

## This is a self-archived version of an original article. This version may differ from the original in pagination and typographic details.

Author(s): Pauls, K. Amande M.; Salmela, Elina; Korsun, Olesia; Kujala, Jan; Salmelin, Riitta; Renvall, Hanna

Title: Human sensorimotor beta event characteristics and aperiodic signal are highly heritable

Year: 2024

Version: Accepted version (Final draft)

Copyright: © 2023 the Authors

Rights: <sub>CC BY 4.0</sub>

Rights url: https://creativecommons.org/licenses/by/4.0/

#### Please cite the original version:

Pauls, K. A. M., Salmela, E., Korsun, O., Kujala, J., Salmelin, R., & Renvall, H. (2024). Human sensorimotor beta event characteristics and aperiodic signal are highly heritable. Journal of Neuroscience, 44(5), Article e0265232023. https://doi.org/10.1523/jneurosci.0265-23.2023



**Research Articles | Systems/Circuits** 

# Human sensorimotor beta event characteristics and aperiodic signal are highly heritable

https://doi.org/10.1523/JNEUROSCI.0265-23.2023

Received: 13 February 2023 Revised: 24 October 2023 Accepted: 3 November 2023

Copyright © 2023 Pauls et al.

This is an open-access article distributed under the terms of the Creative Commons Attribution 4.0 International license, which permits unrestricted use, distribution and reproduction in any medium provided that the original work is properly attributed.

This Early Release article has been peer reviewed and accepted, but has not been through the composition and copyediting processes. The final version may differ slightly in style or formatting and will contain links to any extended data.

Alerts: Sign up at www.jneurosci.org/alerts to receive customized email alerts when the fully formatted version of this article is published.

- 1 <u>Title</u>: Human sensorimotor beta event characteristics and aperiodic signal are highly herita-
- 2 ble
- 3
- 4 Short title: Sensorimotor beta & aperiodic signal heritability
- 5
- 6 K. Amande M. Pauls<sup>1,2</sup>, Elina Salmela<sup>3,4</sup>, Olesia Korsun<sup>2,5</sup>, Jan Kujala<sup>6</sup>, Riitta Salmelin<sup>5</sup>
- 7 Hanna Renvall<sup>2,5</sup>
- Department of Neurology, Helsinki University Hospital and Department of Clinical Neurosciences (Neurology), University of Helsinki, 00029 Helsinki, Finland.
- 102.BioMag Laboratory, HUS Medical Imaging Center, Helsinki University Hospital, 00290 Hel-<br/>sinki, Finland
- Organismal and Evolutionary Biology Research Programme, Faculty of Biological and Environmental Sciences, University of Helsinki, 00014 Helsinki, Finland
- 14 4. Department of Biology, University of Turku, 20014 Turku, Finland
- 155.Department of Neuroscience and Biomedical Engineering, School of Science, Aalto University, 02150 Espoo, Finland
- 17 6. Department of Psychology, University of Jyväskylä, 40014 Jyväskylä, Finland
- 18
- 19 Corresponding author email address: amande.pauls@hus.fi
- 20 Pages: 22, Figures: 3, Tables: 2
- 21 abstract: 243 words; introduction: 645 words; discussion: 1689 words

22

- 23 Conflict of interest statement
- 24 AP, ES, OK, JK, RS and HR report no conflict of interest.
- 25

#### 26 Acknowledgements & funding source

We thank all subjects for participating in the study. We acknowledge the following funding sources: AP has received funding from Helsinki University, the Research Council of Finland (grant number 350242) and the Sigrid Jusélius Foundation. ES has received funding from Jenny and Antti Wihuri Foundation and Ella and Georg Ehrnrooth Foundation. OK is funded by the Instrumentarium Science Foundation and Finnish Cultural Foundation. RS has received funding from the Research Council of Finland (grant numbers 315553 and 355407) and the Sigrid Jusélius Foundation. HR has received funding from the Research Council of Finland
 (grant numbers 127401, 321460 and 355409), Paulo Foundation, and the Finnish Cultural
 Foundation.

36

#### 37 Abstract

Individuals' phenotypes, including the brain's structure and function, are largely determined 38 39 by genes and their interplay. The resting brain generates salient rhythmic patterns that can be characterized non-invasively using functional neuroimaging such as magnetoencephalog-40 41 raphy (MEG). One of these rhythms, the somatomotor ('rolandic') beta rhythm, shows inter-42 mittent high amplitude 'events' that predict behavior across tasks and species. Beta rhythm is altered in neurological disease. The aperiodic ('1/f') signal present in electrophysiological re-43 44 cordings is also modulated by some neurological conditions and aging. Both sensorimotor beta and aperiodic signal could thus serve as biomarkers of sensorimotor function. Knowledge 45 46 about the extent to which these brain functional measures are heritable could shed light on 47 the mechanisms underlying their generation. We investigated the heritability and variability of human spontaneous sensorimotor beta rhythm events and aperiodic activity in 210 healthy 48 male and female adult siblings' spontaneous MEG activity. The most heritable trait was the 49 50 aperiodic 1/f signal, with a heritability of 0.87 in the right hemisphere. Time-resolved beta event 51 amplitude parameters were also highly heritable, whereas the heritabilities for overall beta 52 power, peak frequency and measures of event duration remained nonsignificant. Human sen-53 sorimotor neural activity can thus be dissected into different components with variable herita-54 bility. We postulate that these differences partially reflect different underlying signal generating mechanisms. The 1/f signal and beta event amplitude measures may depend more on fixed. 55 56 anatomical parameters, whereas beta event duration and its modulation reflect dynamic char-57 acteristics, guiding their use as potential disease biomarkers.

58

#### 59 Significance statement

The resting brain shows a prominent, highly modulated beta-range rhythm closely linked to sensorimotor function in health and disease. We investigated the heritability of human spontaneous sensorimotor beta rhythm and its different components in a large cohort of 210 siblings' MEG data. We find that particularly beta event amplitude and its variation as well as aperiodic signal characteristics are highly heritable. The study demonstrates that time-resolved electrophysiological measures of spontaneous human sensorimotor brain activity are determined to a significant degree by genes. We discuss the findings in the context of known

X

and postulated structural underpinnings of MEG signal generation, to highlight their transla tional relevance. The findings have clinical implications, *e.g.*, when considering sensorimotor
 beta alterations as biomarkers of neurological disease.

70

#### 71 Introduction

Individuals' phenotypes are largely determined by their genetic blueprint that regulates prop-72 73 erties ranging from cell products (Barroso and McCarthy, 2019) to system-level brain macro-74 structure (Geschwind et al., 2002; Peper et al., 2007). Genetic influences also underlie func-75 tional brain measures which are constant within, but highly variable between individuals. Elec-76 troencephalography (EEG) and magnetoencephalography (MEG) have been successfully ap-77 plied to quantify the heritability and identify genetic determinants of functional brain measures (Van Beijsterveldt et al., 1996; Smit et al., 2006; Koten et al., 2009; Renvall et al., 2012; van 78 79 Pelt et al., 2012).

