The Topological Architecture of Brain Identity

Simone Poetto,^{1,2} Haily Merritt,^{3,4} Andrea Santoro,² Giovanni Rabuffo,⁵ Demian Battaglia,⁶

Francesco Vaccarino,⁷ Manish Saggar,^{8,9} Andrea Brovelli,¹⁰ and Giovanni Petri^{11,12,2,*}

¹Nicolaus Copernicus University, Toruń, Poland

²CENTAI Institute, Turin, Italy

³Program in Cognitive Science, Indiana University, Bloomington, IN, USA

⁴Department of Informatics, Indiana University, Bloomington, IN, USA

⁵Institut de Neurosciences des Systèmes (INS), UMR 1106,

INSERM, Aix-Marseille Université, Marseille, France

⁶Laboratoire de Neurosciences Cognitives et Adaptatives (LNCA),

Faculté de Psychologie, Université de Strasbourg, Strasbourg, France

⁷Dipartimento di Scienze Matematiche, Politecnico di Torino, Turin, Italy

⁸Department of Psychiatry and Behavioral Sciences,

Stanford University School of Medicine, Stanford, CA, USA

⁹Wu Tsai Neurosciences Institute, Stanford University, Stanford, CA, USA

¹⁰Institut de Neurosciences de la Timone (INT), UMR 7289,

CNRS, Aix-Marseille Université, Marseille, France

¹¹NP Lab, Network Science Institute, Northeastern University London, London, UK

¹²Department of Physics, Northeastern University, Boston, USA

(Dated: June 20, 2025)

Accurately identifying individuals from brain activity—functional fingerprinting—is a powerful tool for understanding individual variability and detecting brain disorders. Most current approaches rely on functional connectivity (FC), which measures pairwise correlations between brain regions. However, FC is limited in capturing the higher-order, multiscale structure of brain organization. Here, we propose a novel fingerprinting method based on homological scaffolds, a topological representation derived from persistent homology of resting-state fMRI data. Using data from the Human Connectome Project (n = 100), we show that scaffold-based fingerprints achieve near-perfect identification accuracy ($\sim 100\%$), outperforming FC-based methods (90%), and remain robust across preprocessing pipelines, atlas choices, and even with drastically shortened scan durations. Unlike FC, in which fingerprinting features localize within networks, scaffolds derive their discriminative power from inter-network connections, revealing the existence of individual mesoscale organizational signatures. Finally, we show that scaffolds act as bridges between redundancy and synergy, by balancing redundancy along high-FC border edges with high synergy across the topological voids that the cycles define. These findings establish topological scaffolds as a powerful tool for capturing individual variability, revealing that unique signatures of brain organization are encoded in the interplay between mesoscale network integration and information dynamics.

I. INTRODUCTION

The human brain can be modeled as a complex network of interconnected regions that continuously exchange information [1]. Advances in neuroimaging techniques, particularly functional magnetic resonance imaging (fMRI). have allowed researchers to investigate the properties of these networks in terms of the functional relationships between different regions of the brain [2–5]. Functional connectivity (FC), typically measured as a statistical dependency between the activity of pairs of brain regions [6], has become a foundational tool to characterize large-scale brain organization [6]. Decades of research using FC and related techniques have demonstrated that human brain networks share common architectural principles. Across individuals, networks consistently exhibit efficient information processing pathways [7], heavy-tailed distributions of connection weights, node degrees with the presence of hubs and rich-club organizations [8–10], modular structures [11], and efficient wiring costs [12].

Although these common features are shared between individuals, each brain network carries a unique signature [13], shaped by individual differences in cognitive abilities, experiences, and plasticity. Indeed, subtle but robust differences in brain network organization form the basis for functional brain fingerprinting [14]. In their seminal work [15], Finn and colleagues demonstrated that individuals can be reliably identified based solely on their whole-brain FC profiles. This work sparked broad interest in uncovering subject-level patterns from fMRI and has since expanded to other modalities, including electroencephalography (EEG) [16–19], functional near-infrared spectroscopy (fNIRS) [20], and magnetoencephalography (MEG) [21, 22]. These studies underscore the robustness and cross-modality generality of individual brain connectivity patterns across modalities and cognitive states [23–25]. Moreover, the clinical relevance of fingerprinting is gaining traction: alterations in an individual's connectivity profile over time can have important

^{*} giovanni.petri@nulondon.ac.uk

implications in clinical settings [26], and in early diagnosis of neurodegeneration [27–29].

While effective for this scope, FC has intrinsic limitations: it captures only pairwise statistical associations and typically assumes temporal stationarity. To address these constraints, emerging approaches extend connectivity analyses across multiple dimensions [30]. Edge time series, for instance, track the temporal evolution of connectivity strength between pairs of regions, characterizing subject-specific patterns that vary across time [31– 33]. Other frameworks move beyond pairwise interactions, focusing on higher-order interactions, which involve three or more brain regions [34]. For instance, information-theoretic approaches, such as multivariate information theory and partial entropy decomposition, quantify how information is shared, redundantly or synergistically, across multiple regions [35–37]. Parallel to these methods, topological data analysis (TDA) provides a lens for identifying higher-order structure in brain networks. Techniques such as persistent homology and simplicial complexes track multiregional interactions that persist across scales and time [38–42]. TDA has effectively identified differences in structural connectivity [39], functional connectivity [38, 43], brain representation [44], cortical thickness [45, 46], EEG dynamics [47], and PET images [48, 49].

A tool of particular interest from TDA for brain network analysis is the homological scaffold [38]. This approach uses persistent homology to identify the topologically significant links in a weighted brain network [50]. A homological scaffold is constructed by performing a weighted rank filtration, where edges are added one at a time to an initially empty graph in order of decreasing weight. At each step of the filtration, the graph is promoted to its clique complex-a simplicial complex formed by treating every complete subgraph (clique) as a simplex. This allows the computation of topological features, such as 1-dimensional homology classes (H_1) , which correspond to loops or topological holes in the network. The final scaffold includes all edges that participate in at least one of these cycles, effectively capturing the network's topological backbone. Unlike conventional thresholding, this method preserves essential multiscale topological features and avoids arbitrary cutoffs, enabling a more global view of mesoscale network organization [40], and is akin to a Rips-Vietoris filtration on metric associated to Pearson correlations. Edge weights in the scaffold can be defined either based on their persistence, i.e., how long an edge remains present during the filtration, or by their frequency, i.e., how many H_1 loops they participate in. Importantly, prior work has shown that homological scaffolds effectively distinguish altered brain states — separating psilocybin-treated individuals from placebo [38], as well as highlight different information integration schemes [51]. However, it remains unknown whether such topological sensitivity to subject-specific and state-specific patterns generalizes in the context of robust individual fingerprinting.

Here, we show that homological scaffolds derived from resting-state fMRI not only enhance individual specificity, but achieve near-perfect identification accuracy across sessions using just a fraction of the connections of classical FC. Beyond their fingerprinting power, scaffolds reveal a novel link between brain network topology and information integration [52]: redundant communication concentrates along borders of topological cycles making up the scaffolds, while interactions across the cycles are significantly more synergistic than expected, presenting lower correlations and thus giving origin to the topological void. This interplay underpins uniquely individual patterns of brain organization, positioning scaffolds as a powerful and interpretable framework for brain fingerprinting and biomarker discovery.

2

RESULTS

To investigate individual differences in brain functional organization, we analyzed resting-state fMRI data from 100 unrelated participants, each scanned in two separate sessions as part of the Human Connectome Project (HCP) [53]. FC matrices were constructed for each participant and session by computing Pearson correlations between the time series of 300 brain regions defined by the Schaefer atlas [54] (Fig. 1A).

To test whether topological structure outperforms conventional FC as a subject fingerprint, we apply persistent homology using weighted rank filtration, adding edges in decreasing order of their FC weight [50]. This process tracks the birth and death of topological features, specifically 1-dimensional loops (i.e. the generators of the H_1 homology group). We identify the core topological structure by constructing "frequency scaffolds" [38], which consist of all links participating in at least one homological generator, weighted by the number of cycles they belong to (Fig. 1B). This procedure typically yields sparse networks, retaining on average $\sim 1\%$ of the original links (Fig. 1C and see Methods for a detailed description). For completeness and as comparative benchmarks, we also compute thresholded FC matrices, retaining the top 25% (following standard FC approaches [55]) and 1% strongest connections (to match scaffold's densities). Finally, we measure the fingerprinting performance of each representation. We do this by vectorizing the upper triangle of the adjacency matrix for each representation, and, for every participant, compute the Pearson's correlations between their Session 1 vector and all Session 2 vectors (and vice versa). Following standard approaches [15, 56], a correct identification occurred when a participant's own data yielded the highest cross-session correlation. We repeated this procedure in both directions — Session 1 to Session 2 and Session 2 to Session 1 -and defined overall accuracy as the average identification rate across the two directions (Fig. 1D).



