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The complexity of the relationship between spontaneous brain activity and glucose metabolism

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5	and glucose metabolism
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Abstract

Brain glucose metabolism as assessed by [¹⁸F]FDG positron emission tomography (PET) is expected 37 to be significantly related to resting-state functional MRI (rs-fMRI) activity and functional 38 connectivity (FC), but the underlying coupling model is still incompletely understood. Employing 39 simultaneous acquisitions, we related [¹⁸F]FDG standard uptake value ratio (SUVR) to 50 features 40 41 pertaining to rs-fMRI 1) signal, 2) hemodynamic response, 3) static and 4) time-varying FC, and 5) 42 phase synchronization. To assess which rs-fMRI variables better describe SUVR across regions, we 43 employed a hierarchical approach, identifying the model at population level, and then estimating it on individual data. Multilevel modelling explained around 40% of the SUVR variance, with signal-44 45 related features as the most relevant fMRI variables. When the model was used to characterize 46 between-network variability of the SUVR-fMRI coupling, the ranking changed. We demonstrate that 47 local activity and synchronization are the most important predictors of glucose metabolism, while 48 large-scale FC properties gain importance within specific networks. 49

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Introduction

Brain glucose consumption can be assessed *in vivo* by [¹⁸F]fluorodeoxyglucose positron emission 56 57 tomography ([¹⁸F]FDG PET) acquisitions¹, usually through semi-quantitative proxies like the standard uptake value ratio (SUVR)². As evidenced by [¹⁸F]FDG studies, glucose metabolism 58 59 displays significant regional variability. The reasons behind this heterogeneity in glucose expense, however, remain largely unexplained. Crucially, most of the remarkable metabolic budget of the 60 brain, ~25% of energy in the face of only 2% of body weight, is spent during rest^{3,4}. This 'dark energy' 61 62 of the brain⁵ with unclear functional meaning is expected to be mainly employed for maintaining resting potentials and subthreshold synaptic transmission⁶, since most of the energy budget of a 63 64 neuron is utilized at the level of the synapses, rather than in the neuron's body⁷.

65 The regional differences in brain metabolism are thus likely to be explained by variability in 66 spontaneous activity, which has been extensively explored with blood-oxygen-level-dependent (BOLD) resting-state functional magnetic resonance imaging (rs-fMRI)^{8,9}. In addition to spontaneous 67 68 activity, the functional relationships between activity patterns of different brain regions may relate to glucose consumption as well¹⁰. To this purpose, rs-fMRI can be used to derive the so-called 69 "functional connectivity" (FC), i.e., the statistical relation between BOLD signal fluctuations in 70 71 different brain regions; this approach has led to the identification of a functional architecture of 72 resting-state networks (RSNs) that recapitulate clusters of regions activated for specific functions^{11,12}. 73 FC can be estimated in a static fashion (sFC), but also with time-varying approaches (tvFC), which 74 interpret FC as non-stationary and changing across adjacent time windows¹³; notably, both might 75 prove relevant to metabolic consumption.

Metrics derived from network science can then be used to characterize the topology of sFC and tvFC, with correlations between areas represented as 'edges' connecting 'nodes', and nodes described in terms of their centrality ('hubness'), number of connections, and so on^{14,15}.

While rs-fMRI studies have provided a wide range of information on the properties of spontaneous activity, the physiological underpinnings of these results remain poorly understood, as the BOLD signal arises from a complex combination of cerebral blood volume (CBV), blood flow (CBF) and metabolic rate of oxygen (CMRO₂)^{17,18}, and is indirectly and nonlinearly related to neuronal activity through the hemodynamic response function (HRF)¹⁹; importantly, it is also subjected to significant contamination from systemic modulations, both hemodynamic (heart rate variability, vasomotion etc.) and respiratory (e.g., respiratory volume variability)²⁰.

Building upon the previous considerations, the relationship between the information provided by rsfMRI and [¹⁸F]FDG PET across brain regions needs to be thoroughly investigated with two main
aims.

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First, to better understand the sources of regional metabolic variability. In particular, we might wonder: 1) how much is the 'dark energy' of the brain related to local activity probed by the BOLD signal? 2) how much is instead related to local synchronization of the BOLD signal, i.e., a possible measure of neural population excitatory-inhibitory balance? 3) does inter-regional static synchrony play a more relevant role than more dynamic, time-varying interactions?

94 Second, the biological basis of the BOLD signal needs to be further characterized, an effort that would 95 help turn rs-fMRI into a more specific biomarker; [¹⁸F]FDG PET, being tightly linked to neural 96 activity⁷, despite its typically low temporal resolution, can provide important insight.

97 Some evidence on this relationship has started to emerge from sequential and simultaneous [¹⁸F]FDG PET/fMRI acquisitions²¹. In particular, the mean BOLD signal and the amplitude of its low frequency 98 99 fluctuations (ALFF) have been found to be associated with [¹⁸F]FDG SUVR across voxels²², with stronger correlations in specific brain regions^{23,24}. With regard to the coupling between FC and 100 101 metabolism, moderate associations between SUVR and global FC metrics were detected^{23,25}, with 102 stronger and more consistent correlations for the regional homogeneity (ReHo) of BOLD (up to Pearson's r = 0.8), which is an index of local synchronization^{26,27}. The topology of FC was also found 103 104 to be important, with more central nodes having a stronger relationship between their FC and metabolic consumption²³. In addition, the relationship between SUVR and local and global FC has 105 been described as a power law or exponential model, especially in some specific networks^{25,28}. The 106 variability in the [¹⁸F]FDG-fMRI coupling across networks has been highlighted in multiple previous 107 studies^{23,24}, with demarcation between visual and default mode regions on one side, and frontoparietal 108 109 regions on the other²⁸. When the [¹⁸F]FDG-fMRI coupling is assessed across subjects, however, much lower correlations between SUVR and rs-fMRI variables are detected in many studies^{23,26,27}. In 110 summary, somewhat inconsistent results emerge from the literature, with bivariate spatial correlations 111 112 between [¹⁸F]FDG PET and a handful of BOLD-derived metrics (ALFF, ReHo, voxel-wise FC) ranging from 0 to 0.64 in explained variance (R²), and substantial differences across brain regions 113 and networks, as well as a low correlation across subjects even in simultaneous acquisitions^{23,27}. 114 115 Notably, no study has ever attempted a multivariate integration of a wider range of rs-fMRI features, 116 as well as a multilevel prediction of SUVR both at the population level and at the subject/network 117 level.

118 We set out to fill these gaps in knowledge with a fully data-driven approach using simultaneously

acquired [¹⁸F]FDG PET and rs-fMRI data from two separately published datasets of 26 subjects^{23,31}.

120 After preliminary assessment of 50 rs-fMRI-derived variables, pooled into 5 categories, i.e., 1) signal,

121 2) HRF, 3) sFC, 4) tvFC, and 5) phase coherence $(PC)^{29}$ (see **Table 1** for the list of the features, their

acronyms and a brief description) we set out to address the following questions.

- 123 1. which is the strength of the bivariate association between these rs-fMRI features and SUVR across
- 124 the whole brain? And then, since regions with high vs. low metabolic consumption are expected to
- have quite different structural and functional properties^{1,3,4}, does this coupling change according to
- 126 the ranking of brain nodes based on SUVR?
- 127 2. is it possible to explain group level SUVR variance across regions by combining rs-fMRI features,
- 128 for the first time, into a multiple regression model? Is the group of selected features more populated
- 129 by local or large-scale brain network metrics, and does it account for between-subject variability
- 130 (BSV)³⁰? Finally, which of the previously identified rs-fMRI features are more important to explain
- 131 SUVR when multilevel modelling is performed across RSNs, i.e., which is the between-network
- 132 variability (BNV) of the SUVR-fMRI association?
- 133

134 Table 1 – Extracted rs-fMRI features and their categories

Fifty fMRI-derived variables, divided according to the pool to which they belong: 1) signal, 2) hemodynamic response function (HRF), 3) static functional connectivity (sFC), 4) time-varying functional connectivity (tvFC), 5) phase coherence (PC). See **Supplementary Methods** for full description of the features.

