Metabolism modulates network synchrony in the aging brain

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Abstract

Brain aging is associated with hypometabolism and associated global changes in functional connectivity. Using 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. In particular, we find healthy brains to be poised at a critical point of this communication strength, enabling a balance between segregated (to functional domains) and integrated (between domains) patterns of synchrony. However, one characteristic of criticality is a high sensitivity to small changes. Thus, minute weakening of pairwise communication between regions, as seen in the aging brain, gives rise to qualitatively abrupt changes in 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 mechanism for global changes in synchrony.

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1 1. Significance Statement

The brain is a biological machine that utilizes chemical energy to process 2 information. However, the mechanism by which the brain adapts to resource 3 constraints is poorly understood. This is particularly relevant in the aging brain, for which the ability of neurons to utilize their primary energy source, glucose, is diminished. Here, we provide a data-driven quantitative model for how brain-wide activity patterns are controlled by resource availability. This model shows that the brain is poised at a critical point, past which even minute changes in glucose utilization cause communication across the brain to markedly re-configure. Together, our results suggest that the clinical trajectory of cog-10 nitive changes associated with aging is discontinuous and can be mediated by 11 metabolism. 12

13 2. Introduction

One 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 the constituent regions.

We draw on two types of experimental evidence. First, as established us-19 ing positron emission tomography (PET), older brains show reduced glucose 20 metabolism [1, 2, 3]. Second, as established by functional magnetic resonance 21 imaging (fMRI), aging is associated with weakened functional connectivity (FC), 22 *i.e.* reduced communication (on average) between brain regions [4, 5, 6]. Com-23 bining both observations suggests that impaired glucose metabolism may un-24 derlie changes in FC [1, 7]. Further supporting this link are studies showing 25 disruptions similar to those seen with aging in Type 2 diabetic subjects [8, 9]. 26

In healthy brains, resting-state brain activity (states during which subjects are not engaged in any explicit task) alternates between *segregating* computations to localized functional domains and *integrating* this information across

these domains [7, 10, 11, 12, 13]. The metabolic cost of these activities increase 30 in proportion to the number and length of functional connections between pairs 31 of brain regions [14], making highly-connected (integrated) networks more ener-32 getically costly [10]. Moreover, connections with the highest cost are the first to 33 weaken with age [6, 7, 15]. Thus, it has been hypothesized that declining glucose 34 metabolism in older brains drives the loss of high-cost (integrated) functional 35 activities [14]. Yet the relationship in aging brains, between energetic constraint 36 at the level of individual brain regions and the apparent re-organization of the 37 functional connectome, is still not well understood. 38

Here, we develop a generative model that describes how the probability 39 distribution of FC patterns transforms with changes in global variables (such as 40 age and metabolic activity)[16]. The approach of choice to understand how these 41 changes arise is statistical physics, which interprets the collective properties of 42 complex systems in terms of individual interactions between the underlying 43 parts [17]. In particular, we employ an *Ising model* [18, 19, 20] to describe 44 how pairwise interactions between brain regions give rise to specific profiles of 45 network synchrony, a time-dependent average of the activity over the entire 46 brain [21, 22, 23] 47

While the Ising model provides a general tool for describing the collective 48 properties of complex systems, we adapt it to examine the specific relationship 49 between brain aging and metabolic activity. To achieve this, we re-analyzed 50 two fMRI datasets. The first is the *lifespan* Cam-CAN 3T fMRI resting state 51 dataset of 636 individuals, ranging over ages 18-88 [24]. The second, in which 52 we hold age constant in order to isolate the effects of metabolic activity alone, 53 is the PAgB 7T fMRI within-subject experiment of 12 healthy young adults 54 scanned while on glycolytic and ketogenic *diets* [25]. Ketone bodies decrease 55 the relative free energy of ATP production by 27% as compared to glucose [26]. 56 This additional efficiency of ketone bodies as a metabolite, observed even in 57 healthy subjects, has been shown to increase both cardiac efficiency [26] as well 58 as brain activity [25]. 59

⁶⁰ The significance of this work is three-fold. First, in contrast to the tools

⁶¹ commonly used to study fMRI networks, our approach provides a predictive ⁶² mechanism for how FC patterns change, in qualitatively significant ways, as ⁶³ a function of the average interaction between brain regions [16]. Second, we ⁶⁴ establish a direct link between network synchrony and the relative frequencies of ⁶⁵ integrated (high-cost) and segregated (low-cost) brain activities [10, 14]. Finally, ⁶⁶ we illustrate a precise relationship between differences in FC over the lifespan ⁶⁷ as well as in response to changes in the brain's access to energy.

68 3. Methods

⁶⁹ 3.1. Lifespan and metabolic neuroimaging datasets

To identify how the collective features of fMRI change across the lifespan, we 70 analyzed a large-scale 3T fMRI dataset: the Cambridge Centre for Ageing and 71 Neuroscience stage II (Cam-CAN: ages 18-88, N = 636) [24]. The Cam-CAN 72 study was designed to identify neural correlates of normal aging and provides 73 a roughly uniform coverage of age groups, allowing comparison between groups 74 as well as a wide array of behavioral measures. While the functional MRI imag-75 ing of Cam-CAN stage II included both task and resting state data, we used 76 only resting state data, for which most regions of the brain have roughly similar 77 statistical properties (see Supplementary Fig. 2). To relate these changes to en-78 ergy in the brain, we additionally analyzed 7T fMRI data from the Protecting 79 the Aging Brain (PAgB) database [25]. In a within-subjects experiment, young 80 healthy adults ($N = 12, \mu_{age} = 28 \pm 6.73$ years; 4 female) were scanned at 81 resting state under two conditions: (1) glycolytic, following their standard diet, 82 without fasting; and (2) ketogenic, following a high-fat, moderate-protein, low-83 carbohydrate (< 50 g/day) diet for one week, by which point all participants 84 were in ketosis (> 0.6 mmol/L ketone blood concentration). For details on the 85 glycolytic and ketogenic dietary regimes, as well as validation of their blood 86 values and neurobiological effects as comparable to calorie-matched administra-87 tion of glucose and D- β -hydroxybuterate, see previous work [25]. Studies were 88 approved by the Institutional Review Boards of Cambridge University and Mas-89 sachusetts General Hospital/Partners Healthcare, respectively; all participants 90

⁹¹ provided informed consent.

