Assessing directionality and strength of coupling through symbolic analysis: an application to epilepsy patients

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Inferring strength and direction of interactions from electroencephalographic (EEG) recordings is of crucial importance to improve our understanding of dynamical interdependencies underlying various physiologic and pathophysiologic conditions in the human epileptic brain. We here use approaches from symbolic analysis to investigate—in a time-resolved manner—weighted and directed, short- to long-ranged interactions between various brain regions constituting the epileptic network. Our observations point to complex spatial-temporal interdependencies underlying the epileptic process and their role in the generation of epileptic seizures, despite the massive reduction of the complex information content of multi-day, multi-channel EEG recordings through symbolisation. We discuss limitations and potential future improvements of this approach.
1. Introduction

Epilepsy is one of the most common neurological disorders, second only to stroke, that affects approximately 1% of the world’s population [1, 2]. In about 25% of patients, epileptic seizures—the cardinal symptom of epilepsy—can not be controlled by any available therapies [3–7]. In order to enhance the quality of life for epilepsy patients, there is thus a great need for improved therapeutic possibilities. An epileptic seizure is defined as “a transient occurrence of signs and/or symptoms due to abnormal, excessive or synchronous neuronal activity in the brain” [8, 9]. Epileptic seizures can be divided into two main categories: focal seizures, which appear to originate from a circumscribed brain region (the so called seizure onset zone (SOZ) [10]) and which may or may not remain restricted to this region, and generalized onset seizures, which appear to engage almost the entire brain. These concepts of focal and generalized seizures, however, are being challenged by increasing evidence of seizure onset within a network of brain regions (the so called epileptic network) which led to a new approach to classification of seizures and epilepsies [11, 12]. In an epileptic network, all its constituents can contribute to the generation, maintenance, spread, and termination of seizures as well as to the many pathophysiologic phenomena seen during the seizure-free interval [13–17]. The concept of an epileptic network may also explain the apparent contradictory findings of seizure precursors that are not confined to the SOZ or its immediate surroundings but can be observed in remote or even contralateral brain regions [18–27].

An improved understanding of the epileptic network and its complex dynamics can be achieved with time series analysis techniques that aim at characterising interactions and their properties, namely strength and direction [28–35]. Over the last years, a number of such bivariate analysis techniques have been proposed, ranging from linear to nonlinear ones. Comprehensive overviews concerning these techniques and their applications in diverse fields can be found in [36–43]. Here, we concentrate on information theoretic approaches [44], and particularly on those that utilize symbol sequences [45–47] derived from empirical time series to characterise interactions. For the strength of interaction we will consider the order parameter $\hat{\gamma}$ proposed by Liu [48] and for the direction of interaction the symbolic transfer entropy [49, 50]. Both approaches allow for a robust and computationally fast quantification of the strength and the preferred direction of information flow between time series from observed data.

Previous studies [49, 50] that investigated directed interactions in the human epileptic brain provided evidence that the aforementioned approaches allow one to reliably identify the hemisphere containing the SOZ without observing actual seizure activity. In these studies, however, analyses were restricted to data from patients with mesial temporal lobe epilepsies and to recordings from within the mesial temporal lobes. Here, we extend these studies in several aspects. First, we investigate multi-day (on average about 4 days), multi-channel electroencephalographic data recorded invasively from 11 patients that suffered from seizures with different anatomical onset locations. Second, we investigate—in a time-resolved manner—changes in interdependencies between all sampled brain regions, thus taking into account interactions for a variety of physiologic and pathophysiologic conditions. Third, we estimate both strength and direction of interactions in order to effectively distinguish the various coupling regimes (uncoupled, weak to strong couplings) that may be observable in the human epileptic brain and in order to avoid misinterpretations [31, 50]. Fourth, we address the question of who is driving preferentially whom in large-scale epileptic brain networks and whether consistent changes in directed interactions can be identified when approaching epileptic seizures.

2. Strength and direction of interaction from symbolic analysis

In the following, we denote with $x_i := x(i \Delta t)$ and $y_i := y(i \Delta t)$, $i = 1, \ldots, N$, time series of observables of systems $X$ and $Y$ recorded simultaneously with sampling interval $\Delta t$. We make use of the seminal work of Bandt and Pompe [46] and derive symbol sequences from reordering
the amplitude values of time series. Let \( l \) and \( m \) denote the embedding delay and embedding dimension, which were chosen appropriately for symbolisation. Then \( m \) amplitude values \( s_i = \{x_i, x_{i+l}, \ldots, x_{i+(m-1)l}\} \) for a given, but arbitrary timestep \( i \) are arranged in ascending order \( \{x_i+(k_1-1)l \leq x_{i+(k_2)-1)l \leq \cdots \leq x_i+(m-1)l\} \) with rank \( k_{ii} \). In case of equal amplitude values the rearrangement is carried out according to the associated index \( k \), i.e., for \( x_i+(k_1-1)l \) we write \( x_i+(k_2-1)l \leq x_i+(m-1)l \) if \( k_1 < k_2 \). This ensures that every \( s_i \) is uniquely mapped onto one of the \( m! \) possible permutations, and a permutation symbol is defined as

\[
\hat{x}_i := (k_{i1}, k_{i2}, \ldots, k_{im}).
\]  

The symbol sequence \( \hat{y}_i \) for system \( Y \) is defined in complete analogy.

A robust method to estimate the strength of interaction from symbol sequences was proposed by Liu [48]. It is based on consistent changing tendencies of temporal permutation entropies \( H_X(w_\eta) = \sum_i p(\hat{x}_i) \log p(\hat{x}_i) \), where the sum runs over all possible symbols. \( H_Y(w_\eta) \) is defined in complete analogy. In order to estimate these changing tendencies, symbol sequences \( \hat{x}_i \) and \( \hat{y}_i \), \( i=1, \ldots, N \), are split into \( N_\eta \) segments \( w_\eta \) with \( \eta = 1, \ldots, N_\eta \) consisting of \( N_w \) datapoints, and permutation entropies \( H_X \) and \( H_Y \) are estimated for each segment \( w_\eta \). \( H_X(w_\eta) \) and \( H_Y(w_\eta) \) can be expected to be similar if some functional relationship—in the sense of generalized synchronization [51]—exists between states of \( X \) and \( Y \). The changing tendency for a time series from system \( X \) can be quantified with

\[
S_X(w_\eta) = \begin{cases} 1 & \text{if } H_X(w_{\eta+1}) > H_X(w_\eta) \\ -1 & \text{else} \end{cases}
\]  

and in complete analogy for a time series from system \( S_Y \). A measure for the strength of interaction between systems \( X \) and \( Y \) can then be defined as

\[
\hat{\gamma} = \frac{1}{N_\eta} \sum_{\eta=1}^{N_\eta} S_X(w_\eta) S_Y(w_\eta).
\]  

This index will be around zero for independent time series and close to unity for generalized synchronization. Note that \( \hat{\gamma} \) might attain slightly negative values.

