Whole-Brain Propagation Delays in Multiple Sclerosis, a Combined Tractography-Magnetoencephalography Study

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Two structurally connected brain regions are more likely to interact, with the lengths of the structural bundles, their widths, myelination, and the topology of the structural connectome influencing the timing of the interactions. We introduce an in vivo approach for measuring functional delays across the whole brain in humans (of either sex) using magneto/electroencephalography (MEG/EEG) and integrating them with the structural bundles. The resulting topochronic map of the functional delays/velocities shows that larger bundles have faster velocities. We estimated the topochronic map in multiple sclerosis patients, who have damaged myelin sheaths, and controls, demonstrating greater delays in patients across the network and that structurally lesioned tracts were slowed down more than unaffected ones. We provide a novel framework for estimating functional transmission delays in vivo at the single-subject and single-tract level.

Key words: brain criticality; brain dynamics; brain networks; conduction velocities; magnetoencephalography; multiple sclerosis

Significance Statement
This article provides a straightforward way to estimate patient-specific delays and conduction velocities in the CNS, at the individual level, in healthy and diseased subjects. To do so, it uses a principled way to merge magnetoencephalography (MEG)/electroencephalography (EEG) and tractography.

Introduction
Higher-level cognitive functions might be emergent and depend from the regulated interaction of multiple brain areas (Stam, 2014). Hence, network theory has been used to describe such large-scale functional interactions within the brain (Bullmore and Sporns, 2009), as well as the underlying structural connections (Watts and Strogatz, 1998; Barabasi and Albert, 1999). From the functional standpoint, most studies used functional magnetic resonance imaging (fMRI), thereby focusing on the slow-evolving activity (typically in the timescale of seconds) and using assumptions of stationarity. The presence of time-averaged spatial patterns of correlated and anti-correlated activity was consistently reported (Fox et al., 2005), which also related to the structural topology (Honey et al., 2009; Tewarie et al., 2019). Per contra, brain activities do not exclusively generate periodic patterns, but also aperiodic ones emerge on the large scale (Zalesky et al., 2014). Structural topology imposes constraints on the evolution of brain dynamics, as shown by the fact that the probability of two brain regions activating sequentially is proportional to the coupling intensity along the brain tract linking them (Sorrentino et al., 2021b). However, structural topology alone is necessary, but not sufficient, to generate the observed dynamics, as delays, and noise, are also key elements (Niebur, 1991; Atay et al., 2004; Ghosh et al., 2008; Deco et al., 2009; Cabral et al., 2014; Petkoski and Jirsa, 2019; Tewarie et al., 2019). The time it takes two gray matter regions to sequentially activate depends...
on multiple factors, such as the distance separating them, the properties of the white matter bundle (structural tract) linking them (such as its diameter and myelination), and the overall topology of the network embedding them (Petkoski et al., 2018). For this reason, estimating a comprehensive map of the delays poses great challenges, and most modeling studies assume homogeneous velocities of conduction (typically ranging from 2 to 10 m/s; Sanz-Leon et al., 2015; Breakspear, 2017). Hence, the delays are conceptualized as purely dependent on distance (Ghosh et al., 2008; Petkoski et al., 2018; Petkoski and Jirsa, 2019). This is a suboptimal simplification, since we know (e.g., from routine examinations such as visual evoked potentials) that the velocities in highly myelinated, thick tracts can reach up to 150 m/s (Purves et al., 2001). Combining magnetoencephalography (MEG) and tractography one can track the beginning and the spread of a regional perturbation (Sorrentino et al., 2021b). Building on this framework, we hypothesized that the time it takes a signal to go from one region to another might be a candidate proxy for the delay of transmission across any given structural tract. Shorter delays might relate, among other factors, to diameter and myelination. This should result in perturbations spreading with velocities that are not constant but, rather, rise as a function of the length of the white-matter tracts (given that longer tracts are typically also the thicker and more myelinated ones). Conversely, we expected the width of the distribution of the delays to be much narrower than what would be expected from the length of the white matter bundles alone (since the longer tracts are hypothesized to be the fastest). We further reasoned that, if our measurements are related to myelination, then lesions in the myelin sheath should increase the delays. To test this hypothesis, we used source-deconstructed MEG signals to estimate the delays, defined as the average time it takes a perturbation generated in region $i$ to spread to region $j$, and then estimated the lengths of the white matter bundles in a cohort of 18 patients affected by multiple sclerosis and 20 healthy subjects using tractography. We used the delay in the subsequent recruitment of brain regions as a proxy for the conduction delay of an impulse between regions and estimated the corresponding velocities by dividing the tract lengths by the corresponding delays. First, we explored the distributions of the delays and velocities in healthy humans. Then, we evaluated the effect of the damage to the myelin sheath on the delays. To further verify the whole workflow, we applied the same procedure to a publicly available dataset based on combined electroencephalogram (EEG) recordings and tractography, and we were able to extract subject-specific delays and, thus, velocities, which had statistics that were similar to those obtained via MEG. The findings were further verified on an independent MEG/Diffusion Tensor Imaging (DTI) dataset, using a different parcellation and a different source-reconstruction algorithm.