Pools	rs-fMRI Variables
	med-BOLD: median of the BOLD time series
	MAD-BOLD: median absolute deviation (MAD) of the BOLD time series
	skew-BOLD: skewness of the BOLD time series
	ApEn-BOLD: approximate entropy (ApEn) of the BOLD time series
al	rApEn-BOLD: range ApEn of the BOLD time series
Signal	AR-BOLD : reflection coefficient of the first-order autoregressive AR(1) model fit to BOLD time series
\mathbf{N}	ALFF: amplitude of low frequency fluctuations (ALFF) of BOLD time series
	ReHo: regional homogeneity of BOLD time series
	MAD-ReHo: MAD of the time-varying ReHo (tvReHo)
	CV-ReHo: CV% of tvReHo
	peaks-BOLD: number of BOLD pseudo-events
	peak-HRF: height of HRF peak
	hrf-DEG: degree (DEG) of HRF correlation matrix
Γ τ ι	hrf-STR: strength (STR) of HRF correlation matrix
HRF	hrf-CC: clustering coefficient (CC) of HRF correlation matrix
	hrf-BC: betweenness centrality (BC) of HRF correlation matrix
	hrf-EC: eigenvector centrality (EC) of HRF correlation matrix
	hrf-LE: local efficiency (LE) of HRF correlation matrix
	hrf-GE: global efficiency (GE) of HRF correlation matrix
	s-DEG: DEG of sFC
	s-STR: STR of sFC
U	s-CC: CC of sFC
sFC	s-BC: BC of sFC
•1	s-EC: EC of sFC
	s-LE: LE of sFC
	s-GE: GE of sFC

	mdiff-DEG : temporal median of the absolute value of 1 st order differentials (mdiff) of DEG time series
	mdiff-STR: mdiff of STR time series
	mdiff-CC: mdiff of CC time series
	mdiff-BC: diff of BC time series
	mdiff-EC: mdiff of EC time series
	mdiff-LE: mdiff of LE time series
	mdiff-GE: mdiff of GE time series
	CV-DEG: coefficient of variation (CV%) of DEG time series
U	CV-STR: CV% of STR time series
tvFC	CV-CC: CV% of CC time series
tv	CV-BC: CV% of BC time series
	CV-EC: CV% of EC time series
	CV-LE: CV% of LE time series
	CV-GE: CV% of GE time series
	SampEn-DEG: sample entropy (SampEn) of DEG time series
	SampEn-STR: SampEn of STR time series
	SampEn-CC: SampEn of CC time series
	SampEn-BC: SampEn of BC time series
	SampEn-LE: SampEn of LE time series
	SampEn-GE: SampEn of GE time series
	med-LEig: median of the Leading Eigenvector (LEig)'s time series
\mathbf{C}	MAD-LEig: MAD of LEig time series
Ď	CV-LEig: CV% of LEig time series
	mdiff-LEig: mdiff of LEig time series

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Results

143 Feature extraction and preliminary evaluation of rs-fMRI variables

144 The flowchart describing the preprocessing and preliminary analysis of the [¹⁸F]FDG PET and rs-

145 fMRI data is shown in **Figure 1** (see the **Methods** section for details).

146 The $[^{18}F]FDG$ PET variable of interest is the SUVR, which was extracted for every region of the 147 Schaefer cortical atlas³² (200 parcels, supplemented by 18 subcortical regions³³) in each subject, and

Schaefer cortical atlas³² (200 parcels, supplemented by 18 subcortical regions³³) in each subject, an
will be considered as the dependent variable in every modelling approach from here onward.

The 50 rs-fMRI variables, extracted at the single-subject level and *a priori* subdivided into 5 pools, are reported in **Table 1**: the signal pool (1) contains features related to the basic statistics of the BOLD

151 time series (median, variance, skewness), its complexity, its low-frequency fluctuations (ALFF), local

152 coherence (ReHo) and high-amplitude events (peaks-BOLD); in the HRF pool (2), then, we placed

153 the amplitude of the HRF peak, calculated using a blind deconvolution method³⁴, and the HRF

154 correlation structure across regions described by means of graph properties; the sFC pool (3)

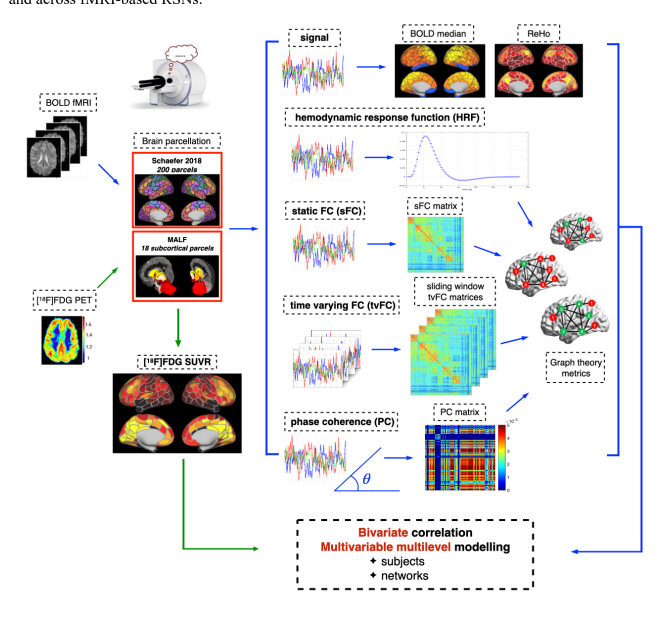
155 characterizes FC calculated across the entire fMRI scan with graph theory metrics; the tvFC pool (4)

assesses graph metrics' temporal variability across sliding windows¹³; finally, the PC pool (5)

157 characterizes FC as coherence of BOLD phase²⁹.

158 Figure 1 – Flowchart of rs-fMRI and [¹⁸F]FDG PET processing, feature extraction and analysis.

- 159 Both rs-fMRI time series and [¹⁸F]FDG SUVR data were parceled using the Schaefer cortical atlas
- 160 (200 ROIs) and 18 subcortical ROIs. The parcel-wise rs-fMRI data were used to extract fifty features
- representative of five pools, i.e., 1) signal, 2) HRF, 3) sFC, 4) tvFC, 5) PC. The PET-fMRI coupling was investigated using bivariate correlation and multivariable multilevel modelling across subjects
- 162 was investigated using bivariate correlation163 and across fMRI-based RSNs.



168 The Spearman's correlation matrix between the 50 rs-fMRI variables at group median level (i.e., by 169 taking the parcel-wise median value of each feature across subjects) was computed (Figure 2a), in order to assess the relationships between the extracted features and their degree of redundancy: the 170 clustering into 5 pools provided by a priori knowledge was fairly consistent with the observed 171 172 correlation structure, with signal, HRF and sFC features (upper block) being clearly distinguished from tvFC features (lower block), which they are negatively correlated with, and PC variables 173 174 demonstrating lower correlation with the rest. However, it was also noticeable that strong correlations between many variables were present, especially for the tvFC pool, and that a feature selection step 175 176 was going to be necessary to use these variables in a numerically sound multivariable model of SUVR: the condition number $\kappa(X)$, which quantifies the level of correlation between predictors in a 177 multiple regression context (i.e., their multicollinearity), was high ($\kappa(X) = 70.58$), way beyond the 178 acceptability range³⁵, and this is known to result in unstable and unreliable models (see **Methods**). 179

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182 SUVR vs. rs-fMRI: bivariate relationships

Before moving to the multiple regression framework, we began by investigating bivariate associations between SUVR and the extracted rs-fMRI variables at the group level, in the so-called naïve average data approach (NAD), as done by many previous studies^{23,25,26}; here, however, a much wider range of fMRI-derived variables was explored. Many significant spatial associations between SUVR and rs-fMRI features were detected across the 218 analyzed regions, as assessed through Spearman's rank correlation (p = 0.05 significance level) with false discovery rate (FDR) multiple comparison correction³⁶. The correlation coefficients are reported in **Figure 2b**.

190

192 Figure 2 – Bivariate correlations among rs-fMRI variables, and between rs-fMRI variables and

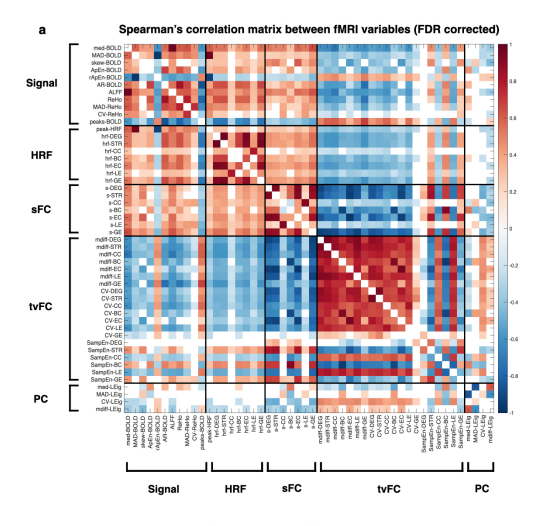
193 SUVR. The pattern of Spearman's correlations (FDR-corrected, non-significant values shown in

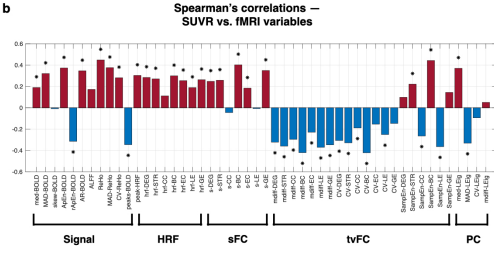
194 white) among rs-fMRI features, assessed at the group level and divided according to the pool to which

they have been assigned (1) signal, 2) HRF, 3) sFC, 4) tvFC, 5) PC), is shown in (a). The rs-fMRI 195 196

features are tested for association with group median SUVR across 218 brain regions (b) via

197 Spearman's correlations (significant values after FDR correction are indicated with an asterisk).





The strongest *positive* associations were with 1) ReHo ($\rho = 0.45$, p < 0.001), 2) s-BC ($\rho = 0.4$, p < 0.001), and 3) SampEn-BC ($\rho = 0.44$, p < 0.001), i.e., respectively 1) a measure of local synchronization of BOLD, 2) a sFC graph metric, betweenness centrality (BC), which describes a node in terms of its *global* connections in a graph, and 3) a measure of temporal complexity of the BC time series. The strongest *negative* correlations were mdiff-BC ($\rho = -0.42$, p < 0.001) and CV-BC ($\rho = -0.42$, p < 0.001) in the tvFC pool, both measures of temporal variability of BC.