Symbolic transfer entropy (STE) [49, 50] has been proposed as a computational fast and robust method to quantify the direction of interaction between coupled systems. It is based on Schreiber’s transfer entropy [52] that allows one to distinguish effectively driving and responding elements and to detect asymmetry in the interaction of systems. STE overcomes some of the limitations of previous techniques that aim at estimating transfer entropy from time series, such as requiring fine tuning of parameters and being highly sensitive to noise contributions [53–55]. More specifically, STE makes use of relative frequencies of symbols to estimate the transition probabilities required to quantify the deviation from the generalised Markov property, \( p(x_i|x_{i-1}, y_{i-1}) = p(x_i|x_{i-1}) \), where \( p(\cdot|\cdot) \) denotes the conditional transition probability density. With transfer entropy this deviation is formulated as a Kullback-Leibler entropy between \( p(x_i|x_{i-1}, y_{i-1}) \) and \( p(x_i|x_{i-1}) \).

With given symbol sequences \( \hat{x}_i \) and \( \hat{y}_i \), symbolic transfer entropy is then defined as

\[
\hat{T}_{Y \rightarrow X} = \sum_{\eta} p(\hat{x}_i, \hat{x}_{i-1}, \hat{y}_{i-1}) \log \frac{p(\hat{x}_i|\hat{x}_{i-1}, \hat{y}_{i-1})}{p(\hat{x}_i|\hat{x}_{i-1})}.
\]  

where the sum runs over all symbols. \( \hat{T}_{Y \rightarrow X} \) is defined in complete analogy, and the directionality index \( \hat{T} := \hat{T}_{X \rightarrow Y} - \hat{T}_{Y \rightarrow X} \) allows one to quantify the preferred direction of information flow between systems \( X \) and \( Y \). \( \hat{T} \) attains positive values for unidirectional coupling with \( X \) as the driver, negative values for \( Y \) driving \( X \), and \( \hat{T} \approx 0 \) for symmetric bidirectional coupling.

Since its invention, STE has been used to study interactions in various disciplines ranging from quantum [56], plasma [57], and laser physics [58] via neurology [59], cardiology [60] and anesthesiology [61–64] to the neurosciences [65]. In addition, extensions have been proposed that
allow one to estimate the dominating direction of information flow even from short or transient signals [65]. A recent modification, termed partial symbolic transfer entropy [66, 67], extends STE for multivariate time series and can help distinguish between direct and indirect interactions.

Estimating interaction properties with information-theoretic measures is prone to biases and statistical errors which depend on the method used and on the characteristics of the data [44, 68–71]. For both, the order parameter \( \hat{\gamma} \) and the directionality index \( \hat{T} \), asymptotic distributions are unknown, and in the following we apply averaging procedures (both in time and space) to decrease the variance of estimation error and consider both aspects of interaction, strength and direction, to minimise the risk for misinterpretations and to effectively distinguish the various coupling regimes.

Based on previous investigations that employed bivariate time series analysis techniques to characterise both strength and direction of interactions in studies on coupled model systems [31, 49, 50, 72, 73], we expect the order parameter \( \hat{\gamma} \) and the directionality index \( \hat{T} \) to depend on the coupling strength \( \kappa \) in some driver–responder system as follows (cf. Fig. 1): \( \hat{\gamma} \) increases monotonously with increasing \( \kappa \) until \( \hat{\gamma} = 1 \) for fully synchronised systems. This dependency thus correctly reflects an increasing strength of interaction due to the increased coupling. On the other hand, \( \hat{T} \) correctly detects the direction of interaction for some intermediate values of \( \kappa \) only. Moreover, \( \hat{T} \approx 0 \) for uncoupled (\( \kappa = 0 \) and \( \hat{\gamma} = 0 \)) or for fully synchronised systems (\( \hat{\gamma} = 1 \)). Distinguishing these coupling regimes thus requires checking the strength of interaction. When investigating interacting systems using empirical data, the coupling strength is usually not known a priori, but we expect the aforementioned dependencies to hold (at least approximately) for such data.

3. Measuring strength and direction of interactions in the human epileptic brain

We retrospectively analysed strength and direction of interactions between various brain regions by applying the aforementioned methods to multi-channel, multi-day electroencephalographic (EEG) data recorded intracranially in 11 patients (4 women, 7 men; mean age at onset of epilepsy 10.5 years, range 0–32 years; mean duration of epilepsy 28 years, range 5–53 years). These data had been recorded during the presurgical evaluation of intractable focal seizures with different anatomical onset locations. The patients had signed informed consent that their clinical data might be used and published for research purposes. The study protocol had previously been approved by the ethics committee of the University of Bonn. EEG data were recorded from, on average, 48 sites (range 21–73; cf. Fig. 2), band-pass-filtered between 1–45Hz and sampled at
200 Hz (sampling interval $\Delta t = 5$ ms) using a 16 bit analog-to-digital converter. Recordings lasted, on average, 98.6 h (range 25.4–206.2 h) during which seven seizures/patient (range 1–21) were captured. In five patients, seizures originated from the mesial temporal lobe, in two patients from the lateral temporal lobe, and in four patients from the frontal lobe. Postoperatively, all patients were seizure-free [74].

As a compromise between the statistical accuracy for the calculation of interaction indices and approximate stationarity [75], we divided the data into a sequence of non-overlapping segments of 20.48 s duration (corresponding to 4096 data points). This allowed us to calculate $\hat{\gamma}$ and $\hat{T}$ for each combination of pairs of recording sites in a time-resolved manner. If not stated otherwise, all calculations were performed with $m = 5$, $l = 3$, $N_w = 2048$, and $N_u = 204$ [76, 77].

Using knowledge concerning location and extent of the SOZ, which is defined by the electrode contacts showing initial seizure activity [10], we assigned all electrode contacts to three location categories:

- **focal** ($f$): electrode contacts located within the SOZ (on average for all patients 13.0 % of all contacts, varying between 5.0 % and 20.8 %);
- **neighbour** ($n$): electrode contacts not more than two contacts distant to those from category $f$ (on average 9.1 %, varying between 2.6 % and 14.9 %);
- **other** ($o$): all remaining electrode contacts (on average 77.9 %, varying between 66.2 % and 87.7 %).
This allowed us to concisely analyze interactions between brain regions and to perform inter-subject comparisons, despite the high variability in location and extent of the individual SOZ, which necessitated an electrode implantation tailored to the individual patient.

Since the order parameter $\hat{\gamma}$ is a symmetric and the directionality index $\hat{T}$ an antisymmetric measure, we will present our findings on the strength resp. direction of interactions from six combination categories (f-f, f-n, n-n, n-o, o-o; the direction of interaction is encoded in the sign of $\hat{T}$).

