A fundamental question in science is how complex dynamics can be characterized by measuring the interactions within a distributed system. To address this question, many approaches have been developed to measure different types of pairwise interactions from dynamical data. For example, in neuroimaging, functional connections between pairs of brain regions are quantified through statistical correlations, which mark changes in human behaviors and differ in neurological diseases. In Earth system science, pairwise causal models have been used to infer mechanistic drivers of natural processes, from the influence of sea-surface temperature on sardine and anchovy populations, to the atmospheric drivers of air circulation. Furthermore, economic analysts have studied the co-integration of paired non-stationary time series—including stock-market indices and their associated future contracts—to infer a statistically significant coupling for building econometric models.

As illustrated schematically in Fig. 1a, the common goal of these studies is to extract meaningful pairwise relationships from multivariate time series (MTS): sets of observations taken regularly over time. In the age of big data, the scientific problems that are studied in diverse disciplinary contexts—from genomics to astronomy—require novel ways to quantify a statistical dependency between two time series, it remains common practice to manually select a single method with minimal comparisons against alternatives. For instance, Pearson correlation remains the most commonly used tool for measuring pairwise relationships in neuroimaging and Earth system science, despite rather restrictive (and often unsatisfied) assumptions that the data are serially independent and normally distributed. Fortunately, many more sophisticated and powerful algorithms have been developed, including those where dependencies are lagged in time (for example,
these methods remain disconnected from one another. In this work, we unify this wealth of interdisciplinary scientific knowledge, empirically connecting previously disjoint methodological traditions to yield a unified set of tools for quantifying interactions in complex dynamics.

As diverse scientific methods for quantifying pairwise interactions have never been compared at scale, there remain many unanswered questions. Are all of these methods capturing unique information on the interactions occurring within a time-varying system? Is there synergy between complementary approaches that, when combined, tells us more about the underlying system than any single method can? Or, do interesting redundancies exist between techniques used across disciplines that hint at a common theoretical underpinning?

Cross-correlation\(^6\), may be misaligned (for instance, dynamic time warping\(^10\)), or where the knowledge of one variable improves the predictability of another (for example, Granger causality\(^11\)).

In this Article, we represent algorithms that measure interactions between pairs of time series as real-valued summary statistics, that is, statistics of pairwise interactions (SPIs). Figure 1b illustrates the diverse theoretical tools and types of interactions covered in the scientific literature on SPIs, from covariance (the foundation of statistics and machine learning) to convergent cross-mapping\(^3\) (developed to infer causal effect in complex ecosystems); however, as the theory underlying statistical interactions between pairs of time series has been developed largely independently across disciplinary contexts,
that has not previously been identified? Following previous highly comparative studies of univariate time series and graphs, this work addresses these questions by simultaneously evaluating hundreds of different SPIs directly from data. Our empirical approach to this problem involves first assembling a comprehensive annotated library of computational methods for quantifying pairwise interactions from data (summarized in Fig. 1b), and then analyzing their behavior across a large and diverse library of MTS (Fig. 1c).

As shown in Fig. 1b, our annotated library of SPIs was organized into six broad categories on the basis of their underlying theory: ‘basic’, ‘distance similarity’, ‘causal indices’, ‘information theory’, ‘spectral’ and ‘miscellaneous’ (see ‘Library of 237 SPIs’ section in the Methods for more details, and Supplementary Note 1 for a full list of SPIs with descriptions and references). To understand the behavior of these SPIs on data, we constructed a library of 1,053 diverse MTS, curated with the aim of capturing the main classes of systems and dynamics that are studied across scientific disciplines, including synchronization, spatiotemporal chaos, wave propagation, criticality and phase transitions. As depicted in Fig. 1c, our MTS library contains 505 synthetic MTS generated from a range of mathematical models, including coupled maps, ordinary differential equations and partial differential equations, and 548 diverse real-world MTS assembled from public databases such as geophysical, medical and financial data (see ‘Library of 1,053 MTS’ section in the Methods for further details, and Supplementary Note 2 for a full list of included MTS).

Results
Organizing pairwise interactions by their empirical behavior
Having assembled diverse libraries of methods (237 SPIs) and data (1,053 MTS), we aimed to analyze how similarly the different SPIs behave on the data. To achieve this, we developed an empirical similarity index, \( R(0 \leq R \leq 1) \), that captures the relationship between any two SPIs by comparing their output when applied to all 195,112 pairwise interactions present across all 1,053 MTS. This index is derived from the average absolute Spearman correlation between a pair of SPIs when applied to all pairs of processes in all datasets (see ‘Quantifying similarity between SPIs using the empirical similarity index’ section of the Methods for details). The minimum value, \( R = 0 \), indicates a pair of maximally distinct SPIs (with uncorrelated behavior on all datasets), whereas the maximum, \( R = 1 \), indicates a pair of SPIs that are perfectly correlated on all datasets (that is, they behave as simple monotonous transformations of one another); thus, a pair of SPIs with a high \( R \) reflects broadly similar behavior across MTS containing very different types of structures, and therefore acts as a suitable candidate index of empirical similarity.

Using the dissimilarity measure, \( D = 1 - R \), we organized all 237 SPIs using hierarchical clustering, yielding the dendrogram shown in Fig. 2b. This presents a data-driven, structured representation of a diverse literature that allows us to probe and interpret relationships between scientific methods at multiple levels. We focus our analysis here on a 14-module resolution (with modules labeled M1–M14), which captures important methodological connections between groups of SPIs with similar behavior on data. As summarized in Fig. 2a, these fourteen modules group common conceptual and theoretical approaches to measuring interactions between pairs of time series, demonstrating the ability of our empirical approach to meaningfully organize the interdisciplinary literature.

In addition to grouping similar types of methods into modules, we found that different high-level conceptual formulations of dynamical interactions were recapitulated in the relationships between modules. For example, modules M3–M6 contain distinct types of SPIs (including Granger causality, directed information and integrated information), all of which capture statistical dependencies between two time series by considering the context of their past. This idea that observable interactions are predicated (or, from a statistical standpoint, conditioned) on the history of a process was first proposed by the Wiener–Granger theories of causality and feedback, specifically by measuring how one time series might improve the self-predictability of another. Our results group SPIs on the basis of this underlying theoretical formulation, due to their characteristic behavior on data. Other types of SPIs (that do not predicate on the self-predictability of a process) also display distinctive behavior, including measures of contemporaneous relationships (the correlation coefficients of M14), dependencies that account for temporal lags (the coherence measures of M12), or temporal dilation and shifts (dynamic time warping and related methods in M10).

