the pooled variability of the APTH (see Materials and Methods; Fig. 4C, inset) in control and NR inactivation conditions. The median APTH variability across

experiments was not significantly changed when NR was inactivated (control median: 9.62, min/max: 8.98/10.60, NR inactivation median: 9.02, min/max: 8.67/13.73, Kolmogorov-Smirnov test, $p = 0.4428$), as shown in Figure 4C. This reveals the fact that, although bursting is reduced, the time extent to which neurons in mPFC fire is maintained. However, the UP state duration was slightly shorter (control: 0.55 s, NR inactivation: 0.48 s,

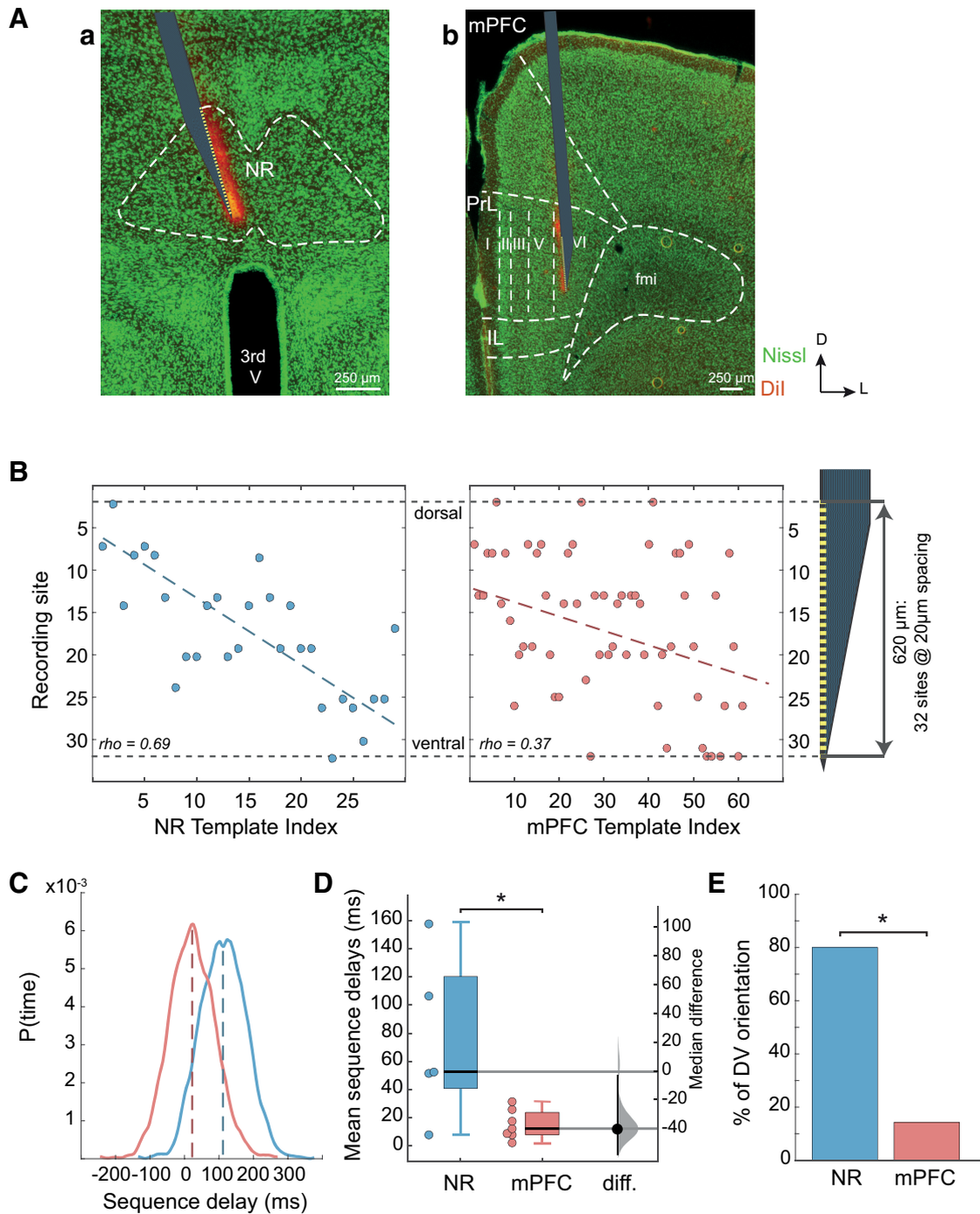


Figure 3. Spatial organization of NR and mPFC cell assemblies. **A**, Representation of the position of a linear silicon probe with 32 recording sites in (**Aa**) the NR and (**Ab**) mPFC. NR and mPFC contours (and layers) are delimited by the white dashed line over the green fluorescent Nissl staining. fmi, Forceps minor of the corpus callosum; third V, third ventricle; PrL, prelimbic area; IL, infralimbic area. Red-orange staining represents the Dil that was deposited at the back of the silicon probe before insertion. D, Dorsal; L, lateral. **B**, Relationship between anatomic dorsoventral location of NR (left) and mPFC (right) neurons (location defined by the site of the probe recording the maximum amplitude of the action potentials) and the