Materials and Methods

Participants

The participants (of either sex) were recruited at the outpatient clinic of the Institute for Diagnosis and Cure Hermitage Capodimonte. The diagnosis of multiple sclerosis was made according to the revised 2017 McDonald criteria (Thompson et al., 2018a). Exclusion criteria were age <18 years, clinical relapse and/or steroid use in the three months before the study, inability to understand and complete "patient reported outcomes" and cognitive evaluation, or inability to undergo the MRI scan. All patients underwent a neurologic clinical examination, Expanded Disability Status Scale (EDSS) scoring, the Symbol Digit Modalities Test (SDMT) to measure cognitive impairment, the Fatigue Severity Scale (FSS), and the Beck Depression Inventory (BDI). The controls for the MS cohort were selected from among the caregivers and spouses of the patients. Genetic relatives were not allowed as controls. The subjects for the second, larger healthy cohort were selected as described previously (Sorrentino et al., 2021b). The independent EEG/DTI dataset is described in (Schirner et al., 2018). The demographics and main clinical and radiologic features of the MS cohort are summarized in Table 1. The study was approved by the local Ethics Committee (Prot.n.93C.E./Reg. n.14-17OSS).

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<th>Table 1. Features of the multiple sclerosis cohort</th>
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MRI acquisition and processing

Each MRI scan was performed immediately after the MEG recording on the same MRI scanner (1.5 Tesla, Signa, GE Healthcare). Analyzed sequences included echo-planar imaging for DTI reconstruction (TR/TE 12 000/95.5 ms, voxel 0.94 × 0.94 × 2.5 mm$^3$, 32 equally spaced diffusion-sensitizing directions, 5 B0 volumes) and 3D-FLAIR volume for WM lesion segmentation (TR/TE 7 000/145/1919 ms, echo train length 170, 212 sagittal partitions, voxel size 0.52 × 0.52 × 0.80 mm$^3$). Preprocessing of the diffusion MRI data were conducted using the software modules provided in the FMRIB software library (FSL; http://fsl.fmrib.ox.ac.uk/fsl). All diffusion MRI datasets were corrected for head movements and eddy current distortions using the "eddy_correct" routine (Smith et al., 2004), rotating diffusion sensitizing gradient directions accordingly (Leemans and Jones, 2009), and a brain mask was obtained from the B0 images using the Brain Extraction Tool routine (Smith, 2002). A diffusion-tensor model was fitted at each voxel, and fiber tracts were generated over the whole brain by deterministic tractography using the Fiber Assignment by Continuous Tracking (FACT) algorithm implemented in Diffusion Toolkit (angle threshold 45°, spline-filtered, masking by the FA maps thresholded at 0.2). Two cortical study-specific ROI datasets were obtained by masking the ROIs available in the AAL atlas (Tzourio-Mazoyer et al., 2002) and in an MNI space-defined volumetric version of the Desikan–Killiany–Tourville (DKT) region of interest (ROI) atlas (Klein and Tourville, 2012) using the gray matter (GM) probability map available in the Statistical Parametric Mapping software package (SPM), thresholded at 0.2. This was done to limit the endpoints of the fibers to cortical and adjacent juxtacortical white matter voxels in the subsequent ROI-based analysis of the tractography data. The analysis was replicated twice to ascertain how robust the method was when used with specific brain parcellations. To obtain the corresponding patient-specific ROI sets, each participant’s FA volume was spatially normalized (Friston et al., 1995) to the FA template provided by FSL using SPM12, and the resulting normalization matrices were inverted and applied to the two ROI sets. Additionally, for each subject, the MS lesion map was obtained by segmenting the 3D-FLAIR volume using the lesion prediction algorithm (Schmitt et al., 2012) implemented in the Lesion Segmentation Tool (LST toolbox version 3.0.0; www.statistical-modeling.de/lst/html) for SPM. The 3D-FLAIR volume was then co-registered to the EPI of the patient (Ashburner and Friston, 1997), and the coregistration matrix was applied to the corresponding WM lesion volume, which was resampled by nearest-neighbor interpolation, thus obtaining the patients’ lesion masks, which would be coregistered with the DTI volume. Finally, for each patient the average length of the fibers connecting each pair of ROIs and whether the voxels crossed by those fibers had MS lesions were calculated separately for the AAL and DKT ROI sets, using an in-house routine written in Interactive Data Language (IDL, Harris Geospatial Solutions). To this end, each tract length was computed, for each subject, as the mean of