Ten of the fourteen modules are homogeneous, containing methods that are derived from similar underlying theories (as indicated by the color of the category labels in Fig. 2bii). Of these ten homogeneous modules, six of them (M4, M7, M8, M9, M11 and M13) comprise SPIs for measuring one specific type of pairwise interaction, differing in either: their specific algorithmic implementation (for example, M13 contains both the Engle–Granger and the Johansen tests for measuring co-integration); extracted summary statistics (for example, M8 contains both the mean and maximum of the wavelet-based phase lag index); or parameter settings (for example, M4 contains SPIs that use five different estimation techniques for transfer entropy). The remaining four homogeneous modules (M1, M2, M6 and M12) comprise methods with very similar theoretical underpinnings; for example, M12 contains many SPIs for measuring undirected interactions via Fourier transformations, such as the magnitude and the imaginary part of the coherence. Of particular interest are the four heterogeneous modules (M3, M5, M10 and M14), which mix SPIs from different literature categories, revealing interesting connections between different theoretical bases for quantifying pairwise dependencies between time series. Although M3 contains a mix of SPIs based on information theory (six labeled ‘information theory’ measures and one, information-geometric conditional independence, labeled as a ‘causal index’), the remaining three modules establish interesting connections between the behavior of seemingly disparate SPIs on MTS data. Three networks that are derived from these modules are plotted in Fig. 2c–e and are investigated in detail below.

As illustrated in Fig. 2c, module M5 contains a mix of two different types of methods: the first includes five linear estimators for integrated information (such as geometric integrated information, \( \Phi_r \) (ref. 22), phi star, \( \Phi^* \) (ref. 16) and stochastic interaction), whereas the second includes 16 estimators for Granger causality in both the time and frequency domains. Although Granger causality and integrated information theory were developed in very different contexts (for example, Granger’s investigations into causality between economic time series in 1969, versus Tononi’s recent integrated information theory (\( \Phi \) of consciousness)), our analysis reveals that all SPIs in this module nevertheless behave similarly on data (with an empirical similarity index, \( R = 0.52 \), which lies in the 95th percentile for all \( R \) values; see Supplementary Fig. 1c). Recent results have indeed shown that Granger causality can be formulated as the information-theoretic measure, transfer entropy, and can thus be grouped under the same information-geometric framework as integrated information theory. However, it was not known whether these information-theoretic SPIs behave similarly in practice and, as such, their relationship was not widely recognized. Module M5 thus demonstrates an important confirmation of our empirical approach in being able to recapitulate emerging theory and unify scientific tools for understanding interacting processes.

Module M10, shown in Fig. 2d, highlights striking connections between three conceptually distinct types of methods: (1) dynamic time warping (DTW), which was developed in the data-mining community to quantify the similarity between two (potentially shifted and dilated) audio signals; (2) cross-spectral phase-based measures—the maximum phase coherence, and the mean and maximum phase-locking value—which were developed to examine frequency-specific
synchronization in neuroimaging data; and (3) the maximum cross-correlation, a classical statistical technique for correlating two time series at different temporal lags. All of these SPIs capture time-lagged interactions between two processes, but in slightly different ways: the maximum cross-correlation finds the highest fixed-lag match, DTW extends this idea by optimizing the distortion of the time axis to best match potentially misaligned time series, and the cross-spectral measures account for time lags in terms of phase differences. Module M10 thus reveals previously unreported connections between diverse approaches to capturing associations between pairs of potentially unaugmented time series, indicating a common conceptual basis for methods developed and applied across disciplines—whether they are measuring synchronization between neuroelectric recordings or recognizing speech from audio signals.

Finally, we discuss module M14, which groups 66 SPIs from all literature categories except for ‘spectral’ (see Fig. 1b for categories). This module recapitulates some theoretical relationships that have already been established, such as the equivalence between linear-Gaussian mutual information and absolute correlation (with a maximum similarity of $R = 1$). To highlight some previously unreported relationships, we focus on a demonstrative submodule (shown in Fig. 2e) comprising 17 SPIs from the ‘causal indices’, ‘distance similarity’ and ‘miscellaneous’ literature categories. We first note the tight cluster of SPIs, labeled ‘i’ in Fig. 2e, that were developed independently in two different domains: distance correlation-based methods from the statistics community, and kernel-based methods from the machine-learning community. This cluster first highlights a recent finding that distance correlation and the Hilbert–Schmidt independence criteria...
(HSIC, a kernel-based method)\textsuperscript{30} are equivalent when computed using certain distance kernels\textsuperscript{1}; our results suggest that similar theoretical connections can be established between the other SPIs of the cluster (including the Heller–Heller–Gorfine test \textsuperscript{32} and multiscale graph correlation\textsuperscript{33}). Second, we find that the distance- and kernel-based statistics display strikingly similar behavior as common implementations of the convergent cross-mapping (CCM) algorithm, which was originally developed for inferring causality in complex ecosystems\textsuperscript{3}. Convergent cross-mapping aims to measure the causal effect of one time series on another by the ability of the second to reconstruct the first using a nearest-neighbor approach. The fact that these methods behave so similarly on MTS data indicates that the well-studied techniques of phase-space reconstruction (used in the CCM algorithm) have a correspondence to the nonlinear kernel-estimation techniques from the statistics and machine learning communities. These observed connections also have important practical ramifications; for example, our results suggest candidate proxy algorithms to substitute for the computationally expensive CCM, which could yield major computational efficiencies and enable new applications on larger datasets.