FIG. 1. Schematic of scaffold computation and fingerprinting workflow. (A) Resting-state fMRI scans from the 100 unrelated subjects of the Human Connectome Project are used to compute individual functional connectivity matrices from two sessions (~10 days apart). (B) Conceptual illustration of homological scaffold construction. Edges are progressively added in decreasing weight order, and those participating in topological cycles (H_1 features) form the scaffold. (C) Comparison between a functional connectivity matrix thresholded to retain 1% of the strongest connections (left) and the corresponding scaffold (right), which achieves similar sparsity while preserving topologically informative links. (D) Visual representation of the functional fingerprinting process: an individual's test session brain connectivity matrix is compared against a database of re-test session matrices to identify the correct match based on the highest similarity.

Homological scaffolds outperform functional connectivity in identifying subjects

To compare the homological scaffold with FC, we compared the similarity distributions — measured via Pearson correlation — between pairs of scans from the same individual and those from different individuals (Fig. 2A). The scaffold representation shows a markedly clearer separation between the within-subject and between-subject similarity distributions, as quantified by a significantly higher Cohen's d effect size compared to FC (Fig. 2B). To strengthen this observation, we also compute the identification success rate. We find that full FC and its sparsified versions (25% and 1% thresholds) plateau around 90%. Homological scaffolds instead achieve near-perfect identification with a $\sim 100\%$ success rate across the cohort (Fig.2A, Section I and Table S1).

Next, we investigate the robustness of these findings. First, we assess the impact of common fMRI preprocessing choices on scaffold performance (Sec. II and Table S4). Overall, we find that fingerprinting capacity of scaffolds is weakly affected by different preprocessing steps. Indeed, the only step to significantly impact performance is low-pass filtering, the omission of which substantially degraded accuracy (from 99% to 83.5% using Glasser atlas data). Conversely, other common preprocessing steps, like global signal regression (GSR), high-pass filtering, and the choice of scan acquisition order (left-right vs.





FIG. 2. Scaffolds outperform traditional functional connectivity in capturing individual-specific brain patterns. (A) Intersession similarity heatmaps and corresponding identification success rates for full Functional Connectivity (FC) (Success Rate = 90.5%), FC thresholded at 25% (Success Rate = 90.0%), FC thresholded at 1% (Success Rate = 87.5%), and Scaffolds (Success Rate = 100.0%). (B) Violin plots display the distributions of intra-subject (within-subject) and inter-subject (between-subject) Pearson correlations for the four representations shown in (A). Cohen's d values (ES) quantify the separation between these distributions: FC full (ES=3.17), FC th=25% (ES=3.37), FC th=1% (ES=3.78), and Scaffolds (ES=7.46). (C) Boxplots compare intra-subject and inter-subject correlation distributions for FC and Scaffolds across different brain atlases: AAL 89 (FC ES=1.37, Scaffolds ES=2.09), Schaefer 100 (FC ES=2.12, Scaffolds ES=2.72), Shen 268 (FC ES=2.99, Scaffolds ES=4.75), and Glasser 360 (FC ES=4.58, Scaffolds ES=8.04), with their respective Cohen's d effect sizes indicated. (D) Line graph showing the identification success rate (%) for Functional Connectivity and Scaffolds as a function of different fMRI window widths (from 100 to 1100 TRs). Standard deviation for scaffolds is also shown. This demonstrates that scaffold-based representations maintain high discriminability even with limited data, outperforming FC across all window scan lengths.

right-left phase encoding) have minimal or slightly detrimental effects on scaffold identifiability, which remained at or near 100% in most conditions.

Second, we evaluate fingerprinting performance for

different brain atlases, including anatomical (AAL89 [57, 58]) and functional parcellations of varying resolutions (Schaefer100/300 [54], Shen268 [59], Glasser360 [60]) (Fig. 2C, Sec. IA, Table S2). Across all atlases, scaffolds consistently outperform the corresponding 1% thresholded FC matrix (as well as the 25% thresholded and full FC, see SI). Identifiability scores strongly depended on atlas resolution, increasing from 43% (AAL89) and 64.5% (Schaefer100) with scaffolds to 100% (Schaefer300) and 99.5% (Glasser360). As also shown in previous studies [61], this suggests that higher resolution parcellations better capture the individual-specific details.

Finally, we examine how scaffold performance changes with shorter scan lengths. Specifically, we progressively shorten fMRI data from 1200 TRs (~ 14.4 minutes) to 50 TRs (~ 0.6 minutes) and compute the identification success rate at each step to quantify identification accuracy (Fig. 2D, see also methods for details, and Fig. S2 for results on effect size). Across all window lengths, scaffolds again consistently outperform thresholded FC, maintaining a high effect size comparable to that achieved using the full FC timeseries, even for windows as short as 100 TRs. This indicates that scaffolds capture stable individual traits even at shorter time scales or limited scan time. In Sec. III we also provide a comparison of fingerprinting capacity within- and across-sessions, showing that within the same session the fingerprinting capacity at short timescales of FC and scaffolds is comparable, supporting the idea that scaffolds captures preserved functional traits, while FC better captures transient states.

Scaffold edges are distributed across the brain

Having established the superior accuracy and robustness of scaffolds, we next turn to: what makes them so effective? Specifically, we examine where individualspecific information is concentrated within these networks. To do this, we calculated the differential power for each edge in both the FC (1% thresholded) and scaffold representations, following Finn et al. [15]. This metric identifies edges that are highly stable within an individual across sessions but highly variable between individuals. Following previous work [15], we selected the top 0.2% of edges with the highest differential power, considering these as the most critical for fingerprinting.

We then examined the distribution of these highimportance edges across 7 canonical resting-state networks [62]. A striking difference emerged (Fig. 3A-B): for FC, the vast majority (72%) of important edges were within individual networks (intra-network connections). In contrast, for the scaffold, the majority (56%) of important edges spanned between different networks (internetwork connections). This suggests that while FC-based identity is reflected in local functional network properties, scaffold-based identity relies on the integration across networks, reflecting a mesoscale organization.

To directly test this hypothesis, we repeated the finger-

printing analysis using only intra-network edges or only inter-network edges for FC and scaffolds (Fig. 3C, Section IB, Table S3). For FC, intra-network edges alone supported relatively high identification rates (e.g., 73%) using Default Mode Network edges), often comparable to full-network performance. This reinforces the view that FC-based identity signatures are localized within specific brain systems. In contrast, scaffold-based identification suffered when restricted to intra-network edges. with performance dropping to 10-50% across networks (maximally 60% for the Somatomotor network). Strikingly, using only the inter-network scaffold edges restored performance to nearly full accuracy ($\sim 98\%$), highlighting again that the key individual-specific information in scaffolds lies in the links between networks. Scaffolds thus appear to capture an inherently mesoscale, distributed signature of brain organization.

Scaffold edges balance synergy and redundancy

Motivated by the mesoscale nature of scaffolds and their definition through topological cycles, we investigated the information-theoretic properties of scaffold edges. Specifically, we examined whether edges forming "holes" — the defining structures of the scaffold – carry distinct information-processing roles. Following prior works [52], we adopted the integrated information decomposition (phi-ID) framework [63] to compute subjectspecific synergy and redundancy matrices for every pair of brain regions. As expected, we found strong correlation between FC and redundancy, while synergy showed no significant relationship with FC (Fig. 4A). This confirms that FC primarily captures redundant information shared between regions.

Our topological analysis distinguishes between edges forming the scaffold borders (those participating in the H_1 generators, typically characterized by high FC weights by the construction of the filtration) and the *internal* edges spanning the topological voids, whose relatively lower weights allow the cycles to form and persist. Given the correlation between FC and redundancy and the filtration construction, scaffold border edges exhibit high redundancy. The critical question is how synergy behaves with respect to topological cycles. We hypothesize that internal edges, despite their lower correlation (and redundancy), might support greater synergistic interactions, acting as integrative "bridges" across topological holes (Fig. 4B).