92 3.2. MRI acquisition

The Cam-CAN lifespan dataset includes multiple imaging modalities (T1 93 and T2-weighted images, diffusion-weighted images, BOLD EPI images during 94 tasks of three varying levels of cognitive demand, MEG images during two sep-95 arate cognitive loads and magnetisation-transfer images). Of these, the resting 96 state BOLD EPI fMRI was the focus of our analysis (full dataset documentation 97 at [24]). The Cam-CAN functional imaging was done at 3T field strength over 8 98 min 40 s. The neuroimaging experiments of Cam-CAN study were conducted in 99 Cambridge, UK at the Medical Research Council Cognition and Brain Sciences 100 Unit (MRC-CBSU). Specifics of the BOLD EPI imaging protocol included: TR 101 $= 1970 \text{ ms}, \text{TE} = 30 \text{ ms}, \text{flip angle} = 78^{\circ}, \text{voxel size} = 3 \times 3 \times 4.44 \text{ mm}, \text{slices} =$ 102 32, number of measurements = 261. The PAgB metabolic dataset was acquired 103 at ultra-high-field (7T) field strength at the Athinoula A. Martinos Center for 104 Biomedical Imaging. Imaging included whole brain BOLD, field map, and T1-105 weighted structural (MEMPRAGE) images. BOLD images were acquired using 106 a protocol quantitatively optimized, using a dynamic phantom, for detection-107 sensitivity to resting state networks [27]: SMS slice acceleration factor = 5, R 108 = 2 acceleration in the primary phase encoding direction (48 reference lines) 109 and online GRAPPA image reconstruction, TR = 802 ms, TE = 20 ms, flip 110 angle = 33° , voxel size = $2 \times 2 \times 1.5$ mm, slices = 85, number of measurements 111 = 740 in each resting state session, for a total acquisition time of 10 minutes. 112 Field map images were acquired using the following parameters: TR = 723 ms, 113 $TE1 = 4.60 \text{ ms}, TE2 = 5.62 \text{ ms}, \text{ flip angle} = 36^{\circ}, \text{ voxel size} = 1.7 \times 1.7 \times 1.5$ 114 mm, slices = 89, for a total acquisition time of 3 min 14 s. The whole-brain 115 T1-weighted structural volumes were acquired using a conventional multi-echo 116 MPRAGE (MEMPRAGE) sequence with 1 mm isotropic voxel size and four 117 echoes with the following protocol parameters: TE1 = 1.61 ms, TE2 = 3.47118 ms, TE3 = 5.33 ms, TE4 = 7.19 ms, TR = 2530 ms, flip angle = 7° , with R = 119 2 acceleration in the primary phase encoding direction (32 reference lines) and 120

online GRAPPA image reconstruction, for a total volume acquisition time of 6
 min 3 s.

123 3.3. MRI pre-processing

Lifespan dataset pre-processing was conducted in the FMRIB Software Li-

¹²⁵ brary (FSL; https://fsl.fmrib.ox.ac.uk/fsl/fslwiki/) and FreeSurfer

(https://surfer.nmr.mgh.harvard.edu/): anatomical images were skull-stripped 126 using FreeSurfer and co-registered to Montreal Neurological Institute (MNI) 127 templates and mean functional images using FLIRT (part of FSL). Functional 128 images were motion and fieldmap-corrected (using MCFLIRT and epidewarp), 129 brain-extracted (using BET), and co-registered to MNI templates using trans-130 formations learned through the anatomical image. Motion parameter as well as 131 tissue segmentation-extracted white-matter and CSF confounds (using FAST) 132 were regressed out at ROI-level time series extraction stage using nilearn package 133 (https://nilearn.github.io) [28]. Metabolic dataset pre-processing used Statisti-134 cal Parametric Mapping 12 135

(SPM12; https://www.fil.ion.ucl.ac.uk/spm/software/spm12/) was used as in 136 our previous studies conducted at the same acquisition parameters [29], [25]. 137 Anatomical images (MEMPRAGE) were normalized to MNI templates using 138 unified segmentation and registration. Images of each individual participant 139 were realigned to account for head movements, and field map-corrected (using 140 epidewarp.fsl) for geometric distortions caused by the magnetic field inhomo-141 geneity. Following normalization, structural images were probabilistically seg-142 mented into three tissues: grey matter, white matter, and cerebral spinal fluid. 143 We did not apply spatial smoothing or global signal regression to pre-processing 144 of either dataset. For all datasets, voxelwise data were parceled into the Willard 145 498 functional regions of interest (ROI) [30] corresponding entirely to grey mat-146 ter voxels. 147

148 3.4. Ising model

Here we use the principle of maximum entropy [18, 20, 23] to build the minimally biased probability distribution of N binary (+1 or -1) node weights,

> ¹⁵¹ { \hat{v}_i } satisfying fixed constraints on the mean (0) and variance, Var(s) of the ¹⁵² global property, $s({\hat{v}_i}) = N^{-1} \sum_{i=1}^N \hat{v}_i$ (synchrony). This is given by [23]:

$$P(\{\hat{v}_i\}) = Z^{-1} e^{N^2 \lambda s(\{\hat{v}_i\})^2}$$
(1)

where Z is the partition function and normalizes the distribution. λ represents 153 the average node-to-node interaction strength and is the basic mechanistic quan-154 tity of our model (Fig. 1a, left). Small values of λ describe networks in which 155 interactions between nodes are weak and in which the node weights are inde-156 pendent of each other. In contrast, large values of λ describe networks in which 157 interactions between nodes are strong and node activities are highly correlated 158 (Fig. 1a, right). A given value of synchrony s may be obtained in many 159 different ways; *i.e.*, it is *degenerate* (Fig. 1b). In other words, since there 160 are $\binom{N}{N(1+s)/2}$ different ways to have $s = N^{-1} \sum_i \hat{v}_i$, we find that the total 161 probability P(s) of different synchronies is: 162

$$P(\{\hat{v}_i\}) = Z^{-1} \binom{N}{N(1+s)/2} e^{N^2 \lambda s(\{\hat{v}_i\})^2}$$
(2)