8808 · J. Neurosci., November 23, 2022 · 42(47):8807–8816 Sorrentino et al. · The Topochronic Map of the Human Brain
the physical distances covered by the fibers composing the tract. These in
turns were calculated as the sum of the Euclidean distances between the
fiber subsequent turning points, including the two extremities, which are
the output of the FACT algorithm.

**MEG preprocessing**

MEG preprocessing and source reconstruction were performed as previ-
ously described (Sorrentino et al., 2018). Preprocessing and source
reconstruction operations were carried out using the FieldTrip Toolbox
(Oostenveld et al., 2011). Each participant underwent a MEG recording,
composed of both eyes-closed resting-state segments of 3’30” each. Four
anatomic coils were applied on the head of each participant and their
position was recorded along with the position of four head anatomic
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position was recorded along with the position of four head anatomic

**Source reconstruction**

Signal time series were reconstructed using both the AAL and DKT
atlases (Tzourio-Mazoyer et al., 2002; Gong et al., 2009), which consist
of 116 and 84 ROIs, respectively. The reconstruction took place using
the volume conduction model proposed by Nolte (Nolte, 2003). Based
on the native MRIs of each subject, the linearily constrained minimum
variance (LCMN; Hillebrand et al., 2016) beamformer was applied to
reconstruct the signal sources based on the centroids of each ROI

(Chialvo, 2010). Source reconstruction

**Neuronal avalanches and branching parameter**

To study the dynamics of brain activity, we based our analysis on “neuronal avalanches.” First, the time series
for each ROI was downsamped to 512 Hz and discretized by calculating the z score; then the positive and
negative excursions beyond a threshold were identified (Fig. 1). The main results reported here refer to a thresh-
hold equal to three standard deviations (|z| = 3), but thresholds of 2.5 and 3.5 were also tested. A neuronal
avalanche began when at least one ROI went above the threshold (|z| > 3) and ended when all the ROIs were
below the threshold (Beggs and Plenz, 2003; Shriki et al., 2013). Before proceeding with the analyses, we binned
the data, to ensure that we captured any critical dynamics, if present. To estimate the suitable time bin length, for
each subject, for each neuronal avalanche, and for each time bin duration, the branching parameter $\sigma$ was
estimated (Harris, 1964; Haldeman and Beggs, 2005). In fact, systems operating at criticality typically display a
branching ratio of $\sim 1$ (Chialvo, 2010). The branching ratio was calculated as the geometrically averaged (over
all the time bins) ratio of the number of events (activations) between the subsequent time bin (descendants)
and that in the current time bin (ancestors) and then averaging the avalanche-specific branching ratio over all
the avalanches (Bak et al., 1987). More specifically:

$$\sigma_i = \frac{\sum_{j=1}^{\text{events } j} \left( \frac{\text{events } j+1}{\text{events } j} \right)}{\text{delay } j+1}$$

where $\sigma_i$ is the branching parameter of the $i^{th}$ avalanche in the data-
set, $N_{\text{bin}}$ is the total number of bins in the $i^{th}$ avalanche, $\text{events } j$ is the total number of events active in the $j^{th}$ bin, and $N_{\text{real}}$ is the total number of avalanches in the dataset. We tested bins from 1 to 5 and picked three for further analysis.

**Estimation of delay matrices and velocity matrices**

The delays were estimated for each avalanche. The procedure is schema-
tized in Figure 1. In an avalanche, from the moment region $i$ activated, we recorded how long it took region $j$ to activate. These are what we con-
sidered to be delays. Hence, for each avalanche we obtained a matrix, in
which the rows and columns represented brain regions and the entries
contained the delays. We then averaged across all the avalanches belong-
ing to one subject, obtaining an average $i^{th}$ delay. The average was per-
formed disregarding zero entries, since each avalanche-specific matrix is
very sparse. With this procedure, a subject-specific delay matrix was built.
Averaging across subjects (again discarding zero entries) yielded a
group-specific matrix. Note that it can happen that no delay estimation
is available for a specific edge in a specific subject, since the esti-
mation depends on the fact that an avalanche propagated across the
edge. Finally, we obtained a velocity matrix, dividing the tract lengths,
obtained from MRI, by the corresponding delays. Velocities were only
obtained for the edges where a direct structural connection was present.

**Experimental design and statistical analyses**

To build null models to test the functional delays estimates, we random-
ized the temporal order of the avalanches without changing the spatial
structure. To this end we built a surrogate dataset where the avalanche
time points were randomly shuffled, but the regions recruited at any

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**Figure 1.** Top left, The structural connectome. From the tractography, we obtained an estimate of each tract length. Averaging across subjects provided a group-level estimate. Top right, Source-reconstructed MEG time-series. The time

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(Hillebrand et al., 2016). ROIs belonging to the cerebel-
lum were excluded because of the low reliability of their
source reconstruction (Lardone et al., 2018), for a total
of 90 ROIs in the AAL atlas and 84 ROIs in the DKT
atlas.
given time point were fixed. By doing this, the time-structure of the recruitment of regions was disrupted, but the purely spatial component was retained. After the permutations, the average delays were again computed for each edge. The procedure was repeated 1000 times, yielding a hundred surrogate delays for each edge. We then used the delays estimated from the random surrogates to compute functional velocities (dividing the length of the structural edges by the surrogate delay). Finally, we computed the growth of the velocities as a function of the length of the structural tracks and compared this with the observed distribution. To test differences between the distribution of the delays in the healthy subjects and patients, we used the Kolmogorov–Smirnov (KS) test. To perform the edge-wise comparison of the delays in the healthy versus lesioned edges in MS patients, we used permutation testing (Nichols and Holmes, 2002). In short, we tested the null-hypothesis that lesions in the edges would not have an impact on the delays. First, we calculated the average edge-wise difference between the delays in each patient and the average delay in the corresponding edge in the controls. Then, we randomly selected a subset of the differences in the delays with the same size as the number of lesioned edges and computed its average. We repeated this procedure 1000 times, building a distribution of the differences in the delays that are to be expected by randomly drawing a subsample of the edges. Finally, we compared this distribution to the observed difference in the lesioned edges to obtain the probability of observing the data under the hypothesis that the edge-specific lesions would not slow down the functional transmission.