80

The brain generates 'background' electrical activity with salient rhythmic, but also arrhythmic 81 82 patterns during wakeful resting. One of the prominent spontaneous rhythms is the somatomo-83 tor (rolandic) beta rhythm (Hari and Salmelin, 1997) that is observed across several mamma-84 lian species (Haegens et al., 2011; Feingold et al., 2015; Sherman et al., 2016). It is modulated 85 by perceptual and cognitive functions, including tactile processing (Pfurtscheller et al., 2001; Haegens et al., 2011), motor function (Salmelin and Hari, 1994; Feingold et al., 2015), action 86 perception (Hari et al., 1998; Babiloni et al., 2002) and attention (Van Ede et al., 2011; Sacchet 87 et al., 2015). Beta band activity is modulated over time, manifesting in intermittent high ampli-88 89 tude 'events' (Feingold et al., 2015; Jones, 2016) relevant for behavior: In the sensorimotor 90 cortex, beta event rate predicts behavior across tasks and species (Shin et al., 2017). Both 91 beta power and beta events are altered in neurological conditions affecting motor function, 92 such as genetically determined Unverricht-Lundborg disease (Silén et al., 2000), stroke (Laaksonen et al., 2012) and Parkinson's disease (Vinding et al., 2020; Pauls et al., 2022). 93

94

Besides rhythmic, or periodic, components, MEG power spectra also contain aperiodic ('1/f') components (He, 2014). These two are important to disentangle as they are probably generated by different neural mechanisms. Aperiodic signal is believed to represent excitation-inhibition balance (Gao et al., 2017), and it is modulated, *e.g.*, by brain maturation (McSweeney et al. 2021; Hill et al. 2022), aging (Voytek et al., 2015; Wilson et al., 2022) and several neurological and psychiatric conditions (Molina et al., 2020; Ostlund et al., 2021; Semenova et al.,

X

101 2021). Cortical beta rhythm (Laaksonen et al., 2012; Pauls et al., 2022) and aperiodic activity 102 (Helson et al., 2023) both relate to clinical symptoms, show good or excellent test-retest reli-103 ability (Pauls et al., 2023, bioRxiv), and thus have potential as diagnostic or prognostic bi-104 omarkers.

105

106 Interpretability of rhythmic and aperiodic neural signals is important for both research and 107 clinical diagnostic applications. MEG signal arises from spatial and temporal summation of 108 underlying neuronal activity (Buzsáki et al., 2012). Structure and function are closely related: 109 e.g., peak oscillation frequency decreases with increasing cortical thickness and processing 110 hierarchy (Mahjoory et al., 2020). Decoding the structure-function-genetics relationship of M/EEG signal generation could help understand signals' individuality and their degradation in 111 112 neurological diseases, raising their value as diagnostic tools: M/EEG may detect pathology 113 before observable structural changes in neurological disorders (Terry et al., 1991). Heritability reflects the contribution of genetic vs. environmental factors to the differences observed be-114 115 tween individuals, and the quantification of the heritability of neural signals can thus lead to 116 insights of the biology behind the measurable phenotypes (Visscher et al., 2008). Beta and 117 other frequency bands' global spectral power is heritable (Van Baal et al., 1996; Van Beijsterveldt et al., 1996; Smit et al., 2005; Salmela et al., 2016); the beta power variability has been 118 119 linked to a GABA<sub>A</sub> receptor locus (Porjesz et al., 2002). Heritability of time-resolved beta 120 events, however, has not been investigated.

121

We investigated the heritability and variability of time-resolved human cortical sensorimotor beta rhythm and aperiodic activity using healthy adult siblings' spontaneous MEG data. We propose that knowledge about the relative heritability of different neural components of sensorimotor activity can shed light on the underlying generating mechanisms and help interpret changes observed in, *e.g.*, patient populations with sensorimotor dysfunction.

127

#### 128 Materials and methods

#### 129 Subjects

130 210 Finnish-speaking siblings from 100 families participated in the study (8 families with three 131 siblings, 1 family with four; 148 females [mean  $\pm$  SD age 29  $\pm$  10 years, range 18-60 years], 132 62 males [30  $\pm$  9 years, range 19-52 years]; 206 right-handed, three ambidextrous, one left-133 handed). None of the participants had a history of neurological or psychiatric disorders. The study was approved by the Hospital District of Helsinki and Uusimaa ethics committee, and all
 participants gave their written informed consent to participate.

#### 136 MEG recordings

Spontaneous cortical activity was recorded in a magnetically shielded room with a 306-channel Vectorview neuromagnetometer (Elekta Oy, Helsinki, Finland) that contains 204 planar gradiometers and 102 magnetometers. Head positioning was measured at the beginning of the measurement. Three minutes of data was collected while participants were resting with their eyes open (REST), as well as while they clenched both hands alternatingly about once per second, self-paced, keeping the eyes open (MOT). The MEG signals were band-pass filtered at 0.03–200 Hz and sampled at 600 Hz.

#### 144 MEG signal processing and beta event extraction

For suppressing external artifacts, MEG data were preprocessed using the signal space sep-145 146 aration method (SSS, (Taulu and Simola, 2006)) implemented in MaxFilter software (MEGIN 147 Oy, Helsinki, Finland). Individual MEG recordings were transferred to one subject's head space using a signal space separation based head transformation algorithm (Taulu et al., 148 149 2004), implemented in MaxFilter. Further signal processing was done using MNE-python version 0.22 (Gramfort et al., 2013). After band-pass filtering the data to 2-48 Hz with a one-pass, 150 151 zero-phase, non-causal FIR filter (MNE firwin filter design using a Hamming window), power 152 spectral density (PSD) was calculated using Welch's method (MNE's psd welch function) with 153 a non-overlapping Hamming window and 1024-point Fast Fourier Transformation (FFT).

The subsequent analysis steps are illustrated in **Figure 1**. The data analysis was performed on the 204 gradiometer signals. First, a channel pair with the highest spectral peak in the beta range ('the peak channel pair') was selected from the region of interest (ROI) of 15 gradiometer channel pairs per hemisphere centered over the sensorimotor cortices, and the frequency at the power peak noted ('peak beta frequency') (see **Figure 1A**). In order to quantify PSD at each recording site, we computed the vector sum of the two orthogonally oriented planar gradiometers at each sensor location ('vector PSD'):

161 PSD<sub>vector</sub> =  $\sqrt{(PSD_{ch1}^2 + PSD_{ch2}^2)}$ 

The resulting 15 vector-sum PSDs per hemisphere were then decomposed into a periodic and aperiodic component using FOOOF (Donoghue et al., 2020). FOOOF models the power spectrum as a combination of two distinct functional processes: an aperiodic component, reflecting 1/f like characteristics (exponential decay with an offset and an exponent), and a variable number of periodic components (putative oscillations), as peaks rising above the aperiodic 167 component. After subtraction of the aperiodic component, the remaining periodic component
 168 was plotted for all 15 vector-sum PSDs for both REST and MOT conditions in the frequency
 169 range of 14-30 Hz. The resulting plots were visually inspected by two observers (AP and OK)
 170 to manually select the beta signal frequency modulated most by MOT compared to REST.