In summary, our empirical approach to unifying a large and disjoint literature of methods for characterizing pairwise interactions in time series has allowed us to: (1) capture the commonalities in
Leveraging diverse methods to address scientific problems

Our results above illustrate the rich diversity of scientific methods for quantifying pairwise interactions. This diversity suggests that, when quantifying pairwise interactions for a given application, there is potential to make comparisons across the scientific literature of SPIs to: (1) select the best-performing SPI in an unbiased, data-driven way; and (2) leverage a synergistic combination of multiple complementary SPIs to better capture complex underlying interactions in MTS. Here we provide a simple demonstration of this strategy (referred to as ‘highly comparative’ due to the broad methodological comparison involved) on three MTS classification problems using three open datasets: (1) the smartwatch activity dataset (Fig. 3a), where the aim is to classify one of four behavioral states (walking, running, resting, or playing badminton) from six-sensor smartwatch MTS (comprising a three-axis accelerometer and three-axis gyroscope)\(^1\); (2) the electroencephalogram (EEG) state dataset (Fig. 3d), where the aim is to distinguish positive versus negative slow cortical potentials from single-subject EEG data (originally used to move a cursor up or down on a computer screen)\(^2\); and (3) the functional magnetic resonance imaging (fMRI) film dataset (Fig. 3g), where the aim is to classify resting and film-watching conditions from fMRI data\(^3\) (see ‘Classification case studies’ section in the Methods for additional details on each dataset). To investigate the performance of different SPIs on these tasks, we represented each MTS as a set of features corresponding to all pairwise interactions between its constituent processes and compared their classification performance using a linear support vector machine (SVM) with cross-validation (see ‘Classification section’ in the Methods for details).

We observed a wide range of SPI performances in each case study, ranging from null performance up to high and statistically significant performance: 27–92% accuracy on the smartwatch activity dataset (Fig. 3b), 48–69% accuracy on the EEG state dataset (Fig. 3e) and 41–95% accuracy on the fMRI film dataset (Fig. 3h). Many SPIs displayed significant classification performance on each dataset (permutation test, Bonferroni-corrected \(p<0.05\)): 213 SPIs for the smartwatch activity dataset, 165 SPIs for the EEG state dataset, and 67 SPIs for the fMRI film dataset. This wide range of observed SPI performance on all three datasets demonstrates the crucial importance of selecting an SPI that is able to capture the relevant types of interactions underlying a given dataset.

To understand the types of interactions that characterize the labeled classes of MTS, we analyzed and interpreted the highest-performing SPIs on each dataset (see Supplementary Data 2 for the full results). To provide a simple demonstration of this process, here we focus on the top-performing individual SPI in each case study. In the smartwatch activity dataset, the top-performing SPI was causally conditioned entropy (CCE) using a Kozachenko–Leonenko estimator (92% accuracy; SPI label ‘cco_kozachenko’; see Supplementary Note 1.4.7 for details). Its high performance is driven in part by its ability to capture the coupling from the wrist’s z-rotation to its x-translocation, which differs strongly between running and walking, as shown in Fig. 3c. This tells us that wrist rotation in the z-direction is more informative of subsequent elbow movement in the x-direction (as the elbow moves side to side) in walking than in running.

The top-performing SPI for the EEG state dataset was, as above, a causally conditioned entropy, but using a Gaussian density estimator (69%, SPI label ‘ccgaussian’; see Supplementary Note 1.4.7 for details). Its performance was driven in part by its ability to capture the increased coupling from EEG channel 2 to 1 (from near the right ear to near the left ear) in cortical negativity versus positivity states, as shown in Fig. 3f. Other statistics designed to capture directed information flow (in a way that includes instantaneous interactions in the presence of feedback) also performed well on this task, including directed information with a Gaussian density estimator (69% accuracy; SPI label ‘ddtf_multi-taper_mean_fs=1_fmin=0–25_fmax=0–5’; see Supplementary Note 1.5.9) and the Hilbert–Schmidt Independence Criterion (66%, SPI label ‘hsic’; see Supplementary Note 1.2.5). Of the classical methods for quantifying EEG connectivity, some are recapitulated as high performers by our data-driven analysis, including mean directed coherence across various frequency bands (all 67% accuracy; see Supplementary Note 1.5.10), whereas others exhibited surprisingly low accuracy, such as algorithmic variants of partial directed coherence (between 55% and 64%).

Finally, for the fMRI film dataset, the top-performing SPI was regression error-based causal inference (95% accuracy; SPI label ‘reci’; see Supplementary Note 1.3.4). Its high performance is driven in part by its ability to capture the stronger coupling from the control network to the ventral attention network during film-watching compared to rest (Fig. 3i). The dominant way of measuring coupling in whole-brain fMRI is to use the Pearson correlation coefficient (annotated in Fig. 3h), which exhibits strong and statistically significant classification accuracy on this problem (86%); however, our data-driven approach highlights 30 alternative SPIs with higher performance (88% to 95%; compare with Supplementary Data 2). These high-performing methods include alternative types of covariance (for example, minimum covariance determinant, 93% accuracy, SPI label ‘cov_MinfCovDet’; see Supplementary Note 1.1.1) and precision estimates (for example, using Ledoit–Wolf shrinkage, 91%, SPI label ‘prec_LedoitWolf’; see Supplementary Note 1.1.2) that better deal with non-Gaussian bivariate distributions. Other high performers include information-theoretic SPIs (such as conditional entropy, joint entropy and mutual information using Gaussian density estimators, all of which exhibited 91% accuracy; see Supplementary Note 1.4.1–1.4.3) and directed SPIs which distinguish asymmetric coupling (for example, the top-performing regression error-based causal inference, ‘reci’). Compared with the typically subjective process of selecting an appropriate method to analyze a given dataset, the highly comparative approach demonstrated here highlights the most useful scientific methods automatically, facilitating interpretable understanding of the conceptual formulations of pairwise dependencies that drive successful performance.

As different types of systems involve different types of interactions between measured processes, we expected different SPIs to perform well across the three datasets. Indeed, we found that an SPI with high performance on one problem does not imply its high performance on other problems. In particular, some SPIs performed well on only a single dataset; for example, dynamic time warping with an Itakura parallelogram (SPI label ‘dtw_constraint–itakura’; see Supplementary Note 1.2.7), which was a top performer on the fMRI film dataset (91%), but showed weaker performance on the smartwatch activity dataset (78%) and null performance on the EEG state dataset (53%). Yet some SPIs did perform strongly across all three datasets, such as the cross distance correlation (SPI label ‘dcorr_maxlag–10’; see Supplementary Note 1.2.2), which ranked among the top 10 SPIs for all three problems (90%, 66% and 91%, for the smartwatch activity, EEG state and fMRI film datasets, respectively). Moreover, different algorithmic variants of causally conditioned entropy were top performers for the smartwatch activity (‘cco_kozachenko’, 92%), EEG state (‘ccgaussian’, 69%) and fMRI film (‘cc_kozachenko_W=0.5’, 90%) datasets (although we note a strong dependence of the density estimation approach on CCE performance). We also found that grouping SPIs based on the fourteen data-driven modules (identified by their similarity of behavior on data, Fig. 2)
better captured their relative performance on these tasks than the six literature categories (from Fig. 1b), as shown in Supplementary Fig. 3, suggesting our modular representation as a useful one for understanding differential SPI performance on a given task.