Results

Delay estimation

In this study, we noninvasively estimated the functional delays in transmission across the network of white matter bundles in vivo. To this end, we combined source-reconstructed magnetoencephalography and tractography. An overview of the pipeline is shown in Figure 1. The upper row of Figure 2 shows the average structural matrix for the controls (i.e., the tract lengths) with the corresponding distributions of the tract lengths on the right. The middle and bottom rows of Figure 2, respectively, show the matrices and distributions of the delays and the velocities. Importantly, the structure of the delay matrix is related to the length of the structural tracts. However, the width of the distribution of the delays is much narrower than what would be expected given constant propagation velocities (see below). When estimating the velocities, we found that a consistent, fat-tailed distribution emerged. As expected, because of the heterogeneity of the tracts, the velocities appeared to be far from homogeneous, ranging from ~2 to ~60 m/s. This range appears to be in accordance with known velocities of major myelin tracts

Data and materials availability

Code is available on GitHub. The MEG data and the reconstructed avalanches are available upon request to the corresponding author, conditional on appropriate ethics approval at the local site. The availability of the data was not previously included in the ethical approval, and therefore data cannot be shared directly. In case data are requested, the corresponding author will request an amendment to the local ethical committee. Conditional to approval, the data will be made available.

Figure 1. A, Upper panel, top, The Topochronic Map of the Human Brain. A, Middle panel, top, Time delays in seconds, log-scale, for each edge connecting two brain regions. The color code conveys the average time it took region j to become active provided region i had been active earlier. Values are in seconds and reported on a log-scale to highlight the texture of the delay matrix, which appears to be strongly correlated to the length matrix. Bottom, Middle panel, top, The Glass-brain provides an overview on the topography of the functional edges with fast functional velocities. In particular, the blue dots represent brain regions, and the edges signify the velocities. Only structural connections with transmission speed higher than 15 m/s are shown. The width of the edge is proportional to its velocity.

Figure 2. A, Top, Track lengths. Rows and columns represent brain regions, and the color code conveys the corresponding length of the tract linking any two regions. Middle, Group average of the delays. Rows and columns represent brain regions, while the color code represents the average time it took region j to become active provided region i had been active earlier. Values are in seconds and reported on a log-scale to highlight the texture of the delay matrix, which appears to be strongly correlated to the length matrix. Bottom, As before, brain regions are represented as rows and columns, while the matrix entries represent the velocities, expressed in meters/s. B, Top, middle, and bottom, Histograms corresponding to the matrices for the delays, lengths, and velocities, respectively.
A Group-level relationship between tract lengths and functional delays. Blue dots represent iso-hemispheric edges, while red dots represent cross-hemispheric edges. B, For each of the 20 healthy controls, the red dot represents the intensity of the relationship between the tract lengths and delays, calculated using Spearman’s correlation coefficient. For each subject, the avalanches were randomized by shuffling the time sequence while preserving the spatial pattern of the active regions within each timestep. Based on these newly randomized avalanches, the edgewise average delay was calculated and then related to the tract lengths. The procedure was repeated 1000 times per subject. The resulting Spearman correlations are represented as blue dots. C, For each subject, the delays were averaged according to the percentile of the corresponding length. Hence, each blue dot represents the average delay across all edges whose length belonged to the (patient-specific) nth percentile. The colored lines show the delays that would be expected if the velocities were homogeneous across all edges. In this case, the growth of the delays followed the growth of the tract lengths. D, The red dots represent the observed velocities (averaged according to the percentile of the tract length). The red line represents the average velocity estimated based on the delays derived from the randomly shuffled avalanches. E, Group-level relationship between the velocities. Blue dots represent iso-hemispheric edges, while red dots represent cross-hemispheric edges.