To test this, we compare the synergy values of internal edges versus border edges. Indeed, we find that internal links displayed significantly higher synergy than border links (Fig. 4C). To confirm this result is not a byproduct of low correlation alone (i.e., an inverse relationship between synergy and redundancy), we compared the synergy of internal edges against three null distributions: (i) one obtained by sampling synergy values randomly from all edges, (ii) a second one obtained by sampling syn-



FIG. 3. Scaffolds rely on inter-network connectivity to enhance individual identification. (A) Heatmaps illustrating the differential power of edges for FC (top) and scaffolds (bottom). In FC, 72% of differential power is concentrated within intra-network connections (diagonal: 0.66), while 28% spans across inter-networks (off-diagonal: 0.17). In contrast, scaffolds distribute differential power more evenly, with 44% within-network (diagonal: 0.46) and 56% across networks (off-diagonal: 0.27), involving all 7 canonical resting-state networks. (B) Circular connectivity diagrams (connectograms) illustrating the location of high differential power edges in FC (top) and Scaffolds (bottom). (C) Box plots comparing identification rates (IdRate) for FCs and Scaffolds when considering only edges across different subnetworks "No subnet") versus using only intra-network edges from specific subnetworks (DA: Dorsal Attention, C: Control, DMN: Default Mode Network, L: Limbic, SA: Salience/Ventral Attention, SM: Somatomotor, V: Visual) or the full matrix. This further confirms that FC heavily relies on within-network connections, while scaffold identity relies on inter-network connections.

ergy values only from edges with FC/redundancy values matched to those of the internal links, and (iii) a third one obtained by sampling synergy values from random non-topological closed cycles (to assess whether the effect emerges from pure geometrical effects [64]). In all cases, the internal scaffold edges showed significantly higher synergy than expected by chance (Fig. 4D).

Together, these findings uncover a novel link between the topological organization of brain networks and their information-processing profiles. Scaffold representations delineate a functional architecture in which redundant communication concentrates along the borders of topological cycles, while synergistic interactions preferentially occupy the voids they enclose. This spatial segregation of redundancy and synergy suggests that scaffolds encode a balance between structured, stable pathways and flexible, integrative processing. Such a balance may underlie the emergence of individual-specific scaffold signatures, capturing distinctive aspects of brain organization that remain hidden to conventional FC analyses.

DISCUSSION

Identifying individuals based on their unique patterns of brain activity and connectivity—brain fingerprinting—has emerged as a powerful paradigm for understanding individual differences in brain organization [14, 15]. While functional connectivity (FC) derived from fMRI has proven remarkably effective for this purpose [15, 65], it primarily captures pairwise linear correlations, potentially overlooking richer, higher-order organizational principles. Here, we introduced an alternative approach leveraging topological data analysis (TDA), specifically homological scaffolds [38], to extract a unique fingerprint based on the persistent topological structure of the functional connectome. Our findings demonstrate that these topological fingerprints not only achieve near-



FIG. 4. Information-Theoretic Properties of Homological loops. (A) Diagram illustrating the relationships between FC, redundancy, and synergy across brain regions. FC is strongly positively correlated with redundancy (ρ =0.86), but only weakly with synergy (ρ =0.23). Redundancy and synergy also show a weak correlation (ρ =0.3). (B) Conceptual illustration distinguishing "boundary links" — edges that form the borders of topological cycles (solid lines) — from internal links that span the interior of the topological cycles (dashed red lines). (C) Boxplots comparing mean synergy (in bits) across different categories of links: Border (true scaffold border links), Internal (true internal links spanning topological holes), Random (randomly selected edges), Random (FC Match) (randomly selected edges matched to FC correlation values of internal links), and Random Cycle Internal (internal edges from randomly selected cycles). Internal links show significantly higher synergy than border links and random selections, suggesting a unique role in integrative information processing within topological cycles.

perfect identification accuracy, but also exhibit superior robustness and reveal distinct organizational features compared to traditional FC-based methods.

The superiority of topological over correlation-based measures aligns with emerging evidence that brain organization is fundamentally shaped by wiring constraints and communication efficiency principles [12, 66]. Topological cycles may represent optimal routing paths that minimize metabolic costs while maximizing information integration—constraints that could vary systematically across individuals due to genetic and developmental factors [67].

The central result of our study is the striking superiority of homological scaffolds over standard FC measures for individual identification. Using test-retest restingstate fMRI data from the Human Connectome Project, scaffolds achieved 100% accuracy in identifying 100 unrelated individuals, significantly outperforming full FC and thresholded FC matrices, which plateaued around 90% (Table S1, Fig. 2A). This advantage was not incidental; it proved remarkably robust across various standard preprocessing choices (Table S4), multiple brain atlases of differing resolutions and modalities (Table S2, Fig. 2C), and perhaps most notably, across substantially shortened time series lengths (Fig. 2D). The ability of scaffolds to maintain high discriminability even with data segments as short as 100 TRs (\sim 72 seconds) suggests they capture highly stable, intrinsic topological traits of an individual's functional brain architecture, potentially offering advantages for clinical applications or studies with limited scan time. This is consistent with recent studies on the temporality of brain fingerprinting, which demonstrated that bursts of "identifiability" can occur even over short time scales [68, 69] (and the additional results re-

ported in Sec. III).

But, why are scaffolds such effective fingerprints? Our results suggest that scaffolds capture fundamentally different aspects of brain organization compared to FC. When examining the edges most critical for identification (using differential power [15]), we found a clear distinction: high-DP FC edges were predominantly located within canonical brain networks (72%), particularly the default mode, control, and visual systems, consistent with previous reports emphasizing the role of frontoparietal and default mode networks in FC-based identifiability [15, 70]. In stark contrast, high-DP scaffold edges were predominantly found between different networks (56%) (Fig. 3a,b). This finding echoes recent work on higher order interactions in brain networks that emphasizes integration across canonical functional systems [71].

Furthermore, restricting the fingerprinting analysis to only intra-network edges severely degraded scaffold performance, whereas using only inter-network scaffold edges largely recovered the full scaffold's near-perfect accuracy (Fig. 3c). This indicates that the individual identifying power of scaffolds resides in their ability to capture the mesoscale distributed organization of brain function – the specific pattern of topological loops and holes formed by interactions across large-scale systems, rather than the localized connectivity strength within those systems. This finding resonates with recent work suggesting that individual differences in cognition are better predicted by between-network rather than within-network connectivity [72, 73]. The inter-network edges captured by scaffolds may reflect individual variations in network integration capacity—a key feature of cognitive flexibility and executive function [74, 75]. This mesoscale topological signature appears to be a more unique and stable individual identifier than the strength of local connections typically emphasized by FC.

Homological scaffolds emerge as a first link between topology and multivariate information theoretic descrip-Confirming previous work [52], we found FC tion. strength strongly correlated with redundant information between regional time series (Fig. 4A). The scaffold construction selects for edges involved in forming topological cycles (H_1 generators). By definition of the weighted rank filtration, edges forming the "borders" of these cycles have high FC weights and, consequently, high redundancy. The intriguing finding emerged when examining the *internal* edges that span the topological voids enclosed by these borders - edges characterized by relatively lower FC and redundancy. These internal edges exhibited significantly higher synergistic information than the border edges, and significantly higher than expected by chance, even when controlling for FC/redundancy levels (Fig. 4C). This suggests a novel link between network topology and information dynamics: the 1-dimensional "holes" revealed by persistent homology are not merely absences of strong pairwise correlation but are structured regions characterized by heightened synergistic interactions among the connections spanning them.

This interplay between redundancy and synergy within the scaffold structure offers a potential explanation for its discriminative power. The border links demonstrate high redundancy, with strong pairwise correlations indicating shared informational content. In contrast, considering pairs of links that span the internal loops reveals high synergy between them, an interaction that emerges despite the weak correlation characterizing the nodes. This spatial segregation of information modes parallels recent findings that synergistic interactions support conscious awareness and cognitive flexibility [52, 71]. The topological cycles may thus delineate functional units where redundant "backbone" connections provide stable information channels, while synergistic "bridges" enable flexible recombination of information—a balance that could be highly individual-specific due to its role in supporting each person's unique cognitive style [76].

An important practical advantage of scaffolds is their sparsity ($\approx 1\%$ of edges), making them computationally efficient for large-scale studies. This efficiency, combined with their robustness to short scan durations, positions scaffolds as a practical tool for clinical implementation where scan time is limited. While strategic study design [77], broader sampling [78], and scan duration [79] are often thought to improve predictive power, these results, in conjunction with others [31], suggest additional routes for identifying individual differences in MRI signal.