Therefore, when λ is small, P(s) is determined by the degeneracy and low syn-163 chrony is most probable. Conversely, when λ is large, P(s) is determined by the 164 interactions between nodes and high synchrony is most probable. In particular, 165 as λ is varied, the relative importance of each of these terms changes. As can be 166 seen in Fig. 1c, this causes P(s) to change from a bimodal (left) to a unimodal 167 (right) distribution. The critical point, λ_c , is the value of λ where this shift 168 happens (i.e. when these two contributions are balanced). Using the standard 169 approximation of the binomial coefficients, P(s) becomes: 170

$$P(s) \approx Z^{-1} \binom{N}{N/2} e^{[\lambda - \frac{1}{2N}]s^2}$$
(3)

Conceptually, when $\lambda < \lambda_c$, P(s) opens downwards like a Gaussian; s = 0is most probable. However, when $\lambda > \lambda_c$, P(s) opens upwards and large values of s (both positive and negative) are probable. When N = 498 (the number of regions), we find that this critical point is $\lambda_c = \frac{1}{2N} = 1.004 \times 10^{-3}$, coinciding with the observed transition between unimodal and bimodal synchrony (**Fig.**

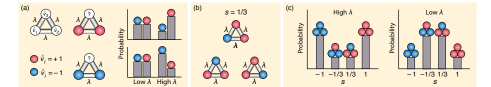


Figure 1 The Ising model predicts network probabilities from interactions between its nodes. (a) The Ising model maps binary variables onto a fully-connected network (left). Each variable (i = 1, 2, ..., N) is a node with binary weight \hat{v}_i (represented by the colors red and blue), and each pair 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,c) The probability of each network is determined by its synchrony (s). (b) Multiple graphs give the same value of synchrony. Since there are 3 ways to have 1 blue node and 2 red nodes, there are 3 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). But, when λ is small, degeneracy wins out and the probability that |s| = 1/3 is large (right).

177 **1c**). To simplify our analysis, now refer to the rescaled interaction Λ : $\Lambda = (\lambda - \lambda)/\lambda$

178 $(\lambda - \lambda_c)/\lambda_c$.

179 3.5. fMRI binarization

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

$$\rho(X,Y) = \frac{Cov(X,Y)}{\sqrt{Var(X)Var(Y)}}.$$
(4)

Here variables (X and Y 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 [31]. Thus, we calculate ρ over each pair of successive time points, reducing Eq. 4 to:

$$\rho^*(X, Y, t) = \frac{\Delta X(t) \Delta Y(t)}{\sqrt{(\Delta X(t))^2 (\Delta Y(t))^2}} = \hat{X}(t) \hat{Y}(t)$$
$$BDM(X, t) = \hat{X}(t)$$
(5)

where \hat{X} and \hat{Y} are the signs of the time derivatives of X and Y respectively 189 and the time-dependent correlation, ρ^* , is their product. This procedure takes 190 our original time series X(t) and produces a simplified, binarized time series 191 $\hat{X}(t)$ (Binarized Derivative Method, BDM). By computing these binarized val-192 ues for long periods of time, we can ask questions about how the probabilities 193 of different sequences (in time) and patterns (over regions) change with dif-194 ferent conditions (such as with age and diet). As validation of this method, 195 we find that this simplified representation preserves fMRI FC patterns across 196 time (Supplementary Fig. 1a) and for different subjects (Supplementary Fig. 197 1b). This approach has two key advantages over previous methods [31, 32]. 198 First, it simplifies complex, many-variable interactions in terms of dynamical 199 patterns of binary (+1 and -1) variables. Second, it is naturally compatible 200 with Ising-like models, which have been shown to be powerful tools in isolating 201 latent relationships within networks of neurons [20, 23]. 202

203 3.6. Model fitting

We then fit the Ising model to our data (Fig. 2). First, we took the fMRI 204 signal $v_i(t)$ for each region i and time t and binarized it using BDM. The model 205 assumes that all nodes have, on average, similar FC strengths. We tested this 206 assumption by computing the total (over all pairs) FC for each region, and we 207 used the subject-averaged (over all diets and ages) FC matrix as our reference 208 (Supplementary Fig. 2). From these signals, we found that most nodes are 209 primarily positively correlated, while a few nodes were primarily negatively cor-210 related with other nodes. For the latter, we flipped $(\hat{v}_i \rightarrow -\hat{v}_i)$ for these regions 211 only in order to satisfy the assumptions of our Ising model (Supplementary Fig. 212 2). For each subject, we then computed the time-dependent synchrony s(t) (each 213 TR is a time point) using the binarized fMRI signals from all (498) regions of the 214 brain. We then took the histogram of s(t) for each subject to get a distribution 215 P(s), giving the variation in synchrony per individual. This was then used to 216 obtain Λ by fitting P(s) to the Ising model Eq. 2. This fit is expressed by the 217 Bayesian posterior distribution $P(\Lambda | \text{Data})$, which captures the relative quality 218 of of our model. We use a uniform (unbiased) prior distribution of Λ ; thus the 219 posterior is computed directly from the likelihood function $\mathcal{L}(\Lambda|\text{Data})$ of our 220 Ising model Eq. 2. In practice, we will summarize this posterior by its peak 221 (the maximum likelihood estimate) and its width (error bars). As fMRI signals 222 are auto-correlated, the data (s(t)) are not fully independent. To compensate 223 for this effect, we consider conservative (0.01 likelihood ratio) error bars for Λ . 224