Statistical validation
First, we compared the delays retrieved from the random time-series to the tract lengths, obtaining 100 Spearman’s rs, to which we compared the observed correlation, \( p < 0.001 \). We then used the random surrogates to compute random velocities, i.e., dividing each edge length by the corresponding delay derived from the random surrogates and, using the same validation scheme, we showed that the random velocities appeared to be more strongly related to the tract lengths than the observed velocities (\( p < 0.001 \)). In other words, when we divided the lengths by the randomized delays, the resulting velocities became a function of the tract length alone. On the other hand, when we divided the tract lengths by the observed delays, the longer tracts appeared to be faster than if the delays were only a function of distance. The mean (red line) and upper and lower bounds (shaded area) of the surrogate velocities derived from surrogate delays (grouped by the percentile of the corresponding tract length) are shown in Figure 3D. The delays derived from the surrogate data led to a higher estimate of the velocities compared with the observed ones. Finally, as shown in Figure 3E, we confirmed, using real data, that the transmission velocities grew as a function of the length of the tracts, such that the longer tracts were also the faster ones. All in all, this part of the analysis showed a finely regulated relationship between the delays and the lengths of the structural tracts, which implies nonhomogeneous functional velocities.

Delay estimation in multiple sclerosis patients
Next, we sought to test our framework in patients affected by multiple sclerosis, which is a prototypical disease in which myelin in the CNS is selectively attacked by the immune system (Thompson et al., 2018b). We excluded patients with severe functional impairment (Expanded Disability Status Scale, <7)) so that we could obtain a picture that is likely to be influenced more by demyelination than by degenerative phenomena (Frischer et al., 2009). We expected to observe greater delays (and lower

(Waxman and Bennett, 1972; Waxman, 1980; Purves et al., 2001). When plotting the distribution of the fastest edges (see glass brains in Fig. 2), it appears that these were not evenly distributed across the brain and that the cross-hemispheric edges were preferentially selected as the fastest ones, a finding which agrees with the literature (Shen et al., 2015). We found that the delays grew as a function of the length of the tracts, as shown by the Spearman correlation between edge length and delays (\( p = 0.35, p < 0.0001 \); Fig. 3A). The results held using Pearson’s correlation (\( p = 0.31, p < 0.0001 \)). This relationship held at the individual level (Fig. 3B). However, while the delays were related to the tract lengths, they were not only determined by the lengths. In fact, the tract lengths ranged across an order of magnitude, whereas the corresponding increase in the delays was only moderate. For comparison, in Figure 3C, we show the expected delays given constant velocities and reveal that the observed delays were remarkably steady despite the difference in the lengths of the structural tracts.

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velocities) in the patient population compared with the controls. As shown in Figure 4A, the average delay per percentile was consistently higher in the multiple sclerosis patients compared with the controls. The empirical cumulative distribution function confirmed the difference in the delays in patients compared with the controls (KS test, \( p < 0.001 \); Fig. 4B). Note that, in this analysis, similar to what we did before, we grouped the delays according to percentile across the whole group. Hence, the average value refers to the average delay of all the edges that fall within a given length percentile, with the percentile defined for each subject. We then moved on to an edge-wise comparison. To this end, for each patient, the average delay of each edge was compared with the average delay of each corresponding edge in the controls. Hence, we obtained the difference between the edge-specific delay and the corresponding average delay in the controls for each patient. To base our findings on a more stable estimate, we considered only edges for which a delay estimate was available for each of the 20 controls (i.e., a total of 1202 edges). Figure 4C shows the distribution of the differences between the delays (patients minus controls). The distribution is not centered around zero, as would be expected if the patients did not have longer delays. Instead, it is evident at the visual inspection that the distribution is heavily skewed toward positive values, implying longer delays in the patients compared with controls. Finally, we investigated edge-specific lesions. In fact, while it is reasonable to expect globally greater delays in the patients compared with the controls, the delays corresponding to edges that were lesioned might be more lengthened compared with the delays corresponding to healthy edges. For each patient, we classified edges as healthy or lesioned, based on the presence or absence of structural damage (i.e., the tracts were considered as lesioned if any of the corresponding voxels was lesioned). Under the null hypothesis that the delays corresponding to structurally lesioned edges do not differ from the delays corresponding to nonlesioned edges, we calculated the average delay difference in random samples of nonlesioned edges (in patients), with the size of the sample equal to the number of lesioned edges. We repeated this procedure 1000 times and compared the obtained distribution to the observed difference in the lesioned edges. The results are shown in Figure 4D. The delays in the lesioned edges slowed more (with respect to the corresponding delays in the healthy population) than the healthy edges. Hence, selecting a subset of edges based on the structural information, we retrieved a difference in the temporal structure of the functional dynamic. This difference would not be expected if the structural damage was unrelated to the delays, as shown by the permutation analysis (\( p < 0.001 \)).

