

Metabolism modulates network synchrony in the aging brain

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Brain aging is associated with hypometabolism and global changes in functional connectivity. Using functional MRI (fMRI), we show that network synchrony, a collective property of brain activity, decreases with age. Applying quantitative methods from statistical physics, we provide a generative (Ising) model for these changes as a function of the average communication strength between brain regions. We find that older brains are closer to a critical point of this communication strength, in which even small changes in metabolism lead to abrupt changes in network synchrony. Finally, by experimentally modulating metabolic activity in younger adults, we show how metabolism alone—independent of other changes associated with aging—can provide a plausible candidate mechanism for marked reorganization of brain network topology.

fMRI | aging | neurometabolism | synchrony | criticality

O ne of the most fundamental questions in neuroscience is how the familiar patterns of collective, brain-wide activity arise from the properties of the constituent neurons and their networks. Here, we study how the brain's global activity patterns change with age and how those changes might arise from the reduced metabolic activity of constituent regions.

We draw on two types of experimental evidence. First, as established using positron emission tomography, older brains show reduced glucose metabolism (1–3). Second, as established by functional MRI (fMRI), aging is associated with weakened functional connectivity (FC; i.e., reduced communication [on average] between brain regions) (4–6). Combining both observations suggests that impaired glucose metabolism may underlie changes in FC (1, 7). Supporting this link are studies showing disruptions similar to those seen with aging in type 2 diabetic subjects (8, 9).

In healthy brains, resting-state brain activity (states during which subjects are not engaged in any explicit task) alternates between segregating computations within localized functional domains and integrating this information across these domains (7, 10–13). The metabolic cost of these activities increases in proportion to the number and length of functional connections between pairs of brain regions (14), making highly connected (integrated) networks more energetically costly (10). Moreover, connections with the highest cost are the first to weaken with age (6, 7, 15, 16). Thus, it has been hypothesized that declining glucose metabolism in older brains drives the loss of high-cost (integrated) functional activities (14, 16). Yet, the relationship between energetic constraints at the level of individual regions and the apparent reorganization of the functional connectome is still not well understood.

Statistical physics, which interprets the collective properties of complex systems in terms of individual interactions between the underlying parts, provides powerful tools for understanding how brain networks reorganize under constraints (17, 18). In particular, we employ an Ising model (19–21) to describe

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how pairwise interactions between brain regions give rise to specific profiles of network synchrony, a time-dependent average of the activity over the brain (22–24). The Ising model is a simple yet powerful tool to capture ordered and disordered phases and the transitions between them. It assumes that two things matter in understanding the collective behaviors of a system of agents (brain regions in this case): the topological relationships among the agents and their strength of interaction. This perspective can provide insight into global changes in FC observed in the data by assuming that brain regions are fully connected with a single average interaction strength.

While the Ising model provides a general tool for describing the collective properties of complex systems, we adapt it to examine the specific relationship between brain aging and metabolic activity. To achieve this, we analyzed four resting-state fMRI datasets. We utilized two large-scale resting-state 3T (Tesla) fMRI datasets to identify aging related effects: the Cambridge Centre for Aging and Neuroscience Stage II Study (Cam-CAN; ages 18 to 88, N = 652) (25) and the Human Connectome Project Aging fMRI dataset (HCP-A; ages 36 to 100, N = 712) (26). To assess metabolic effects, we used two 7T fMRI datasets from the Protecting the Aging Brain fMRI data repository: a Bolus dataset (PAgB Bolus; ages 21 to 48, N = 38) consisting of a within-subject experiment in which individuals were scanned following separate administration of calorie-matched glucose and ketones (D- β -hydroxybutyrate ketone ester [D- β Hb]) and a Diet dataset (PAgB Diet; ages 21 to 43, N = 12) consisting

Significance

How do brains adapt to changing resource constraints? This is particularly relevant in the aging brain, for which the ability of neurons to utilize their primary energy source, glucose, is diminished. Through experiments and modeling, we find that changes to brain activity patterns with age can be understood in terms of decreasing metabolic activity. Specifically, we find that older brains approach a critical point in our model, enabling small changes in metabolic activity to give rise to an abrupt reconfiguration of functional brain networks.