Importantly, when we repeated the analysis within each session (Fig. S3-S4), the gap between scaffolds and FC decreased notably: FC success rates and effect sizes rose to scaffold-like levels, whereas scaffold performance was unchanged. This convergence indicates that FC fingerprints derive a substantial boost from transient, state-dependent co-activation patterns that are naturally aligned within the same scan. By contrast, the scaffold signal—anchored in persistent topological cycles—appears largely insensitive to such momentary fluctuations, capturing instead trait-like organisational features that generalise across days.

Our study naturally presents some limitations. First, the scaffolds were derived from static, time-averaged FC matrices. While the sliding window analysis showed robustness over shorter intervals, exploring scaffolds derived directly from dynamic connectivity or using timeresolved topological methods could reveal further insights into transient brain states and their individual specificity. Second, the interpretation of information-theoretic measures like synergy and redundancy in the context of BOLD fMRI is still an active area of research [52, 80]. While our findings point to a compelling link between topology and synergy, the precise functional meaning requires further investigation. Third, our analysis focused on H_1 homology (loops); exploring higher-dimensional topological features $(H_2, \text{ cavities})$ might uncover additional organizational principles [39].

Future research should investigate the functional relevance of these topological fingerprints. For example,

relevant questions are: How do individual variations in scaffold structure relate to cognitive abilities, personality traits, or behaviour? Can changes in scaffold topology track learning, development, or ageing? Additionally, combining scaffolds with task-based fMRI could reveal how individual topological signatures reconfigure during cognitive demands, potentially linking stable traits to state-dependent flexibility [81, 82]. Furthermore, applying topological fingerprinting to clinical populations could be particularly fruitful. Alterations in brain connectivity are hallmarks of numerous neurological and psychiatric disorders [83, 84]. Recent work has shown altered topological properties in autism [85], schizophrenia [86], and Alzheimer's disease [87]. Our scaffold approach might be particularly sensitive to these alterations given its focus on mesoscale organization, which appears disrupted across multiple psychiatric conditions [88]. Investigating whether scaffold properties are differentially affected in these conditions could provide novel biomarkers for diagnosis, prognosis, or treatment response, potentially offering greater sensitivity than traditional FC measures due to their robustness and mesoscale nature.

In conclusion, this work introduces homological scaffolds as a potent and robust method for brain fingerprinting, significantly surpassing traditional functional connectivity approaches. By capturing the persistent topological structure of brain interactions, particularly the mesoscale arrangement of connections spanning across large-scale networks, scaffolds provide a unique window into individual brain organization. Our findings linking the topological voids within scaffolds to heightened synergistic information suggest a deeper connection between network topology and information processing principles. This work highlights the power of moving beyond pairwise correlations [34] and joining tools from topology and information theory [41, 42] to unravel the complex, individualized architecture of the human brain.

II. METHODS

A. Dataset

We utilized data from 100 unrelated healthy young adults (54 females, 46 males, mean age = 29.1 ± 3.7 years) provided in the Human Connectome Project (HCP) 900-subject data release [53, 89]. The HCP consortium curated this specific subset to ensure individuals were not family relatives, which was crucial for our study to avoid potential confounds related to family structure in identifiability analyses. All participants provided written informed consent according to the HCP protocol, which was approved by the local Institutional Review Board at Washington University in St. Louis. All experiments were performed in accordance with relevant guidelines and regulations.

We focused on the resting-state fMRI data (HCP filenames: rfMRI_REST1 and rfMRI_REST2). These were acquired in separate sessions on two different days. Each session included scans with both left-to-right (LR) and right-to-left (RL) phase-encoding directions to mitigate susceptibility distortions. For all primary analyses, we exclusively utilized the data from the LR phase-encoding direction. To validate this approach, we conducted a verification analysis where we compared the functional connectivity (FC) matrices from the LR scans alone against the mean FC matrices obtained by averaging the results from the LR and RL scans. The results were highly comparable, confirming that using only the LR data provided a reliable measure of functional connectivity representative of the full dataset. Full details on the HCP restingstate acquisition can be found in [90].

B. Preprocessing

The data were processed using the HCP minimal preprocessing pipelines [91]. The preprocessing workflow corrected for gradient distortion, head motion, and B0 field inhomogeneities. Functional data were registered to the individual's T1w structural image and then transformed into MNI152 standard space. All transformations were concatenated and applied in a single step to minimize interpolation blurring, with the final data resampled to 2mm isotropic voxels. To preserve fine-grained spatial detail, no spatial smoothing was applied. Structured noise was removed using ICA-FIX [92, 93], which identifies and removes non-neural signal components. The resulting cleaned time series from the *MNInonlinear* folder served as the primary input for all subsequent analyses.

C. Parcellation

To define network nodes, we applied several standard brain atlases to the preprocessed fMRI data in MNI space. These included: the Automated Anatomical Labeling atlas (AAL, 90 regions) [57], the Schaefer functional parcellations (using resolutions of 100 and 300 regions) [54], the Shen functional parcellation (268 regions) [59], and the Glasser multimodal parcellation (360 cortical regions, to which we added 19 subcortical areas from the HCP release, for a total of 379 regions) [60, 91]. For each participant and each parcellation, regional time series were extracted by averaging the BOLD signal across all voxels within each defined brain region. For analyses involving brain subnetworks, we utilized the 7 canonical resting-state networks defined by Yeo et al. (2011) [62], assigning each cortical parcel from the Schaefer300 atlas to one of these networks.

D. Functional Connectivity (FC)

The conventional functional connectivity, denoted as FC_{ij} , was determined by calculating the Pearson correla-

10

tion coefficient between each pairs i and j of preprocessed and denoised BOLD signal time series corresponding to a different brain region.

Pearson correlation coefficient: The correlation coefficient r_{ij} between two time series $i = \{i_1, ..., i_N\}$ and $j = \{j_1, ..., j_N\}$ is defined as:

$$r_{ij} = \frac{\sum_{k=1}^{N} (i_k - \bar{i})(j_k - \bar{j})}{\sqrt{\sum_{k=1}^{N} (i_k - \bar{i})^2 \sum_{k=1}^{N} (j_k - \bar{j})^2}}$$

where \overline{i} and \overline{j} are the means of the time series i and j, respectively, and N is the number of time points. This calculation results in a symmetric $M \times M$ FC matrix for each subject and session, where M is the number of brain regions in the chosen parcellation.

E. Homological Scaffold

To analyze the topological structure of the weighted networks, we used persistent homology [94]. Specifically, we employed the methodology described in [50].

In this framework, a weighted network is converted into a sequence of unweighted graphs, known as a *filtration*. This is achieved by considering the edge weights as the filtration parameter. We build a sequence of simplicial complexes [30, 50], specifically clique complexes, by ordering the network links by their weights in descending order. The filtration is constructed by creating a series of thresholded graphs, where each step includes all edges with a weight greater than or equal to a given threshold. As the threshold is lowered, more edges are included, and the corresponding clique complex grows.

Within this filtration, we track the birth and death of one-dimensional topological holes, which are the generators of the first homology group (H_1) . A cycle is *born* at a specific filtration value (i.e., edge weight) when a set of edges forms a closed loop. A cycle is said to *die* when it is filled in by 2-simplices (triangles), which occurs when edges are added that connect the vertices of the cycle, effectively turning the hole into a set of complete subgraphs (cliques). The birth (β_g) and death (δ_g) of each H_1 generator, g, are recorded as indices in the filtration sequence.

To summarize the topological information and understand the importance of individual links in forming cyclical structures, we constructed a *frequency homological scaffold*, as introduced in Petri et al. (2014) [38]. The frequency scaffold is a weighted graph where the edge set is composed of all edges that are part of at least one H_1 generator.

The weight of an edge e in the frequency scaffold, ω_e^f , is defined as the total number of distinct H_1 generators to which that edge belongs. This is calculated using the following formula [38]:

$$\omega_e^f = \sum_g \mathbf{1}_{e \in g}$$

where $1_{e \in g}$ is an indicator function that is equal to 1 if edge e is a part of the generator g, and 0 otherwise.

All persistent homology calculations were performed using the Ripserer.jl library [95] in the Julia programming language.

F. Fingerprinting

To assess the ability of different network representations (full FC, thresholded FC, Scaffold) to identify individuals, we implemented a fingerprinting procedure based on cross-session similarity [15]. For each subject and session, the upper triangle of the corresponding connectivity matrix was vectorized.