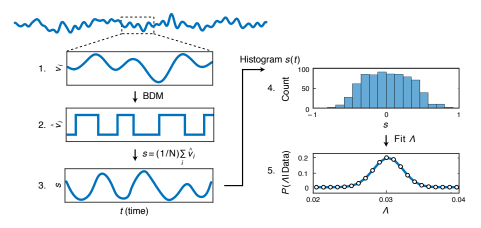


Figure 2 How we obtain the Ising model parameter Λ from fMRI data. [1] shows the fMRI signal $v_i(t)$ from the *i*th brain region (out of 498), as a function of time *t*. [2] We binarize it, to give $\hat{v}_i(t)$. [3] The binarized signals are then averaged over all brain regions, giving that individual's time-dependent synchrony s(t). [4] We then histogram into P(s) the different *s* values over time [4], to express the variations in an individual's synchrony levels. [5] We then find the value of Λ that best fits P(s) for each individual. $P(\Lambda|\text{Data})$ expresses the Bayesian posterior probability (with a uniform prior distribution over Λ) that our data P(s) was generated from an Ising model (Eq. 2) with relative interaction strength Λ .

225 4. Results

To interpret our fMRI data, we developed a generative biophysical approach based on a network *Ising model* [19, 20]. Widely used in physics, the Ising model describes how pairwise interactions among microscopic, binary (± 1) elements give rise to macroscopic behaviors, including correlations ([19], **Fig. 1**). In other words, the Ising model allows us to describe time-dependent variability (probabilities) of different brain states for each subject.

We are particularly interested in the collective (i.e. regionally-averaged) 232 properties of brain activity. In general, collective properties can often be de-233 scribed using mean-field models, where every component of interest is approx-234 imated as being connected to every other component with the same strength 235 [23, 33]. Here the collective property of interest is the observed network syn-236 chrony, s, or the average activity across the 498 Willard Atlas brain regions 237 measured in fMRI experiments [30, 23]. The probability distribution of dif-238 ferent synchronies can then be described by a mean-field Ising model, with a 239

> single average interaction strength (assumed positive) between all pairs of brain 240 regions (see Supplementary Figs. 2 and 3 for further justification). To explore 241 this model, we find the value of the interaction strength, Λ , that best fits the 242 experimentally observed synchrony values for each subject (Fig. 2). Thus, each 243 value of s corresponds to the degree of consensus of a particular network pro-244 duced by the best-fitting Ising model [23]. As further validation for our model, 245 we find that the Ising model, regardless of age and diet, correctly captures the 246 kurtosis of P(s), a higher-order feature that cannot be generally predicted from 247 correlations alone (Supplementary Fig. 3). 248

> Ising models are useful in understanding how changes in smaller-scale prop-249 erties (such as the interactions between brain regions) can give rise to abrupt 250 and qualitatively distinct collective phenomena at larger scales. Much like water 251 at its boiling point, which discontinuously changes from liquid to vapor, these 252 changes occur at an intermediate value of the interaction strength, called the 253 critical point. Here we use Λ to denote the deviation from the critical interaction 254 strength $(\Lambda = 0)$ of the Ising model. Figure 3 illustrates how the distribution 255 of synchronies (with example brain networks shown for comparison) changes as 256 a function of Λ , from unimodal (low synchrony, s = 0, blue) when $\Lambda < 0$ to 257 bimodal (high synchrony, $s = \pm 1$, orange) when $\Lambda > 0$. While both low and 258 high synchrony networks are equally likely at the critical point (Fig. 3, red, 259 $\Lambda = 0$, small changes in Λ lead to large, abrupt changes in this balance. 260

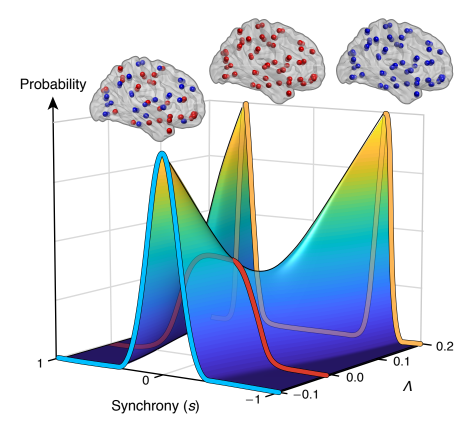


Figure 3 The Ising model applied to brain synchrony. Shown is the probability distribution of different values of synchrony (s) for different values of the dimensionless quantity Λ , reflecting the distance of the actual interaction strength, λ from the critical point λ_c : $\Lambda = (\lambda - \lambda_c)/\lambda_c$. 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 a 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.

> To establish the relationship between synchrony and the occupation proba-262 bilities of specific functional networks, we separately computed the inter-subject 263 average FC matrices during periods of low and high synchrony. During peri-264 ods of low synchrony, functional connections are found to be sparse, favoring 265 connections between local (segregated) networks of regions (Fig. 4a, Seg). In 266 contrast, high synchrony networks are typified by dense connections (integrated) 267 between multiple functional domains across the brain (Fig. 4b, Int) [10]. Con-268 sequently, just as with synchrony (Fig. 3), different values of Λ change the 269 relative time spent in segregated (P_{Seg}) and integrated (P_{Int}) networks (Fig. 270 4c, $R^2 > 0.9$, sigmoidal fit not shown, each colored marker is a subject), inde-271 pendent of age or diet. The time spent in each pattern was computed as the 272 similarity of each subject's FC to the extracted patterns, Int and Seg. When 273 $\Lambda < 0$, low synchronies (i.e. segregated networks) occur more frequently, while 274 the opposite holds when $\Lambda > 0$. In both cases, this balance rapidly shifts at the 275 critical point, $\Lambda = 0$. 276