In Fig. 3 we show, as an example, temporal evolutions of the strength ($\hat{\gamma}$) and the direction ($\hat{T}$) of interactions between various brain regions over a time course of 28 h, during which a seizure was captured. The data were calculated from intracranial EEG recordings of the patient whose electrode implantation scheme is shown in Fig. 2. For only few pairs of recording sites can we observe temporarily increased values of $\hat{\gamma}$. These increases do not appear to coincide with seizure-related activities, but are nevertheless confined to interactions within the SOZ (f-f) and to neighbour–neighbour interactions (n-n). Recognisable indications for directed interactions can be mainly observed for combination categories involving the SOZ, and particularly for interactions between the SOZ and its neighbourhood as well as with other, remote brain regions. These observations indicate the SOZ to be a driving brain region (cf. Refs. [30, 31, 78, 79]) with interactions that are not only local but involve large parts of the epileptic network.

From the data shown in Fig. 3, it can be deduced that, for the most part, indications for directed interactions go along with low to intermediate strengths of interactions, while high values of the latter are accompanied by indications for symmetric bidirectional couplings. Our findings for all pairs of recording sites (see Fig. 4 left) and for all combination categories (see Fig. 4 right) support our expectations mentioned above (see Fig. 1) and underline the importance of investigating both strength and direction of interaction in order to avoid misinterpretations.

We now investigate whether the aforementioned observations of strongest interactions confined to the SOZ and/or its immediate surroundings extend beyond exemplary data. In Fig. 5, we show $\hat{T}$ and $\hat{\gamma}$ for all categories of combinations of recordings sites estimated from the data from all patients. On this group level, strengths of interactions appear to decrease with an increasing distance from the SOZ and its surroundings, which is in line with a number of previous studies [29, 34, 80–88]. If we search, however, for the maximum strength of interactions among all combinations of recording sites in each patient separately, we observe this maximum (and the next 10 lower values) for interactions far off the SOZ (category o-o) in 9 of 11 patients. In one patient, categories f-f and o-o rank almost equally among the ten highest values. In another patient, categories f-f and n-n clearly rank highest among all combinations of recording sites. These observations on a single patient level are in contrast to many of the previous studies mentioned before, and it remains to be shown whether differences can be related to the much lower number of recording sites taken into account before or to possibly different sensitivities of applied estimators for the strength resp. direction of interactions (see, e.g., [29, 41, 89]).

So far we only considered data from the seizure-free interval (inter-ictal state) for our analyses. We now proceed by addressing the question whether changes in directed interactions between brain regions constituting the epileptic network can be identified prior to seizures (pre-ictal state). Our previous investigations [90] indicate that directed interactions between the SOZ and homologous contralateral sites decrease (i.e. from a unidirectional driving to a symmetric bidirectional coupling) during the pre-ictal states in 11 out of 15 patients with mesial temporal lobe epilepsy. In this study, we also employed symbolic transfer entropy but restricted our analyses to intrahippocampal recordings. In the following, we will again assume that a pre-ictal phase of 4 h duration exists [21] and compare the distributions of values of $\hat{T}$ from the pre-ictal periods with those from inter-ictal periods (all data that were recorded at least 4 h prior to and 30 min after a seizure) and taking into account all interactions. In Fig. 6, we show exemplary matrices representing the spatial distribution of averaged strengths and directions of all pairwise interactions during pre-ictal and inter-ictal periods. For both periods, highest values of $\hat{\gamma}$ can be observed for interactions within the SOZ (category f-f; right hippocampal formation; sites...
Figure 3. Exemplary temporal evolutions of the order parameter $\hat{\gamma}$ (upper plot) and the directionality index $\hat{T}$ (lower plot) for interactions between various brain regions (from top to bottom): within the SOZ ($f$–$f$), between SOZ and its neighbourhood ($f$–$n$), between SOZ and other brain areas ($f$–$o$), neighbour–neighbour interactions ($n$–$n$), as well as neighbour–other ($n$–$o$) and other–other interactions ($o$–$o$). The dashed vertical line indicates onset of an epileptic seizure that lasted for approx. 100 s. For readability, time profiles are smoothed using a Hamming window (10 min duration). Time 00 indicates midnight.

TR01–TR10) and its surrounding brain areas (category $n$–$n$; sites TBAR1–TBAR3 and TBPR1–TBPR3) but also for interactions in the ipsilateral neocortex, particularly at parieto-occipital sites (category $o$–$o$; TLR06–TLR08 and TLR14–TLR16). Strongest interactions also involve the most anterior contact in the contralateral hippocampal formation (category $o$–$o$; particularly site TL01) and parieto-occipital sites in the contralateral neocortex (category $o$–$o$; sites TLL07–TLL08).
In line with [91], we observe—in both hippocampal formations—a gap in the inter-regional strength of interactions namely within the entorhinal cortex and within the hippocampus, that indicate the existence of independent rhythms (mostly in the frequency band 0.5–7 Hz) in different subregions of the human medial temporal lobe, produced by autonomous generators. Prominent differences between data from the pre-ictal and the inter-ictal periods are visible for the SOZ, its neighbourhood, homologous sites within the contralateral mesial temporal lobe, and for interactions between inferior and lateral sites from the contralateral temporal cortex. We observe

Figure 4. Left: Contour lines of equal frequency in a bi-dimensional histogram of more than $3.6 \cdot 10^7$ pairs of values of the order parameter $\hat{\gamma}$ and of the directionality index $\hat{T}$ ($\hat{T} \geq 0$) estimated from intracranial EEG data recorded from one patient (cf. Fig. 2) over nine days for all combinations of electrode contacts (seizure-free interval only, i.e. data from a presumed pre-ictal phase (4 h duration), the ictal, and the post-ictal phase (30 min duration) of four seizures were excluded). Right: Temporal and spatial means and standard deviations of $\hat{\gamma}$ and of the absolute value of $\hat{T}$ for all categories of combinations of recordings sites.

Figure 5. Means and standard deviations of patient-wise means of $\hat{\gamma}$ and of the absolute value of $\hat{T}$ for all categories of combinations of recordings sites for the 11 patients.
Figure 6. Averaged strengths (top) and directions (bottom) of all pair-wise interactions during inter-ictal (left) and pre-ictal (middle) periods estimated from an intracranial EEG recording (cf. Fig. 2) that lasted nine days and during which four seizures were captured. Positive (negative) values of $\hat{T}$ indicate that sites listed on the ordinate drive (are being driven by) sites listed on the abscissa. Differences between pre-ictal and inter-ictal data are shown in the right column.

both, a pre-ictal increase and decrease of $\hat{\gamma}$, in line with previous studies that used other, mostly phase-based estimators for the strength of interactions [20, 25, 92–97].