template rank showing a linear correlation for NR neurons but not for mPFC ones in a template experiment. **C**, Distribution of spatial delays (from the beginning to end of a sequence) across the probe from the top-most site to the bottom-most one for mPFC (red) and NR (blue) neurons. **D**, Mean sequence delays (with median difference and 95% CI) across experiments for NR and mPFC. **E**, The percentage of experiments showing a dorsoventral (DV) orientation of sequences was larger in the NR than in the mPFC. (* $p < 0.05$).

Mann–Whitney test, $p < 0.001$), an effect that can be attributed to the extra peak in UP state duration distribution at ~ 0.25 s, which did not exist in control data (Fig. 4D). We then evaluated the outcome of the mPFC sequences in the NR inactivation condition. Inhibition of NR activity reduced the capacity of mPFC to generate reliable sequences compared with the control condition, since only 5.6% of them (min/max: 1.4%/8.4%) were

reliable (compared with control median: 25%, min/max: 5%/30%, Kolmogorov–Smirnov test, $p = 0.0228$; Fig. 4E). Moreover, the fraction of participating neurons in reliable sequences decreased in the inactivation condition (control: 53%; inactivation: 37%; Kolmogorov–Smirnov test, $p < 1e-5$; Fig. 4F). These results support the proposal that NR activity is important for the stability and reliability of mPFC sequences.