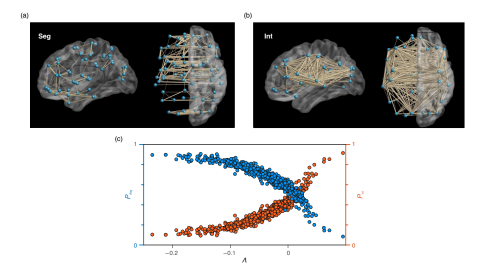


Figure 4 Λ controls the balance between segregated (low s) and integrated (high s) networks. (a,b) Inter-subject average functional connectivity (Pearson Correlation) during low (a) and high (b) synchrony, visualized using the BrainNet Viewer showing the top 10 % of connections [34]. (a) Low synchrony (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 their specific value of Λ) spends in integrated (P_{Int} , orange) and segregated (P_{Seg} , blue) networks. Time spent was calculated from a bivariate regression of the functional connectivity (here over all s, from each subject) with the patterns, Seg and Int. $\Lambda < 0$ 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, $\Lambda = 0$.

Changes in FC with both age and diet can be described by changes in the 278 region-region interaction strength Λ . In particular, we find that Λ significantly 279 decreases with age $(p = 1.7 \times 10^{-38}, N = 636,$ Fig. 5a), suggesting that ag-280 ing is associated with a marked shift from integrated towards more segregated 281 network activities. But, upon switching from a lower-energy glycolytic to a 282 higher-energy ketogenic diet, Λ increases $(p = 1.2 \times 10^{-3}, N = 12, \text{Fig. 5b})$ 283 by about 25% to 50% of the decrease seen over the entire lifespan. Thus, by 284 toggling the relative frequencies of segregated and integrated networks, Λ re-285 flects an average cost of functional activity and, as suggested by our metabolic 286 experiment, the amount of energy available to the brain. Thus one way the 287 brain may conserve energy when this amount of energy available is decreased, 288 such as through aging, is by decreasing Λ . 289

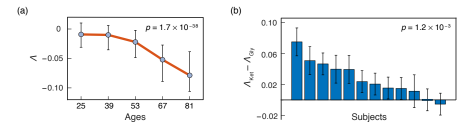


Figure 5 Λ significantly decreases with age (a, $p = 1.7 \times 10^{-38}$) and increases on the higherenergy, ketogenic diet (b, $p = 1.2 \times 10^{-3}$). (a) Each point (as well as the orange curve connecting them) reflects the median best-fit Λ values for each (of 5) equal width (14 years) age groups. Error bars represent the upper and lower quartiles. We used a Spearman-rank Permutation test (N = 636, $\rho(634) = -0.48$) to test significance of the nonlinear relationship between Λ and age. (b) Change in Λ for each subject (N = 12, W = 3) when switching from a lower-energy glucose (glycolytic, Gly) to higher-energy ketone (Ket) metabolism. Error bars reflect a 0.01 likelihood ratio confidence interval. A Wilcoxon 1-sided signed rank test (N = 12) was used to test if ketones significantly increased Λ .

But why would a small change in Λ lead to the dramatic changes in FC seen 291 in older age? Precisely because young healthy brains are poised at the critical 292 point $(\Lambda = 0)$, very small changes in the interaction strength between regions 293 lead to a sharp transition in the ratio of integrated to segregated networks 294 [19, 20]. Figure 6 expresses this in terms of the probability distribution of 205 s, now viewed from the top-down. Here younger brains (green, age 25 ± 7) 296 are near the critical point (black), allowing them to access both high and low 297 synchrony networks. But as Λ (a proxy for energy availability, [14]) decreases, 298 such as observed in older brains (yellow, age 81 ± 7), the probabilities of higher 299 synchrony networks quickly fall to 0. 300

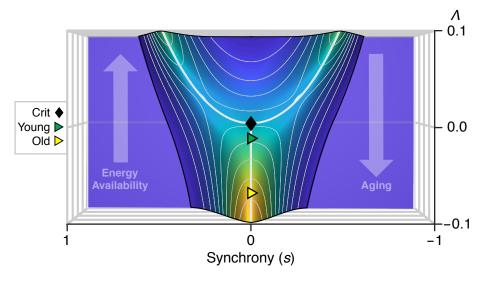


Figure 6 Younger brains are poised at a critical point; this is disrupted by decreasing energy availability. Shown is the probability distribution of synchrony (s) vs Λ , viewed from the top-down. At the critical point ($\Lambda = 0$, black), peak synchrony (indicated by a white line) changes from low (s = 0) towards high (s < 0 and s > 0) values. Near this transition, such as seen in younger brains (Age 25 ± 7 , N = 85, green), both low and high synchrony networks can be accessed. Reducing energy availability causes Λ (through associated decreases in FC, [14]) to decrease. Older brains (Age 81 ± 7 , N = 121, yellow) have smaller Λ and only access low synchrony networks. The plotted triangles correspond to the Λ values centered at ages 25 (Young) and 81 (Old) (**Fig. 5**).

301 5. Discussion

Our results suggest that the principal functional changes associated with 302 aging, in terms of network synchrony, are controlled by an average interaction 303 strength (Λ) between pairs of brain regions. Crucially, unlike graph theoretic 304 features normally used to describe such data [16], Λ encodes how the aging brain 305 rewires. We have also shown that Λ governs a trade-off between low-cost, seg-306 regated and high-cost, integrated activity patterns. Furthermore, as suggested 307 by our findings, we hypothesize that Λ is decreased in older brains to com-308 pensate for glucose hypometabolism. But, because younger brains are poised 309 near a critical point, this compensation results in sharp changes in functional 310 connectivity. 311