If we assume that the direction of interactions can best be resolved at intermediate strengths (see Fig. 1), we accordingly observe high absolute values of $\hat{T}$ particularly for interactions involving categories $f$-$n$, $f$-$o$, and $n$-$o$. During both the pre-ictal and the inter-ictal periods, the SOZ (right hippocampal formation; sites TR01–TR10) appears to persistently drive all other brain regions, while these regions appear to drive structures near the SOZ (sites TBAR1–TBAR3 and TBPR1–TBPR3). Nevertheless, we also observe directed interactions between brain regions that are apparently not associated with the SOZ (category $o$-$o$). As a result, no clear-cut directionality patterns can be observed from the differences between data from the pre-ictal and the inter-ictal periods, although strongest changes in directionality, i.e. more pronounced indications for a unidirectional driving coupling, are confined to brain regions surrounding the SOZ (category $n$-$n$; sites TBAR1–TBAR3 and TBPR1–TBPR3). In Fig. 7, we show the inter-ictal and pre-ictal frequency distributions of indices $\hat{\gamma}$ and $\hat{T}$ for all categories of combinations of recordings sites. Pronounced relative changes (> 20 %) can only be observed for directed interactions between brain regions surrounding the SOZ and all other sampled regions (category $n$-$o$; inter-ictal: $-0.09 \pm 0.18$; pre-ictal: $-0.15 \pm 0.16$) and for directed interactions between the SOZ and its neighbourhood (category $f$-$n$; inter-ictal: $0.26 \pm 0.15$; pre-ictal: $0.33 \pm 0.13$).
Figure 7. Frequency distributions of order parameter $\hat{\gamma}$ (top) and directionality index $\hat{T}$ (bottom) for all categories of combinations of recordings sites estimated from intracranial EEG data (cf. Fig. 2) recorded over nine days. Left: data from inter-ictal periods. Right: data from pre-ictal periods.

Table 1. Number of patients for which an increase, a decrease, or no change (else; changes in amplitude values lower than 5% or undefined relative changes) of $\hat{\gamma}$ and of $\hat{T}$ can be observed pre-ictally as compared to the inter-ictal periods.

| interactions between brain regions | order parameter $\hat{\gamma}$ | directionality index $\hat{T}$ |
|-----------------------------------|-------------------------------|-------------------------------|
|                                   | decrease | increase | else | decrease | increase | else |
| SOZ $\Leftarrow$ neighbour        | 3        | 6         | 2     | 2        | 5         | 4     |
| SOZ $\Leftarrow$ other            | 1        | 5         | 5     | 5        | 3         | 3     |
| neighbour $\Leftarrow$ other      | 0        | 4         | 7     | 3        | 5         | 3     |

In order to investigate whether the aforementioned findings extend beyond exemplary data, we estimated—for each combination category (but excluding those that involve the same location category)—the number of patients for which pre-ictally an increase or a decrease (of at least 5%), or no change of the order parameter and of the directionality index can be observed as compared to the inter-ictal periods (see Tab. 1). For about half the cases, the strength of
interactions between the SOZ, its surrounding and other brain regions (categories \( f-n \) and \( f-o \)) increased during the pre-ictal periods. The strength of interactions between brain regions not involving the SOZ (category \( n-o \)) remained unchanged pre-ictally in the majority of cases. The direction of interactions increased pre-ictally in about half the cases for interdependencies between the SOZ and its surrounding (category \( f-n \)) as well as between brain regions not involving the SOZ (category \( n-o \)). The direction of interactions decreased pre-ictally in about half the cases for interdependencies between the SOZ and other brain regions (category \( f-o \)).

In Tab. 2, we report the number of patients for which \( \hat{T} \) (using \( \hat{\gamma} \) as selection criterion) indicates a preferred direction of interactions (preferentially driving vs. preferentially responding) during pre-ictal and inter-ictal periods. For this purpose we calculated, for each investigated combination category, the temporal and spatial median of \( \hat{T} \) and of \( \hat{\gamma} \) (denoted as \( \langle \hat{T} \rangle \) and \( \langle \hat{\gamma} \rangle \), respectively), and regarded the interaction as preferential driving (responding) if \( \langle \hat{T} \rangle \) was positive (negative) and the accompanying values of \( \langle \hat{\gamma} \rangle \) were from the interquartile range of the respective distribution of values of \( \hat{\gamma} \). We could neither observe a clear-cut indication for a preferential driving nor for a preferential responding between the investigated brain regions. Pre-ictally, the preferential driving of surrounding brain region by the SOZ (category \( f-n \); as seen inter-ictally) appears to be lost, however, the driving of surrounding brain regions by other regions (category \( n-o \)) can be observed more often. Although speculative, these observation probably point to an effective inhibitory surround [98, 99] that prevents the spreading of epileptiform activities during the seizure-free interval.

4. Conclusion

Using approaches from symbolic analysis, we investigated—in a time-resolved manner—changes in strength and direction of short- to long-ranged interactions between brain regions constituting the epileptic network. We here employed the order parameter \( \hat{\gamma} \) and the directionality index \( \hat{T} \) (derived from symbolic transfer entropy), both allowing for a robust and computationally fast quantification of the mentioned interaction properties. We followed previous recommendations [31, 50] to investigate both aspects of interactions in order to avoid misinterpretations and to allow an effective differentiation between various coupling regimes.

Analysing multi-day, multi-channel EEG recordings from 11 epilepsy patients, we could confirm some previous findings concerning the strength of interactions that had been derived with frequency-based, phase-based, state-space-based, or statistical approaches. Summarising these observations, the strength of interactions decreases with an increasing distance from the seizure-onset zone (SOZ) and its surroundings during the seizure-free (inter-ictal) periods, while it exhibits a possibly location-specific increase or decrease during the pre-ictal periods. Our findings obtained from investigating interdependencies between all sampled brain regions, however, indicate that the maximum strength of interactions can be observed for interactions between brain regions far off the SOZ in the majority of cases. It thus remains to be shown whether the dynamics of SOZ can indeed be characterised by an elevated local synchrony or elevated strength of interactions [80–83, 86–88] and whether this property can help to identify the SOZ in the presurgical evaluation. It also remains to be shown whether the location-specific changes of the strength of interactions (increase or decrease) during the pre-ictal periods can be regarded as

| Interactions between brain regions | Driving \((A \rightarrow B)\) | Responding \((A \leftarrow B)\) |
|----------------------------------|----------------|----------------|
| SOZ \(\rightarrow\) neighbour     | 3              | 0              |
| SOZ \(\rightarrow\) other        | 2              | 3              |
| neighbour \(\rightarrow\) other    | 2              | 4              |
precursors of epileptic seizures. Such an investigation would require using statistical methods for testing the significance of the predictive performance of the applied analysis techniques [25, 100–106]. The same holds for the direction of interactions since the unambiguous inference of the direction of interactions remains an unsolved problem, despite some developments to test the significance of directionality estimates [107–110].