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of an independent within-subject experiment of healthy, young adults scanned at resting state while on glycolytic and ketogenic diets (27). Ketone bodies decrease the relative free energy of adenosine triphosphate (ATP) production by approximately 27% as compared with glucose (28). The additional efficiency of ketone bodies as a cellular fuel source, observed even in healthy subjects, has been shown to increase both cardiac efficiency (28) as well as brain activity (27).

The significance of this work is threefold. First, we demonstrate that the Ising model, with a fully connected topology, provides a mechanism for how FC patterns change in qualitatively abrupt ways as a function of the average interaction between brain regions (18). Second, we establish a direct link between network synchrony and the relative frequencies of integrated (high-cost) and segregated (low-cost) brain activities (10, 14). Finally, we suggest a precise relationship between differences in FC over the life span as well as in response to changes in the brain's access to energy.

Methods

Life Span and Metabolic Neuroimaging Datasets. The Cam-CAN life span study was designed to identify neurobiological correlates of normal aging and provides a roughly uniform coverage of age groups, allowing comparison between groups as well as a wide array of behavioral measures. We excluded 10 subjects from analysis due to severe structural atrophy that precluded coregistration to a standard brain template (N = 642, 321 female).

The HCP-A study of healthy adults was comparable with the Cam-CAN dataset in terms of both its goals and design but differed in acquisition parameters and preprocessing methods. Given these differences, we used the HCP-A dataset to test the generalizability of our results with respect to aging.

The PAgB Bolus study consisted of two within-subject conditions time locked on separate days: 1) D- β Hb ketone ester bolus following a 12-h overnight fast (deltaG, TdeltaS Ltd, Thame, UK) and 2) glucose bolus following a 12-h overnight fast, breaking an overnight fast with a glucose drink (Glucose Tolerance Test Beverages; Fisher Scientific, Inc.). The D- β Hb ketone ester was weight dosed for each participant at 395 mg/kg and calorie matched between D- β Hb ketone ester and glucose (detailed documentation of the dataset may be found in ref. 27).

The PAgB Diet study scanned participants under three conditions: 1) standard diet: following their standard diet without fasting; 2) fasting: following their standard diet with an overnight (12-h) fast; and 3) ketogenic diet: following a ketogenic (high-fat, moderate-protein, low-carbohydrate [<50-g/d]) diet for 1 wk, by which point all participants were in ketosis (>0.6 mmol/L ketone blood concentration). Of these, we examined the standard (glycolytic) and ketogenic conditions to test the generalizability of our results in the context of metabolism. All conditions were conducted at resting state. Detailed documentation of the two PAgB metabolic datasets may be found in ref. 27.

MRI Acquisition. The Cam-CAN life span dataset contains multiple imaging modalities, including blood oxygen level-dependent (BOLD) MRI and magnetoencephalography (MEG). The neuroimaging experiments of the Cam-CAN study were conducted in Cambridge, United Kingdom at the Medical Research Council Cognition and Brain Sciences Unit. We focused on the resting-state BOLD echo-planar imaging (EPI) scans collected at 3T field strength over 8 min and 40 s. The BOLD EPI imaging protocol consisted of repetition time (TR) = 1,970 ms, echo time (TE) = 30 ms, flip angle = 78°, voxel size = $3 \times 3 \times 4.44$ mm, slices = 32, and number of measurements = 261. Further acquisition details can be found in ref. 25.

The HCP-A dataset was collected at four different sites while using identical equipment and parameters. Images were acquired at 3T field strength through two separate sessions for each subject. We concatenated the time series from the two sessions within each subject, resulting in a total of 26 min of resting-state fMRI data. The BOLD EPI imaging protocol consisted of TR = 800 ms, TE = 37 ms, flip angle = 52° , voxel size = $2 \times 2 \times 2$ mm, slices = 72, number of measurements = 2×488 per session, and 1,952 volumes in total. Further acquisition details can be found in ref. 29.

The two PAgB metabolic datasets (PAgB Bolus and PAgB Diet) were acquired at the Athinoula A. Martinos Center for Biomedical Imaging.