171 As the manual channel selection may be prone to human observer bias, we compared the 172 inter-rater agreement between two slightly different approaches, conducted independently 173 years apart on the same data. The peak beta band frequencies had previously been extracted 174 by one of the authors (HR) without separating the 1/f aperiodic signal part and by using 175 Welch's method with 4096-point FFT, eight data segments overlapping by 50% and Hamming 176 windowing. When allowing deviation of +/- 3 Hz in the extracted peaks (taken the different FFT 177 sizes and different handling of the aperiodic 1/f component), the two approaches resulted in 178 85% agreement, which is considered good.

Using the manually selected peak frequencies, the periodic components of the 15 vector-sum
PSDs were searched automatically to determine the recording channel with the highest peak

and its frequency (± 1 Hz) for both hemispheres' ROIs, and visually inspected again by AP.

The peak beta frequency and corresponding peak power of the chosen vector-sum PSD, the total beta band power (periodic part of PSD area under curve (AUC) from 14-30 Hz, 1/f component subtracted), as well as the aperiodic component information obtained via FOOOF (offset and exponent chi), were further used in the heritability analysis. All electrophysiological parameters included in the heritability analysis are illustrated in **Figure 1C**.

187 The channel pair and peak beta frequency corresponding to the chosen vector-sum PSD were 188 used for beta burst analysis (see **Figure 1B**). Beta event extraction was carried out similarly 189 to the method described in Pauls et al. (2022): the channel pair's raw unfiltered time series 190 data were downsampled to 200 Hz, high-pass filtered at 2 Hz and decomposed by convolving 191 the signal with a set of complex Morlet wavelets over the frequency range of 7-47 Hz with 1 192 Hz resolution and n\_cycles=frequency/2. The signal was then averaged within the individual narrow-band beta frequency range, *i.e.*, ± 1.5 Hz around the individual peak beta frequency, 193 194 discarding the other frequencies. The vector sum over the two channels' beta band time series 195 was calculated as described above, and the resulting signal was rectified to obtain one beta 196 band amplitude envelope for the channel pair. The envelope was smoothed with a 100-ms 197 FWHM kernel and thresholded at the 75th percentile value. Periods exceeding this threshold 198 for 50 ms or longer were defined as beta events. For event amplitude and event duration, the 199 mean, median, robust maximum (defined as mean of the top 5% values) and standard devia-200 tion values were calculated. Furthermore, events per second (event rate) and event dispersion 201 were calculated similarly to Pauls et al. (2022). Times between beta events were defined as

waiting times. To estimate the variation of waiting times ('event dispersion'), we calculated the coefficient  $C_V$  proposed by Shinomoto et al. (Shinomoto et al., 2005), defined as the waiting times' standard deviation  $\sigma$  divided by their mean  $\mu$ :

205

$$C_V = \frac{\sigma}{\mu}$$

206

All values were calculated for both hemispheres in all subjects (see also Figure 1B).

208 Effect sizes for MEG features were based on Cohen's *d* values for single group designs:

$$209 D = M/S$$

210 where M and S are the mean and the standard deviation of the feature values across subjects

211 (Goulet-Pelletier and Cousineau, 2018).

h<sup>2</sup>= V<sub>genetic</sub>/V<sub>phenotypic</sub>

212 Heritability analysis

Heritability is defined as the proportion of (additive) genetic variance of the total phenotypic variance of a population.

215

216

217

Phenotype heritabilities were calculated using the software program Merlin version 1.1.2 (Abecasis et al., 2002), which employs a variance component approach as detailed by Amos (Amos, 1994). Heritability estimates are calculated based on variance components. The coefficient estimating genetic variance is adjusted by the degree of relationship, which is 0.5 (50% shared genes) in full siblings. The full sibling status of our study individuals has been confirmed by [an earlier] DNA analysis (Renvall et al. 2012).

224

Merlin requires non-negative values for correct interpretation so phenotypes with negative values were multiplied by -1. Such a transformation is standard for the Merlin analysis tool. Correctness of the input data format was checked by the Pedstats program (Wigginton and Abecasis, 2005). As the analysis assumes the studied phenotypes to be normally distributed while many of them were not, we also re-ran the analyses after first correcting the phenotype values' distributions with the inverse normal correction internal to Merlin. As both analyses produced highly concordant results, we report here the results based on the non-correctedvalues.

233

The probability of the observed heritability values being different from zero was assessed by 234 235 permuting the family labels of the study subjects 6000 times and calculating the heritability for 236 each of the permuted datasets. For each phenotype, the number of permutations k where the 237 permuted heritability was higher than the heritability observed in the real data was recorded 238 and used to calculate the one-tailed probability of the observed heritability exceeding zero as 239 k/6000. This permutation scheme may slightly inflate the permuted heritabilities, as it does not 240 explicitly ensure that the permutation does not reproduce any of the original sibships. This may lead to conservative significance estimates. Likewise, to correct for the multiple tests 241 242 performed (n = 30), we performed a Bonferroni correction, which may be overly conservative 243 considering that some of the phenotypes were correlated.

#### 244 Code and data accessibility

These data cannot be made publicly available due to Finnish data protection law. Data can, however, be shared for research collaboration with an amendment to the research ethics permit and a related data transfer agreement. All analysis code is available on GitHub (https://github.com/BioMag/Beta-sibling-study).

#### 249 **Results**

A summary of the beta band phenotypic features (both beta PSD features as well as beta band burst characteristics) is given in **Table 1**. **Figure 2A** shows examples of different beta power spectral phenotypes observed, and **Figure 2B** depicts beta band phenotypes for pairs of siblings. Typical PSD phenotypes were *i*) ones with a narrow peak on either side of 20 Hz, *ii*) a broad band activity typically spanning 15-25 Hz, and *iii*) two distinctive peaks, one typically in the lower beta range (14-20 Hz) and the other in the high beta range (20-30 Hz).