We here considered a spatial average over pre-defined location categories using knowledge concerning location and extent of the SOZ and observed in some patients a preferential driving between some brain regions while in other patients directionality was inverted for the very same brain regions. This ambiguity persisted when comparing data from the inter-ictal and the pre-ictal periods. Due to these inconsistencies we can not yet give a clear-cut answer to the question of who is driving whom in large-scale epileptic brain networks.

There are, by now, only a few studies that investigated directed interaction in the epileptic brain [15, 90, 111–117], and these studies mostly concentrated on seizures or other epileptiform activities (such as epileptic spikes) and/or were mostly restricted to selected recording sites. A comparison of our findings with those obtained in these studies is thus rather constricted.

The inference of directed interactions from empirical data may be limited by a number of influencing factors. Bivariate analysis, as used here, is often not sufficient to reveal the correct interaction structure, i.e. distinguishing direct and indirect interactions. An extension to the concept of symbolic transfer entropy that aims at accounting for the presence of other observed confounding variables has recently been proposed (so called partial symbolic transfer entropy [66, 67]), but its suitability for characterising directed interactions in epileptic brain networks remains to be investigated. It is well known that interactions between and within brain regions may be delayed, with time delays reaching up to 200 ms depending on region and functions [118], and not accounting for such delays may lead to spurious indications for directed interactions. An improved characterisation of such delayed, directed interactions may be achieved with recently proposed analysis techniques that explicitly incorporate time delays in estimates for information transfer [119–123].

In the future, we expect further improvements in the data-driven characterisation of weighted and directed, short- to long-ranged interactions in evolving epileptic brain networks to advance our understanding of the dynamical disease epilepsy, which may guide new developments for individualised diagnosis, treatment, and control.

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References

1 Duncan JS, Sander JW, Sisodiya SM, Walker MC, 2006 Adult epilepsy. Lancet 367, 1087–1100. doi:10.1016/S0140-6736(06)68477-8
2 Guerrini R, 2006 Epilepsy in children. Lancet 367, 499–524. doi:10.1016/S0140-6736(06)68182-8
3 Schuele S, Lüders HO, 2008 Intractable epilepsy: management and therapeutic alternatives. Lancet Neurol. 7, 514–524. doi:10.1016/S1474-4422(08)70108-X
4 Spencer S, Huh L, 2008 Outcomes of epilepsy surgery in adults and children. Lancet Neurol. 7, 525–537. doi:10.1016/S1474-4422(08)70109-1
5 Schramm J, 2008 Temporal lobe epilepsy surgery and the quest for optimal extent of resection: A review. Epilepsia 49, 1296–1307. doi:10.1111/j.1528-1167.2008.01604.x
6 Kwan P, Schachter SC, Brodie MJ, 2011 Drug-resistant epilepsy. N. Engl. J. Med. 365, 919–926
7 Perucca E, Tomson T, 2011 The pharmacological treatment of epilepsy in adults. Lancet Neurol. 10, 446–456. doi:10.1016/S1474-4422(11)70047-3
8 Fisher RS, van Emde Boas W, Blume W, Elger CE, Genton P, Lee P, Engel Jr J, 2005 Epileptic seizures and epilepsy: definitions proposed by the International League Against Epilepsy (ILAE) and the International Bureau for Epilepsy (IBE). *Epilepsia* **46**, 470–472. doi:10.1111/j.0013-9580.2005.66104.x

9 Engel Jr J, 2006 Report of the ILAE Classification Core Group. *Epilepsia* **47**, 1558–1568. doi:10.1111/j.1528-1167.2006.00215.x

10 Rosenow F, Lüders H, 2001 Presurgical evaluation of epilepsy. *Brain* **124**, 1683–1700. doi:10.1093/brain/124.9.1683

11 Berg AT, Scheffer IE, 2011 New concepts in classification of the epilepsies: Entering the 21st century. *Epilepsia* **52**, 1058–1062. doi:10.1111/j.1528-1167.2011.03101.x

12 Lehertez K, Ansmann G, Bialonski S, Dickten H, Geier C, Porz S, 2014 Evolving networks in the human epileptic brain. *Physica D* **267**, 7–15. doi:10.1016/j.physd.2013.06.009

13 Bragin A, Wilson CL, Engel Jr J, 2000 Chronic epileptogenesis requires development of a network of pathologically interconnected neuron clusters: a hypothesis. *Epilepsia* **41** (Suppl. 6), S144–S152. doi:10.1111/j.1528-1157.2000.tb01573.x

14 Barlow SS, Mammarella E, 2002 Neural networks in human epilepsy: Evidence of and implications for treatment. *Epilepsia* **43**, 219–227. doi:10.1046/j.1528-1157.2002.26901.x

15 D’Alessandro M, Vachtsevanos G, Esteller R, Echauz J, Cranstoun S, Worrell G, Parish L, Litt B, 2005 A multi-feature and multi-channel univariate selection process for seizure prediction. *Clin. Neurophysiol.* **116**, 506–516

16 Kalitzin S, Velis D, Suffczynski P, Parra J, Lopes da Silva F, 2005 Electrical brain-stimulation paradigm for estimating the seizure onset site and the time to ictal transition in temporal lobe epilepsy. *Clin. Neurophysiol.* **116**, 718–728. doi:10.1016/j.clinph.2004.08.021

17 Le Van Quyen M, Soss J, Navarro V, Robertson R, Chavez M, Baulac M, Martinerie J, 2005 Preictal state identification by synchronization changes in long-term intracranial EEG recordings. *Clin. Neurophysiol.* **116**, 559–568

18 Mormann F, Kreuz T, Rieke C, Andrzejak RG, Kraskov A, David P, Elger CE, 2005 On the predictability of epileptic seizures. *Clin. Neurophysiol.* **116**, 569–587. doi:10.1016/j.clinph.2004.08.025

19 Navarro V, Martinerie J, Le Van Quyen M, Baulac M, Dubois F, Gotman J, 2005 Seizure anticipation: do mathematical measures correlate with video-EEG evaluation? *Epilepsia* **46**, 385–396

20 Federico P, Abbott DF, Brielmann RS, Harvey AS, Jackson GD, 2005 Functional MRI of the pre-ictal state. *Brain* **128**, 1811–1817

21 Meier R, Häussler U, Aertsen A, Deransart C, Depaulis A, Egert U, 2007 Short-term changes in bilateral hippocampal coherence precede epileptiform events. *NeuroImage* **38**, 138–149

22 Feldwisch-Drentrup H, Ihle M, Le Van Quyen M, Teixeira C, Dourado A, Timmer J, Sales F, Navarro V, Schulze-Bonhage A, Schelter B, 2011 Anticipating the unobserved: Prediction of subclinical seizures. *Epilepsy Behav.* **22**, S119–S126. doi:10.1016/j.yebeh.2011.08.023

23 Aarabi A, He B, 2012 A rule-based seizure prediction method for focal neocortical epilepsy. *Clin. Neurophysiol.* **123**, 1111–1122. doi:10.1016/j.clinph.2012.01.014

24 Seyal M, 2014 Frontal hemodynamic changes precede EEG onset of temporal lobe seizures. *Clin. Neurophysiol.* **125**, 442–448. doi:10.1016/j.clinph.2013.09.003

25 Arndt J, Grassberger P, Lüders H, 1999 A robust method for detecting interdependences: application to intracranially recorded EEG. *Physica D* **134**, 419–430
29 Osterhage H, Mormann F, Staniek M, Lehnertz K, 2007 Measuring synchronization in the epileptic brain: A comparison of different approaches. *Int. J. Bifurcation Chaos Appl. Sci. Eng.* 17, 3539–3544.