**Replication in independent datasets**

The results of the delay estimation were tested using an independent dataset based on co-registered MRI and EEG, and all the main findings were confirmed (Fig. 5). Note that the cleaning and source-reconstruction algorithms were different in this dataset as compared with ours, showing robustness to both the technique used and data processing. Furthermore, we replicated the main results using different parcellations (AAL and DKT), varying the z score threshold (2.5 and 3.5 in addition to the 3.0 reported here), varying the binnings (see Materials and Methods), and using yet another source-reconstruction algorithm (i.e., the “residual variance” method). The residual variance is a particular dipole-fit approach that involves the minimization of the signal that remains unexplained by a current source model in which dipoles are assumed to have fixed position and fixed or varying orientation. Conversely to LCMV, there is no linear constraint involved in the minimization process (Grech et al., 2008). Furthermore, a larger MEG/tractography dataset, involving 47 young healthy subjects, was also used to further explore the robustness of our findings. These results were replicated using the AAL atlas. Finally, we varied the minimum size of the avalanches used to compute the delays, from taking all avalanches into account, up to selecting only those longer than 10/15 samples (analyses shown in Fig. 5). Finally, we confirmed that the delay matrices showed convergence as the sample size increased, as a further check on the validity of our results (Fig. 5).

**Discussion**

**The tophochronic map**

In this study, we developed a novel method to quickly and noninvasively obtain an estimate of the time delays that occur when locally generated perturbations spread to other brain regions. Our methodology was able to capture the presence of homogeneous delays across brain regions, although the length of the structural tracts grew according to a fat-tail distribution (Sporns and
If the delays were only dependent on the tract lengths, they should be longer for longer tracts (i.e., it should take longer to cover a greater distance). This effect should be major, given that longer white-matter tracts are roughly one order of magnitude longer than shorter ones (Bullmore and Sporns, 2009). The fact that the delays do not scale with increased tract lengths implies one or more compensatory mechanisms. Such mechanisms might involve varying axonal diameters, myelination, and network effects. In particular, myelination greatly affects the velocity of propagation (Waxman, 1980; Waxman et al., 1980; Horowitz et al., 2015). In fact, given our estimated delays, we found that, in the healthy subjects, the velocities ranged from \( \sim 3 \) to \( \sim 60 \) m/s. This observation is in sharp contrast with the typical simplifying assumptions, made in modeling studies, that velocities are constant, which would be expected to lead to a broad range of delays across the network. The fact that the transmission between the regions connected by the longest edges is faster than those with shorter edges is likely important in terms of the unfolding of the dynamics and should not be surprising considering the experimental results on stimulation, although until this current study solid measurements of personalized links on the whole-brain level had been missing. Hence, noninvasive estimates of the temporal constraints in vivo are highly relevant for modeling individual dynamics. Furthermore, many of the fast edges connect regions that are highly central to the brain network, thus, likely causing reverberations that greatly and nonlinearly impact the average transit time (He et al., 2007). Although we can only speculate, we think that it is possible that homogeneous delays, that is, having perturbations reach wide-spread brain regions or arrive at a focal point from wide-spread regions simultaneously, would be favorable for allowing the brain to have simultaneous access to information from multiple locations across the brain, a concept which fits well into the framework of global workspace theory (Dehaene et al., 1998). This supposition finds support in that white matter damage impairs conscious access in multiple sclerosis patients (Reuter et al., 2009; Rosanova et al., 2012). A reduction in the complexity of the spatiotemporal spreading of such perturbations (referred to as “neuronal avalanches” within the framework of critical dynamics) has been shown to be related to states of reduced consciousness (Tononi and Sporns, 2003) as well as to neurodegeneration (Rucco et al., 2020; Sorrentino et al., 2021a). Importantly, our analyses focused on rare, intermittent, large-scale bursts of activations, which have been consistently observed in human brains (Shriki et al., 2013). The importance of such rapid transients to large-scale brain dynamics is confirmed by recent findings showing that the patterns of functional connectivity are shaped by specific, short moments in time (Zamani Esfahiani et al., 2020). Furthermore, the fact that avalanches preferentially spread along structural tracts indicates that functional delays might be used as a proxy to estimate the velocities across individual tracts (Sorrentino et al., 2021b). As we investigated the topographic distribution of the fast edges, we found that they were spatially distributed nonhomogeneously. Importantly, fast connections seem to be preferentially cross-hemispheric, a finding that is not surprising from a neuroanatomical standpoint, provided that these connections are mediated, for example, via the corpus callosum (Caminiti et al., 2013). While our data were based on broad-band data, the patterns that emerged were not dominated by the occipital or frequency, as it is often the case with M/EEG data. However, we wish to stress that such results might have been biased by the tractography, which may have preferentially estimated tracts in specific anatomic regions (Shen et al., 2015). Hence, further validation is needed to confirm this finding, which should be considered explorative. The replication of our results in an independent MEG dataset as well as in a publicly available multimodal EEG-MRI dataset (Schirmer et al., 2018) indicates the reliability of our findings by increasing the probability that they are not modality specific.