Let $v_{i,s1}$ be the vector for subject *i* in session 1 and $v_{j,s2}$ be the vector for subject *j* in session 2. We computed a similarity matrix **Sim**, where **Sim**_{*ij*} is the Pearson correlation coefficient between $v_{i,s1}$ and $v_{j,s2}$. To identify subject *i* from session 1, we found the subject j^* in session 2 that yielded the maximum similarity: $j^* = \arg \max_j(\mathbf{Sim}_{ij})$. Subject *i* was considered correctly identified if $j^* = i$. This process was repeated to identify subjects from session 2 based on session 1 similarity. The overall success rate was calculated as the average percentage of correctly identified subjects across both identification directions (Session $1 \rightarrow$ Session 2 and Session $2 \rightarrow$ Session 1). We provide the disaggregated values in Table S1.

G. Effect Size

To quantify the discriminability offered by different representations, we calculated the effect size [96], separating the distributions of within-subject and betweensubject similarity scores. Within-subject similarity refers to the correlation between vectors of the same subject across the two sessions (i.e., the diagonal elements of **Sim**, **Sim**_{ii}). Between-subject similarity refers to the correlations between vectors of different subjects across sessions (i.e., the off-diagonal elements **Sim**_{ij} where $i \neq j$). We used Cohen's d as the measure of effect size [97], defined as:

$$d = \frac{\mu_{within} - \mu_{between}}{\sqrt{\frac{(n_{within} - 1)s_{within}^2 + (n_{between} - 1)s_{between}^2}{n_{within} + n_{between} - 2}}}$$

where μ and s^2 are the mean and variance of the similarity scores for the within-subject and between-subject distributions, respectively, and n is the number of scores in each distribution. The denominator represents the pooled standard deviation.

H. Bootstrap Subsampling

To assess the stability and robustness of our findings regarding the fingerprinting success rates for FC and scaf-

11

folds, we employed a bootstrap subsampling procedure. We randomly sampled 80 subjects without replacement. We then calculated the fingerprinting success rate based only on this subset of 80 subjects. This subsampling process was repeated 1000 times, generating a distribution of success rates, which allowed us to estimate the variability of the metric due to subject selection.

I. Sliding Window Analysis

To investigate the temporal dynamics and stability of fingerprinting features, analyses were conducted on various shorter portions of the original 1200 TRs restingstate time series. We explored time series of the following lengths: 1100, 1000, 900, 800, 700, 600, 500, 400, 300, 200, 100, and 50 TRs. For each selected time series length, we employed a sliding window approach. Each time series was partitioned into consecutive, potentially overlapping segments of a fixed length, w. Each window was treated as a separate realization from which an independent FC matrix was derived. For each length we selected a window spanning from the first TRs t = 1 to t = w. The window was then shifted forward by a step size s. A variable sliding parameter s was used, with a minimum value of 100 TRs. This meant that for longer time series, there was an unavoidable overlap, resulting in a maximum number of windows. For example, with a time series length of 1100 TRs, only two windows of that length could be selected: one from 0 to 1100 TRs, and a second from 100 to 1200 TRs. When possible, we aimed to select six windows of each length, minimizing the overlap between them or selecting the most spread-apart windows to ensure broad coverage of the original time series. This iterative process continued until the window encompassed the final portion of the time course, resulting in a collection of connectivity matrices. To compare different window lengths, we created numerous realizations. For each realization, we randomly selected a single window per subject and calculated relevant metrics, such as the effect size and success rate, from the derived FC matrices. We repeated this random selection and metric computation process 100 times to ensure the statistical robustness of our findings.

J. Differential Power

To identify edges most critical for individual identification, we calculated the differential power (DP) for each edge, inspired by the approach of [15]. The differential power metric quantifies the contribution of an edge to fingerprinting by assessing its stability within an individual across sessions relative to its variability across the group. Conceptually, edges with high differential power exhibit connection strengths that are highly consistent for the same person across time but differ substantially between individuals. We used this metric to rank edges and select the top fraction (in this case 0.2%) deemed most important for identification for subsequent analyses, such as computing their densities across canonical brain networks.

K. Integrated Information Decomposition

To explore the information-theoretic properties underlying functional connections, particularly within the scaffold structure, we utilized integrated information decomposition based on the Partial Information Decomposition (PID) framework. Specifically, we employed the 'phi-ID' (Φ ID) formalism [63] to decompose the mutual information (MI) between the time series of pairs of brain regions (X, Y) into distinct information atoms: redundancy (I_{red}), unique information ($I_{unq,X}, I_{unq,Y}$), and synergy (I_{syn}).

$$MI(X;Y) =$$

$$I_{red}(X;Y) + I_{unq,X}(X;Y) + I_{unq,Y}(X;Y) + I_{syn}(X;Y)$$

In particular we adopted the same approach used in [52] to compute the pairwise value of synergy and redundancy from fMRI time series data based on Time Delayed Mutual Information TDMI. We computed the redundancy (I_{red}) and synergy (I_{syn}) components for each pair of regions resulting in subject-specific synergy and redundancy matrices.

L. Synergy Distribution Analysis

We specifically investigated the synergy values associated with the topological structures identified by the scaffold. We compared the distribution of synergy values for edges belonging to the scaffold "borders" (i.e., edges that are part of the frequency scaffold matrix Swith $S_{ij} > 0$) against the synergy values of "internal" edges. Internal edges were conceptually defined as those edges connecting pairs of nodes that lie within a topological "hole" (an H_1 cycle) but are not part of the cycle boundary itself. To verify that the observed higher synergy of the internal scaffold was not due to confounding factors, we compared its distribution to that of randomly selected edges defined in several ways.

- 1. Random: Synergy values were drawn from an equal number of edges selected uniformly at random from all possible edges in the connectome.
- 2. Random FC-Matched: Synergy values were drawn from randomly selected edges, but constrained such that their original FC weights (or redundancy values) were comparable to those of the actual internal edges being analyzed. This controls

12

for potential dependencies between synergy and FC/redundancy.

3. Random Cycle Internal: Synergy values were drawn after selecting a number of random cycles matching the number and lengths of original cycle. This was done to exclude the fact that the higher synergy was given by the structure of the cycle itself, rather than a specific properties of the connectome.

Statistical tests (two-sided t-tests) was used to compare

the synergy distributions between border links, internal links, and the random distributions.

ACKNOWLEDGMENTS

G.P. acknowledges partial support by ERC Consolidator Grant RUNES (Grant no. 101171380) and the MSCA Doctoral Network *BeyondTheEdge*(Grant no. 101120085). A.S. acknowledges funding from the European Union's Horizon 2020 research and innovation programme under the Marie Skłodowska-Curie grant agreement no. 101208090.

- Van Den Heuvel, M. P. & Pol, H. E. H. Exploring the brain network: a review on resting-state fmri functional connectivity. *European neuropsychopharmacology* 20, 519–534 (2010).
- [2] Sporns, O. Networks of the Brain (MIT press, 2010).
- [3] De Vico Fallani, F., Richiardi, J., Chavez, M. & Achard, S. Graph analysis of functional brain networks: Practical issues in translational neuroscience. *Philosophical Transactions of the Royal Society B: Biological Sciences* **369**, 20130521–20130521 (2014). 1406.7391.
- [4] Fornito, A., Zalesky, A. & Bullmore, E. Fundamentals of Brain Network Analysis (Academic Press, 2016).
- [5] Sporns, O. Structure and function of complex brain networks. *Dialogues in Clinical Neuroscience* 15, 247– 262 (2022).
- [6] Friston, K. J. Functional and effective connectivity in neuroimaging: a synthesis. *Human brain mapping* 2, 56–78 (1994).
- [7] Sporns, O. & Zwi, J. D. The small world of the cerebral cortex. *Neuroinformatics* 2, 145–162 (2004).
- [8] Van Den Heuvel, M. P. & Sporns, O. Rich-club organization of the human connectome. *Journal of Neuro*science **31**, 15775–15786 (2011).
- [9] Hagmann, P. et al. Mapping the structural core of human cerebral cortex. PLoS biology 6, e159 (2008).
- [10] Betzel, R. F. & Bassett, D. S. Specificity and robustness of long-distance connections in weighted, interareal connectomes. *Proceedings of the National Academy of Sciences* **115**, E4880–E4889 (2018).
- [11] Sporns, O. & Betzel, R. F. Modular brain networks. Annual review of psychology 67, 613–640 (2016).
- [12] Bullmore, E. & Sporns, O. The economy of brain network organization. Nature Reviews Neuroscience 13, 336–349 (2012).
- [13] Gratton, C. et al. Functional brain networks are dominated by stable group and individual factors, not cognitive or daily variation. Neuron 98, 439–452 (2018).
- [14] Miranda-Dominguez, O. et al. Connectotyping: model based fingerprinting of the functional connectome. PloS one 9, e111048 (2014).
- [15] Finn, E. S. et al. Functional connectome fingerprinting: identifying individuals using patterns of brain connectivity. Nature neuroscience 18, 1664–1671 (2015).
- [16] Fraschini, M. et al. The effect of epoch length on estimated EEG functional connectivity and brain network

organisation. *Journal of Neural Engineering* **13**, 036015 (2016).