In general, it can be noted that positive associations emerged for the majority of the signal-based, 206 207 HRF and sFC-related features, while tvFC metrics displayed a consistent and never previously 208 reported negative association with SUVR (Figure 2b). Notable exceptions amongst signal-based features are rApEn-BOLD ($\rho = -0.31$, p < 0.001), a measure of BOLD signal complexity, and peaks-209 210 BOLD ($\rho = -0.34$, p < 0.001), which quantifies the number of signal peaks exceeding one standard 211 deviation from the baseline: both exhibited negative relationships with SUVR. Amongst tvFC 212 features, SampEn-BC ($\rho = 0.44$, p < 0.001) shows a strong positive coupling with SUVR, in contrast 213 to the behavior of the other tvFC metrics. Interestingly, the dynamics of local synchronization, i.e., 214 MAD-ReHo and CV-ReHo, displays a positive association with SUVR as well.

215 Overall, it can be noted that the detected spatial correlations were at best moderate.

216

217 SUVR-fMRI associations are strengthened in low SUVR nodes

As the relationship between [¹⁸F]FDG PET and rs-fMRI could be spatially heterogeneous, 218 219 Spearman's correlations were also re-evaluated across groups of nodes selected according to linearly 220 increasing percentiles of the SUVR distribution, i.e., by progressively retaining the highest SUVR values, from the 1st up to the 85th percentile, as well as linearly decreasing percentiles of the same 221 222 distribution, i.e., by retaining the lowest SUVR values, from the 100th to the 15th (Supplementary 223 Figure S1). The purpose of the analysis was to verify whether SUVR-fMRI associations would be 224 strengthened in high SUVR nodes or, conversely, in low SUVR nodes, as SUVR provides a clear 225 ranking of brain regions, that is expected to be related to crucial structural and functional properties, 226 e.g., neuron-to-glia ratio, richness in neuroreceptors, excitatory-inhibitory activity^{1,3,4}.

Spearman's correlations (p = 0.05 significance level, FDR-corrected) between parcel-wise SUVR and all 50 rs-fMRI features (*rows*) are shown in **Figure 3a**, for each of the threshold levels along the SUVR distribution (*columns*). Going towards nodes with high SUVR (*right side of* **Figure 3a**) does not lead to any relevant effect for most features (except for three measures): therefore, hardly any strengthening of SUVR-fMRI relationships is detected in high SUVR nodes.

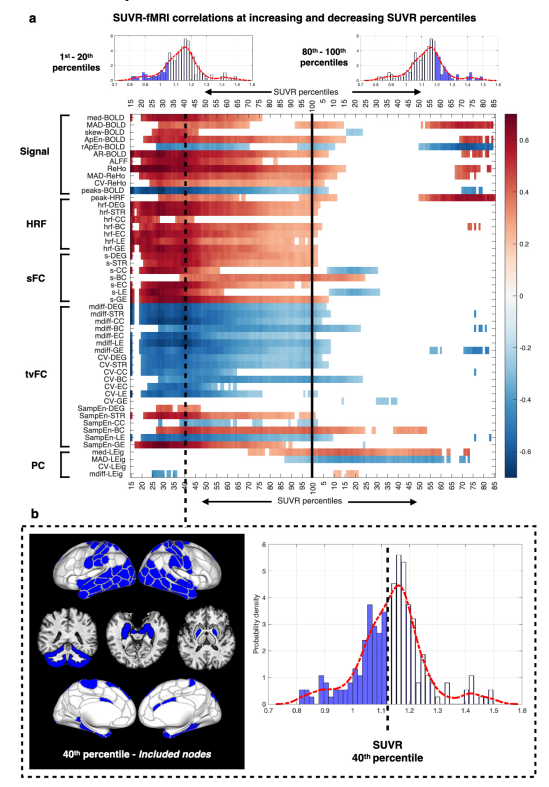
232 Interestingly, however, a marked increase in many of the bivariate associations can be observed by

233 selecting nodes with low values of SUVR (*left side of* Figure 3a), with highly significant correlations

even after FDR correction.

235 Figure 3 – The SUVR-*fMRI* correlation changes strongly in low SUVR nodes.

Spearman's correlations (FDR-corrected, non-significant values shown in white) between SUVR and all fifty rs-fMRI features (*y axis*) across nodes selected according to linearly increasing (*x axis - right*) and decreasing (*x axis - left*) percentiles of SUVR (**a**). The dashed black line shows the percentile with maximum correlation across features (i.e., nodes in the $1^{st} - 40^{th}$ percentile range). The histogram (**b**), on the right, highlights the chosen percentile in the SUVR distribution and the range of percentiles included in the correlation. The brain regions shown on the left, plotted on the cortical surface and subcortex, are the parcels over which correlations are assessed.



We then identified the percentile threshold corresponding to the highest total correlation value across features: the spatial pattern of the 87 nodes below the 40th percentile of the SUVR distribution is shown in **Figure 3b.** These parcels, where the SUVR-fMRI association is emphasized, mainly belong to temporal/limbic areas (including hippocampus), sensorimotor cortices, and subcortical regions, such as cerebellum and globus pallidus (**Supplementary Figure S1**).

This finding suggests the presence of *nonlinear* relationships between [¹⁸F]FDG SUVR and most rsfMRI features: tighter and more linear associations are present across a limited range of low SUVR nodes, with weaker coupling as SUVR gets higher. This nonlinear association was thus further characterized (see **Supplementary Results**), expanding on previous studies which were focused on specific networks and features^{25,28}.

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255 SUVR vs. rs-fMRI: multivariable multilevel model

We then set out to assess which combination of rs-fMRI features was better able to explain SUVR 256 257 across brain regions with multiple regression and multilevel modelling, in a fully data-driven way. In 258 multilevel modelling, the model structure is usually known, or selected at the lower level, i.e., at the 259 individual level³⁰. However, as significant BSV in the SUVR-fMRI association is expected, 260 especially for the rs-fMRI features, we chose to identify the model predictors at the population level (NAD approach), thus exploiting the denoising properties of averaging. The model structure selected 261 262 at the group median level was then used for multilevel modelling across subjects to characterize the 263 BSV of the SUVR-fMRI association, as shown in (Figure 4, top), trying to fully capitalize on the fact that [¹⁸F]FDG and rs-fMRI data were acquired in the same subjects. 264

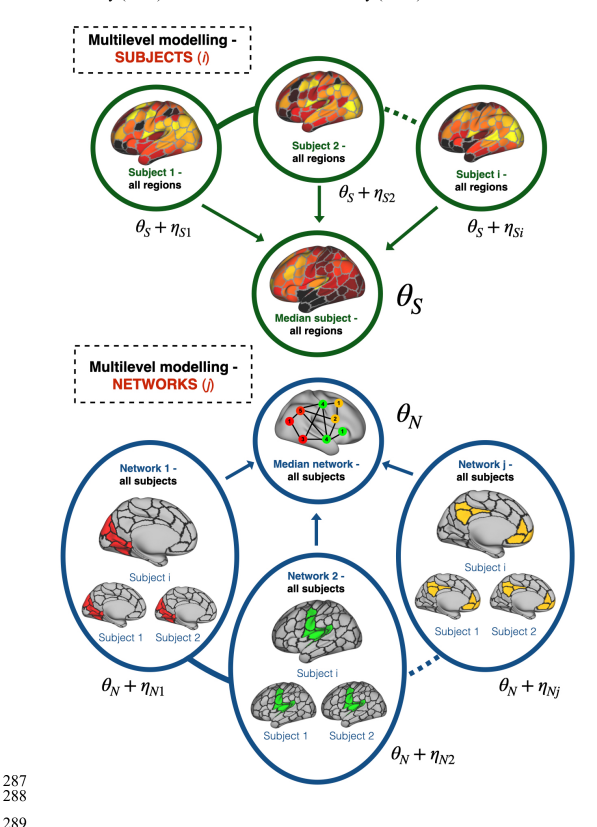
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266 <u>Maximum explanatory power of SUVR:</u>

To assess the maximum explanatory power provided by the fMRI-derived features, we began by fitting an ordinary least squares (OLS) multiple regression model employing *all* the available features in a log-linear form (i.e., exploring the relationship between SUVR and the log-transformed rs-fMRI explanatory variables), to account for the detected nonlinearity (**Supplementary Results**): it can in fact be noted that the log-linear pairwise relationship between SUVR and each of the rs-fMRI features outperformed to the linear model in many occasions (62%). From now we will call this log-linear model.

The OLS model had an R^2 value of 0.62: the maximum explanatory power thus reaches up to around 60% of the SUVR variance, without fully saturating the variance despite marked overparameterization (i.e., 50 rs-fMRI predictors). Due to the high number of predictors and the presence of multicollinearity, the precision of numerous parameter estimates (expressed as coefficients of variation, $CVs\%)^{37}$ was low, as expected (**Supplementary Figure S3c**).

280 Figure 4 – Multilevel modelling approach for subjects and networks. The multivariable model 281 structure defined at the population level for the SUVR-fMRI relationship was used in a hierarchical 282 model. The multilevel approach was first applied to individual data with subjects (S) as grouping 283 variables (S, top), then with networks (N, bottom): fixed effects θ_S and θ_N describe the population 284 parameters, and random effects η_{Si} and η_{Ni} describe how much each subject *i* or network *j* deviates from the population fixed effects for each of the selected parameters, i.e., the between-subject 285 286 variability (BSV) and between-network variability (BNV) of the SUVR-fMRI model.