30 Osterhage H, Mormann F, Wagner T, Lehnertz K, 2007 Measuring the directionality of coupling: phase versus state space dynamics and application to EEG time series. *Int. J. Neural. Syst.* 17, 139–148.

31 Osterhage H, Mormann F, Wagner T, Lehnertz K, 2008 Detecting directional coupling in the human epileptic brain: Limitations and potential pitfalls. *Phys. Rev. E* 77, 011914.

32 Prusseit J, Lehnertz K, 2008 Measuring interdependences in dissipative dynamical systems with estimated Fokker-Planck coefficients. *Phys. Rev. E* 77, 041914.

33 Wendling F, Bartolomei F, Senhadji L, 2009 Spatial analysis of intracerebral electroencephalographic signals in the time and frequency domain: identification of epileptogenic networks in partial epilepsy. *Phil. Trans. Roy. Soc. A* 367, 297–316. doi:10.1098/rsta.2008.0220.

34 Andrzejak RG, Chicharro D, Lehnertz K, Mormann F, 2011 Using bivariate signal analysis to characterize the epileptic focus: The benefit of surrogates. *Phys. Rev. E* 83, 046203. doi:10.1103/PhysRevE.83.046203.

35 Andrzejak RG, Kreuz T, 2011 Characterizing unidirectional couplings between point processes and flows. *EPL* 96, doi:10.1209/0295-5075/96/50012.

36 Pikovsky AS, Rosenblum MG, Kurths J, 2001 *Synchronization: A universal concept in nonlinear sciences*. Cambridge, UK: Cambridge University Press. doi:10.1017/CBO9780511755743.

37 Boccaletti S, Kurths J, Osipov G, Valladares DL, Zhou CS, 2002 The synchronization of chaotic systems. *Phys. Rep.* 366, 1–101. doi:10.1016/S0370-1573(02)00137-0.

38 Pereda E, Quian Quiroga R, Bhattacharya J, 2005 Nonlinear multivariate analysis of neurophysiological signals. *Prog. Neurobiol.* 77, 1–37. doi:10.1016/j.pneurobio.2005.10.003.

39 Gourevitch B, Le Bouquin-Jeannes R, Faucon G, 2006 Linear and nonlinear causality between signals: methods, examples and neurophysiological applications. *Biol. Cybern.* 95, 349–369. doi:10.1007/s00422-006-0098-0.

40 Marwan N, Romano MC, Thiel M, Kurths J, 2007 Recurrence plots for the analysis of complex systems. *Phys. Rep.* 438, 237–329. doi:10.1016/j.physrep.2006.11.001.

41 Lehnertz K, Bialonski S, Horstmann MT, Krug D, Rothkegel A, Staniek M, Wagner T, 2009 Synchronization phenomena in human epileptic brain networks. *J. Neurosci. Methods* 183, 42–48. doi:10.1016/j.jneumeth.2009.05.015.

42 Friedrich R, Peinke J, Sahimi M, Tabar MRR, 2011 Approaching complexity by stochastic methods: From biological systems to turbulence. *Phys. Rep.* 506, 87–162. doi:10.1016/j.physrep.2011.05.003.

43 Lehnertz K, 2011 Assessing directed interactions from neurophysiological signals—an overview. *Physiol. Meas.* 32, 1751–1724. doi:10.1088/0967-3334/32/11/R01.

44 Hlaváčková-Schindler K, Paluš M, Vejmelka M, Bhattacharya J, 2007 Causality detection based on information-theoretic approaches in time series analysis. *Phys. Rep.* 441, 1–46. doi:10.1016/j.physrep.2006.12.004.

45 Hao BL, 1989 *Elementary Symbolic Dynamics and Chaos in Dissipative Systems*. Singapore: World Scientific.

46 Bandt C, Pompe B, 2002 Permutation entropy: A natural complexity measure for time series. *Phys. Rev. Lett.* 88, 174102.

47 Daw C, Finney C, Tracy E, 2003 A review of symbolic analysis of experimental data. *Rev. Sci. Instrum.* 74, 915–930.

48 Liu Z, 2004 Measuring the degree of synchronization from time series data. *Europhys. Lett.* 68, 19–25.

49 Staniek M, Lehnertz K, 2008 Symbolic transfer entropy. *Phys. Rev. Lett.* 100, 158101.

50 Staniek M, Lehnertz K, 2009 Symbolic transfer entropy: inferring directionality in biosignals. *Biomed. Tech.* 54, 323–328.