It is important to note that aging and ketosis each exerts independent sys-312 temic effects that need to be considered in interpreting the results. For exam-313 ple, older subjects often have cardiovascular changes that affect neurovascular 314 coupling[35] and thus, by extension, the blood oxygen level dependent (BOLD) 315 response measured by fMRI. Likewise, ketosis has systemic effects, such as di-316 uresis and thus lowered blood pressure, as well as reduced cellular need for 317 oxygen, all of which also could theoretically affect BOLD. However, there are 318 several reasons to suspect that these alternative mechanisms are not the sole 319 causal influence of shifts in Λ . First, to minimize the primary cause of neu-320 rovascular confounds, the lifespan dataset specifically excluded individuals with 321 cardiovascular disease, including cerebral ischeaemia [36]. Moreover, while the 322 impact of arteriosclerosis in reducing the dynamic range of BOLD could reduce 323 signal/noise and therefore reduce the strength of measured connections over-324 all, it would not discriminate between integrated versus segregated networks 325 and the transitions between them. Second, shifts in λ were observed not only 326 in the aging dataset, but also in the dietary dataset, the latter of which in-327 328 cluded only younger individuals and thus eliminated systemic aging effects as a variable. Third, systemic (non-metabolic) effects of ketosis, such as reduced 329 cerebral blood pressure and reduced need for oxygen, should decrease BOLD ac-330

> tivation, while we have previously shown ketosis to increase BOLD activation, both in our dietary dataset as well as an independent dataset in which ketosis was achieved by administering exogenous D- β -hydroxybuterate[25]. Nevertheless, dissociating metabolic from more systemic influences of aging and ketosis is one important direction for our future research.

> The metabolic cost of connectivity is known to reflect both signaling along 336 axons as well as between synapses. As such, Λ may reflect the average synaptic 337 connectivity across the brain, as suggested by recent evidence linking global 338 resting state fMRI fluctuations to synaptic activity [21]. Indeed, synaptic con-339 nections weaken with age [37, 38] and are particularly vulnerable to metabolic 340 disruptions [39, 40, 41, 42]. However, the fact that age was associated with a 341 reduced probability of integrated activities (with longer connections) in favor 342 of segregated activities (with shorter connections) suggests that the metabolic 343 cost of axon conductance may also play a key role. Long-range connections 344 are known to be disproportionately diminished not only with age [15] but also 345 epilepsy [43], the latter of which commonly shows improvement with ketosis. 346

> That brains at their presumed peak of functionality should be poised so close 347 to a critical point of synchrony may reflect an evolutionary selective advantage. 348 Criticality is not only a widely-observed feature of neural activity [44, 45, 46], 340 but also enables the broadest range of functional patterns while also achieving 350 maximum sensitivity to external drivers (e.g. sensory stimuli) [19, 47]. Some 351 recent work suggests that signatures resembling criticality may be generic fea-352 tures of systems with many unobserved variables [48]. However, if this were the 353 case, one would find these signatures in both younger and older brains, which 354 is not consistent with our findings. 355

> In conclusion, the Ising model provides a data-driven generative model for how the brain adapts to resource constraints, such as progressive glucose hypometabolism in aging brains. By simply shifting the balance between integration and segregation away from the critical point, the brain is able to modulate its fuel efficiency without the need to invest in new synaptic connections [7, 14]. Thus toggling Λ reflects an optimal strategy for the brain, enabling

362 the smoothest adaptation for the smallest energetic cost. At the same time,

- $_{\tt 363}$ $\,$ the brain's protective strategy in conserving energy may produce discontinuous
- ³⁶⁴ trajectories for cognitive changes associated with aging, both in terms of di-
- ³⁶⁵ minished sensitivity to sensory stimuli (as predicted by shifts from criticality)
- ³⁶⁶ as well as cognitive processing associated with flexibility in switching between
- ³⁶⁷ both segregated and integrated networks.

368 Data and code availability

- Lifespan fMRI data are publicly available from Cam-CAN [24]. Metabolic
- ³⁷⁰ fMRI data are located at Data Archive for the Brain Initiative (DABI:
- ³⁷¹ https://dabi.loni.usc.edu/explore/project/42) in the Protecting the Aging Brain
- ³⁷² (PAgB), Project 1926781 repository. Additional details (including links to cus-
- ³⁷³ tom MATLAB and Python codes used in the processing and analyses of data)
- ³⁷⁴ can be found at http://www.lcneuro.org/software-and-instrumentation.
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380 CRediT authorship contribution statement

LRMP designed the experiments. AA pre-processed the data. CW did the modeling and designed the methods used. CW, LRMP, and KD wrote and revised the paper.

- 384 Declarations of interest
- 385 None.

386 References

³⁸⁷ [1] Lisa Mosconi. Glucose metabolism in normal aging and alzheimers disease:

methodological and physiological considerations for pet studies. Clinical
 and translational imaging, 1(4):217–233, 2013.

³⁹⁰ [2] Manu S Goyal, Andrei G Vlassenko, Tyler M Blazey, Yi Su, Lars E Couture,

³⁹¹ Tony J Durbin, Randall J Bateman, Tammie L-S Benzinger, John C Morris,

- ³⁹² and Marcus E Raichle. Loss of brain aerobic glycolysis in normal human
- aging. Cell metabolism, 26(2):353–360, 2017.