- 290 Assessment of a parsimonious and informative group level multivariable model:
- 291 Multiple feature selection approaches (eleven methods) were then explored. The results of each and
- the reason for the choice of the optimal approach are detailed in **Supplementary Results**.
- 293 The chosen feature selection process was performed in two stages. First, a sign-constrained non-
- 294 negative least squares (NNLS) estimator³⁸ was employed; then, the NNLS estimates were refined
- with a second stage of feature selection with elastic net regression³⁹. The reached solution was optimal
- in comparison with the other ten methods, in terms of both goodness of fit ($R^2 = 0.411$) and precision
- of the estimates (CVs% $\mu \pm \sigma = 66.73 \pm 17.79$ %). The selected rs-fMRI predictors are: 1) ApEn-
- 298 BOLD, 2) rApEn-BOLD, 3) ReHo, 4) CV-ReHo, 5) peaks-BOLD, 6) hrf-LE, 7) s-BC, 8) CV-BC, 9)
- 299 med-LEig.
- The first five predictors belong to the signal and local synchronization pool, while the other four to the remaining groups of rs-fMRI features, suggesting only an *indirect* relationship with the largescale network connectivity measures. Notably, most of the identified rs-fMRI predictors were chosen with high consistency across the employed feature selection methods, which highlights the robustness of their association with SUVR (**Supplementary Figure S4, Supplementary Figure S5**).
- 305

306 <u>Multivariable multilevel model across subjects:</u>

307 The hierarchical modelling framework was then applied to the individual data using the identified 308 predictors, to characterize the BSV of the SUVR-fMRI association. The log-linear model identified at the group level was re-estimated using a linear mixed-effect (LME) approach³⁰. The fixed-effect 309 310 (θ_s) parameter estimates, which represent the equivalent of the parameters estimated at the group 311 median level, are reported in (Figure 5a) with their standard errors (SEs). To get an accurate ranking of the most relevant predictors in explaining SUVR, the estimated θ_s were ordered by their relative 312 313 contribution to the model using dominance analysis (DA)⁴⁰ (Figure 5b). In terms of general dominance (see Methods), at the top was ReHo (48% of the total R²), followed by peaks-BOLD 314 (19%), CV-BC (11.74%), CV-BC (10.50%), s-BC (8.02%), ApEn-BOLD (3.67%), med-Leig 315 316 (2.60%), hrf-LE (1.47%), rApEn-BOLD (0.02%). Notably, the features belonging to the signal pool collectively accounted for 76.17% of the hierarchical model R². The random effects (η_{si}) describe 317 318 the deviation from the group value of the parameters for a specific subject *i*, i.e., how much the 319 parameters of each subject *i* are distant from the group-level estimates θ_s (Supplementary Figure 320 S7b). In this case we found that the BSV in the SUVR-fMRI association is clearly non-negligible. In 321 fact, the group-level θ_s estimates are very close to those obtained using the NAD approach, 322 confirming the adequacy of the average approach in describing the relationship between the variables. However, as expected, the R² of the overall model, i.e., considering BSV, was lower and equal to 323 324 0.245, due to the capability of the multilevel mixed-effect approach to keep into account both

- between- and within-subject variability. The R^2 values of the subject-level models are reported in
- **Supplementary Figure S7a**, and they display high variability (from 0.05 to 0.45).
- The median across subjects of the model's residuals v_{Si} , which highlight how well the SUVR of each region is explained by the identified model, can be visualized in **Figure 5c**. Notably, high positive

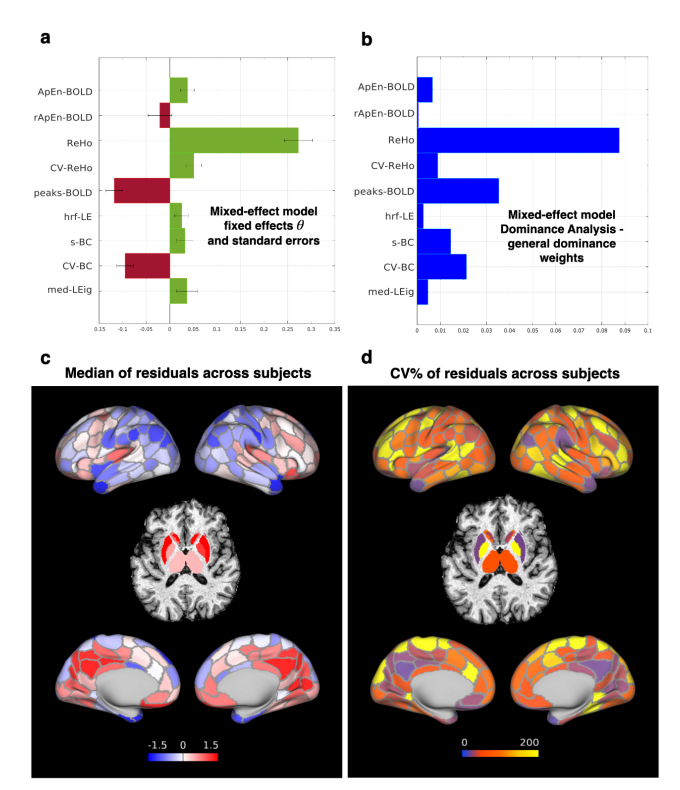
region is explained by the identified model, can be visualized in **Figure 5c**. Notably, high positive values are present in posteromedial cortex (posterior cingulate cortex (PCC) in particular) and

330 subcortex (putamen): these areas identify nodes with high SUVR values which are not satisfactorily

- 331 explained by the available rs-fMRI features. Importantly, this deficiency in explanatory power is
- 332 highly consistent across subjects, as evidenced by the low variability (CVs%) of the residuals in those
- areas (Figure 5d).

Figure 5 – Multivariable multilevel modelling of SUVR across subjects. The multivariable loglinear model predictors chosen at the group NAD level are shown. Parameter estimates and standard errors for the fixed effects θ_s , which represent the parameters that best explain SUVR across regions at the group level (a). The relative importance weights produced by dominance analysis (DA), highlighting the proportion of the multivariable multilevel model R² explained by each predictor (general dominance) (b). Across-subject median (c) and CVs% (d) of weighted residuals v_{Si} of the multilevel model, plotted on the brain cortex and subcortex.

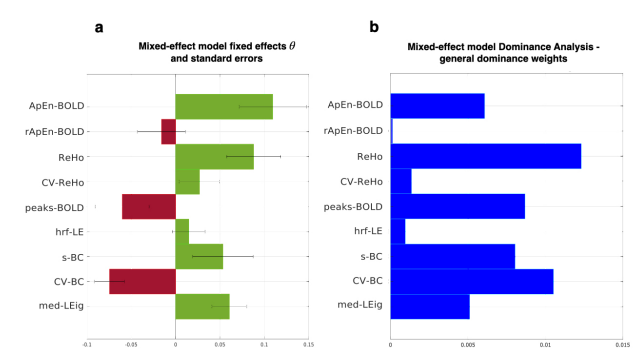
342



345 SUVR vs. rs-fMRI: multilevel model across networks

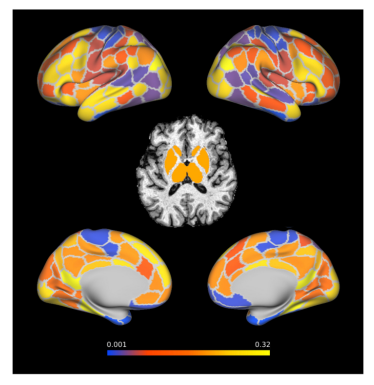
- Finally, the log-linear model with the 9 selected rs-fMRI predictors was tested in its ability to describe the expected network-level variability of the SUVR-fMRI association (BNV), i.e., by considering only the parcels referring to specific brain networks. RSNs were here grouped according to the Schaefer' functional atlas in its 17-RSN partition³², supplemented by 18 subcortical anatomical regions³³. A multilevel modelling framework was again employed, but with RSNs as the grouping
- 351 factor for individual level data, instead of subjects (Figure 4, *bottom*).
- 352 The fixed effects θ_N and their SEs for the between-network model are reported in (Figure 6a). ReHo
- 353 and peaks-BOLD are still highly relevant parameters in describing the SUVR across networks,
- 354 together with ApEn-BOLD and CV-BC; rApEn-BOLD and hrf-LE, instead, lose importance, and
- 355 their fixed effect θ_N becomes irrelevant (with a SE range crossing the zero-line). To confirm the
- 356 ranking, DA was performed in this context as well: ReHo was still the most important predictor in
- terms of general dominance (explaining 23.24% of the model's R²), followed by CV-BC (19.85%),
- 358 peaks-BOLD (16.39%), s-BC (15.19%), ApEn-BOLD (11.46%), med-LEig (9.65%), CV-ReHo
- 359 (2.55%), hrf-LE (1.80%), rApEn-BOLD (0.13%) (Figure 6b).
- 360 Notably, the R² of model prediction considering network-wise estimates is markedly lower than when
- 361 subjects are used to cluster nodes. As shown in (Figure 6c), the single RSNs are highly heterogeneous
- in terms of model R², ranging from around 0 to 0.32, with an overall prediction with $R^2 = 0.147$.