Relative to investigating individual SPIs one at a time, we finally aimed to investigate the value of drawing on multiple SPIs simultaneously. We developed a combined representation of the pairwise dependence structures captured by all SPIs, allowing us to simultaneously represent each MTS using a large and diverse set of pairwise dependency structures (through feature concatenation, as described in the ‘Classification’ section of the Methods). Although this approach represents each MTS in a much higher-dimensional space than the individual SPI representation analyzed above (with associated challenges for robust classifier design), we expected it to outperform the individual best SPI on datasets involving multiple types of interactions, such that simultaneously leveraging multiple SPIs provides complementary and useful information about class differences. Relative to the top-performing individual SPI, our combined SPI approach improved classification accuracy on the smartwatch activity dataset (to 96%) and the EEG state dataset (to 71%), shown as vertical red lines in Fig. 3b, e. On the fMRI film dataset, it yielded slightly lower performance (91%) than the top individual SPI, ‘reci’ (95%), suggesting that the associated interactions are well-captured by a single, well-chosen SPI on this dataset (that is, multiple SPIs do not provide an advantage sufficient to overcome the challenges of fitting a classifier in a higher-dimensional space). By simultaneously drawing on a wide range of SPIs, the simple statistical approach demonstrated here can quantify multiple complementary types of interactions from MTS data (and is likely to yield improved accuracy through optimization; see ‘Discussion’ section).

Discussion

Our empirical organization of SPIs (Fig. 2b) identifies connections between a rich literature of diverse methods for quantifying interactions between time series, and thereby highlights fruitful directions for future research to consolidate, extend and develop new theory. These results were obtained using a similarity index, R, that averages across a large and diverse range of 1,053 MTS but—as the behavior of an SPI depends on the types of interactions present in a dataset—the resulting relationships between SPIs depend on the MTS we chose to include. As a simple illustrative example, consider the case that our MTS library only contained data generated by lag-1 vector autoregressive models, VAR(1). Then the unique behavior of more sophisticated SPIs (such as those capturing nonlinear dependence or causal coupling on longer time lags) would not be observed, as data containing those types of interactions would be absent. Consequently, the variety of behaviors exhibited by our library of SPIs would be reduced, and the modular structure evident in the dendrogram of Fig. 2b would become far more homogeneous. In constructing the MTS data library used in this work, we thus aimed to be as even-handed and comprehensive as possible in sampling from diverse systems, which has been sufficient to yield a meaningful and useful representation of the interdisciplinary literature on SPIs. However, as any such finite sample is incomplete and relies on subjective decisions in its construction, future work may explore the dataset-dependent similarity of SPI pairs in detail to construct more nuanced organizations of the literature. This would provide new understanding of how different mechanisms (and hence empirical dependency structures) play out in different classes of complex systems. Future work may also revisit the simple methodological choices made here, including our decisions to: (1) quantify the similarity of a pair of SPIs using a single number (R); (2) represent the resulting relationships in one dimension (as a dendrogram); and (3) analyze the resulting dendrogram at a single selected resolution (a 14-module decomposition). For example, more complex unsupervised methods, such as overlapping community-detection methods, could reveal interesting new relationships between SPIs at different resolution scales.

As different time-varying systems contain different statistical relationships between their elements (for example, those that capture instantaneous or time-delayed responses, linear or nonlinear interactions, are conditioned on the past, allow for non-constant delays, or infer directional coupling), our results demonstrate that a highly comparative approach can be used to detect the most informative types of SPIs for capturing the relevant types of interactions underlying a given dataset. Unlike many machine-learning approaches to MTS classification, which can be challenging to interpret, this approach connects scientists to interpretable theory that shapes understanding of the most important types of pairwise interactions in a dataset, following recent undertakings to unify interdisciplinary literatures on summary statistics (‘features’) for univariate time series and complex networks. This broad, comprehensive methodological comparison stands in contrast to a more conventional approach in which the data analyst manually selects a method—a practice that leaves open the possibility that alternative methods may provide clearer interpretation, better performance, or computational efficiencies. The three classification case studies analyzed here provide a simple demonstration of the procedure, automatically highlighting high-performing SPIs from across the literature and providing interpretable understanding of the relevant interactions in each dataset. We observed a wide range of performance in all cases, highlighting the importance of careful selection of SPIs for a given application. In the EEG state dataset, for example, our analysis flagged high-performing SPIs consistent with common methodological practice in EEG analysis (thus recapitulating existing domain knowledge), and others with surprisingly poor performance, as well as flagging novel high-performing SPIs as promising candidates for methodological innovation in the field. Future work investigating which types of SPIs are best suited to which types of problems will yield new insights into the interactions underlying different complex systems, and is likely to uncover additional novel applications of SPIs to new problems.

Although the ability to compare across the full library of SPIs is powerful, it comes at a computational cost, particularly for larger MTS datasets in which each MTS is sampled over many time points and processes. The high redundancy between many groups of SPIs (Fig. 2), together with the preliminary indications from case studies that some SPIs may exhibit generally high performance, suggest the potential for major efficiency gains by constructing a high-performing and minimally redundant reduced set of SPIs. Beyond a simple approach of selecting a representative SPI from each of the 14 modules, a more systematic approach could compare SPI performance across a wide range of representative MTS tasks, for example, following recent work reducing a large pool of thousands of interdependent univariate time-series features to just 22 representatives. There is also much scope for statistical optimization in applying any given set of SPIs to real-world problems, beyond the simple choices made here. In particular, performance could be optimized by tackling the challenges of high-dimensionality and strong interdependence of SPIs using dimensionality reduction, regularization or feature-selection methods to find small but highly explanatory combinations of complementary SPIs tailored to a given dataset. Given the methodological freedoms involved in highly comparative time-series analysis, pre-registration should be considered for transparency and to avoid bias (for example, ref. 49).