394	[3]	Christian-Alexandre Castellano, Scott Nugent, Nancy Paquet, Sébastien
395		Tremblay, Christian Bocti, Guy Lacombe, Helene Imbeault, Eric Turcotte,
396		Tamas Fulop, and Stephen C Cunnane. Lower brain 18f-fluorodeoxyglucose
397		uptake but normal 11c-acetoacetate metabolism in mild alzheimer's disease
398		dementia. Journal of Alzheimer's Disease, 43(4):1343–1353, 2015.
399	[4]	Jessica R Andrews-Hanna, Abraham Z Snyder, Justin L Vincent, Cindy
400		Lustig, Denise Head, Marcus E Raichle, and Randy L Buckner. Disruption
401		of large-scale brain systems in advanced aging. Neuron, 56(5):924–935,
402		2007.
403	[5]	Linda Geerligs, Remco J Renken, Emi Saliasi, Natasha M Maurits, and
404		Monicque M Lorist. A brain-wide study of age-related changes in functional
405		connectivity. <i>Cerebral cortex</i> , 25(7):1987–1999, 2015.
406	[6]	Sophie Achard and Ed Bullmore. Efficiency and cost of economical brain
407		functional networks. PLoS Comput Biol, 3(2):e17, 2007.
408	[7]	Ed Bullmore and Olaf Sporns. The economy of brain network organization.
409		Nature Reviews Neuroscience, 13(5):336–349, 2012.
410	[8]	Louis A Profenno, Anton P Porsteinsson, and Stephen V Faraone. Meta-
411		analysis of alzheimer's disease risk with obesity, diabetes, and related dis-
412		orders. Biological psychiatry, 67(6):505–512, 2010.
413	[9]	Daihong Liu, Lihua Chen, Shanshan Duan, Xuntao Yin, Wu Yang, Yanshu
414		Shi, Jiuquan Zhang, and Jian Wang. Disrupted balance of long-and short-
415		range functional connectivity density in type 2 diabetes mellitus: A resting-
416		state fmri study. Frontiers in Neuroscience, 12:875, 2018.
417	[10]	Andrew Zalesky, Alex Fornito, Luca Cocchi, Leonardo L Gollo, and Michael
418		Breakspear. Time-resolved resting-state brain networks. Proceedings of the
419		National Academy of Sciences, 111(28):10341–10346, 2014.
420	[11]	Karl J Friston. Modalities, modes, and models in functional neuroimaging.
421		Science, 326(5951):399–403, 2009.

- ⁴²² [12] Olaf Sporns. Network attributes for segregation and integration in the ⁴²³ human brain. *Current opinion in neurobiology*, 23(2):162–171, 2013.
- ⁴²⁴ [13] Danielle Smith Bassett and ED Bullmore. Small-world brain networks. *The neuroscientist*, 12(6):512–523, 2006.
- [14] Dardo Tomasi, Gene-Jack Wang, and Nora D Volkow. Energetic cost of
 brain functional connectivity. *Proceedings of the National Academy of Sciences*, 110(33):13642–13647, 2013.
- ⁴²⁹ [15] Dardo Tomasi and Nora D Volkow. Aging and functional brain networks.
 ⁴³⁰ Molecular psychiatry, 17(5):549–558, 2012.
- [16] Richard F Betzel and Danielle S Bassett. Generative models for network
 neuroscience: prospects and promise. Journal of The Royal Society Inter face, 14(136):20170623, 2017.
- ⁴³⁴ [17] Réka Albert and Albert-László Barabási. Statistical mechanics of complex
 ⁴³⁵ networks. *Reviews of modern physics*, 74(1):47, 2002.
- [18] Gasper Tkacik, Elad Schneidman, II Berry, J Michael, and William
 Bialek. Spin glass models for a network of real neurons. arXiv preprint
 arXiv:0912.5409, 2009.
- [19] Thierry Mora and William Bialek. Are biological systems poised at critical values.
 (a) Cality? Journal of Statistical Physics, 144(2):268–302, 2011.
- ⁴⁴¹ [20] Elad Schneidman, Michael J Berry, Ronen Segev, and William Bialek.
 ⁴⁴² Weak pairwise correlations imply strongly correlated network states in a
 ⁴⁴³ neural population. *Nature*, 440(7087):1007–1012, 2006.
- [21] Marieke L Schölvinck, Alexander Maier, Q Ye Frank, Jeff H Duyn, and
 David A Leopold. Neural basis of global resting-state fmri activity. Pro-
- $_{446}$ ceedings of the National Academy of Sciences, 107(22):10238–10243, 2010.
- Li Zhao, David C Alsop, John A Detre, and Weiying Dai. Global fluctua tions of cerebral blood flow indicate a global brain network independent of

systemic factors. Journal of Cerebral Blood Flow & Metabolism, 39(2):302–

- 450 312, 2019.
- ⁴⁵¹ [23] Gašper Tkačik, Olivier Marre, Dario Amodei, Elad Schneidman, William
 ⁴⁵² Bialek, Michael J Berry, et al. Searching for collective behavior in a large
 ⁴⁵³ network of sensory neurons. *PLoS computational biology*, 10(1), 2014.
- ⁴⁵⁴ [24] Jason R Taylor, Nitin Williams, Rhodri Cusack, Tibor Auer, Meredith A
 ⁴⁵⁵ Shafto, Marie Dixon, Lorraine K Tyler, Richard N Henson, et al. The cam⁴⁵⁶ bridge centre for ageing and neuroscience (cam-can) data repository: struc⁴⁵⁷ tural and functional mri, meg, and cognitive data from a cross-sectional
 ⁴⁵⁸ adult lifespan sample. *Neuroimage*, 144:262–269, 2017.
- Lilianne R Mujica-Parodi, Anar Amgalan, Syed Fahad Sultan, Botond Antal, Xiaofei Sun, Steven Skiena, Andrew Lithen, Noor Adra, Eva-Maria
 Ratai, Corey Weistuch, et al. Diet modulates brain network stability, a
 biomarker for brain aging, in young adults. *Proceedings of the National*Academy of Sciences, 117(11):6170–6177, 2020.
- ⁴⁶⁴ [26] Kiyotaka Sato, Y Kashiwaya, CA Keon, N Tsuchiya, MT King, GK Radda,
 ⁴⁶⁵ B Chance, K Clarke, and RL Veech. Insulin, ketone bodies, and mitochon⁴⁶⁶ drial energy transduction. *The FASEB Journal*, 9(8):651–658, 1995.
- ⁴⁶⁷ [27] Daniel J DeDora, Sanja Nedic, Pratha Katti, Shafique Arnab, Lawrence L
 ⁴⁶⁸ Wald, Atsushi Takahashi, Koene RA Van Dijk, Helmut H Strey, and Lil⁴⁶⁹ ianne R Mujica-Parodi. Signal fluctuation sensitivity: An improved metric
 ⁴⁷⁰ for optimizing detection of resting-state fmri networks. *Frontiers in Neu-*⁴⁷¹ roscience, 10:180, 2016.
- ⁴⁷² [28] Alexandre Abraham, Fabian Pedregosa, Michael Eickenberg, Philippe
 ⁴⁷³ Gervais, Andreas Mueller, Jean Kossaifi, Alexandre Gramfort, Bertrand
 ⁴⁷⁴ Thirion, and Gaël Varoquaux. Machine learning for neuroimaging with
 ⁴⁷⁵ scikit-learn. *Frontiers in neuroinformatics*, 8:14, 2014.