51 Rulkov NF, Sushchik MM, Tsimring LS, Abarbanel HDI, 1995 Generalized synchronization of chaos in directionally coupled chaotic systems. *Phys. Rev. E* 51, 980–994.
52 Schreiber T, 2000 Measuring information transfer. Phys. Rev. Lett. 85, 461–464
53 Kaiser A, Schreiber T, 2002 Information transfer in continuous processes. Physica D 166, 43–62
54 Verdes PF, 2005 Assessing causality from multivariate time series. Phys. Rev. E 72, 026222
55 Lungarella M, Pitti A, Kuniyoshi Y, 2007 Information transfer at multiple scales. Phys. Rev. E 76, 056117. doi:10.1103/PhysRevE.76.056117
56 Kowalski AM, Martin MT, Plastino A, Zunino L, 2010 Information flow during the quantum-classical transition. Phys. Lett. A 374, 1819–1826. doi:10.1016/j.physleta.2010.02.037
57 Melzer A, Schella A, 2014 Symbolic transfer entropy analysis of the dust interaction in the presence of wakefields in dusty plasmas. Phys. Rev. E 89, 041103. doi:10.1103/PhysRevE.89.041103
58 Nian-Qiang L, Wei P, Lian-Shan Y, Bin L, Ming-Feng X, Yi-Long T, 2012 Quantifying information flow between two chaotic semiconductor lasers using symbolic transfer entropy. Chin. Phys. Lett. 29, 030502. doi:10.1088/0256-307X/29/3/030502
59 Blain-Moraes S, Mashour GA, Lee H, Huggins JE, Lee U, 2013 Altered cortical communication in amyotrophic lateral sclerosis. Neurosci. Lett. 543, 172–176. doi:10.1016/j.neulet.2013.03.028
60 Jun W, Zheng-Feng Y, 2012 Symbolic transfer entropy-based premature signal analysis. Chin. Phys. B 21, 018702. doi:10.1088/1674-1056/21/1/018702
61 Ku SW, Lee U, Noh GJ, Jun IG, Mashour GA, 2011 Preferential inhibition of fronto-to-parietal feedback connectivity is a neurophysiologic correlate of general anesthesia in surgical patients. PLoS One 6, e25155. doi:10.1371/journal.pone.0025155
62 Jordan D, et al., 2013 Simultaneous electroencephalographic and functional magnetic resonance imaging indicate impaired cortical top-down processing in association with anesthetic-induced unconsciousness. Anesthesiology 119, 1031–1042. doi:10.1097/ALN.0b013e31829103f5
63 Martin M, Kranz TA, Wagner T, Lehnerzt K, 2011 Inferring directional interactions from transient signals with symbolic transfer entropy. Phys. Rev. E 83, 011919. doi:10.1103/PhysRevE.83.011919
64 Kugiumtzis D, 2013 Partial transfer entropy on rank vectors. Eur. Phys. J.-Spec. Top. 222, 401–420. doi:10.1140/epjst/e2013-01849-4
65 Papana A, Kyrtou C, Kugiumtzis D, Diks C, 2013 Simulation study of direct causality measures in multivariate time series. Entropy 15, 2635–2661. doi:10.3390/e15072635
66 Hahs DW, Pethel SD, 2011 Distinguishing anticipation from causality: Anticipatory bias in the estimation of information flow. Phys. Rev. Lett. 107, 128701. doi:10.1103/PhysRevLett.107.128701
67 Barnett L, Bossemaier T, 2012 Transfer entropy as a log-likelihood ratio. Phys. Rev. Lett. 109, 138103. doi:10.1103/PhysRevLett.109.138105
68 Smirnov DA, 2013 Spurious causalties with transfer entropy. Phys. Rev. E 87, 042917. doi:10.1103/PhysRevE.87.042917
69 Haruna T, Nakajima K, 2013 Permutation complexity and coupling measures in hidden markov models. Entropy 15, 3910–3930. doi:10.3390/e15093910
70 Paluš M, Vejmelka M, 2007 Directionality of coupling from bivariate time series: How to avoid false causalties and missed connections. Phys. Rev. E 75, 056211
71 Waddell J, Dzakpasu R, Booth V, Riley B, Reasor J, Poe G, Zachowski M, 2007 Causal entropies–A measure for determining changes in the temporal organization of neural systems. J. Neurosci. Methods 162, 320–332
72 Engel Jr J, van Ness PC, Rasmussen TB, Ojemann LM, 1993 Outcome with respect to epileptic seizures. In J Engel Jr, ed., Surgical Treatment of the Epilepsies, 609. New York: Raven Press
75 Blanco S, Garcia H, Quian Quiroga R, Romanelli L, Rosso OA, 1995 Stationarity of the EEG series. IEEE Eng. Med. Biol. 4, 395–399. doi:10.1109/51.395321
76 Cao Y, Tung W, Gao JB, Protopopescu VA, Hively LM, 2004 Detecting dynamical changes in time series using the permutation entropy. Phys. Rev. E 70, 046217
77 Staniek M, Lehnertz K, 2007 Parameter selection in permutation entropy measurements. Int. J. Bifurcation Chaos Appl. Sci. Eng. 17, 3729
78 Paluš M, Komárek V, Hrncíř Z, Štěrbová K, 2001 Synchronization and information flow in EEGs of epileptic patients. Med. Biol. Mag. 20, 65–71
79 Osterhage H, Bialonski S, Staniek M, Schindler K, Wagner T, Elger CE, Lehnertz K, 2008 Bivariate and multivariate time series analysis techniques and their potential impact for seizure prediction. 189–208. Berlin: Wiley-VCH
80 Ben-Jacob E, Boccaletti S, Pomyalov A, Procaccia I, Towle VL, 2007 Detecting and localizing the foci in human epileptic seizures. Chaos 17, 043113
81 Lai YC, Frei MG, Osorio I, Huang L, 2007 Characterization of synchrony with applications to epileptic brain signals. Phys. Rev. Lett. 98, 108102
82 Schevon CA, et al., 2007 Cortical abnormalities in epilepsy revealed by local EEG synchrony. NeuroImage 35, 140–148
83 Ortega GJ, Menendez de la Prida L, Sola RG, Pastor J, 2008 Synchronization clusters of interictal activity in the lateral temporal cortex of epileptic patients: Intraoperative electrocorticographic analysis. Epilepsia 49, 269–280
84 Zaveri HP, Pincus SM, Goncharova II, Duckrow RB, Spencer DD, Spencer SS, 2009 Localization-related epilepsy exhibits significant connectivity away from the seizure-onset area. NeuroReport 20, 891–895
85 Warren C, Hu S, Stead M, Brinkmann BH, Bower MR, Worrell GA, 2010 Synchrony in normal and focal epileptic brain: The seizure onset zone is functionally disconnected. J. Neurophysiol. 104, 3530–3539
86 Bettus G, et al., 2011 Interictal functional connectivity of human epileptic networks assessed by intracerebral EEG and BOLD signal fluctuations. PLoS ONE 6, e20071. doi:10.1371/journal.pone.0020071
87 Ortega GJ, Peco IH, Sola RG, Pastor J, 2011 Impaired mesial synchronization in temporal lobe epilepsy. Clin. Neurophysiol. 122, 1106–1116. doi:10.1016/j.clinph.2010.11.001
88 Palmigiano A, Pastor J, Garcia de Sola R, Ortega GJ, 2012 Stability of synchronization clusters and seizurability in temporal lobe epilepsy. PLoS ONE 7, e41799. doi:10.1371/journal.pone.0041799
89 Ansari-Asl K, Senhadji L, Bellanger JJ, Wendling F, 2006 Quantitative evaluation of linear and nonlinear methods characterizing interdependencies between brain signals. Phys. Rev. E 74, 031916
90 Lhernzt K, Krug D, Staniek M, Glüsenkamp D, Elger CE, 2011 Preictal directed interactions in epileptic brain networks. 265–272. Boca Raton, FL, USA: CRC Press
91 Mormann F, Osterhage H, Andrzejak RG, Weber B, Fernandez G, Fell J, Elger CE, Lehnertz K, 2008 Independent delta/theta rhythms in the human hippocampus and entorhinal cortex. Front. Hum. Neurosci. 2, 3. doi:10.3389/neuro.09.003.2008
92 Mormann F, Lehnertz K, David P, Elger CE, 2000 Mean phase coherence as a measure for phase synchronization and its application to the EEG of epilepsy patients. Physica D 144, 358–369. doi:10.1016/S0167-2789(00)00087-7
93 Mormann F, Andrzejak RG, Kreuz T, Rieke C, David P, Elger CE, Lehnertz K, 2003 Automated detection of a preseizure state based on a decrease in synchronization in intracranial electroencephalogram recordings from epilepsy patients. Phys. Rev. E 67, 021912. doi:10.1103/PhysRevE.67.021912
94 Winterhalder M, Schelter B, Maiwald T, Brandt A, Schad A, Schulze-Bonhage A, Timmer J, 2006 Spatio-temporal patient-individual assessment of synchronization changes for epileptic seizure prediction. Clin. Neurophysiol. 117, 2399–2413
95 Kuhlmann L, Freestone D, Lai AL, Burkitt AN, Fuller K, Grayden D, Seiderer L, Vogrin S, Mareels IMY, Cook MJ, 2010 Patient-specific bivariate-synchrony-based seizure prediction for short prediction horizons. Epilepsy Res. 91, 214–231. doi:10.1016/j.epilepsyres.2010.07.014
Wang L, Wang C, Fu F, Yu X, Guo H, C CX, Jing X, Zhang H, Dong X, 2011 Temporal lobe seizure prediction based on a complex Gaussian wavelet. *Clin. Neurophysiol.* 122, 656–663. doi:10.1016/j.clinph.2010.09.018