Figure 6 – Multilevel SUVR modelling across networks – parameter estimates and explained variance. The nine features chosen at the group level are shown. Parameter estimates and standard errors for the fixed effects θ_N , which represent the parameters that best explain SUVR across regions in an average network (a). Relative importance weights produced by dominance analysis (DA) in terms of the proportion of the between-network model R² explained by each predictor (general dominance) (b). Network-wise R² values, representing the percentage of SUVR variance explained by the mixed-effect model at the network level, plotted on the cortical surface and subcortex (c).



С

Network-wise model R²



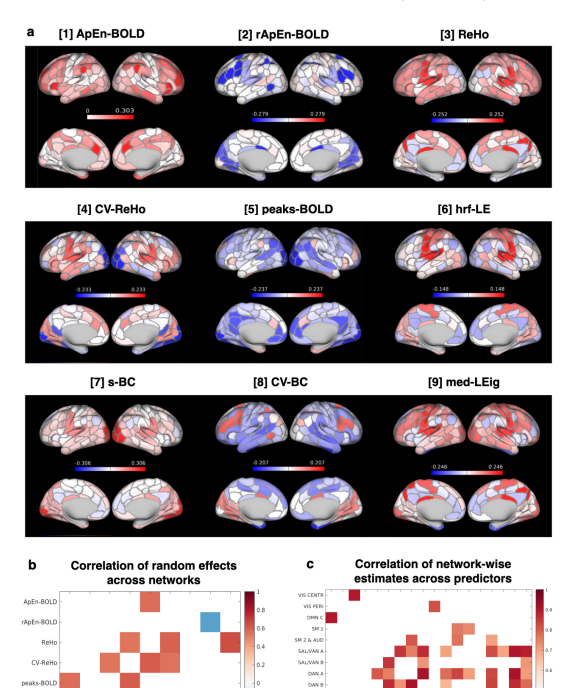
373 The BNV of the SUVR-fMRI association is measured by the random effects η_{Ni} for each network, with some RSNs displaying significant distance from the model estimates θ_N of the "average-374 network". To better assess this variability, the nine rs-fMRI predictors' parameter estimates ψ_{Ni} (i.e., 375 sum of fixed effects θ_N and random effects η_{Nj} for every network j) were plotted (Figure 7a). We 376 377 can observe that most predictors included in the multivariable model display heterogeneity across 378 networks in their relationship with SUVR, with either positive or negative associations depending on 379 the specific RSN, which cannot be captured by the average situation described by the fixed effects 380 θ_N of (Figure 6a). Some predictors show notably consistent spatial patterns, and therefore, to assess their similarity, the correlation between their random effects was evaluated across networks (Figure 381 382 7b). Notably, although uncorrected for multiple comparisons, significant correlations (p < 0.05) can 383 be found between the patterns of ReHo, CV-ReHo, hrf-LE and med-LEig, with strong positive 384 weights for somatomotor network B (SM(B)) and also control network (CTR(C)). Another interesting 385 pattern emerges for CV-BC, which displays both positive (CTR(A), VIS(B)) and negative weights 386 (TEMP/PAR, LIMBIC(A), SAL/VAN(A), DMN(B)).

387 Finally, the network-wise ψ_{Nj} values were correlated across predictors, to assess how similar the 388 RSNs were to one to another in terms of their SUVR-fMRI coupling (Figure 7c). When considering 389 only significant correlations (p < 0.05, uncorrected), an interesting pattern emerges: some RSNs are 390 fairly isolated from the rest of the brain in their SUVR-fMRI association pattern (e.g. DMN(A), 391 DMN(C), VIS(A), VIS(B), SM(A), CTR(A)), with only 1-2 significant correlations with other RSNs; 392 other RSNs, instead, have many significant correlations, and thus are similar to many other networks 393 in their SUVR-fMRI coupling (SAL/VAN(A), DAN(A), DAN(B), CTR(B), CTR(C), DMN(B), 394 LIMBIC(B), SUB).

396 Figure 7 – Multilevel SUVR modelling across networks – multivariable network-level estimates.

Individual network parameter estimates (ψ_{Nj} , sum of fixed effects θ_N and random effects η_{Nj} , which 397 398 describe the variability from the fixed effect for each RSN *j*), plotted on the brain surface for each 399 predictor (a). Correlation matrix (non-significant values in white, p = 0.05, uncorrected) of the nine 400 predictors' random effects η_{Ni} across RSNs (b). Correlation matrix (non-significant values in white, p = 0.05, uncorrected) of the RSNs' parameter estimates ($\psi_{Nj} = \theta_N + \eta_{Nj}$) across predictors (c). 401

402



CTR A

CTR B

CTR C

TEMP/PAR

DMN A

DMN B

LIMBIC A

LIMBIC B

SUE

SM 1

DMN (

CTR E

DMN MBIC

CTR CTR (

DAN

-0.2

-0.4

-0.6

-0.8

ned-LEig

403

hrf-LE

s-BC

CV-BC

med-LEig

-BOLD

ADEN-BOLD

eaks-BOLI

404

Discussion

In this work, we thoroughly investigated the spatial relationship between a wide range of features
extracted from rs-fMRI and simultaneously acquired [¹⁸F]FDG PET, while also accounting for the
variability across subjects (i.e., BSV) and networks (i.e., BNV) in this relationship.

408

409 Relationship between SUVR and rs-fMRI through bivariate correlation

In addition to the rs-fMRI variables that have already been associated to SUVR, i.e., ALFF, ReHo,
sFC strength^{22,23,25}, we extended our assessment to a wide variety of previously unexplored features,
such as time-varying functional connectivity (tvFC) and HRF-related variables.

To our knowledge, the relationship between [¹⁸F]FDG metabolism and FC temporal variability has never been tested before. While it is established that regions with stronger static FC tend to have higher cerebral blood flow⁴¹ and higher energy metabolism²⁵, possibly reflecting the fact that they are also more strongly connected anatomically⁴² (with recent work highlighting that structural connectivity graph properties are positively associated with [¹⁸F]FDG SUVR^{43,44}), the tvFC coupling with glucose metabolism is not established.

419 We found that tvFC (as captured by graph theory metrics' temporal variability) has a moderate-strong 420 negative association with SUVR. The interpretation of this finding can be supported by knowing that 421 sFC and tvFC graph metrics are negatively correlated, as clearly shown by the correlation matrix 422 among rs-fMRI predictors (Figure 2a) and that the higher the strength of a static correlation across 423 the entire rs-fMRI acquisition, the lower its temporal variability across time windows from the same acquisition⁴⁵. However, examination of the relationships between tvFC and sFC graph metrics with 424 SUVR suggests that a different pattern of correlations emerges, with tvFC-SUVR associations not 425 426 being just the *inverse* of the sFC findings (Figure 2b). Similarly to our findings with [¹⁸F]FDG, tvFC has been previously linked to L-[1-11C]leucine PET, with regions having strong protein turnover 427 428 displaying less temporal variability of their graph properties⁴⁶.

Interestingly, this negative association stands out when compared to CV-ReHo, which is instead positively associated with SUVR, implying that the dynamics of local synchronization vs. global FC may have different neurobiological underpinnings: this is in line with a rs-fMRI study which showed that, unlike with FC, static ReHo and time-varying ReHo are *positively* correlated⁴⁷.

Additionally, a strong negative relationship is found between SUVR and the number of BOLD pseudo-events (peaks-BOLD), which is related to the interpretation of the BOLD signal as a point process, with sparse neural events governing its dynamics⁴⁸. While puzzling at first, one interpretation might come from considering that higher local oxygen consumption by active neurons is associated with decreased positive BOLD fluctuations⁴⁹, and therefore the higher the number of BOLD peaks

438 and extreme events, the lower the oxidative metabolism and SUVR might be in that region.

439 We then examined how these relationships would be modulated by selecting parcels according to their ranking in the SUVR distribution. In order to better probe the spatial relationship between SUVR 440 and rs-fMRI, which is heterogeneous across the brain^{23,28}, we chose to explore the changes in 441 correlations selecting nodes from the SUVR standpoint, instead of according to FC properties^{24,25}. 442 443 Interestingly, nodes with progressively higher SUVR, which are expected to be the richest in terms 444 of receptor density, local activity and inter-regional communications^{1,3,4}, did not show different 445 relationships between metabolism and rs-fMRI, but the correlations with most rs-fMRI features 446 became significantly stronger when considering nodes with progressively lower SUVR (Figure 3).

447 This finding suggests that only in nodes with lower metabolism is the [¹⁸F]FDG-fMRI relationship emphasized, implying the presence of a nonlinear association for most of the rs-fMRI features, not 448 449 just for the previously explored metrics^{25,28}, with high SUVR nodes remaining unexplained by the 450 available features. This nonlinear association was also assessed through model selection, with either 451 an exponential, a power law or a log-linear relationship attributed to the majority (86%) of the 452 evaluated bivariate associations (see Supplementary Results). The nonlinearity of the coupling 453 between glucose consumption and BOLD is partly expected: 1) known nonlinearity exists in the associations between BOLD and neuronal activity¹⁸, to which glucose metabolism is instead linearly 454 455 related⁷; 2) nonlinear models such as power laws are common in biology, and in particular in metabolic budget²⁵; 3) the $[^{18}F]FDG$ coupling with local and global FC^{25,28} has been previously 456 described with a power law within specific areas; 4) nonlinear relationships between cerebral blood 457 458 flow and glucose metabolism have been reported⁵⁰.