### Implications in neurologic diseases and multiple sclerosis

As explained, myelination is believed to greatly influence conduction velocity and, hence, to modulate delays. To test this, we
measured the delays in multiple sclerosis patients. As expected, the delays were greater in multiple sclerosis patients than in matched healthy controls. Selecting MS patients at a fairly early stage should make the role of demyelinating lesions more prominent than the role of degeneration (Stadelmann et al., 2011). However, this was not quantified and remains a potential source of confounds that needs further investigation. To provide edge-specific information about the time delays, we used subject-specific lesion masks to separate lesioned edges from nonlesioned ones in the patients. We focused the analysis on the delays to avoid potential biases induced by unreliable length estimates because of the demyelinating lesions (Delettre et al., 2019). We observed, for both the lesioned and nonlesioned edges, that functional delays increased in the multiple sclerosis patients, in accordance with the hypothesis that damage to the myelin would provoke longer delays. This also provides support for the claim that our measurements are related to temporal patterns of delays that are imposed on the overall structural connectome. The fact that avalanches propagate more slowly along lesioned edges shows the relevance of the direct pathways in determining the edge-specific delays given the expected relationship between the structural integrity of the tract and the velocity of propagation along it. However, it is important to stress that the role of network effects cannot be easily disentangled (Bullmore and Sporns, 2009). In fact, the transmission in patients was slower even across unaffected tracts. This might be interpreted as an expression of the fact that the delays likely depend from a combination of both direct and indirect paths through which a perturbation can potentially travel between two regions. In this sense, one does not expect two regions that are linked by a healthy edge that is embedded in a diseased network to communicate as quickly as two regions that are also linked by a healthy edge as well as embedded in a healthy network. Other contributing factors may include the erroneous classification of damaged but subthreshold edges as healthy.

Cross-dataset validation and limitations

The strength of this study is that the tract lengths and the delays were measured using EEG, using a different preprocessing pipeline, applying the robustness of our findings. Similar results were obtained of the perturbations and the brain parcellation, again showing the potential to improve virtual, personalized in-silico brain models (Hashemi et al., 2020). Importantly, the fact that delay estimation can be obtained from EEG, a widely available technique, allows nearly every facility to include subject-specific delays into personalized brain models. Furthermore, large cohorts will be needed to get normative data from large, stratified cohorts, to create priors to be included in personalized models, in case subject-specific estimations are not available. In conclusion, we proposed a simple method for combining multimodal imaging within the framework of statistical mechanics, to derive subject-specific topochronic maps of large-scale brain dynamics in both healthy and diseased populations.

References