- [17] Kong, W., Wang, L., Xu, S., Babiloni, F. & Chen, H. EEG Fingerprints: Phase Synchronization of EEG Signals as Biomarker for Subject Identification. *IEEE Ac*cess 7, 121165–121173 (2019).
- [18] Demuru, M. & Fraschini, M. EEG fingerprinting: Subject-specific signature based on the aperiodic component of power spectrum. *Computers in Biology and Medicine* **120**, 103748 (2020).
- [19] Miri, M. et al. Brain fingerprinting using eeg graph inference. In 2023 31st European Signal Processing Conference (EUSIPCO), 1025–1029 (IEEE, 2023).
- [20] Rodrigues, J. d. S., Ribeiro, F. L., Sato, J. R., Mesquita, R. C. & Júnior, C. E. B. Identifying individuals using fNIRS-based cortical connectomes. *Biomedical Optics Express* 10, 2889–2897 (2019).
- [21] da Silva Castanheira, J., Orozco Perez, H. D., Misic, B. & Baillet, S. Brief segments of neurophysiological activity enable individual differentiation. *Nature Communications* 12, 5713 (2021).
- [22] Sareen, E. et al. Exploring MEG brain fingerprints: Evaluation, pitfalls, and interpretations. *NeuroImage* 240, 118331 (2021).
- [23] Ricchi, I. et al. Spine-prints: Transposing brain fingerprints to the spinal cord. bioRxiv 2025–05 (2025).
- [24] Tolle, H. M. et al. The unique neural signature of your trip: Functional connectome fingerprints of subjective psilocybin experience. Network Neuroscience 8, 203– 225 (2024).
- [25] Luppi, A. I. et al. General anaesthesia decreases the uniqueness of brain functional connectivity across individuals and species. Nature Human Behaviour 1–18 (2025).
- [26] Sorrentino, P. et al. Clinical connectome fingerprints of cognitive decline. NeuroImage 238, 118253 (2021).
- [27] Romano, A. et al. The progressive loss of brain network fingerprints in Amyotrophic Lateral Sclerosis predicts clinical impairment. *NeuroImage: Clinical* 35, 103095 (2022).
- [28] Troisi Lopez, E. et al. Fading of brain network fingerprint in parkinson's disease predicts motor clinical impairment. Human Brain Mapping 44, 1239–1250 (2023).
- [29] Stampacchia, S. *et al.* Fingerprints of brain disease: connectome identifiability in alzheimer's disease. *Com*-

munications Biology 7, 1169 (2024).

- [30] Battiston, F. et al. Networks beyond pairwise interactions: Structure and dynamics. Physics reports 874, 1–92 (2020).
- [31] Cutts, S. A., Faskowitz, J., Betzel, R. F. & Sporns, O. Uncovering individual differences in fine-scale dynamics of functional connectivity. *Cerebral cortex* 33, 2375– 2394 (2023).
- [32] Betzel, R. F., Cutts, S. A., Greenwell, S., Faskowitz, J. & Sporns, O. Individualized event structure drives individual differences in whole-brain functional connectivity. *NeuroImage* 252, 118993 (2022).
- [33] Jo, Y., Faskowitz, J., Esfahlani, F. Z., Sporns, O. & Betzel, R. F. Subject identification using edge-centric functional connectivity. *NeuroImage* 238, 118204 (2021).
- [34] Battiston, F. et al. The physics of higher-order interactions in complex systems. Nature Physics 17, 1093–1098 (2021).
- [35] Clauw, K., Stramaglia, S. & Marinazzo, D. Higherorder mutual information reveals synergistic subnetworks for multi-neuron importance. arXiv preprint arXiv:2211.00416 (2022).
- [36] Varley, T. F., Pope, M., Grazia, M., Joshua & Sporns, O. Partial entropy decomposition reveals higher-order information structures in human brain activity. *Pro*ceedings of the National Academy of Sciences **120**, e2300888120 (2023).
- [37] Puxeddu, M. G., Pope, M., Varley, T. F., Faskowitz, J. & Sporns, O. Leveraging multivariate information for community detection in functional brain networks. *bioRxiv* 2024–07 (2024).
- [38] Petri, G. et al. Homological scaffolds of brain functional networks. Journal of The Royal Society Interface 11, 20140873 (2014).
- [39] Sizemore, A. E. et al. Cliques and cavities in the human connectome. Journal of computational neuroscience 44, 115–145 (2018).
- [40] Sizemore, A. E., Phillips-Cremins, J. E., Ghrist, R. & Bassett, D. S. The importance of the whole: topological data analysis for the network neuroscientist. *Network Neuroscience* 3, 656–673 (2019).
- [41] Santoro, A., Battiston, F., Petri, G. & Amico, E. Higher-order organization of multivariate time series. *Nature Physics* 19, 221–229 (2023).
- [42] Santoro, A., Battiston, F., Lucas, M., Petri, G. & Amico, E. Higher-order connectomics of human brain function reveals local topological signatures of task decoding, individual identification, and behavior. *Nature Communications* 15, 10244 (2024).
- [43] Chung, M. K., Lee, H., DiChristofano, A., Ombao, H. & Solo, V. Exact topological inference of the restingstate brain networks in twins. *Network Neuroscience* 3, 674–694 (2019).
- [44] Giusti, C., Pastalkova, E., Curto, C. & Itskov, V. Clique topology reveals intrinsic geometric structure in neural correlations. *Proceedings of the National Academy of Sciences* **112**, 13455–13460 (2015).
- [45] Chung, M. K., Bubenik, P. & Kim, P. T. Persistence diagrams of cortical surface data. In International Conference on Information Processing in Medical Imaging, 386–397 (Springer, 2009).
- [46] Pachauri, D., Hinrichs, C., Chung, M. K., Johnson, S. C. & Singh, V. Topology-based kernels with application to inference problems in alzheimer's disease. *IEEE*

transactions on medical imaging **30**, 1760–1770 (2011).

- [47] Poetto, S. & Duch, W. Classification of schizophrenia eeg recording using homological features. In 2024 International Joint Conference on Neural Networks (IJCNN), 1–8 (IEEE, 2024).
- [48] Lee, H., Kang, H., Chung, M. K., Kim, B.-N. & Lee, D. S. Persistent brain network homology from the perspective of dendrogram. *IEEE transactions on medical imaging* **31**, 2267–2277 (2012).
- [49] Lee, H., Chung, M. K., Kang, H., Kim, B.-N. & Lee, D. S. Discriminative persistent homology of brain networks. In 2011 IEEE international symposium on biomedical imaging: from nano to macro, 841–844 (IEEE, 2011).
- [50] Petri, G., Scolamiero, M., Donato, I. & Vaccarino, F. Topological strata of weighted complex networks. *PloS* one 8, e66506 (2013).
- [51] Lord, L.-D. et al. Insights into brain architectures from the homological scaffolds of functional connectivity networks. Frontiers in systems neuroscience 10, 85 (2016).
- [52] Luppi, A. I. et al. A synergistic core for human brain evolution and cognition. Nature Neuroscience 25, 771– 782 (2022).
- [53] Van Essen, D. C. *et al.* The wu-minn human connectome project: an overview. *Neuroimage* 80, 62–79 (2013).
- [54] Schaefer, A. et al. Local-global parcellation of the human cerebral cortex from intrinsic functional connectivity mri. Cerebral cortex 28, 3095–3114 (2018).
- [55] Bassett, D. S. & Sporns, O. Network neuroscience. Nature neuroscience 20, 353–364 (2017).
- [56] Amico, E. & Goñi, J. The quest for identifiability in human functional connectomes. *Scientific reports* 8, 8254 (2018).
- [57] Tzourio-Mazoyer, N. et al. Automated Anatomical Labeling of Activations in SPM Using a Macroscopic Anatomical Parcellation of the MNI MRI Single-Subject Brain. NeuroImage 15, 273–289 (2002).
- [58] Rolls, E. T., Huang, C.-C., Lin, C.-P., Feng, J. & Joliot, M. Automated anatomical labelling atlas 3. *NeuroImage* 206, 116189 (2020).
- [59] Shen, X., Tokoglu, F., Papademetris, X. & Constable, R. T. Groupwise whole-brain parcellation from restingstate fMRI data for network node identification. *NeuroImage* 82, 403–415 (2013).
- [60] Glasser, M. F. *et al.* A multi-modal parcellation of human cerebral cortex. *Nature* **536**, 171–178 (2016).
- [61] Abbas, K. *et al.* Tangent functional connectomes uncover more unique phenotypic traits. *iScience* 26, 107624 (2023).
- [62] Thomas Yeo, B. T. et al. The organization of the human cerebral cortex estimated by intrinsic functional connectivity. Journal of Neurophysiology 106, 1125– 1165 (2011).
- [63] Mediano, P. A. et al. Towards an extended taxonomy of information dynamics via integrated information decomposition. arXiv preprint arXiv:2109.13186 (2021).
- [64] Varley, T. F., Mediano, P. A., Patania, A. & Bongard, J. The topology of synergy: linking topological and information-theoretic approaches to higherorder interactions in complex systems. arXiv preprint arXiv:2504.10140 (2025).
- [65] Airan, R. D. *et al.* Factors affecting characterization and localization of interindividual differences in func-

tional connectivity using mri. Human brain mapping **37**, 1986–1997 (2016).