459

460 *The multivariable multilevel model*

461 To our knowledge, this is the first study to investigate the [¹⁸F]FDG-fMRI coupling using a 462 multivariable approach, attempting to identify the best subset of metrics, among a wide range of 463 fMRI-derived variables, to explain SUVR variability across regions. Moreover, to fully capitalize on 464 the fact that PET and fMRI data were acquired in the same subjects, we employed a multilevel 465 modelling approach, with the selection of the best features performed at the group (higher) level, and modelling performed at the individual (lower) level, to characterize the between-subject variability 466 467 of the SUVR-fMRI association (Figure 4, top). The selected model consisted of nine rs-fMRI 468 variables (Figure 5 a, b) which represented all pools of features: signal (ApEn-BOLD, rApEn-469 BOLD, peaks-BOLD, ReHo, CV-ReHo), HRF (hrf-LE), sFC (s-BC), tvFC (CV-BC), PC (med-Leig). 470 The strongest predictors are related to the BOLD signal and its local synchronization properties 471 (peaks-BOLD, ReHo), which consistently emerged as relevant across all feature selection methods 472 (Supplementary Results). The fact that the SUVR-fMRI spatial coupling is emphasized when local

- BOLD variables are involved might reflect the interplay between excitatory and inhibitory neural
 populations⁵¹, which regulate CBF, a main ingredient in many fMRI-related features^{17,18,52,53}.
- 475 Overall, the explanatory power provided by BOLD rs-fMRI reached a 40% of the SUVR variance at
- the group level (24% across subjects). Zones of polarization in the model residuals emerged in subcortical (putamen), posteromedial (PCC), and lateral frontal regions, which could mainly be attributed to outliers with higher metabolism (**Figure 5c**), which are poorly explained by the available rs-fMRI features in a consistent manner across subjects (**Figure 5d**). These results point to the idea that the BOLD signal and FC, even though related to CBV, CMRO₂ and CBF^{17,18}, reflect the metabolic architecture established by [¹⁸F]FDG SUVR only partially, even in simultaneously acquisitions, and that rs-fMRI FC and its graph metrics cannot be considered a proxy of glucose
- 483 metabolism.

484 Moreover, the individual model R^2 values were variable across subjects, highlighting the fact that the 485 SUVR-fMRI relationship displays significant between-subject variability, with subjects whose 486 BOLD signal and FC architecture are more related to SUVR, and others that have hardly any 487 relationship.

- Next, we used the multilevel modelling approach and the identified predictors to characterize
 between-network variability, exploiting the fMRI-derived RSNs to group the individual data in a
 network-by-network fashion (Figure 4, *bottom*).
- Importantly, the rs-fMRI predictors selected for the between-subject model proved to still be relevant
 for evaluating the between-network SUVR-fMRI association, but their ranking, as assessed by
 dominance analysis, changed noticeably (Figure 6b), with static and dynamic FC features (CV-BC
 in particular) gaining importance in the model.
- 495 Moreover, when the network-wise effects are considered, significant positive and negative 496 associations emerge for each of the nine predictors (Figure 7a). These patterns of predictors have 497 some degree of similarity across networks, with a cluster of RSNs (subcortical, limbic, salience, 498 dorsal attention etc.) being highly correlated, which implies they have a similar SUVR-fMRI 499 multivariable association pattern (Figure 7c). Other networks, instead, seem to be more isolated in 500 their SUVR-fMRI coupling (default mode, visual, somatomotor etc.), possibly reflecting their 501 enrichment in high SUVR nodes (DMN, VIS) that are more difficult to explain using fMRI features 502 (Figure 5c).
- 503 These findings add to and enrich previous work highlighting between-network variability in the 504 SUVR-fMRI association through bivariate associations^{23,28}.

505 Notably, the regions where SUVR-fMRI *bivariate* correlations are higher (Figure 3) seem to fall into

506 networks with fairly high R^2 values in the *multivariable* model (Figure 6c), confirming that the

507 SUVR-fMRI coupling is emphasized in these regions. It is also important to underline that marked

regional heterogeneity has also been described in the coupling between [¹⁸F]FDG SUVR and local

509 CBF⁵⁴, which is likely to underlie some of the variability detected here due to CBF contributions to

- 510 rs-fMRI variables.
- 511

512 *Limitations*

A comprehensive understanding of the relationship between [¹⁸F]FDG PET and rs-fMRI will require 513 simultaneous assessment of other features, such as CBF and CMRO₂, ideally probed by [¹⁵O]H₂O 514 PET⁵⁵ and [¹⁵O]O₂ PET⁵⁶ respectively, which would thoroughly describe the effects of 515 516 hemodynamics and oxidative and non-oxidative metabolism, possibly bridging the gap between these 517 measures. Additionally, while the dataset employed here consists of standard rs-fMRI acquisitions 518 (single-echo, TR of 2s, voxel size 3-4 mm, duration ~10 minutes), the BOLD-[¹⁸F]FDG coupling 519 would likely be improved by more advanced rs-fMRI methods (e.g. multi-echo imaging⁵⁷), recordings 520 of physiological variables (respiratory volume, end-tidal CO_2 and heart rate)²⁰, and regression of the CBF contribution⁵² out of the BOLD signal and local coherence features. 521

For what concerns [¹⁸F]FDG PET, it must also be remembered that SUVR, which was employed here 522 as well as in all the literature on [18F]FDG-fMRI coupling^{22,23,25,27,28}, may offer a biologically 523 confounded view of glucose consumption: SUVR is in fact a semi-quantitative and relative index, 524 which results from interactions between the rate constants of the [¹⁸F]FDG compartmental model, 525 526 i.e., K_1 (ml/cm³/min), describing tracer uptake through the blood-brain barrier, k_2 (min⁻¹), describing its efflux into the venous blood, and k₃ (min⁻¹), quantifying the phosphorylation rate of the hexokinase 527 in neurons and glia^{1,2}. There is therefore the possibility that some of the contribution of CBF to the 528 529 ¹⁸F]FDG-fMRI coupling comes from SUVR, which is in fact highly correlated with the early, CBFrelated frames of [¹¹C]PiB PET in healthy controls⁵⁹, as well as with PET-derived CBF estimates⁵⁴. 530 531 It is therefore likely that PET kinetic modelling will help disentangle the biological processes 532 underlying both BOLD rs-fMRI and static PET estimates.

533

534 Conclusion

535 In conclusion, we thoroughly investigated for the first time the spatial relationship between [¹⁸F]FDG 536 SUVR and a wide range of features derived from rs-fMRI (pooled into 1) signal, 2) HRF, 3) sFC, 4) 537 tvFC and 5) PC-based features) using simultaneous PET/fMRI data. Selection of low SUVR parcels led to a strengthening of SUVR-fMRI associations, implying the presence of a nonlinear relationship 538 539 for many features. Moreover, a novel multivariable multilevel modelling framework was employed 540 to identify the best subset of rs-fMRI predictors able to explain regional SUVR variance, highlighting 541 that predictors based on the BOLD signal and its local synchronization (ReHo and BOLD pseudoevents, in particular) are the ones that are more tightly related to [¹⁸F]FDG SUVR across brain 542

- 543 regions. This suggests a local contribution of CBF that should be tested for and, possibly, regressed
- 544 out from the BOLD signal.
- Notably, the overall explanatory power provided by rs-fMRI on the regional metabolic variability did not exceed 40% of the variance at the group level, with significant variability across subjects. When multilevel modelling of the SUVR-fMRI coupling was carried out across networks, the selected predictors were still relevant for description of RSN metabolism, but noticeable variability across networks was present: new positive and negative associations emerged, and sFC and tvFC network
- 550 features gained importance. In conclusion, SUVR variability across parcels is only partly expression
- 551 of brain network organization described by rs-fMRI.
- 552
- 553

- 554 List of abbreviations:
- 555 ALFF, amplitude of low frequency fluctuations
- 556 ApEn, approximate entropy
- 557 BC, betweenness centrality
- 558 BNV, between-network variability
- 559 BOLD, blood oxygen level dependent
- 560 BSV, between-subject variability
- 561 CBF, cerebral blood flow
- 562 CBV, cerebral blood volume
- 563 CMRO₂, cerebral metabolic rate of oxygen
- 564 CTR, control network
- 565 CV, coefficient of variation
- 566 DAN, dorsal attention network
- 567 DMN, default mode network
- 568 FC, functional connectivity
- 569 FDG, fluorodeoxyglucose
- 570 FDR, false discovery rate
- 571 fMRI, functional magnetic resonance imaging
- 572 HRF, hemodynamic response function
- 573 LE, local efficiency
- 574 LIMBIC, limbic network
- 575 LME, linear mixed-effect
- 576 MAD, median absolute deviation
- 577 med-LEig, median of Leading Eigenvectors
- 578 OLS, ordinary least squares
- 579 NAD, naïve average data
- 580 NNLS, non-negative least squares
- 581 PC, phase coherence
- 582 PET, positron emission tomography
- 583 rApEn, range approximate entropy
- 584 ReHo, regional homogeneity
- 585 rs-fMRI, resting-state fMRI
- 586 RSN, resting-state network
- 587 SAL/VAN, salience/ventral attention network
- 588 SE, standard error