Images were collected at 7T field strength with identical parameters for the two datasets. Resting-state BOLD images were 10 min long each and involved a protocol quantitatively optimized, using a dynamic phantom, for detection sensitivity to resting-state networks (30, 31). The BOLD EPI imaging protocol consisted of TR = 802 ms, TE = 20 ms, flip angle = 33° , voxel size = $2 \times 2 \times 1.5$ mm, slices = 85, and number of measurements = 740 in each resting-state session. Further acquisition details can be found in ref. 27.

MRI Preprocessing. Aside from the HCP-A data, which were provided in an already preprocessed form, we preprocessed all fMRI data using fMRIPrep (20.0.6), which is based on Nipype 1.4.2 (32, 33). First, the T1-weighted (T1w) structural images were corrected for intensity nonuniformity and skull stripped. Brain tissue segmentation of cerebrospinal fluid (CSF), white matter, and gray matter was then performed on the brain-extracted T1w images. Subsequently, volume-based spatial normalization to one standard space (ICBM 2009c Nonlinear Asymmetric template) was carried out through nonlinear registration.

For each of the BOLD runs found per subject, images were skull stripped and motion and slice-time corrected. A deformation field to correct for susceptibility distortions was then estimated based on fMRIPrep's field mapless approach. The BOLD images were then coregistered to the T1w structural images using a boundary-based registration with nine degrees of freedom and resampled into standard space. To control for nonneuronal systemic effects associated with aging and ketosis, white matter and CSF, but not global, signals along with six motion regressors were regressed out using the nilearn package (https://nilearn.github.io).

Preprocessing of the fMRI images from the HCP-A dataset up to this point involved a different procedure. We accessed these images in an already preprocessed state, as per the steps described in refs. 34 and 35.

Finally, for all four datasets, the cleaned voxel space time series were band-pass filtered to include the neuronal frequencies (0.01 to 0.1 Hz) and were parceled into the Willard 498 functional regions of interest.

fMRI Binarization. In order to access the time-dependent network properties of our data, we first binarize the fMRI time series. This method simplifies the time series while preserving their FC patterns. In particular, the Pearson correlation $\rho(v_i, v_j)$ is widely used to estimate FC between arbitrary pairs of variables (v_i, v_i) :

$$\rho(\mathbf{v}_i, \mathbf{v}_j) = \frac{\operatorname{Cov}(\mathbf{v}_i, \mathbf{v}_j)}{\sqrt{\operatorname{Var}(\mathbf{v}_i)\operatorname{Var}(\mathbf{v}_j)}}.$$
[1]

Here, variables (v_i and v_j for example) are the nodes of a graph, and ρ is the weight of the edge between them. However, these connection strengths often change over time (36). To capture these changes, ρ is calculated over each pair of successive time points (i.e., a window length of one), reducing Eq. 1 to

$$\rho^*(\mathbf{v}_i, \mathbf{v}_j, t) = \frac{\Delta \mathbf{v}_i(t) \Delta \mathbf{v}_j(t)}{\sqrt{(\Delta \mathbf{v}_i(t))^2 (\Delta \mathbf{v}_j(t))^2}} = \hat{\mathbf{v}}_i(t) \hat{\mathbf{v}}_j(t)$$
$$BDM(\mathbf{v}_i, t) = \hat{\mathbf{v}}_i(t),$$
[2]

where \hat{v}_i and \hat{v}_j are the signs of the time derivatives of v_i and v_j , respectively, and the time-dependent correlation, ρ^* , is their product. Thus, the binarized derivative method (BDM) takes a raw signal $v_i(t)$ and binarizes it $\hat{v}_i(t)$.

While the binarization $\hat{v}_i(t)$ provides a simplified representation of the raw time series $v_i(t)$, it still preserves FC (*SI Appendix*, Fig. S1). For each subject in the life span dataset, we selected a random interval of fixed length (the window size), computed the region–region correlation matrix for both raw and binarized time series (Eq. 1), and then, computed the correlation between both FC estimates. These estimates were highly correlated, regardless of the number of time points (the averaging window) used (*SI Appendix*, Fig. S1A) and the age of the subject (*SI Appendix*, Fig. S1B) (window size = 260 TR), suggesting that BDM can be used to simplify functional signals while preserving their connectivity patterns. More broadly, the BDM approach has two key advantages over previous methods (36, 37). First, it simplifies complex, many-variable interactions in terms of dynamical patterns of binary (+1 and -1) variables. Second, it is naturally compatible with lsing models, which have been shown to be powerful tools in isolating latent relationships within networks of neurons (21, 24).