			Left hemisphere				Right hemisphere				
PSD characteristics		mean	median	std	range	mean	median	std	range		
peak beta frequend	cy (Hz)	19.7	19.3	3.0	14.1-25.8	19.8	19.3	3.1	14.1-29.3		
peak beta power (fT/cm) <sup>2</sup>		276	144	374	15-3231	133	70	173	5-1251		
total beta band power (periodic)		2979	1686	3318	164-16284	1093	619	1322	26-9652		
1/f component exponent		1.03	1.01	0.19	0.34-1.76	1.07	1.05	0.18	0.67-1.66		
1/f component offs	et	-22.95	-22.99	0.37	-23.7521.75	-23.28	-23.32	0.34	-24.1922.25		
Beta event characteristics		mean	median	std	range	mean	median	std	range		
duration (ms)	mean	256.9	248.0	49.4	181.7-498.2	265.2	253.7	50.0	182.0-454.0		
	median	199.0	195.0	32.3	152.5-420.0	201.4	195.0	34.9	150.0-355.0		
	standard deviation	198.4	182.4	63.2	99.8-487.5	213.0	198.3	67.4	67.3-531.7		
	robust maximum	858.0	788.6	257.4	439.5-2145.0	912.1	852.2	263.9	382.7-2135.0		
amplitude (ft/cm)	mean	325	285	165	104-994	221	188	113	73-654		
	median	301	260	155	93-984	203	173	105	70-618		
	standard deviation	86	71	48	21-264	62	52	34	14-202		
	robust maximum	564	486	286	174-1467	390	335	196	114-1128		
event rate (1/s)		1.00	1.00	0.16	0.50-1.36	0.97	0.98	0.16	0.55-1.37		
dispersion		1.14	1.05	0.44	0.65-5.59	1.16	1.06	0.44	0.41-5.58		

#### **Table 1 – PSD (beta & 1/f) and beta event descriptives**

Parameters used in the heritability analysis. Peak frequency – frequency between 14-30 Hz most modulated by hand movement; peak power – PSD amplitude at peak frequency; total beta band power
(periodic) – total AUC from 14-30 Hz of the periodic part of the signal (1/f signal component subtracted);
1/f component chi – exponential decay coefficient and offset describing 1/f (aperiodic) signal component. Beta event characteristics: robust maximum – mean of top 5 % values; burst rate – number of
bursts/recording time; dispersion – stdev(inter-burst intervals)/mean(inter-burst intervals).

Heritability results are shown in **Table 2**. Overall, the right-hemispheric parameters were more heritable than the left-hemispheric ones. The right hemisphere's 1/f aperiodic exponent and offset were significantly heritable (exponent h<sup>2</sup>=0.87, offset h<sup>2</sup>=0.69). Measures of beta burst amplitudes were also significantly heritable (range of significant heritability values h<sup>2</sup> of 0.28-0.81). Notably, of the beta burst amplitude measures, the measures reflecting the dynamic range (beta event amplitude maximum and its standard deviation) were most highly heritable. Apart from the peak beta power with moderate effect size in both hemispheres (Cohen's d 0.74-0.77), all effect sizes were either large (Cohen's d > 0.80) or very large (Cohen's d > 1.2).

		Left her	misphere		Right h	Right hemisphere		
PSD characteristics		h²	р	n sig. (/6000)	h²	р	n sig. (/6000)	
peak beta frequ	ency	0.45	0.0047	28	0.41	0.0103	62	
peak beta powe	r	0.28	0.0648	389	0.58	0.0072	43	
total beta band power (periodic)		0.49	0.0068	41	0.44	0.0157	94	
1/f component	f component exponent*		0.0035	21	0.87	0.0000*	0	
1/f component	0.35	0.0258	155	0.69	0.0000*	0		
Beta event characteristics		h²	р	n sig. (/6000)	h²	р	n sig. (/6000)	
duration	mean	0.45	0.1350	810	0.36	0.0222	133	
	median	0.28	0.1338	803	0.40	0.0172	103	
	standard deviation	0.49	0.2495	1497	0.32	0.0412	247	
	robust maximum	0.47	0.2383	1430	0.33	0.0372	223	
amplitude	mean*	0.35	0.0060	36	0.75	0.0002*	1	
	median*	0.45	0.0110	66	0.72	0.0002*	1	
	standard deviation*	0.28	0.0005*	3	0.81	0.0000*	0	
	robust maximum*	0.49	0.0007*	4	0.79	0.0000*	0	
event rate		0.47	0.0543	326	0.38	0.0137	82	
dispersion		0.35	0.2850	1710	0.00	1.0000	6000	

276 277 Table 2. Heritability h<sup>2</sup> of the oscillatory phenotypes calculated by Merlin. The nominal probability 278 that the heritability differs from zero is calculated from an empirical distribution based on 6000 permutations of the sibship statuses/family IDs of the subjects. The variables and values that are significant 279 280 after a Bonferroni correction for multiple testing are given in bold.

281

#### 282 Discussion

To our knowledge, this is the first study investigating the heritability of spontaneous time-re-283 284 solved sensorimotor beta event dynamics and aperiodic neural activity. Time-resolved beta 285 event amplitude parameters were highly heritable, whereas the heritabilities for peak fre-286 guency and measures of event duration were not significantly different from zero. Interestingly, 287 the most heritable trait was the aperiodic 1/f exponent, with a heritability of 0.87 in the right 288 hemisphere. Overall, the right-hemispheric phenotypic traits were more heritable than the lefthemispheric ones. 289

290

#### 291 Heritability of MEG/EEG traits including beta oscillatory activity

292 Heritability of electrophysiological traits has been little investigated to date. In twin studies, 293 EEG alpha, beta, theta and delta range peak frequencies (Van Beijsterveldt et al., 1996), oc-294 cipital alpha power and peak frequency at rest (Smit et al., 2006), as well as MEG visual task-295 related gamma peak frequency (van Pelt et al., 2012) have been found to be highly heritable. 296 We have previously demonstrated that auditory evoked fields' amplitude (Renvall et al., 2012) 297 as well as occipital resting-state alpha oscillatory activity (Salmela et al., 2016) are heritable 298 in siblings, and that MEG power spectral features at rest allow identification of sibling relation-299 ship (Leppäaho et al., 2019). These MEG traits were associated with certain genetic loci / genomic regions (Renvall et al., 2012; Salmela et al., 2016; Leppäaho et al., 2019) but it is
 likely that most functional brain traits are controlled polygenetically. Furthermore, functional
 connectivity in theta, alpha and beta bands as measured with MEG appears progressively
 more similar as the strength of genetic relationship increases (Colclough et al., 2017).

304

#### 305 MEG signal generative mechanisms and possible relation to heritability

306 MEG measures magnetic fields arising from the temporal and spatial summation of electric 307 currents occurring in the underlying brain tissue (Buzsáki et al., 2012). The measured raw 308 signal time series can be summarized in different ways, e.g., as power spectral density. Re-309 duction in global beta power can result from various changes in the neuronal signalling, such 310 as smaller amplitude beta oscillation events, or fewer or shorter beta oscillation events without 311 simultaneous changes in amplitude. Thus, decomposing beta power into components gives additional information about the underlying neural processing. We postulate that these MEG 312 dynamical measures reflect different aspects of MEG signal generation. The upper panel of 313 **Figure 3** schematically summarizes factors that contribute to the generation of MEG signals, 314 and the lower panel indicates how those factors may relate to the functional parameters ad-315 316 dressed in this study.