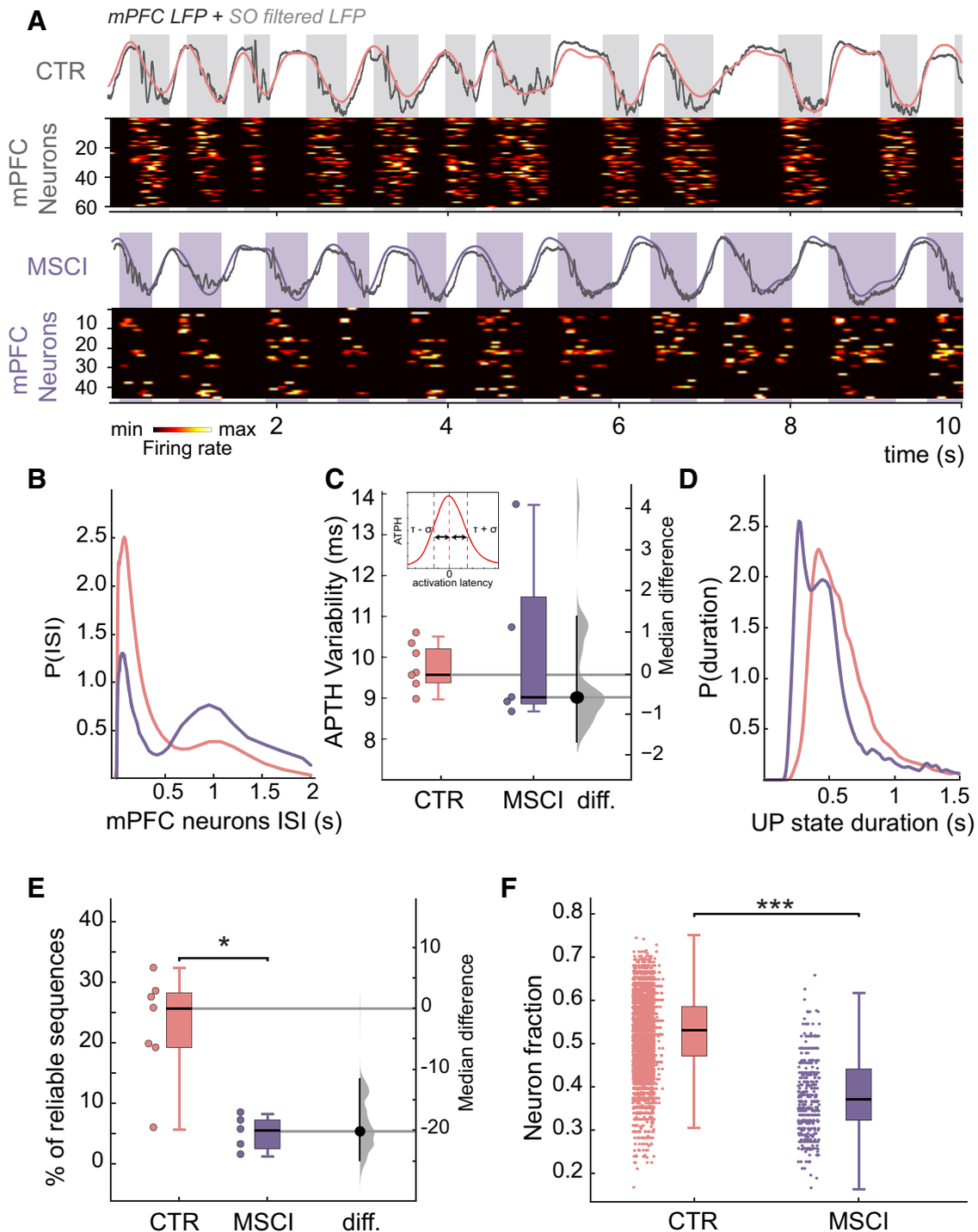


Figure 4. Inactivation of NR impairs reliable sequential activation of mPFC neurons at the beginning of the UP state. **A**, Template heat maps of normalized mPFC unit activities during NR inactivation condition (bottom) inducing less stable cell assemblies compared with control condition (top). The mPFC LFP appears above the heat maps in gray with the corresponding [0.5 2] Hz filtered trace (pink and purple), and the UP states marked by the gray and purple boxes. **B**, Interspike interval distribution of mPFC neurons grouped data for control (pink) and NR inactivation conditions (purple), showing that a decrease in burst firing when the NR is inactivated. **C**, Grouped data with median difference and 95% CI of the variability of the APTH showing no significant change between control (CTR) and NR inactivation conditions (MSCI). Inset, The APTH quantifies the lag distribution with respect to the population activity peak for each neuron. **D**, Distribution of the duration of the UP state in control (pink) and NR inactivation (purple) condition, where a peak at short values indicates the emergence of shorter UP states. **E**, The number of reliable sequences found in mPFC was decreased when the NR was inactivated (MSCI) compared with the control (CTR) condition (grouped data, $n = 5$ experiments). **F**, The fraction of mPFC neurons participating to reliable sequences was decreased when the NR was inactivated. ($*p < 0.05$, $***p < 0.001$).

The UP state of the SOs poorly modulates the activity of HPC neurons and the formation of timely organized cell assemblies. However, sequential activation of HPC neurons consistently occurs during SPW-Rs (Buzsáki, 2015). We therefore investigated the consequences of NR inactivation on HPC sequences

during SPW-Rs. NR inactivation did not affect HPC neurons firing rate (control average: 1.07 ± 0.05 Hz, NR inactivation average: 1.08 ± 0.05 Hz, t test, Cohen's $d = 0.019$, $p = 0.97$; Fig. 5A,E). In addition, NR inactivation did not alter the frequency of SPW-R occurrence (median control: 0.087 Hz, min/max: 0.043/