Motivation for the Ising Model. Here, we provide the Ising model of the network synchrony s and demonstrate it on a simple system of three



Fig. 1. The Ising model predicts network probabilities from interactions between its nodes. (*A*) The Ising model maps binary variables onto a fully connected network (*Left*). Each variable (i = 1, 2, ..., N) is a node with binary weight \hat{v}_i (represented by the colors red and blue), and each pair of nodes is connected by an edge with weight λ . Here, we show the example of N = 3. The value of λ (> 0) describes the average interaction strength between nodes; the larger λ is, the more likely the unknown value of \hat{v}_3 is to be similar to its neighbors (*Right*). (*B* and *C*) The probability of each network is determined by its synchrony (*s*). (*B*) Multiple graphs give the same value of synchrony. Since there are three ways to have one blue node and two red nodes, there are three different graphs that give s = 1/3 (red minus blue divided by N = 3). This degeneracy effectively triples the probability of s = 1/3. (*C*) The probability distribution of s given by the Ising model is a function of λ and degeneracy. When λ (interaction) is large, the probability that |s| = 1 is large (*Left*). However, when λ is small, degeneracy wins out, and the probability that |s| = 1/3 is large (*Right*).

interacting regions. Although first developed to describe ferromagnetism, the Ising model can be more generally derived using maximum entropy (19, 21, 24). For a set of binary nodes, such as what is provided by BDM, synchrony *s*(*t*) is defined mathematically as

$$s(t) = N^{-1} \sum_{i=1}^{N} \hat{v}_i(t),$$
 [3]

where $\hat{v}_i(t)$ is the value of the *i*th binarized signal at time *t* (24). The Ising model of the distribution of synchronies *P*(*s*) (as derived in *SI Appendix, Derivation of the Ising Model of fMRI Network Synchrony*) is given by

$$P(s) = Z^{-1} {N \choose N(1+s)/2} e^{N^2 \lambda s^2}$$
$$Z = \sum_{s=-1}^{s=-1} {N \choose N(1+s)/2} e^{N^2 \lambda s^2}.$$
[4]

Here, the normalizing constant Z is called the partition function, a familiar quantity in statistical physics (20). To account for the multiple ways a particular value of synchrony can occur, P(s) also contains the binomial prefactor, $\binom{N}{N(1+s)/2}$ (Fig. 1). In statistical physics, P(s) is equivalent to a well-known model of ferromagnets called the Curie–Weiss (or fully connected) Ising model (38). Conceptually, Var(s) expresses the average correlation across our N nodes. As a result, λ represents the average node-to-node interaction strength (over N^2 pairs) and is the basic mechanistic quantity of our model (Fig. 1 A, Left). Small values of λ describe correlational networks in which interactions between nodes are weak and in which the

node weights are independent of each other. In contrast, large values of λ describe networks in which interactions between nodes are strong and node activities are highly correlated (Fig. 1 *A*, *Right*). As described earlier, however, a given value of synchrony *s* may be obtained in many different ways (i.e., it is degenerate) (Fig. 1*B*). Therefore, when λ is small, *P*(*s*) is determined by the degeneracy, and low synchronies are most likely. Conversely, when λ is large, *P*(*s*) is determined by the interactions between nodes, and higher synchronies are more likely. In particular, as λ is varied, the relative importance of each of these terms changes. As can be seen in Fig. 1*C*, this causes *P*(*s*) to change from a bimodal (Fig. 1 *C*, *Left*) to a unimodal (Fig. 1 *C*, *Right*) distribution. For the Ising model, this shift occurs at the critical point $\lambda_c = \frac{1}{2N}$ (38). To simplify our future analysis, we will now refer only to the rescaled interaction Λ : $\Lambda = (\lambda - \lambda_c)/\lambda_c$.