317

#### 318 What underlies the heritability of beta event amplitude?

319 We postulate that the MEG beta event amplitude reflects relatively fixed anatomical factors 320 summarized in Figure 3 (upper panel, left). Pyramidal cells are neocortex' most abundant cell 321 type. Synaptic currents and their state-dependent modulation are the main determinants of 322 intra- and extracellular field strength, and their spatial summation is governed by pyramidal 323 cell morphology, cortical microstructure and layering, as well as synaptic input density 324 (Buzsáki et al., 2012). Beta event amplitudes are probably crucially dependent on these mi-325 crostructural properties: While both temporal and spatial superposition determine event am-326 plitude, especially the amplitude's dynamic range is limited by local cortical microstructure. 327 Interestingly, in the current study, event amplitudes' dynamic range measures (standard devi-328 ation, maximum) were most strongly heritable.

- Brain anatomical traits such as cortical thickness (Geschwind et al., 2002; Schmitt et al., 2014)
- and cortical myelination (Schmitt et al., 2020) have previously been shown to be heritable. By
- 332 late adolescence, differences in cortical thickness in the sensorimotor regions are largely due

to heritable factors, whereas environmental factors play only a weak role (Schmitt et al., 2014).
Thus, throughout development, sensorimotor cortical structure appears increasingly governed
by the underlying genetics.

336

Both beta peak amplitude and the power at the beta band (which is determined by the amplitude, number and duration of individual beta events) appeared more heritable in the right than left hemisphere. Our result is in agreement with earlier studies that have found cortical morphology/volume to be more genetically controlled in the right than left hemisphere in righthanded individuals (Geschwind et al., 2002); functional studies point in the same direction (Smit et al., 2006).

343

#### 344 Why are event duration parameters not similarly heritable?

In the current cohort, measures of beta event duration were not significantly heritable. Tem-345 346 poral summation of neural events, which determines the timing and duration of beta events, 347 arises from the interplay between several brain areas, their connections and relative input 348 timings and strength (Figure 3, top panel, right). Important cortical pyramidal cell afferent in-349 puts originate from other adjacent pyramidal cells (intrinsic input) (Lorente de No, 1949), cor-350 tico-cortical connections (Kandel et al., 2000) and thalamic connections, including connections 351 from sensory organs, and from other cortical areas ('higher-order' thalamic input) (Sherman et al., 2016; Mo and Sherman, 2019). Computational models suggest that sensory induced 352 353 beta events are generated by synchronous bursts of excitatory synaptic drive to superficial and deep cortical layers, with asymmetry in the respective input strengths (Jones et al., 2009; 354 Sherman et al., 2016; Neymotin et al., 2020): The stronger the superficial input, the more 355 356 prominent is the beta activity (Sherman et al., 2016). Experimental data are compatible with 357 this model (Sherman et al., 2016; Bonaiuto et al., 2021; Law et al., 2022). Thus, beta event 358 timing and duration appear to depend on the timing and strength of inputs from several differ-359 ent cortical and subcortical input sources.

360

Network resonance could also play a role in beta event generation: In a dopamine-depleted state, cortical beta events are associated with increased synchrony between EEG/ECoG cortical activity and basal ganglia spiking activity (Cagnan et al., 2019). In animal models of parkinsonism, high cortical beta synchrony can be generated by changing the relative timings between thalamic and cortico-cortical inputs (Reis et al., 2019). Hence, network resonant properties could contribute to temporal summation at least in some disease states, but possibly in
a dopamine-dependent fashion also in healthy brains.

368

369 Thus, compared to spatial summation, temporal summation relies on more individual factors 370 and their interplay (e.g., network structural and functional properties), making heritability more 371 multifactorial and thus less likely to show heritability in the present analysis. Methodological 372 factors could also contribute to the lack of heritability: signal-to-noise ratio of the recordings 373 affects event duration more than event amplitude measures. Finally, the resting-state beta 374 event duration could be a randomly fluctuating parameter, governed by stochastic events and 375 their timing. These explanations, however, seem less likely given the outlined experimental evidence, as well as our test-retest reliability results (Pauls et al. 2023, bioRxiv). 376

377

#### 378 Why is the aperiodic signal component heritable?

379 Aperiodic signal components were the most heritable of the investigated parameters in the 380 present study. The aperiodic signal is closely related to anatomical microstructure: Cortical 381 pyramidal cells and their dendritic morphology and density are believed to be the most im-382 portant determinants of the mammalian cortical 1/f signal observed with MEG (Lindén et al., 2010; Buzsáki et al., 2012). The 1/f signal is thought to stem from passive dendrite filtering 383 properties (Halnes et al., 2016) but it is also modulated in an activity-dependent way (Petter-384 385 sen et al., 2014). It has been shown to be affected by brain maturation (McSweeney et al. 2021; Hill et al. 2022) and aging (Voytek et al., 2015; Wilson et al., 2022) as well as neurolog-386 387 ical (Semenova et al., 2021) and psychiatric diseases (Ostlund et al., 2021). Furthermore, 1/f 388 reflects the attentional state (Waschke et al., 2021) and may contribute to integration of signals 389 over longer periods of time (Maniscalco et al., 2018). Thus, the signal's relative stability over 390 extended periods of time, and its close relationship to cortical microstructure may explain the high heritability. 391

392

#### 393 Stability of beta events and aperiodic activity - a prerequisite for clinical use

Movement-related beta suppression and rebound at the sensorimotor cortices show excellent test-retest stability over weeks in EEG recordings (Espenhahn et al., 2017). Similarly, beta rhythm modulation after tactile and proprioceptive stimulation was recently demonstrated to be highly reproducible in healthy subjects within a year (Illman et al., 2022). In an independent 398 cohort of 50 healthy subjects measured twice during wakeful resting, both the aperiodic power 399 spectral features as well as several beta event characteristics showed good to excellent test-400 retest stability (Pauls et al., 2023, bioRxiv). Recordings of 2-3 minutes of resting state data 401 were sufficient to obtain stable results for most parameters, speaking for their feasibility in 402 clinical settings. In the future, the heritability of dynamic oscillatory activity also outside the 403 somatosensory cortices could be addressed. This would, however, likely require automated 404 approaches which, in turn, might be more prone to signal-to-noise variations than the partly 405 manual phenotyping applied here.

406

#### 407 <u>Limitations</u>

As the analysis assumes normal distribution of the phenotypes, the fact that many of the phenotypes were non-normally distributed may have decreased the statistical power of the study. The permutation procedure adopted for testing the significance of the heritability values should, however, correct for any inflation of the heritabilities caused by the non-normality. The analyses were additionally conducted with the internal normality correction functionality of Merlin, resulting in values qualitatively similar to (although slightly more significant than) those based on the non-corrected data presented here.