Zheng Y, Wang G, Li K, Bao G, Wang J, 2014 Epileptic seizure prediction using phase synchronization based on bivariate empirical mode decomposition. *Clin. Neurophysiol.* 125, 1104–1111. doi:10.1016/j.clinph.2013.09.047

Prince DA, Wilder J, 1967 Control mechanisms in cortical epileptogenic foci. "surround" inhibition. *Arch. Neurol.* 16, 194–202

Elger CE, Speckmann EJ, 1983 Penicillin induced epileptic foci in the motor cortex: vertical inhibition. *Electroencephalogr. Clin. Neurophysiol.* 56, 604–622

Andrzejak RG, Mormann F, Kreuz T, Rieke C, Kraskov A, Elger CE, Lehnhertz K, 2003 Testing the null hypothesis of the nonexistence of a preseizure state. *Phys. Rev. E* 67, 010901(R)

Andrzejak RG, Chicharro D, Elger CE, Mormann F, 2009 Seizure prediction: Any better than chance? *Clin. Neurophysiol.* 120, 1465–1478. doi:10.1016/j.clinph.2009.05.019

Winterhalder M, Maiwald T, Voss HU, Aschenbrenner-Scheibe R, Timmer J, Schulze-Bonhage A, 2003 The seizure prediction characteristic: A general framework to assess and compare seizure prediction methods. *Epilepsy Behav.* 3, 318–325

Kreuz T, Andrzejak RG, Mormann F, Stögbauer H, Elger CE, Lehnhertz K, Grassberger P, 2004 Measure profile surrogates: A method to validate the performance of epileptic seizure prediction algorithms. *Phys. Rev. E* 69, 061915

Schelter B, Winterhalder M, Maiwald T, Brandt A, Schad A, Schulze-Bonhage A, Timmer J, 2006 Testing statistical significance of multivariate time series analysis techniques for epileptic seizure prediction. *Chaos* 16, 013108

Wong S, Gardner AB, Krieger AM, Litt B, 2007 A stochastic framework for evaluating seizure prediction algorithms using hidden Markov models. *J. Neurophysiol.* 97, 2525–2532

Snyder DE, Échauz J, Grimes DB, Litt B, 2008 The statistics of a practical seizure warning system. *J. Neural. Eng.* 5, 392–401

Porta A, Furlan R, Rimoldi O, Pagani M, Malliani A, van de Borne P, 2002 Quantifying the strength of the linear causal coupling in closed loop interacting cardiovascular variability signals. *Biol. Cybern.* 86, 241–251. doi:10.1007/s00422-001-0292-z

Thiel M, Romano MC, Kurths J, Rolfs M, Kliegl R, 2006 Twin surrogates to test for complex synchronisation. *Europhys. Lett.* 75, 535–541

Romano MC, Thiel M, Kurths J, Mergenthaler K, Engbert R, 2009 Hypothesis test for synchronization: Twin surrogates revisited. *Chaos* 19, 015108. doi:10.1063/1.3072784

Faes L, Porta A, Nollo G, 2010 Testing frequency-domain causality in multivariate time series. *IEEE Trans. Biomed. Eng.* 57, 1897–1906. doi:10.1109/TBME.2010.2042715

Chavez M, Martinerie J, Le Van Quyen M, 2003 Statistical assessment of nonlinear causality: application to epileptic EEG signals. *J. Neurosci. Methods* 124, 113–128

Smirnov DA, Bodrov MB, Velazquez JLP, Wennberg RA, Bezruchko BP, 2005 Estimation of coupling between oscillators from short time series via phase dynamics modeling: Limitations and application to EEG data. *Chaos* 15, 024102

Wenke C, van Drongelen W, Kohnman M, He B, 2009 Identification of epileptogenic foci from causal analysis of ECoG interictal spike activity. *Clin. Neurophysiol.* 120, 1449–1456. doi:10.1016/j.clinph.2009.04.024
118 Nunez PL, 1995 *Neocortical Dynamics and Human EEG Rhythms*. Oxford, UK: Oxford University Press

119 Frenzel S, Pompe B, 2007 Partial mutual information for coupling analysis of multivariate time series. *Phys. Rev. Lett.* **99**, 204101. doi:10.1103/PhysRevLett.99.204101

120 Pompe B, Runge J, 2011 Momentary information transfer as a coupling measure of time series. *Phys. Rev. E* **83**, 051122. doi:10.1103/PhysRevE.83.051122

121 Runge J, Heitzig J, Petoukhov V, Kurths J, 2012 Escaping the curse of dimensionality in estimating multivariate transfer entropy. *Phys. Rev. Lett.* **108**, 258701. doi:10.1103/PhysRevLett.108.258701

122 Runge J, Heitzig J, Marwan N, Kurths J, 2012 Quantifying causal coupling strength: A lag-specific measure for multivariate time series related to transfer entropy. *Phys. Rev. E* **86**, 061121. doi:10.1103/PhysRevE.86.061121

123 Dickten H, Lehnertz K, 2014 Identifying delayed directional couplings with symbolic transfer entropy. *Phys. Rev. E* **90**, 062706. doi:10.1103/PhysRevE.90.062706