Beyond the applications to MTS classification presented here, the ability to represent the coupling structure in a MTS using a large and diverse set of pairwise interactions could form the foundation for tackling myriad statistical inference problems, including regression, clustering, anomaly detection, and causal network inference. For example, current methods for inferring networks of causal interactions from MTS data range from heuristic approaches based on thresholding a set of pairwise dependencies defined by a given SPI through to full statistical inference. Rather than manually selecting an SPI for this purpose, the ability to compare diverse SPIs provides the flexibility to...
capture different types of underlying interactions that could form the basis of improved network-inference algorithms. Such an extension may also go beyond point estimates of pairwise dependence strengths (considered here) towards assessing statistical significance through comparison to an appropriate null model\(^{34,35}\). Future work could also explore comprehensive statistical representations of MTS using more diverse properties than just the set of pairwise interactions, by also incorporating properties of univariate dynamics of individual system components\(^{36}\), system-level structure in the full set of pairwise interactions represented as a network\(^{37}\) and higher-order interactions\(^{38}\).

The flexible and extendable software accompanying this work, pyspi\(^{2}\) (see ‘Code Availability’ section), facilitates the application of our highly comparative approach to quantifying pairwise interactions to myriad applications. Through careful framing of systems-level inference\(^{39}\), it has the potential to highlight unexpected and powerful methodological approaches to quantifying interaction patterns in time-varying systems. The data library of over 1,000 MTS provided with this work\(^{40}\) (see ‘Data Availability’ section) is another resource that will allow researchers to characterize the behavior of their computational methods on a comprehensive range of real-world and simulated systems, addressing issues associated with only testing new complex systems methods on idealized datasets\(^{41}\). In summary, this work demonstrates the utility of an empirical approach to unifying diverse complex dynamical systems and their methods of analysis, providing insights and tools for scientific discovery.

Methods

**Comprehensive scientific libraries of methods and data**

**Library of 237 SPIs.** This work takes an empirical approach to understanding the relationships between diverse scientific methods for quantifying pairwise interactions between time series, formulated as an SPI. The first step in this approach was to construct a comprehensive, annotated library of SPIs. Here we introduce a library of 237 SPIs, organized into six broad categories based on their underlying theory: ‘basic’ (such as co-variance, Kendall’s \(\tau\) (ref. 59) and cross-correlation); ‘distance similarity’ (for example, distance correlations\(^{42}\), kernel-based independence tests\(^{30,32}\) and dynamic time warping\(^{10}\)); ‘causal indices’ (including additive noise models and convergent cross-mapping\(^{43}\)); ‘information theory’ (such as Granger causality\(^{44,45}\), note that Granger causality is included in the information theory category as it is equivalent to transfer entropy evaluated with a Gaussian estimator\(^{46}\), transfer entropy\(^{47}\) and integrated information\(^{48}\)); ‘spectral’ (derived from Fourier or wavelet transformations, for example, coherence magnitude, phase-locking value\(^{49}\) and spectral Granger causality\(^{50}\)); and ‘miscellaneous’ (for example, co-integration and model fits). A full list of SPIs, along with descriptions and references, is presented in Supplementary Note 1.

**Library of 1,053 MTS.** To understand how each SPI behaves on different types of multivariate time-series data, we constructed a library of 1,053 diverse model-generated and real-world MTS. Our aim in assembling these data was to capture the main classes of systems and dynamics that are studied across scientific disciplines, including synchronization, spatiotemporal chaos, wave propagation, criticality and phase transitions. Our library contains SOS synthetic MTS generated from mathematical models, including: uncorrelated and correlated noise (for example, Cauchy and normally distributed noise and Brownian motion); coupled maps (such as vector autoregression\(^{51}\) and coupled map lattices\(^{52}\)); coupled ordinary differential equations (including Kuramoto oscillators\(^{53}\), Hodgkin–Huxley and Wilson–Cowan networks\(^{54}\)); and partial differential equations (namely, wave equations, in which processes are embedded in physical space). It also contains 548 diverse real-world MTS from public databases across: geophysics (such as earthquake seismograms and atmospheric processes); medicine (for example, heartbeat sonograms, fMRI data and EEGs); physiology (including accelerometer and gyroscope readings for sports and basic motions); and finance (for instance, exchange rates and stock prices), among others. Each MTS comprises between 500 and 40 processes, and between 100 and 2,000 observations—characteristics that were designed to match many real-world datasets. Across all MTS in our library, we have a total of 195,112 pairwise interactions that we used to evaluate the SPIs. Descriptions of the full set of MTS analyzed here are presented in Supplementary Note 2.

**Quantifying similarity between SPIs using the empirical similarity index**

Our main aim in this work was to assess the similarity of behavior of any two SPIs across a diverse range of MTS. In this section we describe the empirical similarity index, \(R\), used to capture the similarity between the behavior of all pairs of SPIs across our diverse library of 1,053 MTS.

This index measures the monotonic relationship between a pair of SPIs across a large number of datasets that have different numbers of processes (and thus a different number of pairwise interactions). For an \(M\)-variate time series, \(z = (z_1, \ldots, z_M)\), we first consider the \(M \times M\) matrix of pairwise interactions (MPI), constructed by evaluating a given SPI for all pairs of \(M\) processes within the MTS. The scalar values of the MPI, \(\mathbf{s} = (s_{ij}) \in \mathbb{R}^{M \times M}\), where the \((i, j)\) entries of the matrix, \(s_{ij} = s(z_i, z_j)\), denotes an SPI evaluated on the \(i\)th and \(j\)th time series, \(z_i\) and \(z_j\), respectively. In general, undirected statistics (for which \(s_{ij} = s_{ji}\) such as Kendall’s \(\tau\)) yield symmetric MPIs, whereas directed statistics (such as transfer entropy, which have \(s_{ij} \neq s_{ji}\)) yield asymmetric MPIs. Furthermore, some SPIs are signed (for example, correlation coefficients are within \([-1, 1]\)), whereas others are unsigned (for example, the distance correlation is within \([0, 1]\)). To compare all methods appropriately, we converted signed SPIs to their absolute value, such that all statistics increase with the strength of dependency between \(z_i\) and \(z_j\) (for example, we analyze the magnitude of the covariance rather than its sign). Some example MPIs are shown in Supplementary Fig. 1a. Having computed MPIs for all SPIs on all 1,053 MTS, we computed the empirical similarity index between each pair of SPIs via a two-step process (depicted in Supplementary Fig. 1b). First, we defined the similarity of a pair of SPIs, \(k\) and \(l\), on a given dataset, \(d\). We did this by computing MPIs for \(k\) and \(l\) (yielding \(S_k\) and \(S_l\)), respectively, and then computing the absolute value of the Spearman’s rank correlation coefficient between each of their (off-diagonal) entries, \(|r_{kl}|\). The resulting correlation value, \(|r_{kl}|\), thus captures the strength of the monotonic relationship between the output of the two SPIs on the dataset \(d\). After repeating this computation for all MTS in our library, we calculated the empirical similarity index, \(R\), as the average of \(|r_{kl}|\) across all datasets. Although \(R\) provides a useful scalar summary of the similarity between a pair of SPIs, it is important to note that some pairs of SPIs have quite a wide distribution of scores, \(|r_{kl}|\), across datasets, indicating that their yield highly correlated outputs on some MTS (high \(|r_{kl}|\)) but not on others (low \(|r_{kl}|\)). See Supplementary Note 3.1 for more details, including a detailed visual breakdown of \(|r_{kl}|\).