415

Any measurement noise contributes to the phenotypic variability, thus reducing estimated heritability. The effect sizes calculated here did not suggest a systematic effect of signal-to-noise ratio on the observed heritabilities: for example, the effect size for event duration was higher than the effect size for event amplitude. Furthermore, in our recent study (Pauls et al., 2023, bioRxiv) the test-retest reliability of somatomotor beta activity was not directly related to relative heritabilities observed in the current study. Thus, the observed heritability differences do not solely reflect differences in the signal reliability nor the signal-to-noise ratio.

423

#### 424 Conclusion

We here show that the human sensorimotor beta and aperiodic cortical activity can be dissected into highly heritable and non-heritable components. We postulate that the different heritabilities reflect, in part, different underlying signal generating mechanisms and their weighting in the generation of different signal characteristics. In combination with increased information resulting from the time-resolved beta signal decomposition, the results generate an interesting framework to interrogate and interpret M/EEG data both in healthy subjects as

- 431 well as patient populations. This framework also increases the potential of whole-brain elec-
- 432 trophysiology measures, such as beta band activity, as disease biomarkers.

#### 433 **References**

- Abecasis GR, Cherny SS, Cookson WO, Cardon LR (2002) Merlin Rapid analysis of
   dense genetic maps using sparse gene flow trees. Nat Genet 30:97–101.
- Amos CI (1994) Robust variance-components approach for assessing genetic linkage in
   pedigrees. Am J Hum Genet 54:535–543.
- Babiloni C, Babiloni F, Carducci F, Cincotti F, Cocozza G, Del Percio C, Moretti DV, Rossini
  PM (2002) Human cortical electroencephalography (EEG) rhythms during the observation of simple aimless movements: A high-resolution EEG study. Neuroimage 17:559–
  572.
- Barroso I, McCarthy MI (2019) The Genetic Basis of Metabolic Disease. Cell 177:146–161
  Available at: https://pubmed.ncbi.nlm.nih.gov/30901536/.
- Bonaiuto JJ, Little S, Neymotin SA, Jones SR, Barnes GR, Bestmann S (2021) Laminar dynamics of high amplitude beta bursts in human motor cortex. Neuroimage 242 Available
  at: https://pubmed.ncbi.nlm.nih.gov/34407440/.
- Buzsáki G, Anastassiou CA, Koch C (2012) The origin of extracellular fields and currentsEEG, ECoG, LFP and spikes. Nature Reviews Neuroscience 13:407–420 Available at:
  https://pubmed.ncbi.nlm.nih.gov/22595786/.
- Cagnan H, Mallet N, Moll CKE, Gulberti A, Holt AB, Westphal M, Gerloff C, Engel AK, Hamel
  W, Magill PJ, Brown P, Sharott A (2019) Temporal evolution of beta bursts in the parkinsonian cortical and basal ganglia network. Proc Natl Acad Sci U S A 116:16095–16104.
- Colclough GL, Smith SM, Nichols TE, Winkler AM, Sotiropoulos SN, Glasser MF, Van Essen
   DC, Woolrich MW (2017) The heritability of multi-modal connectivity in human brain ac tivity. Elife 6 Available at: https://pubmed.ncbi.nlm.nih.gov/28745584/.
- Donoghue T, Haller M, Peterson EJ, Varma P, Sebastian P, Gao R, Noto T, Lara AH, Wallis
   JD, Knight RT, Shestyuk A, Voytek B (2020) Parameterizing neural power spectra into
   periodic and aperiodic components. Nat Neurosci 23:1655–1665.
- 459 Espenhahn S, de Berker AO, van Wijk BCM, Rossiter HE, Ward NS (2017) Movement-re460 lated beta oscillations show high intra-individual reliability. Neuroimage 147:175–185.
- Feingold J, Gibson DJ, Depasquale B, Graybiel AM (2015) Bursts of beta oscillation differentiate postperformance activity in the striatum and motor cortex of monkeys performing
  movement tasks. Proc Natl Acad Sci U S A 112 Available at:
  http://dx.doi.org/10.1073/pnas.1517629112.
- Gao R, Peterson EJ, Voytek B (2017) Inferring synaptic excitation/inhibition balance from
   field potentials. Neuroimage 158:70–78.
- Geschwind DH, Miller BL, DeCarli C, Carmelli D (2002) Heritability of lobar brain volumes in
   twins supports genetic models of cerebral laterality and handedness. Proc Natl Acad Sci
   U S A 99:3176–3181.

- 470 Goulet-Pelletier J-C, Cousineau D (2018) A review of effect sizes and their confidence inter-471 vals, Part 1: The Cohen's d family. Tutor Quant Methods Psychol 14:242–265.
- Gramfort A, Luessi M, Larson E, Engemann DA, Strohmeier D, Brodbeck C, Goj R, Jas M,
  Brooks T, Parkkonen L, Hämäläinen M (2013) MEG and EEG data analysis with MNEPython. Front Neurosci 7 Available at: https://pubmed.ncbi.nlm.nih.gov/24431986/.
- Haegens S, Nácher V, Hernández A, Luna R, Jensen O, Romo R (2011) Beta oscillations in
  the monkey sensorimotor network reflect somatosensory decision making. Proc Natl
  Acad Sci U S A 108:10708–10713.
- Halnes G, Mäki-Marttunen T, Keller D, Pettersen KH, Andreassen OA, Einevoll GT (2016)
  Effect of Ionic Diffusion on Extracellular Potentials in Neural Tissue. PLoS Comput Biol
  12 Available at: https://pubmed.ncbi.nlm.nih.gov/27820827/.
- Hari R, Forss N, Avikainen S, Kirveskari E, Salenius S, Rizzolatti G (1998) Activation of hu man primary motor cortex during action observation: A neuromagnetic study. Proc Natl
   Acad Sci U S A 95:15061–15065.
- Hari R, Salmelin R (1997) Human cortical oscillations: A neuromagnetic view through the
  skull. Trends in Neurosciences 20:44–49 Available at: https://pubmed.ncbi.nlm.nih.gov/9004419/.
- He BJ (2014) Scale-free brain activity: Past, present, and future. Trends in Cognitive Sciences 18:480–487 Available at: https://pubmed.ncbi.nlm.nih.gov/24788139/.
- Helson P, Lundqvist D, Vinding MC, Kumar A (2023) Cortex-wide topography of 1/f-expo nent in Parkinson's disease. bioRxiv:2023.01.19.524792.
- Hill AT, Clark GM, Bigelow FJ, Lum JAG, Enticott PG (2022) Periodic and aperiodic neural
   activity displays age-dependent changes across early-to-middle childhood. Dev Cogn
   Neurosci 54:101076.
- 494 Illman M, Laaksonen K, Jousmäki V, Forss N, Piitulainen H (2022) Reproducibility of
   495 Rolandic beta rhythm modulation in MEG and EEG. J Neurophysiol 127:559–570.
- Jones SR (2016) When brain rhythms aren't "rhythmic": implication for their mechanisms and
   meaning. Current Opinion in Neurobiology 40:72–80 Available at: https://pub med.ncbi.nlm.nih.gov/27400290/.
- Jones SR, Pritchett DL, Sikora MA, Stufflebeam SM, Hämäläinen M, Moore CI (2009) Quantitative analysis and biophysically realistic neural modeling of the MEG mu rhythm:
  Rhythmogenesis and modulation of sensory-evoked responses. J Neurophysiol
  102:3554–3572.
- Kandel ER, Schwartz JH, Jessell TM (2000) The Anatomical Organization of the Central
   Nervous System. In: Principles of Neural Science, pp 317–336.
- Koten JW, Wood G, Hagoort P, Goebel R, Propping P, Willmes K, Boomsma DI (2009) Ge netic contribution to variation in cognitive function: An fMRI study in twins. Science
   323:1737–1740.
- Laaksonen K, Kirveskari E, Mäkelä JP, Kaste M, Mustanoja S, Nummenmaa L, Tatlisumak