- [66] Avena-Koenigsberger, A., Misic, B. & Sporns, O. Communication dynamics in complex brain networks. *Nature Reviews Neuroscience* 19, 17–33 (2018).
- [67] Fornito, A., Zalesky, A. & Breakspear, M. The connectomics of brain disorders. *Nature Reviews Neuroscience* 16, 159–172 (2015).
- [68] Van De Ville, D., Farouj, Y., Preti, M. G., Liégeois, R. & Amico, E. When makes you unique: Temporality of the human brain fingerprint. *Science Advances* 7, eabj0751 (2021).
- [69] Horien, C. et al. Considering factors affecting the connectome-based identification process: Comment on waller et al. NeuroImage 169, 172–175 (2018).
- [70] Hu, D. et al. Existence of functional connectome fingerprint during infancy and its stability over months. *Journal of Neuroscience* 42, 377–389 (2022).
- [71] Varley, T. F., Pope, M., Grazia, J. F. & Sporns, O. Multivariate information theory uncovers synergistic subsystems of the human cerebral cortex. *Communications Biology* 6, 451 (2023).
- [72] Bertolero, M. A., Yeo, B. T. & D'Esposito, M. The modular and integrative functional architecture of the human brain. *Proceedings of the National Academy of Sciences* **112**, E6798–E6807 (2015).
- [73] Cohen, J. R. & D'Esposito, M. The segregation and integration of distinct brain networks and their relationship to cognition. *Journal of Neuroscience* 36, 12083– 12094 (2016).
- [74] Shine, J. M. et al. The dynamics of functional brain networks: integrated network states during cognitive task performance. Neuron 92, 544–554 (2016).
- [75] Pedersen, M., Zalesky, A., Omidvarnia, A. & Jackson, G. D. Multilayer network switching rate predicts brain performance. *Proceedings of the National Academy of Sciences* **115**, 13376–13381 (2018).
- [76] Mediano, P. A. et al. Greater than the parts: a review of the information decomposition approach to causal emergence. Philosophical Transactions of the Royal Society A 380, 20210246 (2022).
- [77] Kang, K. et al. Study design features increase replicability in brain-wide association studies. Nature 1–9 (2024).
- [78] Marek, S. *et al.* Reproducible brain-wide association studies require thousands of individuals. *Nature* **603**, 654–660 (2022).
- [79] Ooi, L. Q. R. et al. Mri economics: Balancing sample size and scan duration in brain wide association studies. bioRxiv (2024).
- [80] Stramaglia, S. et al. Synergetic and redundant information flow detected by unnormalized granger causality: Application to resting state fmri. *IEEE Transactions* on Biomedical Engineering 63, 2518–2524 (2016).
- [81] Braun, U. et al. Dynamic reconfiguration of frontal brain networks during executive cognition in humans. *Proceedings of the National Academy of Sciences* 112, 11678–11683 (2015).
- [82] Salehi, M. *et al.* There is no single functional atlas even for a single individual: Functional parcel definitions change with task. *NeuroImage* **208**, 116366 (2020).
- [83] Greicius, M. Resting-state functional connectivity in neuropsychiatric disorders. *Current opinion in neurol*ogy **21**, 424–430 (2008).

- [84] Fornito, A., Zalesky, A. & Breakspear, M. The connectomics of brain disorders. *Nature Reviews Neuroscience* 16, 159–172 (2015).
- [85] Watanabe, T. & Rees, G. Brain network dynamics in high-functioning individuals with autism. *Nature Communications* 8, 16048 (2017).
- [86] Zhang, W. et al. Disrupted brain connectivity patterns in patients with schizophrenia. Schizophrenia Research 233, 44–53 (2021).
- [87] Daianu, M. et al. Rich club analysis in the alzheimer's disease connectome reveals a relatively undisturbed structural core network. *Human Brain Mapping* 36, 3087–3103 (2015).
- [88] Crossley, N. A. *et al.* The hubs of the human connectome are generally implicated in the anatomy of brain disorders. *Brain* 137, 2382–2395 (2014).
- [89] Van Essen, D. C. *et al.* The human connectome project: a data acquisition perspective. *Neuroimage* 62, 2222– 2231 (2012).
- [90] Smith, S. M. et al. Resting-state fmri in the human connectome project. Neuroimage 80, 144–168 (2013).
- [91] Glasser, M. F. *et al.* The minimal preprocessing pipelines for the human connectome project. *Neuroim*age 80, 105–124 (2013).
- [92] Salimi-Khorshidi, G. et al. Automatic denoising of functional mri data: combining independent component analysis and hierarchical fusion of classifiers. *Neuroim*age **90**, 449–468 (2014).
- [93] Griffanti, L. et al. Ica-based artefact removal and accelerated fmri acquisition for improved resting state network imaging. Neuroimage 95, 232–247 (2014).
- [94] Edelsbrunner, H., Harer, J. et al. Persistent homologya survey. Contemporary mathematics 453, 257–282 (2008).
- [95] Čufar, M. Ripserer. jl: flexible and efficient persistent homology computation in julia. *Journal of Open Source Software* 5, 2614 (2020).
- [96] Sullivan, G. M. & Feinn, R. Using effect size—or why the p value is not enough. *Journal of graduate medical* education 4, 279–282 (2012).
- [97] Cohen, J. Statistical power analysis for the behavioral sciences (routledge, 2013).
- [98] Bertolero, M. A., Yeo, B. T. & D'Esposito, M. The modular and integrative functional architecture of the human brain. *Proceedings of the National Academy of Sciences* **112**, E6798–E6807 (2015).
- [99] Cohen, J. R. & D'Esposito, M. The segregation and integration of distinct brain networks and their relationship to cognition. *Journal of Neuroscience* 36, 12083– 12094 (2016).
- [100] Saggar, M. et al. Towards a new approach to reveal dynamical organization of the brain using topological data analysis. *Nature Communications* 9, 1399 (2018).
- [101] Betzel, R. F. et al. The modular organization of human anatomical brain networks: Accounting for the cost of wiring. Network Neuroscience 1, 42–68 (2018).
- [102] Seitzman, B. A. et al. Trait-like variants in human functional brain networks. Proceedings of the National Academy of Sciences 116, 22851–22861 (2019).
- [103] Horien, C., Shen, X., Scheinost, D. & Constable, R. T. The individual functional connectome is unique and stable over months to years. *NeuroImage* 189, 676–687 (2019).
- [104] Faskowitz, J. et al. Connectome topology of mammalian

brains and its relationship to taxonomy and phylogeny. *Frontiers in neuroscience* **16**, 1044372 (2023).