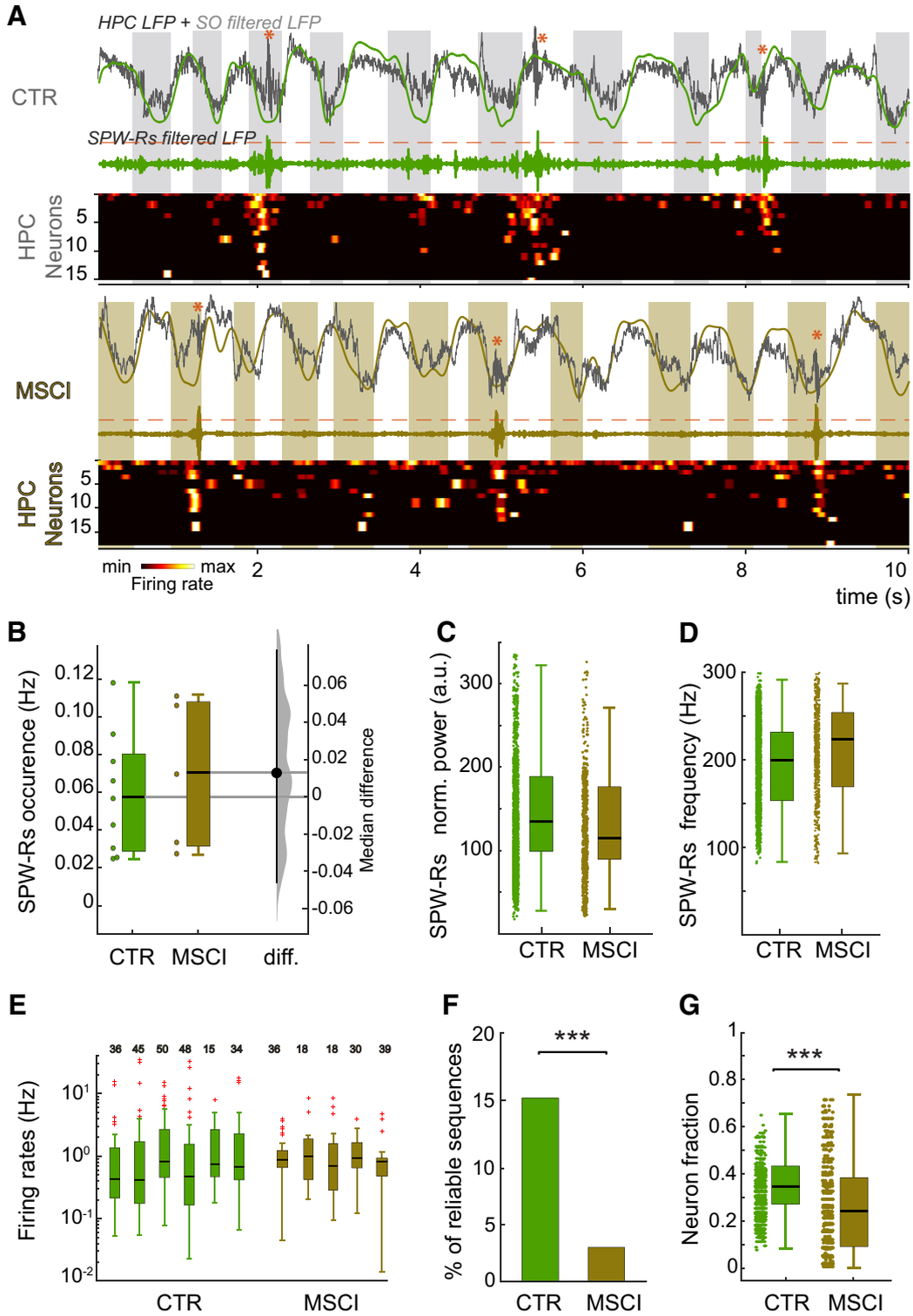


Figure 5. Inactivation of NR impairs reliable sequential activation of HPC neurons during SPW-Rs. **A**, Template heat maps of normalized HPC unit activities in control (CTR) and NR inactivation (MSCI) condition (bottom of each panel). Top, The HPC LFP (gray) and the corresponding [0.5–2] Hz filtered trace (green and khaki). Gray and khaki boxes represent the UP states. Orange stars represent the SPW-Rs, which are visible in the [80–200] Hz filtered LFP (bottom). **B**, SPW-R occurrence did not change when NR was inactivated (grouped data, $n = 5$ experiments; $n = 913$ SPW-R events in CTR, $n = 149$ SPW-R events in MSCI). **C**, NR inactivation did not change the normalized power (a.u., Arbitrary unit) or the **(D)** inner frequency of SPW-Rs. **E**, The median firing rates of HPC neurons were not affected by NR inactivation. Red crosses represent outlier values. The numbers on the top of each statistical box indicate the number of recorded neurons in the session. **F**, The proportion of reliable cell assemblies during SPW-Rs was considerably reduced when the NR was inactivated. **G**, The fraction of reliably participating HPC neurons to sequences during SPW-R events was decreased by NR inactivation. (***) $p < 0.001$.