**Classification case studies**

**Datasets.** Smartwatch activity dataset. The smartwatch activity dataset is derived from the Basic Motions problem in the University of East Anglia (UEA) MTS classification repository\(^{44}\). Each MTS includes six sensors—a three-axis accelerometer and a three-axis gyroscope—recorded for 10 s at 10 Hz, yielding 1,000 time points. There are 20 MTS in each of the four classes (resting, walking, running or playing badminton) for a total of 80 MTS in the dataset. This dataset was recently analyzed in a large MTS classification challenge\(^{45}\) in which the data was split 50%–50% into training and test subsets with 30 stratified repeats. The baseline classifier (based on dynamic time warping) achieved 95.25% accuracy; the best algorithm (HIVE–COTE) achieved 100% accuracy\(^{46}\).
The EEG state dataset corresponds to the SelfRegulationSCPI problem in the UEA MTS classification repository and was originally published in work by Birbaumer and colleagues. Electrical activity was measured from six EEG channels in one participant as they were instructed to move a cursor up or down on a computer screen by generating negative or positive slow cortical potentials, respectively. The physical placement of these EEG electrodes is depicted in Fig. 3d. Cortical activity (measured in microvolts) was recorded for 3.5 s at 256 Hz, yielding 896 time points in each channel per MTS, with 282 negativity trials and 279 positivity trials. This dataset was analyzed in work by Ruiz et al., assessing accuracy using 30 stratified train–test splits of 268 training and 293 test samples. In the original comparison, the baseline DTW algorithm achieved an accuracy of 81.8% and the best algorithm (TapNet) achieved 95.68% (ref. 33).

The fMRI film dataset is derived from a functional connectome fingerprinting study examining individual signatures of cortical activity in $N = 29$ individuals at rest or while watching a film. In this dataset, blood-oxygen-level-dependent signals were recorded in 114 parcellated cortical regions with a repetition time of 813 ms (a sampling rate of 1.23 Hz) for either 1,200 frames (resting), or between 952 and 1,000 frames (film-watching). Each trial type (rest versus film-watching) was repeated four times per participant. Here we analyzed pre-processed data obtained from work by Betzel and co-workers. For consistency and simplicity, we examined the first rest and first film-watching session per participant; across all participants and trials, only the first 947 frames contain real data, and thus we restricted our analysis to this time range. To retain a comparable number of processes as the first two classification case studies, we averaged blood-oxygen-level-dependent signals from the 114 original brain regions into the seven functional networks from Yeo et al., as depicted in Fig. 3g.

For all three case studies, our simple approach to SPI-based classification involved computing the MPIs ($6 \times 6$ for the smartwatch activity and EEG state datasets, and $7 \times 7$ for the fMRI film dataset) for each z-scored MTS and repeating for each SPI. We then used the elements of these matrices as features for a linear SVM classifier. Note that we used the most recent version of pyspi (v.0.4.0) to compute SPIs for the classification case studies, which included some improved implementations of some SPIs. Features were extracted from each SPI differently for directed and undirected SPIs (see Supplementary Data 1); we used the upper triangular entries as features for undirected SPIs (for which the corresponding MPIs symmetri, whereas we used all non-diagonal elements as features for directed SPIs. As a preprocessing step, for each case study, we removed any SPI that had invalid entries (due to numerical issues) in any of the MPIs, or gave constant results across all MTS (see Supplementary Data 3 for a list of omitted SPIs). This yielded a set of 228 SPIs for the smartwatch activity problem, 219 SPIs for the EEG state problem and 227 SPIs for the fMRI film problem. For the analysis involving combining all SPIs into a single classifier, this yielded a total of 4,755 features for the smartwatch activity dataset, 4,659 features for the EEG state dataset and 6,743 features for the fMRI film dataset.

The linear SVM was implemented using default settings from Python’s scikitlearn as part of a classification pipeline that involved z-score feature normalization (fitted on training data and applied to unseen test data). The very simple methodological choices made here allowed us to focus on demonstrating the key conceptual types of analyses made possible by drawing on a diverse set of SPIs, aiding transparency while acknowledging that more complicated statistical methodologies are likely to improve the classification performance quoted here. We implemented 30 class-stratified train–test splits for cross-validation with the same proportions implemented in ref. 33 using the StratifiedShuffleSplit function for the smartwatch activity and EEG state datasets. As there are $N = 29$ individuals in the fMRI film dataset, we implemented leave-one-individual-out cross-validation, such that each classifier was trained with the rest and film scans of $N = 28$ participants, and tested on the rest and film fMRI scans of the left-out participant.

We measured classification performance using total accuracy for all three case studies. Statistical significance was estimated using a permutation testing approach whereby 100 null models were fitted (using randomly shuffled class labels) and evaluated using the same cross-validation classification procedure described above. The observed classification performance for each SPI was then compared with the pooled null distribution of all SPIs, yielding $P$-values that were then adjusted for multiple comparisons by controlling the family-wise error at 0.05 using the method of Bonferroni. The performance metric for the union of all SPIs was similarly compared with its corresponding null permutation distribution to yield a single $P$-value per classification problem.

**Data availability**

The full database of 1,053 diverse real-world and simulated MTS analyzed here is available on the Zenodo repository at: https://doi.org/10.5281/zenodo.711947 (ref. 58). This resource could be used to test scientific methods on a diverse sample of MTS. Time-series data used in the three case studies are from open sources, as described in the ‘Classification case studies’ section in the Methods. Source data are provided with this paper.