- [105] Puxeddu, M. G. et al. Relation of connectome topology to brain volume across 103 mammalian species. Plos Biology 22, e3002489 (2024).
- [106] Termenon, M., Jaillard, A., Delon-Martin, C. & Achard, S. Reliability of graph analysis of resting state fmri using test-retest dataset from the human connectome project. *Neuroimage* 142, 172–187 (2016).
- [107] Van De Ville, D., Farouj, Y., Preti, M. G., Liégeois, R. & Amico, E. When makes you unique: temporality of the human brain fingerprint. *Science advances* 7, eabj0751 (2021).
- [108] Baracchini, G. et al. covstatis: a multi-table technique for network neuroscience. arXiv preprint arXiv:2403.14481 (2024).
- [109] Wang, X. et al. Decomposition of individual-specific and individual-shared components from resting-state functional connectivity using a multi-task machine learning method. Neuroimage 238, 118252 (2021).
- [110] Mokhtari, F., Laurienti, P. J., Rejeski, W. J. & Ballard, G. Dynamic functional magnetic resonance imaging connectivity tensor decomposition: A new approach to analyze and interpret dynamic brain connectivity. *Brain connectivity* 9, 95–112 (2019).
- [111] Xiao, J. et al. A spatio-temporal decomposition framework for dynamic functional connectivity in the human brain. NeuroImage 263, 119618 (2022).
- [112] Sahoo, D., Satterthwaite, T. D. & Davatzikos, C. Hierarchical extraction of functional connectivity components in human brain using resting-state fmri. *IEEE Transactions on Medical Imaging* 40, 940–950 (2020).
- [113] Hannum, A., Lopez, M. A., Blanco, S. A. & Betzel, R. F. High-accuracy machine learning techniques for functional connectome fingerprinting and cognitive state decoding. *Human Brain Mapping* 44, 5294–5308 (2023).
- [114] Caputi, L., Pidnebesna, A. & Hlinka, J. Promises and pitfalls of topological data analysis for brain connectivity analysis. *NeuroImage* 238, 118245 (2021).

1

Supplementary Information: The Topological Architecture of Brain Identity

I. ROBUSTNESS OF IDENTIFIABILITY

To assess the identifiability of each subject across scanning sessions, we computed the Pearson correlation between the vectorized upper triangles of the connectivity matrices (or scaffolds) from Session 1 and all subjects' matrices in Session 2, and vice versa. Identification was considered correct when a subject's own scan yielded the highest similarity across sessions. We computed success rates in both directions—Session 1 \rightarrow Session 2 and Session 2 \rightarrow Session 1—and report the final identifiability as the average of the two. This bidirectional approach provides a robust estimate of cross-session consistency in individual-specific brain signatures.

A. Robustness across atlases

Across all parcellations, *homological scaffolds* consistently outperform their density-matched (1%) thresholded FC counterparts, yielding higher identification success rates and larger within- vs. between-subject effect sizes (Table S1, Fig. S1, Table S2). For the Schaefer 300 atlas (Table S1), scaffolds achieve perfect identification in both sessions, while full FC performance plateaus around 90%. These results confirm that scaffolds retain strong subject-specific features even in extremely sparse representations.

Performance across different parcellation schemes (Table S2) further supports the robustness of scaffold-based fingerprinting. While scaffolds consistently outperform 1% thresholded FC for all atlases, their absolute accuracy varies with parcellation resolution. For coarse atlases like AAL-89 and Schaefer-100, scaffold performance is relatively modest (43% and 64.5%, respectively), but this is likely due to the small number of regions limiting the number and diversity of topological cycles. As the resolution increases (e.g., Shen 268, Schaefer 300, Glasser 360), scaffold performance rapidly rises, reaching near-perfect identifiability (99.5–100%). This trend supports the idea that the discriminative power of scaffolds emerges from capturing mesoscale topological features, which become increasingly expressive at higher resolutions.

B. Localization of identifying edges

To explore where the most discriminative edges are located, we also computed identifiability success rates when restricting the analysis to within-network edges only (Table S3). As expected from prior work, FC-based identifiability is highest within canonical networks such as the Default Mode (73%) and Somatomotor (62%) systems. In contrast, scaffold-based performance drops substantially when limited to intra-network connections (e.g., 33% in DMN, 50% in SMN), confirming that their fingerprinting power does not rely on localized connectivity. Rather, as discussed in the main text, scaffold identity signatures are distributed across inter-network edges and capture subject-specific integration patterns across the entire brain. This supports a fundamentally different mechanism for individual differentiation, one that emphasizes topological integration over local coherence.

		FC full	FC 25%	FC 1%	Scaffolds
Session 1	L	89%	90%	87%	100%
Session 2	2	92%	90%	88%	100%

TABLE S1. Identifiability (success rates) for the Schaefer 300 atlas with full time series.

	FC 1%	Scaffolds
AAL89	40.5%	43%
Schaefer100	44%	64.5%
Shen268	85%	94.5%
Schaefer300	87.5%	100%
Glasser360	92.5%	99.5%

TABLE S2. Identifiability (success rates) for the various atlases of 1% thresholded FC vs. scaffolds.



FIG. S1. Similarity matrices for different atlases and different FC thresholds versus scaffolds. From top to bottom, atlas and number of regions: AAL (89), Glasser (360), Schaefer (100), Shen (268).

	\mathbf{FC}	Scaffolds
Frontoparietal	40%	21%
Default	73%	33%
DorsalAtt	25%	10%
Limbic	24%	10%
VentralAtt	31%	19%
Somatomotor	62%	50%
Visual	50%	33%

TABLE S3. Success rate performance when considering within-network connections only.



FIG. S2. Effect sizes of intra- vs inter-subject similarities. We show here results for FC (1%) and scaffolds as a function of window width, complementary to Fig. 2D.

4

II. EFFECTS OF PREPROCESSING

Table S4 reports the impact of several standard preprocessing steps on fingerprinting accuracy across different network representations. Homological scaffolds consistently outperform all functional connectivity (FC) benchmarks, including full, 25%, and 1% thresholded matrices. In most conditions, scaffolds achieve near-perfect identification (99–100%), demonstrating remarkable robustness to changes in preprocessing. Notably, scaffold performance remains high whether only LR phase encoding is used or LR/RL runs are averaged, and is largely unaffected by the inclusion or omission of global signal regression (GSR) or high-pass filtering. The only substantial performance drop is observed when low-pass filtering is omitted: in this setting, scaffold accuracy decreases to 83.5%, though it still exceeds all FC-based methods under the same condition. This suggests that low-pass filtering helps suppress high-frequency noise that may interfere with the topological cycle formation central to scaffold construction. By contrast, FC performance is more variable and systematically lower across all preprocessing variants. These findings highlight the superior stability of scaffold-based fingerprinting under typical sources of preprocessing variability, making it especially promising for robust subject identification across heterogeneous fMRI pipelines.

	FC full	FC 25%	FC 1%	Scaffolds
LR (Schaefer)	90.5%	90%	87.5%	100%
Mean LR/RL	82%	82%	81.5%	99.5%
GSR (Glasser)	94%	95.5%	92.5%	99.5%
no GSR	84.5%	82.5%	81%	99%
low pass filter (Glasser)	84.5%	82.5%	81%	99%
no low-pass	58.5%	58.5%	42.5%	83.5%
no high-pass (Schaefer)	89.5%	89.5%	88.5%	100%
high-pass	90.5%	93.5%	87%	93.5%

TABLE S4. Effects of Preprocessing effects on identifiability (success rates).

III. TRAIT VS. STATE SENSITIVITY IN SCAFFOLD AND FC REPRESENTATIONS

In addition to cross-session identifiability, we examined fingerprinting performance within a single scanning session (i.e., intrasession comparisons). Results are reported in Figures S3. In this setting, both functional connectivity (FC) and scaffold-based representations show improved success rates and larger effect sizes relative to intersession analyses, as expected due to reduced temporal variability. However, we observe that –for short timescales– the gap between FC and scaffold performance narrows substantially: the effect sizes and success rates for FC increase and become more comparable to those of scaffolds (Fig. S4).

This convergence suggests an important interpretive distinction between the two representations. The scaffold, by design, captures the persistent topological structure of functional interactions—features that are stable over time and thus likely reflect individual traits in brain organization. FC, in contrast, appears more sensitive to transient fluctuations in co-activation patterns that may reflect cognitive or physiological states present during scanning. As a result, while scaffolds maintain high identifiability across sessions (trait-like stability), FC gains relative discriminability when constrained to a single session where state-related variability is minimized.

This interpretation aligns with the emerging view that functional fingerprinting reflects a mixture of stable, trait-like architecture and more dynamic, state-driven reconfigurations. Scaffolds may therefore provide a complementary lens on brain individuality, emphasizing temporally invariant mesoscale integration patterns less influenced by momentary fluctuations in cognitive or physiological state.



FIG. S3. Intrasession fingerprinting performance. Comparison of identifiability success rates for FC and scaffold representations when computed within-session (left) and across-sessions (right).



FIG. S4. Ratio of scaffold vs FC effect sizes, intra- and inter-session as a function of the window length. We see that for short time scales within the same sessions the ratio between effect sizes for scaffolds and FC is quite small, signaling a comparable capacity of FC and scaffolds to fingerprint.