0.11 Hz, NR inactivation: 0.049 Hz, min/max: 0.022/0.111 Hz, Kolmogorov–Smirnov test, $p = 0.8254$; Fig. 5B). Similarly, neither their power (mean normalized power control: 157 ± 2 , NR inactivation: 144 ± 5 , t test, $p = 0.295$; Fig. 5C) nor their inner frequency (mean frequency control: 193 ± 1 Hz, NR inactivation: 213 ± 3 Hz, t test, $p = 0.084$; Fig. 5D) was modified. This result shows that NR inactivation does not change either SPW-R properties or global HPC cell firing. In contrast, the number of reliable sequences found within SWP-Rs was largely reduced when the NR was inactivated (control median: 15%, NR inactivation median: 3%, z test, $p < 0.001$; Fig. 5F). In this condition, the remaining cell assemblies recruited a significantly lower number of neurons (control median: 37.5%, min/max: 12.5%/80%, NR inactivation median: 27.8%, min/max: 0%/74.4%, Kolmogorov–Smirnov test, $p < 0.001$; Fig. 5G). These findings suggest that the NR activity has a causal role in the sequential organization of neuronal firing in the HPC during SPW-Rs.

Discussion

In this study, we show that four-fifths of NR neurons fire within robustly spatially and temporally organized cell assemblies at UP state onset; and that NR activity influences cell assemblies' stability in mPFC at UP state onset and in HPC during SPW-Rs. These results further support the concept that the NR is a key functional hub in memory networks involving the mPFC and the HPC.

The sequential activation of neuronal assemblies is believed to constitute a core feature of information processing in the brain (Hebb, 1949; Buzsáki, 2010). Cell assemblies are found in the archicortex (Lee and Wilson, 2002; Harris et al., 2003; Dragoi and Buzsáki, 2006; Pastalkova et al., 2008; Villette et al., 2015; Malvache et al., 2016), cortical areas (Kenet et al., 2003; MacLean et al., 2005; Ferezou et al., 2006; Euston et al., 2007; Luczak et al., 2007; Luczak and Maclean, 2012), as well as in striatum (Lansink et al., 2009). They constitute a way to code/encode/store information (Maass, 2016; Kitamura et al., 2017). During NREM sleep, cortical activity is dominated by the sequential activation of cortical neurons at the onset of the UP state, while in the HPC the sequential activation mostly occurs during SPW-Rs (Sirota et al., 2003; Battaglia et al., 2004; Peyrache et al., 2009; Maingret et al., 2016; Khodagholy et al., 2017). Similar sequential firing occurs during the SOs measured during anesthesia (Luczak et al., 2007), supporting the view that such brain state shares many features with NREM sleep (Tung and Mendelson, 2004; Isomura et al., 2006; Clement et al., 2008; Quilichini et al., 2010; Hutt, 2011; Ferraris et al., 2018). Whether sequences represent internally generated representations (Pastalkova et al., 2008), preconfigured cell assemblies (Dragoi and Tonegawa, 2011; Liu et al., 2019) or a functional template of offline replay in the framework of memory consolidation (Lee and Wilson, 2002; Buzsáki, 2015; Pfeiffer, 2020) still remains to be elucidated. Our work demonstrates that sequences can also be recorded in a thalamic nucleus. This property appears to be specific to the NR, on the basis of recordings of other thalamic nuclei recorded in the vicinity. A systematic study of all thalamic nuclei should now be performed, to determine whether this property is specific to the NR. In the present work, we used anesthesia conditions; therefore, our results should also be validated during long sleep sessions. It is, however, noteworthy that anesthesia is likely to affect the network dynamics and neuronal firing. As shown by Luczak et al. (2007), anesthesia provides a regular alternation of UP and DOWN, whereas natural sleep displays much briefer DOWN states, resulting in highly asymmetric field events. They, however, show that

neuronal firing patterns at UP state onset bear clear similarities between anesthesia and natural sleep, arguing that the results obtained here are informative about natural brain dynamics.