**Code availability**

Accompanying this paper is an extendable Python-based package called pyspi, which includes implementations of all 237 SPIs. This software allows users to compare the behavior of an interdisciplinary literature on methods for quantifying interactions between pairs of time series. Furthermore, code and instructions for reproducing the results and figures presented in this work can be found on the Zenodo repository at: https://doi.org/10.5281/zenodo.8027702 (ref. 67).

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**Author contributions**

N.T. and B.D.F. conceived of the project. O.M.C. developed the software and large data repositories and performed the main analysis. A.G.B. performed the classification case study analyses. B.D.F. supervised the project with input from J.T.L. and N.T. O.M.C. and B.D.F. wrote the paper, with input from all other co-authors.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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Software and code

Data analysis

The pysipr software package was developed by us; it accompanies this work. Classification analyses were performed using pysipr v0.4.0: https://github.com/DynamicsAndNeuralSystems/pysipr (https://doi.org/10.5281/zenodo.8223340). A docker image for pysipr is available here: https://hub.docker.com/r/arianguyen/pysipr

Analysis and visualizations were done using a combination of R (v4.3.0) and python (v3.9.7).

Full code and data to reproduce the presented analyses are in https://github.com/DynamicsAndNeuralSystems/pysipr_paper_code (https://doi.org/10.5281/zenodo.8027702).

Python package versions are as follows: pandas (v1.5.2), numpy (v1.22.4), scikit-learn (v0.24.1), pyarrow (v10.0.1), scipy (v1.7.3), statsmodels (v0.12.1), pyyaml (v5.4), tqdm (v4.502), nitime (v0.9), hypopo (v0.2.1), pyEDM (v1.9.3.0), jupyter (v1.2.0), scikitimage (v0.8.0), dill (v0.3.2), spectral-connectivity (v0.1.4.27), torch (v1.10.0), cdt (v0.5.23), oct2py (v5.2.0), tlearn (v0.5.2), mne (v0.23.0), seborn (v0.11.0).

R package versions are as follows: tidyverse (v2.0.0), glue (v1.6.2), reticulate (v1.28), icesTAF (v4.2.0), cowplot (v1.1.1), plyr (v1.8.8).

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Data analyzed in this study were both generated via numerical simulation and assembled from publicly available sources. See Supplementary Section 2 for full details and sources. The full set of multivariate time-series datasets analyzed are available in the following Zenodo repository: https://zenodo.org/record/7118947.

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| Sample size | Our interdisciplinary library of 1053 diverse multivariate time series aimed to be as comprehensive as possible. This large collection of MTS was curated with the aim of capturing the main classes of systems and dynamics that are studied across scientific disciplines, including synchronization, spatiotemporal chaos, wave propagation, criticality, and phase transitions. Our aim was to encompass a sufficiently diverse sample of the types of data analyzed in the scientific literature as comprehensively as possible within practical constraints, such that the empirical similarity index evaluated in this work constitutes a meaningful measure of the behavioral similarity of two statistics of pairwise interaction (SIs).
| Data exclusions | We included MTS that contained between 5–40 processes and between 100–2000 observations. We set this range both for practical purposes (avoiding large calculation times required of larger datasets), and because these characteristics match those of real-world datasets that are commonly analyzed.
| Replication | This study aims to understand the relationship between different algorithms for quantifying statistical dependence between a pair of time series. We note in the Discussion that, since the behavior of an SPI depends on the types of interactions present in a dataset, the organization we obtain depends on the data included in this set of MTS. Please see the relevant paragraph in Discussion for additional discussion of this point, including why we might expect different structures to be found when analyzing data obtained from different types of systems.
| Randomization | Cross-validation and permutation-based statistics were applied to the classification case studies, as described in Methods: ‘Classification’.
| Blinding | Randomization is not applicable to this study as it does not involve an experimental design that involves measuring data from participants.
| Blinding | Blinding is not applicable to this study as it does not involve an experimental design that involves measuring data from participants.
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Palaeontology and Archaeology

Specimen provenance
Provide provenance information for specimens and describe permits that were obtained for the work (including the name of the issuing authority, the date of issue, and any identifying information). Permits should encompass collection and, where applicable, export.

Specimen deposition
Indicate where the specimens have been deposited to permit free access by other researchers.
Dating methods

If new dates are provided, describe how they were obtained (e.g. collection, storage, sample pretreatment and measurement), where they were obtained (i.e. lab name), the calibration program and the protocol for quality assurance OR state that no new dates are provided.

☐ Tick this box to confirm that the raw and calibrated dates are available in the paper or in Supplementary Information.

Ethics oversight

Identify the organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or guidance was required and explain why not.

Note that full information on the approval of the study protocol must also be provided in the manuscript.

### Animals and other research organisms

Policy information about studies involving animals: ARRIVE guidelines recommended for reporting animal research, and Sex and Gender in Research

| Laboratory animals | For laboratory animals, report species, strain and age OR state that the study did not involve laboratory animals. |
|--------------------|--------------------------------------------------|
| Wild animals       | Provide details on animals observed in or captured in the field; report species and age where possible. Describe how animals were caught and transported and what happened to captive animals after the study (if killed, explain why and describe method; if released, say where and when) OR state that the study did not involve wild animals. |
| Reporting on sex   | Indicate if findings apply to only one sex; describe whether sex was considered in study design, methods used for assigning sex. Provide data disaggregated for sex where this information has been collected in the source data as appropriate; provide overall numbers in this Reporting Summary. Please state if this information has not been collected. Report sex-based analyses where performed, justify reasons for lack of sex-based analysis. |
| Field-collected samples | For laboratory work with field-collected samples, describe all relevant parameters such as housing, maintenance, temperature, photoperiod and end-of-experiment protocol OR state that the study did not involve samples collected from the field. |
| Ethics oversight   | Identify the organization(s) that approved or provided guidance on the study protocol, OR state that no ethical approval or guidance was required and explain why not. |

Note that full information on the approval of the study protocol must also be provided in the manuscript.