Model Fitting. We next fit the Ising model to our data (Fig. 2). First, we took the fMRI signal $v_i(t)$ for each region *i* and time *t* and binarized it using BDM. The model assumes that all regions have, on average, similar FC strengths. We tested this assumption by computing the total (over all pairs) FC for each region, and we used the subject-averaged (over Cam-CAN and PAgB Bolus) FC matrix as our reference (*SI Appendix*, Fig. 52). From these signals, we selected a subset of 100 positively correlated brain regions with greatly elevated FC in younger (age < 45) compared with older (age > 64) subjects. As validation, we find that our theory is robust to this choice (*SI Appendix*, Fig. 53). For each subject, we then computed the time-dependent synchrony *s*(*t*) using the binarized fMRI signals from the 100 selected regions of the brain. We then took the histogram of *s*(*t*) for each subject to get a distribution *P*(*s*), giving the variation in synchrony per individual. This was then used to obtain Λ by fitting *P*(*s*) to the Ising model. Here, the



Fig. 2. How we obtain the Ising model parameter Λ from fMRI data. 1) The fMRI signal $v_i(t)$ from the *i*th brain region (of 100) as a function of time *t*. 2) We binarize it to give $\hat{v}_i(t)$. 3) The binarized signals are then averaged over all brain regions, giving that individual's time-dependent synchrony s(t). 4) We then histogram into P(s) the different *s* values over time to express the variations in an individual's synchrony levels. 5) We find the value of Λ that best fits P(s) for each individual. $P(\Lambda | Data)$ expresses the Bayesian posterior probability (with a uniform prior distribution over Λ) that our data P(s) were generated from an Ising model (Eq. 4) with relative interaction strength Λ .



Fig. 3. The Ising model applied to brain synchrony. Shown is the probability distribution of synchrony (s) for different values of the dimensionless quantity Λ , reflecting the distance of the actual interaction strength, λ from the critical point λ_c : $\Lambda = (\lambda - \lambda_c)/\lambda_c$. For $\Lambda < 0$ (weak interactions), there is a single unimodal population having a peak at s = 0 (blue line). For $\Lambda > 0$ (strong interactions), the population is bimodal, with peaks at $s \gg 0$ and $s \ll 0$ (orange line). Above each peak is an example network; nodes are brain regions, and colors are states (red +1, blue -1). $\Lambda = 0$ defines the critical point, where s = 0 changes from a minimum to a maximum and P(s) rapidly changes (red line). At the critical point, low- and high-synchrony networks are equally probable.

precision of the fit is a function of the Bayesian posterior distribution $P(\Lambda|\text{Data})$. We use a uniform (unbiased) prior distribution of Λ ; thus, the posterior is computed directly from the likelihood function $P(\Lambda|\text{Data}) = \prod_t P(s(t); \Lambda)$, with $P(s; \Lambda) = P(s; \lambda)$ given by Eq. 4. In practice, we will summarize this posterior by its peak (the maximum likelihood estimate) and its width (error bars). Small relative error bars indicate high confidence in the fitted values of Λ . For the Ising model, the maximum likelihood estimator for Λ satisfies

$$\frac{1}{T}\sum_{t} s(t)^2 = \sum_{s=-1}^{s=1} s^2 P(s; \Lambda),$$
[5]

where T is the number of time points used. To account for autocorrelation within the fMRI time series (which effectively reduces the number of independent samples available for each subject), we consider 0.01 like-lihood ratio error bars for Λ . This approximately corresponds to one SE with a maximum synchrony autocorrelation time of 10 TR (i.e., at minimum, 10% of the samples are uncorrelated) or three SEs if all points are independent.

Network Identification and Analysis. To establish the relationship between Λ and brain network reorganization, we isolated specific functional patterns associated with different values of synchrony. The binarized data were combined across all Cam-CAN subjects (N = 642) and classified into two categories of synchrony: low (s = 0) and high (|s| > 1/2). Time points corresponding to intermediate values of synchrony were excluded (at this stage only) in order to obtain a clear separation between groups. Network connectivity was computed (separately for each group) as the Pearson correlation between each pair of brain regions. During periods of low synchrony, functional connections were found to be sparse, favoring connections between local (segregated) networks of regions. In contrast, periods of high synchrony were characterized by dense connections (integrated) between multiple functional domains across the brain (10). Reflecting these differences, we further refer to these networks as Seg and Int, respectively.