The way mPFC and HPC sequences are generated remains poorly understood. Our results demonstrate that NR activity is a key regulator of mPFC and HPC sequence stability. Interestingly, previous (Ferraris et al., 2018) and present results demonstrate that the NR does not seem to regulate oscillatory patterns, as NR inactivation does not change the frequency of SOs, γ oscillations, and SPW-Rs. It is yet important to note that our NR manipulations indicate a causal role for NR activity, but not necessarily for the sequential organization of its neurons. However, the NR activity appears to organize the synchronization of γ oscillations between the HPC and the mPFC (Ferraris et al., 2018) as well as the sequential activation of cell assemblies in the mPFC during SOs or in the HPC during SPW-Rs. The NR is ideally located for such fine-tuning of cell activity, as it is bidirectionally connected to the mPFC and HPC (Vertes, 2006; Cassel et al., 2013; Varela et al., 2014). The fact that NR sequences always precede mPFC ones at UP state onset suggests that NR cells may directly drive mPFC cells. In keeping with this proposal, NR neurons have an excitatory action on HPC and mPFC by modulating/activating both interneurons and principal cells (Dolleman-Van der Weel et al., 1997; Dolleman-Van der Weel and Witter, 2000; Di Prisco and Vertes, 2006), their axons innervating the stratum lacunosum moleculare of CA1 of mainly the ventral and intermediate portions of the HPC (Wouterlood et al., 1990; Vertes et al., 2006), and the infralimbic, prelimbic, and anterior cingulate cortices of the mPFC with fibers terminating densely in layer 1 and layers 5/6 of these regions (Hoover and Vertes, 2012; Varela et al., 2014). However, the influence of NR activity on sequences during hippocampal SPW-Rs is more difficult to explain as SPW-Rs occur at variable times after UP state onset (Sirota et al., 2003; Battaglia et al., 2004). This rather suggests that the NR is an important hub in a wider network of networks, from which oscillations can emerge.

The presence of assemblies specifically in the NR is quite remarkable. Since NR neurons are involved in reference memory consolidation (Loureiro et al., 2012) and in spatial memory (Jankowski et al., 2014, 2015; Ito et al., 2015; Cholvin et al., 2018), NR assemblies may constitute an activity template used to organize information at the beginning of the UP state. Such a mechanism may enable the transmission of information to target areas in a packet-based manner (Luczak and Maclean, 2012), a default activity in a default mode (Sanchez-Vives and Mattia, 2014; Luczak et al., 2015).

Another remarkable feature of NR assemblies is their dorsoventral organization, which suggests a precise topological organization in terms of afferences and efferences despite the fact that NR does not have a layered organization (Jones, 1985; Bokor et al., 2002; Van der Werf et al., 2002; Dolleman-van der Weel et al., 2019). The NR probe was inserted in the brain with an angle to properly target the NR (Fig. 3Aa). As a result, the bottom-most sites were located more medially than the top-most ones. Although the mediolateral distance was $<250 \mu\text{m}$ between the top and bottom sites, we cannot fully rule out that the spatial organization we describe is mediolateral rather than dorsoventral (or both). Luczak et al. (2007) have shown that local sequences in the cortex are independent of global activity patterns, such as traveling waves, which argues in favor of local computation. However, the NR sequences described here are identified within a smaller spatial scale ($<250 \times 620 \mu\text{m}$ here vs $160 \times 1400 \mu\text{m}$ in Luczak et al. (2007)), indicating that those NR sequences might lie somewhere between entities of global patterns and of local

networks. The origin of the dorsoventral organization of these sequences remains to be elucidated. While NR outputs to the subiculum are topographically organized along the dorsoventral axis, no study has reported such precise topological organization of NR inputs (Dolleman-Van Der Weel and Witter, 1996). Only differences involving the rostrocaudal axis have been so far reported (Cassel et al., 2013), as well as a preferential targeting of NR fibers to prelimbic and infralimbic deep layers (5 and 6) and superficial layer 1 in mPFC (Berendse and Groenewegen, 1991; Vertes, 2004; Hoover and Vertes, 2012). The fact that we could not find any dorsoventral pattern in mPFC sequences does not mean that there is none to be found as, given the spatial organization of the mPFC, we could only target one layer. There is also no available information on the local connectivity among NR neurons, except a caudal to rostral pathway (Dolleman-Van der Weel et al., 1997). The NR includes different cell types, but the lack of specific molecular markers prevents, so far, a proper optogenetic investigation (Bokor et al., 2002; Walsh et al., 2017; Dolleman-van der Weel et al., 2019).

In conclusion, our results further support the concept that the NR plays a key role as an anatomic and functional hub between the mPFC and the HPC. The influence it exerts on mPFC and HPC information packets suggests that it strongly participates in the organization of information in both regions but also in the transfer of information from the HPC to the mPFC. Its internal organization allows the genesis of information packet sequences, which may represent similar features as those coded in the mPFC and the HPC.

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