- 509 T, Forss N (2012) Effect of afferent input on motor cortex excitability during stroke re-
- 510 covery. Clin Neurophysiol 123 Available at:
- 511 http://dx.doi.org/10.1016/j.clinph.2012.05.017.
- Law RG, Pugliese S, Shin H, Sliva DD, Lee S, Neymotin S, Moore C, Jones SR (2022)
   Thalamocortical Mechanisms Regulating the Relationship between Transient Beta
   Events and Human Tactile Perception. Cereb Cortex 32:668–688.
- Leppäaho E, Renvall H, Salmela E, Kere J, Salmelin R, Kaski S (2019) Discovering heritable modes of MEG spectral power. Hum Brain Mapp 40:1391–1402.
- Lindén H, Pettersen KH, Einevoll GT (2010) Intrinsic dendritic filtering gives low-pass power
   spectra of local field potentials. J Comput Neurosci 29:423–444.
- Lorente de No R (1949) Cerebral Cortex. Architecture, intracortical connections, motor projections. In: Physiology of the Nervous System, 3rd ed. (Fulton JF, ed), pp 288–330.
  New York: Oxford University Press.
- 522 Mahjoory K, Schoffelen JM, Keitel A, Gross J (2020) The frequency gradient of human rest-
- ing-state brain oscillations follows cortical hierarchies. Elife 9 Available at:
   http://dx.doi.org/10.7554/ELIFE.53715.
- Maniscalco B, Lee JL, Abry P, Lin A, Holroyd T, He BJ (2018) Neural integration of stimulus
   history underlies prediction for naturalistically evolving sequences. Journal of Neurosci ence 38:1541–1557.
- McSweeney M, Morales S, Valadez EA, Buzzell GA, Fox NA (2021) Longitudinal age- and
   sex-related change in background aperiodic activity during early adolescence. Dev
   Cogn Neurosci 52:101035.
- Mo C, Sherman SM (2019) A sensorimotor pathway via higher-order thalamus. Journal of
   Neuroscience 39:692–704.
- Molina JL, Voytek B, Thomas ML, Joshi YB, Bhakta SG, Talledo JA, Swerdlow NR, Light GA
   (2020) Memantine Effects on Electroencephalographic Measures of Putative Excita tory/Inhibitory Balance in Schizophrenia. Biological Psychiatry: Cognitive Neuroscience
   and Neuroimaging 5:562–568.
- Neymotin SA, Daniels DS, Caldwell B, McDougal RA, Carnevale NT, Jas M, Moore CI,
  Hines ML, Hämäläinen M, Jones SR (2020) Human neocortical neurosolver (HNN), a
  new software tool for interpreting the cellular and network origin of human MEG/EEG
  data. Elife 9 Available at: https://pubmed.ncbi.nlm.nih.gov/31967544/.
- 541 Ostlund BD, Alperin BR, Drew T, Karalunas SL (2021) Behavioral and cognitive correlates of
  542 the aperiodic (1/f-like) exponent of the EEG power spectrum in adolescents with and
  543 without ADHD. Dev Cogn Neurosci 48 Available at: https://pub544 med.ncbi.nlm.nih.gov/33535138/.
- Pauls KAM, Korsun O, Nenonen J, Nurminen J, Liljeström M, Kujala J, Pekkonen E, Renvall
  H (2022) Cortical beta burst dynamics are altered in Parkinson's disease but normalized
  by deep brain stimulation. Neuroimage 257 Available at: https://pub-
- 548 med.ncbi.nlm.nih.gov/35569783/.

- Pauls KAM, Nurmi P, Ala-Salomäki H, Renvall H, Kujala J, Liljeström M (2023) Human sen sorimotor resting state beta events and 1/f response show good test-retest reliability. bi oRxiv Available at: http://dx.doi.org/10.1101/2023.08.16.553499.
- Peper JS, Brouwer RM, Boomsma DI, Kahn RS, Hulshoff Pol HE (2007) Genetic influences
   on human brain structure: A review of brain imaging studies in twins. Human Brain Map ping 28:464–473 Available at: https://pubmed.ncbi.nlm.nih.gov/17415783/.
- Pettersen KH, Lindén H, Tetzlaff T, Einevoll GT (2014) Power Laws from Linear Neuronal
   Cable Theory: Power Spectral Densities of the Soma Potential, Soma Membrane Cur rent and Single-Neuron Contribution to the EEG. PLoS Comput Biol 10 Available at:
   https://pubmed.ncbi.nlm.nih.gov/25393030/.
- Pfurtscheller G, Krausz G, Neuper C (2001) Mechanical stimulation of the fingertip can in duce bursts of β oscillations in sensorimotor areas. J Clin Neurophysiol 18:559–564.
- Porjesz B et al. (2002) Linkage disequilibrium between the beta frequency of the human
   EEG and a GABAA receptor gene locus. Proc Natl Acad Sci U S A 99:3729–3733.
- Reis C, Sharott A, Magill PJ, van Wijk BCM, Parr T, Zeidman P, Friston KJ, Cagnan H
   (2019) Thalamocortical dynamics underlying spontaneous transitions in beta power in
   Parkinsonism. Neuroimage 193:103–114.
- Renvall H, Salmela E, Vihla M, Illman M, Leinonen E, Kere J, Salmelin R (2012) Genome wide linkage analysis of human auditory cortical activation suggests distinct loci on chro mosomes 2, 3, and 8. Journal of Neuroscience 32:14511–14518.
- Sacchet MD, LaPlante RA, Wan Q, Pritchett DL, Lee AKC, Hämäläinen M, Moore CI, Kerr
   CE, Jones SR (2015) Attention drives synchronization of alpha and beta rhythms be tween right inferior frontal and primary sensory neocortex. Journal of Neuroscience
   35:2074–2082.
- Salmela E, Renvall H, Kujala J, Hakosalo O, Illman M, Vihla M, Leinonen E, Salmelin R,
  Kere J (2016) Evidence for genetic regulation of the human parieto-occipital 10-Hz
  rhythmic activity. Eur J Neurosci 44:1963–1971.
- Salmelin R, Hari R (1994) Spatiotemporal characteristics of sensorimotor neuromagnetic
   rhythms related to thumb movement. Neuroscience 60:537–550.
- Schmitt JE, Neale MC, Fassassi B, Perez J, Lenroot RK, Wells EM, Giedd JN (2014) The
  dynamic role of genetics on cortical patterning during childhood and adolescence. Proc
  Natl Acad Sci U S A 111:6774–6779.
- Schmitt JE, Raznahan A, Liu S, Neale MC (2020) The genetics of cortical myelination in
  young adults and its relationships to cerebral surface area, cortical thickness, and intelligence: A magnetic resonance imaging study of twins and families: Genetics of Cortical
  Myelination, Area, Thickness, and Intelligence. Neuroimage 206 Available at:
  https://pubmed.ncbi.nlm.nih.gov/31678229/.
- Seedat ZA, Quinn AJ, Vidaurre D, Liuzzi L, Gascoyne LE, Hunt BAE, O'Neill GC, Pakenham
  DO, Mullinger KJ, Morris PG, Woolrich MW, Brookes MJ (2020) The role of transient
  spectral "bursts" in functional connectivity: A magnetoencephalography study. Neuroimage 209 Available at: http://dx.doi.org/10.1016/j.neuroimage.2020.116537.