### Clinical data

Policy information about clinical studies

All manuscripts should comply with the ICMJE guidelines for publication of clinical research and a completed CONSORT checklist must be included with all submissions.

| Clinical trial registration | Provide the trial registration number from ClinicalTrials.gov or an equivalent agency. |
| Study protocol             | Note where the full trial protocol can be accessed OR if not available, explain why. |
| Data collection            | Describe the settings and locales of data collection, noting the time periods of recruitment and data collection. |
| Outcomes                   | Describe how you pre-defined primary and secondary outcome measures and how you assessed these measures. |

### Dual use research of concern

Policy information about dual use research of concern

### Hazards

Could the accidental, deliberate or reckless misuse of agents or technologies generated in the work, or the application of information presented in the manuscript, pose a threat to:

| No | Yes |
|----|-----|
| ☒ Public health | ☐ |
| ☒ National security | ☐ |
| ☒ Crops and/or livestock | ☐ |
| ☒ Ecosystems | ☐ |
| ☒ Any other significant area | ☐ |
Experiments of concern

Does the work involve any of these experiments of concern:

| No | Yes |
|----|-----|
| ☒ | ☐  | Demonstrate how to render a vaccine ineffective
| ☒ | ☐  | Confer resistance to therapeutically useful antibiotics or antiviral agents
| ☒ | ☐  | Enhance the virulence of a pathogen or render a nonpathogen virulent
| ☒ | ☐  | Increase transmissibility of a pathogen
| ☒ | ☐  | Alter the host range of a pathogen
| ☒ | ☐  | Enable evasion of diagnostics/detection mechanisms
| ☒ | ☐  | Enable the weaponization of a biological agent or toxin
| ☒ | ☐  | Any other potentially harmful combination of experiments and agents

ChIP-seq

Data deposition

☐ Confirm that both raw and final processed data have been deposited in a public database such as GEO.

☐ Confirm that you have deposited or provided access to graph files (e.g. BED files) for the called peaks.

Data access links

May remain private before publication.

For "Initial submission" or "Revised version" documents, provide reviewer access links. For your "Final submission" document, provide a link to the deposited data.

Files in database submission

Provide a list of all files available in the database submission.

Genome browser session

Provide a link to an anonymized genome browser session for "Initial submission" and "Revised version" documents only, to enable peer review. Write "no longer applicable" for "Final submission" documents.

Methodology

Replicates

Describe the experimental replicates, specifying number, type and replicate agreement.

Sequencing depth

Describe the sequencing depth for each experiment, providing the total number of reads, uniquely mapped reads, length of reads and whether they were paired- or single-end.

Antibodies

Describe the antibodies used for the ChIP-seq experiments; as applicable, provide supplier name, catalog number, clone name, and lot number.

Peak calling parameters

Specify the command line program and parameters used for read mapping and peak calling, including the ChIP, control and index files used.

Data quality

Describe the methods used to ensure data quality in full detail, including how many peaks are at FDR 5% and above 5-fold enrichment.

Software

Describe the software used to collect and analyze the ChIP-seq data. For custom code that has been deposited into a community repository, provide accession details.

Flow Cytometry

Plots

Confirm that:

☐ The axis labels state the marker and fluorochrome used (e.g. CD4-FITC).

☐ The axis scales are clearly visible. Include numbers along axes only for bottom left plot of group (a ‘group’ is an analysis of identical markers).

☐ All plots are contour plots with outliers or pseudocolor plots.

☐ A numerical value for number of cells or percentage (with statistics) is provided.

Methodology

Sample preparation

Describe the sample preparation, detailing the biological source of the cells and any tissue processing steps used.

Instrument

Identify the instrument used for data collection, specifying make and model number.
**Magnetic resonance imaging**

**Experimental design**

- **Design type**: Indicate task or resting state; event-related or block design.
- **Design specifications**: Specify the number of blocks, trials or experimental units per session and/or subject, and specify the length of each trial or block (if trials are blocked) and interval between trials.
- **Behavioral performance measures**: State number and/or type of variables recorded (e.g. correct button press, response time) and what statistics were used to establish that the subjects were performing the task as expected (e.g. mean, range, and/or standard deviation across subjects).

**Acquisition**

- **Imaging type(s)**: Specify: functional, structural, diffusion, perfusion.
- **Field strength**: Specify in Tesla
- **Sequence & imaging parameters**: Specify the pulse sequence type (gradient echo, spin echo, etc.), imaging type (EPI, spiral, etc.), field of view, matrix size, slice thickness, orientation and TE/TR/flip angle.
- **Area of acquisition**: State whether a whole brain scan was used or define the area of acquisition, describing how the region was determined.
- **Diffusion MRI**: Used [ ] Not used [ ]

**Preprocessing**

- **Preprocessing software**: Provide detail on software version and revision number and on specific parameters (model/functions, brain extraction, segmentation, smoothing kernel size, etc.).
- **Normalization**: If data were normalized/standardized, describe the approach(es): specify linear or non-linear and define image types used for transformation OR indicate that data were not normalized and explain rationale for lack of normalization.
- **Normalization template**: Describe the template used for normalization/transformation, specifying subject space or group standardized space (e.g. original Talairach, MNI305, ICBM152) OR indicate that the data were not normalized.
- **Noise and artifact removal**: Describe your procedure(s) for artifact and structured noise removal, specifying motion parameters, tissue signals and physiological signals (heart rate, respiration).
- **Volume censoring**: Define your software and/or method and criteria for volume censoring, and state the extent of such censoring.

**Statistical modeling & inference**

- **Model type and settings**: Specify type (mass univariate, multivariate, RSA, predictive, etc.) and describe essential details of the model at the first and second levels (e.g. fixed, random or mixed effects; drift or auto-correlation).
- **Effect(s) tested**: Define precise effect in terms of the task or stimulus conditions instead of psychological concepts and indicate whether ANOVA or factorial designs were used.
- **Specify type of analysis**: Whole brain [ ] ROI-based [ ] Both [ ]
- **Statistic type for inference**: Specify voxel-wise or cluster-wise and report all relevant parameters for cluster-wise methods.
- **Correction**: Specify the type of correction and how it is obtained for multiple comparisons (e.g. FWE, FDR, permutation or Monte Carlo).
Models & analysis

| n/a | Involved in the study |
|-----|-----------------------|
| ☑   | Functional and/or effective connectivity |
|     | Graph analysis |
| ☑   | Multivariate modeling or predictive analysis |

**Functional and/or effective connectivity**

We compared over 230 measures of functional connectivity in the “fMRI film” case study.

**Graph analysis**

Report the dependent variable and connectivity measure, specifying weighted graph or binarized graph, subject- or group-level, and the global and/or node summaries used (e.g. clustering coefficient, efficiency, etc.).

**Multivariate modeling and predictive analysis**

Specify independent variables, features extraction and dimension reduction, model, training and evaluation metrics.