- 589 sFC, static FC
- 590 SM, somatomotor network
- 591 STR, strength
- 592 SUB, subcortical regions
- 593 SUVR, standard uptake value ratio
- 594 TEMP/PAR, temporo-parietal network
- 595 tvFC, time-varying FC
- 596 VIS, visual network
- 597
- 598
- 599
- 600

601

Methods

602 603

604 **Data and Imaging protocols**

The dataset includes 26 healthy subjects from two studies: 11 subjects (8 males; 52.2 ± 10.4 years), 605 hereby referred to as dataset A (Munich)²³, and 15 subjects (6 males; 64.7 ± 7.9 years), i.e., dataset B 606 (Naples)³¹. Subjects were scanned in *eves open* condition while looking at a fixation cross. Exclusion 607 608 criteria included a history of psychiatric or neurological disorders, use of psychoactive drugs or treatment with CNS-active medications, pregnancy, and MR-related contraindications. The subjects 609 provided their informed written consent according to the Code of Ethics of the World Medical 610 Association 611 and the Institutional Review Board and Ethics Committee at the Technische Universität München, for dataset A, and the SDN Foundation, for dataset B. Both 612 centers simultaneously collected [¹⁸F]FDG PET and rs-fMRI data accompanied by a structural MR 613 614 image on two identical Biograph mMR 3T scanners (Siemens Healthcare, Erlangen, Germany) equipped with the standard-supply head-neck coil (12-channel). 615

616 *Dataset A:* MRI data consisted in a structural magnetization prepared rapid acquisition gradient echo 617 (MPRAGE) T1-weighted (T1w) image (TR/TE = 2300/2.98 ms, FA = 9°, 1 mm isotropic voxel size 618 with 0.5 mm gap), 300 volumes of T2*-weighted gradient-echo echo-planar imaging (GE-EPI) with 619 TR/TE = 2000/30 ms and voxel size of 3 mm isotropic (0.6 mm inter-slice gap). PET acquisition 620 consisted in a saturated list mode (10 min duration), started 30 minutes post-injection and 621 reconstructed with voxel size of $3.7 \times 2.3 \times 2.7$ mm³.

- *Dataset B*: MRI data consisted in a similar T1- MPRAGE structural image and 240 volumes of GE EPI for rs-fMRI with 4 mm isotropic voxel and TR/TE = 1920/32 ms. Simultaneous PET/fMRI
 measurements started 30 min post-injection, and PET images were acquired in sinogram mode for 15
- 625 min with reconstruction voxel size of $1.12 \times 1.12 \times 2.0$ mm³.
- The interested reader should refer to the respective papers^{23,31} for more detailed information on each
 dataset.
- 628
- 629

630 Data pre-processing

631 All subjects were identically pre-processed to obtain local metabolism information from [¹⁸F]FDG

632 PET data, and BOLD-based measures from rs-fMRI data, employing a pipeline similar to the Human

- 633 Connectome Project (HCP) minimal preprocessing pipeline⁶⁰ with the addition of PET processing.
- 634

635 Structural imaging pre-processing

Structural T1w images were N4 bias field-corrected⁶¹, skull-stripped, and segmented into grey matter 636 (GM), white matter (WM) and cerebrospinal fluid. The brain cortex was delineated with Freesurfer 637 (recon-all volume and surface reconstruction pipelines)⁶², obtaining pial and GM-WM interface 638 639 surfaces. Manual editing was performed to correct for surface delineation errors. Generated surfaces were resampled over the fs LR mesh provided by Conte69 atlas (symmetric-hemisphere mesh of 32k 640 nodes) to obtain aligned cortical surfaces for each subject. The Schaefer functional atlas³² was used 641 to parcellate cortical surfaces into 200 parcels, grouped according to Yeo's 17 RSNs scheme¹² into 642 643 Central Visual (VIS(A)), Peripheral Visual (VIS(B)), Somatomotor A (SM(A)), Somatomotor B 644 (SM(B)), Temporal Parietal (TP), Dorsal Attention A (DAN(A)), Dorsal Attention B (DAN(B)), 645 Salience/Ventral Attention A (VAN(A)), Salience/Ventral Attention B (VAN(B)), Control A (CTR(A)), Control B (CTR(B)), Control C (CTR(C)), Default Mode A (DMN(A)), Default Mode B 646 647 (DMN(B)), Default Mode C (DMN(C)), Limbic A (L(A)) and Limbic B (L(B). The cortical regions 648 were supplemented by 18 subcortical regions extracted from Freesurfer (bilaterally: Caudate, 649 Putamen, Accumbens, Pallidum, Amygdala, Hippocampus, Thalamus, Ventral diencephalon, Cerebellar cortex) and delineated in single-subject space employing the Multi-Atlas Label Fusion 650 651 (MALF) method³³. Parcels corresponding to subcortical regions were assigned to the Subcortical (SUB) group. 652

653

654 PET data pre-processing

[¹⁸F]FDG PET images, after normalization to injected dose and subject's body weight, were linearly
 resampled in T1w space with FSL's *flirt*⁶³ and intensity-normalized by the whole-brain average
 uptake⁶⁴:

658
$$SUV = \frac{[{}^{18}\text{F}]\text{FDG concentration [MBq/ml]}}{\frac{\text{injected dose [MBq]}}{\text{body weight [kg]}}}$$
 $SUVR = \frac{SUV_{\text{target}}}{SUV_{\text{reference}}}$

659 SUVR maps were then parcellated in the same way as the rs-fMRI data, and parcel-wise SUVR was 660 computed as the median value of the vertices inside a region. All pre-processing steps avoided any 661 further spatial smoothing on both [¹⁸F]FDG and rs-fMRI data.

- 662
- 663 Functional MRI data pre-processing

The first four rs-fMRI volumes were discarded to avoid non-equilibrium magnetization effects. The remaining volumes were corrected for slice timing difference by realigning them to the median volume, using FSL's *mcflirt*⁶³. A template EPI volume was obtained with *antsBuildTemplate*⁶⁵ from realigned rs-fMRI data and used to estimate an affine transform (*flirt*, FSL), subsequently employed to map main tissue segmentations obtained from the pre-processed T1w image to the native EPI

- 669 space. Nuisance signals consisted in motion traces and their first order derivatives complemented by 670 the first five temporal principal components, obtained after principal component analysis of WM and CSF EPI signals, explaining 70% and 50% of the average variance across subjects⁶⁶, which were 671 regressed out from all brain voxels in native EPI space⁶⁷. Regression residuals were finally resampled 672 673 first to the T1w space and then on top of the mid-thickness cortical surface mesh with Connectome Workbench⁶⁸. Finally, the BOLD signal was high-pass filtered with a cut-off of 0.008 Hz. No low-674 pass filter was applied, as the higher frequency components (0.1-0.25 Hz) of BOLD are likely to 675 provide relevant neural information⁶⁹. The vertex-wise BOLD signal was parcellated according to the 676 677 Schaefer cortical atlas and the supplementary subcortical MALF parcels as previously described.
- 678

679 rs-fMRI feature extraction

Feature extraction as well as subsequent analyses were performed in MATLAB (ver. 2020a, TheMathworks, Natick, MA).

682 50 different features were extracted from the BOLD signal, either at the vertex or the parcel level.

683 The extracted features were chosen as descriptors of different aspects of the BOLD 1) signal, 2) HRF,

684 3) sFC, 4) tvFC, and 5) PC. A list of the features and their acronyms is reported in **Table 1**. A detailed 685 description of the features and how they were extracted can be found in the **Supplementary**

- Methods. The extracted features were then employed first in a bivariate correlation analysis against SUVR, and then in a multivariable multilevel modelling procedure to verify how much SUVR variance could be explained across the whole brain at the group level, as well as accounting for individual level information.
- 690

691 Bivariate analysis of the metabolism-fMRI relationship

692 rs-fMRI features vs. glucose metabolism across all brain regions

The bivariate relationship between node-wise SUVR and rs-fMRI properties was assessed at the group level (naïve average data approach, NAD), employing the region-wise median values across subjects for SUVR and each of the 50 extracted features. The association between fMRI-derived features and metabolism across nodes was separately tested via Spearman's rank bivariate correlation (significance level 0.05, corrected for multiple comparisons using the Benjamini-Hochberg FDR approach³⁶).