- 476 [29] Jaime S Ide, Sanja Nedic, Kin F Wong, Shmuel L Strey, Elizabeth A Law-
- 477 son, Bradford C Dickerson, Lawrence L Wald, Giancarlo La Camera, and
 478 Lilianne R Mujica-Parodi. Oxytocin attenuates trust as a subset of more
- general reinforcement learning, with altered reward circuit functional connectivity in males. *Neuroimage*, 174:35–43, 2018.
- [30] Andre Altmann, Bernard Ng, Susan M Landau, William J Jagust, and
 Michael D Greicius. Regional brain hypometabolism is unrelated to regional
 amyloid plaque burden. *Brain*, 138(12):3734–3746, 2015.
- [31] R Matthew Hutchison, Thilo Womelsdorf, Elena A Allen, Peter A Bandettini, Vince D Calhoun, Maurizio Corbetta, Stefania Della Penna, Jeff H
 Duyn, Gary H Glover, Javier Gonzalez-Castillo, et al. Dynamic functional
 connectivity: promise, issues, and interpretations. *Neuroimage*, 80:360–
 378, 2013.
- [32] James M Shine, Oluwasanmi Koyejo, Peter T Bell, Krzysztof J Gorgolewski, Moran Gilat, and Russell A Poldrack. Estimation of dynamic
 functional connectivity using multiplication of temporal derivatives. NeuroImage, 122:399–407, 2015.
- [33] L.D. Landau and E.M. Lifshitz. *Mechanics*. Number v. 1. Elsevier Science,
 1982.
- ⁴⁹⁵ [34] Mingrui Xia, Jinhui Wang, and Yong He. Brainnet viewer: a network
 ⁴⁹⁶ visualization tool for human brain connectomics. *PloS one*, 8(7):e68910,
 ⁴⁹⁷ 2013.
- [35] Mark D'Esposito, Leon Y. Deouell, and Adam Gazzaley. Alterations in
 the bold fmri signal with ageing and disease: a challenge for neuroimaging.
 Nature Reviews Neuroscience, 4(1):9, 2003.
- [36] Meredith A. Shafto, Lorraine K. Tyler, Marie Dixon, Jason R. Taylor,
 James B. Rowe, Rhodri Cusack, Andrew J. Calder, William D. Marslen Wilson, John Duncan, Tim Dalgleish, Richard N. Henson, Carol Brayne,

> Fiona E. Matthews, and C. A. N. Cam. The cambridge centre for ageing and neuroscience (cam-can) study protocol: a cross-sectional, lifespan, multidisciplinary examination of healthy cognitive ageing. *BMC neurology*, 14(1):25, 2014.

- [37] Yuri Geinisman, Leyla de Toledo-Morrell, Frank Morrell, Inna S Persina,
 and Marvin Rossi. Age-related loss of axospinous synapses formed by two
 afferent systems in the rat dentate gyrus as revealed by the unbiased stere ological dissector technique. *Hippocampus*, 2(4):437–444, 1992.
- [38] Thomas C Foster and Christopher M Norris. Age-associated changes in
 ca2+-dependent processes: Relation to hippocampal synaptic plasticity.
 Hippocampus, 7(6):602–612, 1997.
- ⁵¹⁵ [39] Julia J Harris, Renaud Jolivet, and David Attwell. Synaptic energy use and supply. *Neuron*, 75(5):762–777, 2012.
- [40] Simonetta Camandola and Mark P Mattson. Brain metabolism in health,
 aging, and neurodegeneration. *The EMBO journal*, 36(11):1474–1492,
 2017.
- Fei Yin, Alberto Boveris, and Enrique Cadenas. Mitochondrial energy
 metabolism and redox signaling in brain aging and neurodegeneration. An tioxidants & redox signaling, 20(2):353-371, 2014.
- [42] Dimitrios Kapogiannis and Mark P Mattson. Disrupted energy metabolism
 and neuronal circuit dysfunction in cognitive impairment and alzheimer's
 disease. *The Lancet Neurology*, 10(2):187–198, 2011.
- [43] S. Nedic, S.M. Stufflebeam, C. Rondinoni, T.R. Velasco, A.C. dos Santos,
 J.P. Leite, A.C. Gargaro, L.R. Mujica-Parodi, and J.S. Ide. Using network
 dynamic fmri for detection of epileptogenic foci. *BMC Neurology*, 15(1):262,
 2015.

- ⁵³⁰ [44] Luca Cocchi, Leonardo L Gollo, Andrew Zalesky, and Michael Breakspear.
- ⁵³¹ Criticality in the brain: A synthesis of neurobiology, models and cognition.
- ⁵³² Progress in neurobiology, 158:132–152, 2017.
- ⁵³³ [45] Dante R Chialvo. Emergent complex neural dynamics. Nature physics,
 ⁵³⁴ 6(10):744-750, 2010.
- [46] Jens Wilting and Viola Priesemann. 25 years of criticality in neuro scienceestablished results, open controversies, novel concepts. *Current opin- ion in neurobiology*, 58:105–111, 2019.
- [47] John M Beggs. The criticality hypothesis: how local cortical networks
 might optimize information processing. *Philosophical Transactions of* the Royal Society A: Mathematical, Physical and Engineering Sciences,
 366(1864):329-343, 2008.
- [48] David J Schwab, Ilya Nemenman, and Pankaj Mehta. Zipfs law and criticality in multivariate data without fine-tuning. *Physical review letters*,
 113(6):068102, 2014.