To relate the occupation probabilities of these different networks to Λ , we decomposed the FC matrix (M_i , computed over all synchronies) of each subject *i* into a weighted contribution of Seg and Int:

$$M_i = W'_{Seg}$$
Seg + W'_{Int} Int

The weights associated to each subject (W_{seg}^{i} and W_{int}^{i}) were then computed using a least-squares fit. As a preprocessing step, we zeroed the artificial, diagonal elements and then normalized both networks (by dividing by the total correlation across all elements) to account for differences in scale. To compare these weights across subjects and different values of Λ , we only consider the ratio of these weights:

$$P_{Seg} = \frac{W_{Seg}}{W_{Seg} + W_{Int}}, \qquad P_{Int} = 1 - P_{Seg}.$$

For each subject, P_{Seg} represents the relative proportion of FC explained by the Seg network. In other words, P_{Seg} represents the fraction of time spent in segregated networks as compared with integrated networks (P_{Int}).

Results

Here, we analyze the collective properties of fMRI signals and how they change with age and metabolic activity. The collective property of our interest is the observed network synchrony *s*, a measure of the average coordination between the 100 Willard Atlas brain regions selected for our analysis (*Model Fitting*) (24, 39). In general, collective properties can often be described using mean-field models, where every component of interest is approximated as being connected to every other component with the same strength (24, 40). In addition, the Ising model is known to describe how pairwise interactions among microscopic, binary (\pm 1) elements give rise to macroscopic behaviors (20, 21). Thus, to model how the observed frequencies *P*(*s*) of different synchronies vary across experimental conditions, we utilize a mean-field Ising model, with a single average interactions strength (assumed positive) between all pairs of brain regions (Fig. 1; *SI Appendix*, Fig. S2 has further justification).

We find the value of the interaction strength, Λ , that best fits the experimentally observed synchrony values for each subject (Fig. 2). This is equivalent to choosing Λ such that the average value of s^2 predicted by the lsing model matches precisely with the empirical estimate for each subject and condition (Eq. 5 and *Methods*). In addition, we find that this model reasonably captures the full distribution of synchrony observed in the data



Fig. 4. Λ controls the balance between segregated (low *s*) and integrated (high *s*) networks. (*A* and *B*) FC (averaged over all Cam-CAN subjects) during low (*A*) and high (*B*) synchrony, visualized using the BrainNet Viewer showing the top five percentages of connections (41). (*A*) Low synchrony (*s* = 0) reflects segregation (Seg). (*B*) High synchrony (|s| > 1/2) reflects integration (Int). (*C*) The fraction of time each subject (each data point and the specific value of Λ) spends in integrated (P_{Int} ; orange) and segregated (P_{seg} ; blue) networks. $\Lambda < 0$ corresponds to large P_{seg} and small P_{Int} , while $\Lambda > 0$ corresponds to the opposite. The cross-over in *C* occurs at the critical point, $\Lambda = 0$.



Fig. 5. A significantly decreases with age and increases with a ketogenic metabolism compared with a glycolytic metabolism. (A) Each point (as well as the orange curve connecting them) reflects the median best-fit Λ values for each (of five) equal width (14-y) age groups across Cam-CAN subjects. Error bars represent the upper and lower quartiles. We used a Spearman rank permutation test $[N = 642, \rho(640) = -0.53, P = 3.0 \times 10^{-48}]$ to test significance of the nonlinear relationship between Λ and age. (B) Change in Λ for each PAgB Bolus subject after a higher-energy ketone bolus (Ket) compared with a lower-energy glucose (Glu) bolus. Error bars reflect a 0.01 likelihood ratio CI. A Wilcoxon one-sided signed rank test was used to test if ketones significantly increased Λ (N = 38, W = 242, $P = 3.2 \times 10^{-2}$). (C and D) Independent replicates of our findings for both age and metabolism using HCP-A (C) and PAgB Diet (D), respectively.

(*SI Appendix*, Fig. S4). Most importantly, the Ising model predicts the two qualitatively distinct regimes present in the data: one characterized by a unimodal distribution of synchrony and the other by a broad or bimodal distribution of synchrony.