699

700 rs-fMRI features vs. metabolism in specific clusters of nodes

The spatial heterogeneity in the [¹⁸F]FDG PET-fMRI relationship, which has previously been reported^{23,25}, was probed by selecting clusters of nodes with increasingly high or increasingly low SUVR. The threshold level was determined by considering linearly increasing percentiles of the

704 SUVR distribution over all nodes, in the range going from the 1st to 85th percentiles, with step 1 (from 705 218 up to 33 nodes); moreover, in the opposite direction, nodes were selected according to linearly decreasing percentiles of SUVR, from the 100th to the 15th percentile (from 218 down to 33 nodes). 706 707 Selected nodes at every level are reported in the binary matrix in Supplementary Figure S1. For 708 each threshold level, Spearman's correlation between SUVR and all fMRI-derived features was 709 calculated across the selected nodes, and FDR-corrected for multiple comparisons across thresholds 710 and rs-fMRI features (significance level 0.05)³⁶. The absolute values of Spearman's correlation were 711 summed across the 50 fMRI variables for each percentile, to determine which threshold had the 712 maximum correlation across features. 713

714

715 Multivariable modelling of the SUVR-fMRI relationship at the group level

The relationship between SUVR and each of the 50 rs-fMRI properties ($fMRI_{ip}$, for i = 1, ..., 218regions, and p = 1, ..., 50) was then tested with four different bivariate models:

718 1) a linear model,

719

 $SUVR_i = \alpha_p + \beta_p \cdot fMRI_{ip}$

720 2) a mono-exponential model,

 $SUVR_i = \alpha_n \cdot e^{\beta_p \cdot fMRI_{ip}}$

722 3) a power law model,

723

721

 $SUVR_i = \alpha_p \cdot fMRI_{ip}^{\beta_p}$

4) a log-linear model,

725

 $SUVR_i = \alpha_p + \beta_p \cdot \log fMRI_{ip}$

which were compared in terms of their residual sum of squares (RSS). This bivariate model selection
process led to choose the log-linear model for multiple regression modelling (see Supplementary
Results).

At the NAD level, a multiple linear regression approach was employed to verify how much of the group-wise SUVR variance could be explained by the linear combination of different fMRI-based features. The ordinary least squares (OLS) problem was formulated as follows:

732 $y = X\beta + \varepsilon$

where y and ε are $n \times 1$ vectors of the response/dependent variable (i.e., SUVR) and the model error, and $X \in \mathbb{R}^{n \times p}$ is the matrix of p regressors (i.e., log-transformed rs-fMRI predictors), or design matrix. Before performing OLS regression, all predictors were z-scored, i.e., centered and scaled by their standard deviation (SD). The outcome variable, i.e., SUVR, was z-scored as well, so no model intercept needed to be estimated. The solution to the OLS problem was obtained as

$$\hat{\beta} = (X^T X)^{-1} X^T y$$

739 The model design matrix consisted of 50 parameters. The model was formulated as follows:

740 $SUVR_i = \beta_1 \cdot \log fMRI_{i1} + \beta_2 \cdot \log fMRI_{i2} + \dots + \beta_p \cdot \log fMRI_{ip} + \varepsilon_i$

for each observation i = 1, ..., n. The relationships amongst the predictors were evaluated by Spearman's correlation (**Figure 2**), to assess the presence of strong correlations (i.e., multicollinearity). Since high multicollinearity amongst predictors is known to result in lower precision, switched signs of the coefficients, and a lack of statistical significance of the multivariable model³⁵, the ill-conditioning of the design matrix was quantified using the condition number, i.e.,

746
$$\kappa(X) = \frac{\sigma_{\max}(X)}{\sigma_{\min}(X)}$$

747 with $\sigma_{\text{max}}(X)$ and $\sigma_{\text{min}}(X)$ as the highest and lowest singular values of *X*, respectively. As a rule of 748 thumb, $\kappa(X)$ requires attention if higher than 30^{35} .

The OLS fit was obtained with all the rs-fMRI variables and interpreted as the highest possible predictive power that could be extracted from the available features. However, it is well-known that, in the case of overparameterized linear models, OLS is generally not useful, as many CVs%, (i.e., percent error variance divided by the absolute value of the parameter estimates) are too high (CVs% > 100%) and the model is not *a posteriori* identifiable, so it should be rejected³⁷. As discussed, performing feature selection at the individual level would lead to unstable estimates, so we continued to work at the group (i.e., NAD) level.

Eleven feature selection strategies, namely non-negative least squares (NNLS), elastic net regression, hierarchical clustering, stepwise selection, and general-to-specific modelling in different combinations were tested and compared to identify the best group of features to explain SUVR variability at group level. More details are in (Supplementary Figure S4, Supplementary Table S1, Supplementary Methods).

761

762 Full hierarchical modelling of the SUVR-fMRI relationship

As a NAD approach like the one described so far is statistically sound and unbiased only in case of low between-subject variability, a multilevel population modelling approach (mixed-effect model) was employed in order to characterize in a single stage both the group-level (fixed) and individuallevel (random) effects³⁰ contributing to the relationship between the selected rs-fMRI variables and SUVR (**Figure 4**, *top*). First, the link between model and SUVR was described at individual level by the following equation:

769 770 $y_{Si} = F_{Si}(X_{Si}, \psi_{Si})$ $z_{Si} = y_{Si} + v_{Si}$

with y_{Si} as the SUVR model prediction for the *i*th subject (i = 1, ..., m), which is a function of X_{Si} (the fixed-effects design matrix composed by the features extracted from the rs-fMRI data of

subject *i*), and the parameters to be estimated for subject *i*, ψ_{Si} ; z_{Si} is the vector of the measured SUVR data of subject *i* and v_{Si} is the within-subject variability, or residual unexplained variability, assumed to be normally distributed with zero mean and variance σ_i^2 .

Second, at population level, ψ_{Si} was described by a function combining population parameters (or fixed effects, θ_S), and random variability of individual parameters around the population mean (or random effects, η_{Si}), according to the following assumptions:

779 $\eta_{Si} \sim N(0, \Omega_S)$

where η_{Si} is assumed to be Gaussian, with zero mean, independent across individuals and with covariance matrix Ω_S (another population parameter); as a consequence, ψ_{Si} have a normal distribution as well. The matrix Ω_S was assumed to be full.

 $\psi_{Si} = \theta_S + \eta_{Si}$

The intra-individual (first level) model structure was composed by the nine features selected with the NAD approach, here at single-subject level. Data normalization was performed within subjects via zscoring. The inter-individual model (second level) describing the BSV of the parameters was set according to the aforementioned assumptions.

This estimation requires solving the penalized least squares problem, i.e., the penalized weighted
residual sum of squares (PWRSS),

790 $PWRSS(\Omega_{S}, \theta, y_{Si}|Z_{Si}) = WRSS(\Omega_{S}, \theta_{S}, y_{Si}|Z_{Si}) + ||y_{Si}|Z_{Si}||^{2}$

with Z_{Si} as the random-effects design matrix. This nonlinear optimization problem was solved using the restricted maximum likelihood (REML) estimation method⁷⁰. The standard errors (SE) were calculated for each θ_S parameter estimate as the square root of the diagonal of their covariance matrix. The overall and subject-wise multilevel model R² were also evaluated. The residual unexplained variability v_{Si} was evaluated by calculating its median and variability (CV%) across subjects.

The hierarchical modelling approach was also performed across networks (*N*) in order to characterize between-network variability. RSNs were used as the grouping factor instead of subjects in a model formulated as follows:

799

800

$$y_{Nj} = F_{Nj}(X_{Nj}, \psi_{Nj})$$
$$z_{Nj} = y_{Nj} + v_{Nj}$$

801 with *j* as the *j*th network (j = 1, ..., q). Normalization of SUVR and rs-fMRI variables was performed 802 via z-scoring within RSNs.

803 The random effects η_{Nj} and the resulting individual parameters ψ_{Nj} were evaluated in terms of their 804 correlation structure, both across RSNs (1st dimension) and across the nine predictors (2nd dimension), 805 as seen in (**Figure 7b, c**).

807

808 **Relative importance analysis to determine predictor importance**

Relative importance analysis^{40,71} was employed as a supplement to the results of hierarchical 809 modelling. This type of analysis allows to appropriately partition the model's explained variance 810 811 amongst multiple predictors when there is still significant multicollinearity, which makes typical indicators of importance (e.g., standardized regression coefficients) flawed. Dominance analysis 812 (DA), in particular, works by rank-ordering the predictors in term of relative importance by 813 comparing the additional contributions they make to the R² of all possible subset models. Specifically, 814 we assessed the general dominance of the variables, which is established for one predictor over 815 816 another when the average of its conditional contributions over all model sizes is greater than that of 817 the other. The obtained general dominance weights are also measures of relative effect sizes, as they sum to the model R²: the percent contribution to the model R² was therefore calculated and reported. 818 819 While DA was originally proposed for OLS models, it was later extended to multilevel models⁴⁰. In 820 order to apply DA to hierarchical models, a null model with no predictors must be provided, and the 821 slopes of first-level models must be considered fixed even when they are random in the identified 822 model, to simplify dominance evaluation. DA was used to assess the extent to which each selected 823 variable was driving the prediction in the context of the LME models across subjects (S) and across 824 networks (N), as they were still affected by non-negligible multicollinearity. 825

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- 826
- 827

829 Data availability

- The multimodal [¹⁸F]FDG PET and rs-fMRI data used in the present study can be accessed via request to the groups who performed the original studies^{23,31}. Data sharing will be subject to the policies and procedures of the institution where each dataset was collected.
- 833 The codes and processed data that support the conclusions of this research work can be accessed via834 request to the corresponding authors.
- 835

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840

841 Author contributions:

T.V. and A.B. designed the study, performed the analysis, interpretation of the data, and the drafting of the article. E.S. contributed to the preprocessing of the data and to the drafting of the article. M.C. contributed to the interpretation of the data and to the drafting of the article. M.A. designed and collected the data for the original study for dataset B and contributed to revision of the draft. All authors approved the final version of the article to be published.

847

848 **Competing interests:**

- 849 The authors declare no competing interests.
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