Ising models are useful in understanding how changes in smaller-scale properties (such as the interactions between brain regions) can give rise to abrupt and qualitatively distinct collective phenomena at larger scales. Much like water at its boiling point, which discontinuously changes from liquid to vapor, these changes occur at an intermediate value of the interaction strength, called the critical point. Here, we use Λ to denote the deviation from the critical interaction strength ($\Lambda=0$) of the Ising model. Fig. 3 illustrates how the distribution of synchronies (with example brain networks shown for comparison) changes as a function of Λ , from unimodal (low synchrony, s=0; blue) when $\Lambda<0$ to bimodal (high synchrony, $|s| \gg 0$; orange) when $\Lambda>0$. Here, the distributions (which are discrete by nature) are smoothed to simplify the visualization. While both low- and high-synchrony networks are equally likely at the critical point (Fig. 3, red) ($\Lambda=0$), small changes in Λ lead to large, abrupt changes in this balance.

Different values of synchrony also correspond to different functional network patterns (Fig. 4 and *Network Identification and Analysis*). In particular, low-synchrony patterns (near s = 0) reflect short-range, domain-specific (segregated) activities (Fig. 4A). In contrast, high-synchrony patterns (|s| > 1/2) capture integration, or long-range communication between distant regions (Fig. 4B). Consequently, just as with synchrony (Fig. 3), different values of Λ change the relative time spent in segregated (P_{seg}) and integrated (P_{lnt}) networks (in Fig. 4C, each colored marker is a subject). When $\Lambda < 0$, low synchronies (i.e., segregated networks) occur more frequently, while the opposite holds when $\Lambda > 0$. In both cases, this balance rapidly shifts at the critical point, $\Lambda = 0$.

Both age and metabolism-related changes in FC can be described by changes in the region-to-region interaction strength Λ . In particular, we find that Λ significantly decreases with age (Cam-CAN, Spearman rank per-

mutation test, $P = 3.0 \times 10^{-48}$, N = 642) (Fig. 5A), suggesting that aging is associated with a marked shift from integrated toward more segregated network activities. In contrast, upon administration of a more metabolically efficient (higher-energy) ketone bolus compared with a less metabolically efficient (lower-energy) glucose bolus, Λ increases (PAgB Bolus, Wilcoxon one-sided signed rank test, $P = 3.2 \times 10^{-2}$, N = 38) (Fig. 5B) on average by about 10% ($\bar{\Lambda}_{Glu} = 0.082$ to $\bar{\Lambda}_{Ket} = 0.096$) of the decrease seen over the entire life span (28). Thus, by toggling the relative frequencies of segregated and integrated networks, Λ reflects an average cost of functional activity and as suggested by our metabolic manipulation, the energy available to the brain. As aging brains gradually become less efficient in metabolizing glucose, one way that they may conserve energy is by decreasing Λ .

The application of the Ising model to fMRI is dependent on the magnitude, rather than differences, in the functional connections measured from the data. Consequently, such an analysis could be sensitive to differences in data acquisition, preprocessing, and head motion. To assess reproducibility, we thus replicated our life span and metabolic analyses on two independent resting-state datasets: HCP-A (Spearman rank permutation test, $P = 1.2 \times 10^{-22}$, N = 712) (Fig. 5C) and PAgB Diet (Wilcoxon one-sided signed rank test, $P = 8.8 \times 10^{-2}$, N = 12) (Fig. 5D), respectively. Both sets of analyses qualitatively replicated our original findings, suggesting that our original results were robust to differences in processing pipelines and experimental design. Furthermore, we tested and found no relationship between our results and head motion (*SI Appendix*, Fig. S6).

Precisely because aging brains approach the critical point ($\Lambda = 0$), however, very small changes in the interaction strength between regions (Λ) lead to a sharp transition in the ratio of integrated to segregated networks (20, 21). Fig. 6 expresses this in terms of the probability distribution of s, now viewed from the top down. Here, we observe that as brains age, from younger (yellow; age 25 ± 7, N = 86) to older (red; age 81 ± 7, N = 130), Λ [a proxy for metabolism (14)] decreases toward the critical point (black). Consequently, aging induces a sharp reconfiguration of the distribution of synchrony.

Discussion

Our results suggest that disrupted transitioning between integrated vs. segregated networks, as seen with aging, can be quantitatively modeled by an Ising model, in which the range of synchronies observed for each individual is controlled by an average interaction strength (Λ) between pairs of brain regions. We find that Λ governs a trade-off between lower-metabolic cost, segregated and higher-metabolic cost, integrated neural activity patterns (16). Specifically, the observed link between global resting-state FC patterns and synaptic activity, as measured through the local field potential (22), suggests that Λ reflects the average synaptic connectivity across the brain. Indeed, synaptic connections weaken with age (42, 43) and are particularly vulnerable to metabolic disruptions (44–47). Utilizing subjectmatched fMRI and MEG data provided by Cam-CAN, we also find Λ to be



Fig. 6. Older brains approach a critical point, a behavior we have shown to be consistent with decreased metabolism. Shown is the probability distribution of synchrony (s) vs. Λ , viewed from the top down. At the critical point ($\Lambda = 0$; black line), peak synchrony (indicated by a white line) changes from low (s = 0) toward high (|s| > 1/2) values. Decreased metabolic activity causes Λ [through associated decreases in FC (14)] to decrease toward a critical point. Thus, aging from younger (yellow) to older (red) induces a sharp reconfiguration in the distribution of synchrony. Here, the plotted triangles correspond to the Λ values (from Cam-CAN) centered at ages 25 (younger) and 81 (older) (Fig. 5).

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significantly correlated with alpha-band MEG coherence, thus providing a potential, more direct link between our findings and disrupted neural dynamics in the aging brain (*SI Appendix*, Fig. S5) (25, 48). Consistent with our findings, we hypothesize that Λ is decreased in older brains to compensate for encroaching glucose hypometabolism. However, because older brains approach a critical point, even very small metabolic changes will result in large reconfigurations of FC.

Both aging and ketosis exert independent systemic effects that we considered in interpreting our results. For example, older subjects often have cardiovascular changes that affect neurovascular coupling (49) and thus, by extension, the BOLD response measured by fMRI. Likewise, ketosis has systemic effects, such as diuresis and thus, lowered blood pressure, as well as reduced cellular need for oxygen, all of which also could affect BOLD (50). In addition, FC is known to be affected by changes in cerebrovascular reactivity (CVR), which in turn, decreases with age (51). However, there are several reasons to suspect that these alternative mechanisms are not the sole causal influence of shifts in Λ . First, we used a hybrid global signal regression on white matter and CSF voxels, but not gray matter, to minimize the influence of nonneuronal confounds (52, 53). Second, while systemic effects might introduce spurious global correlations across the brain (54-56), they would not be expected to differentiate between integrated and segregated networks. Third, both CVR and MEG alpha-power are modulated by systemic CO₂ (a by-product of cellular metabolism), suggesting a common metabolic mechanism for these age-associated changes (51, 57). Nevertheless, experimentally dissociating metabolic from more systemic influences of aging and ketosis (e.g., by replicating our results using M/EEG or arterial spin labeling in a large-scale life span dataset that also

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experimentally modulates metabolism) is one important direction for future research.

In conclusion, the Ising model provides a mechanism for how the brain adapts to resource constraints, such as age-related glucose hypometabolism. By shifting the balance between integration and segregation toward the critical point, the brain may be able to modulate its fuel efficiency without the need to invest in new synaptic connections (7, 14, 16). Thus, toggling Λ may reflect an optimal strategy for the brain, enabling the greatest adaptation for the smallest energetic cost. At the same time, the brain's protective strategy in conserving energy may produce discontinuous trajectories for cognitive changes associated with aging.

Data Availability. Lifespan fMRI and MEG data are publicly available from Cam-CAN (25). Metabolic fMRI data are located on OpenNeuro (https://openneuro.org/datasets/ds003453 and https:// openneuro.org/datasets/ds003437). fMRI data from the HCP-A dataset are publicly available from ref. 26 and were accessed in an already preprocessed state as described in refs. 34 and 35. Additional details, including links to custom MATLAB used in the analyses of data, can be found at GitHub (https:// github.com/lcneuro/synch).

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