- Semenova U, Popov V, Tomskiy A, Shaikh AG, Sedov A (2021) Pallidal 1/f asymmetry in pa tients with cervical dystonia. Eur J Neurosci 53:2214–2219.
- Sherman MA, Lee S, Law R, Haegens S, Thorn CA, Hämäläinen MS, Moore CI, Jones SR
  (2016) Neural mechanisms of transient neocortical beta rhythms: Converging evidence
  from humans, computational modeling, monkeys, and mice. Proc Natl Acad Sci U S A
  113:E4885–E4894.
- Shin H, Law R, Tsutsui S, Moore CI, Jones SR (2017) The rate of transient beta frequency
   events predicts behavior across tasks and species. Elife 6 Available at:
   http://dx.doi.org/10.7554/eLife.29086.
- Shinomoto S, Miura K, Koyama S (2005) A measure of local variation of inter-spike intervals.
  In: BioSystems, pp 67–72. Biosystems.
- Silén T, Forss N, Jensen O, Hari R (2000) Abnormal reactivity of the ~20-Hz motor cortex
   rhythm in unverricht lundborg type progressive myoclonus epilepsy. Neuroimage
   12:707–712.
- Smit CM, Wright MJ, Hansell NK, Geffen GM, Martin NG (2006) Genetic variation of individ ual alpha frequency (IAF) and alpha power in a large adolescent twin sample. Int J Psy chophysiol 61:235–243.
- 607 Smit DJA, Posthuma D, Boomsma DI, De Geus EJC (2005) Heritability of background EEG 608 across the power spectrum. Psychophysiology 42:691–697.
- Taulu S, Kajola M, Simola J (2004) Suppression of interference and artifacts by the signal
   space separation method. Brain Topogr 16:269–275.
- Taulu S, Simola J (2006) Spatiotemporal signal space separation method for rejecting
   nearby interference in MEG measurements. Phys Med Biol 51:1759–1768.
- Terry RD, Masliah E, Salmon DP, Butters N, DeTeresa R, Hill R, Hansen LA, Katzman R
   (1991) Physical basis of cognitive alterations in Alzheimer's disease: synapse loss is the
   major correlate of cognitive impairment. Ann Neurol 30:572–580.
- Van Baal GCM, De Geus EJC, Boomsma DI (1996) Genetic architecture of EEG power
   spectra in early life. Electroencephalogr Clin Neurophysiol 98:502–514.
- Van Beijsterveldt CEM, Molenaar PCM, De Geus EJC, Boomsma DI (1996) Heritability of
   human brain functioning as assessed by electroencephalosraphy. Am J Hum Genet
   58:562–573.
- Van Ede F, De Lange F, Jensen O, Maris E (2011) Orienting attention to an upcoming tactile
   event involves a spatially and temporally specific modulation of sensorimotor alpha- and
   beta-band oscillations. Journal of Neuroscience 31:2016–2024.
- van Pelt S, Boomsma DI, Fries P (2012) Magnetoencephalography in twins reveals a strong
   genetic determination of the peak frequency of visually induced gamma-band synchroni zation. Journal of Neuroscience 32:3388–3392.
- Vinding MC, Tsitsi P, Waldthaler J, Oostenveld R, Ingvar M, Svenningsson P, Lundqvist D
   (2020) Reduction of spontaneous cortical beta bursts in Parkinson's disease is linked to

- 629 symptom severity. Brain Communications 2 Available at: http://dx.doi.org/10.1093/brain-630 comms/fcaa052.
- Visscher PM, Hill WG, Wray NR (2008) Heritability in the genomics era--concepts and mis conceptions. Nat Rev Genet 9:255–266.
- Voytek B, Kramer MA, Case J, Lepage KQ, Tempesta ZR, Knight RT, Gazzaley A (2015)
   Age-related changes in 1/f neural electrophysiological noise. Journal of Neuroscience
   35:13257–13265.
- Waschke L, Donoghue T, Fiedler L, Smith S, Garrett DD, Voytek B, Obleser J (2021) Modal ity-specific tracking of attention and sensory statistics in the human electrophysiological
   spectral exponent. Elife 10 Available at: https://pubmed.ncbi.nlm.nih.gov/34672259/.
- Wigginton JE, Abecasis GR (2005) PEDSTATS: Descriptive statistics, graphics and quality
   assessment for gene mapping data. Bioinformatics 21:3445–3447.
- 641 Wilson LE, Castanheira J da S, Baillet S (2022) Time-resolved parameterization of aperiodic
- 642 and periodic brain activity. Elife 11 Available at: https://pub-
- 643 med.ncbi.nlm.nih.gov/36094163/.

#### 644 **Figure Captions**

#### 645 Figure 1: Extraction of sensorimotor beta phenotype characteristics.

646 A. Channel selection. A region of interest (ROI) was defined for both hemispheres. The 15 selected 647 gradiometer-channel pairs were combined into 15 vector-sum PSDs (one per channel pair). The peri-648 odic spectral component of the vector-sum PSD was obtained using FOOOF. From these, a peak beta frequency and peak channel pair were selected. B. Beta event extraction. The peak channel pair and 649 650 peak frequency selected in A were used to calculate the channel pair's amplitude envelope. From the raw data, narrow-band filtered data were obtained using wavelet decomposition, and the individual 651 652 channels' band-filtered signals were combined to one amplitude envelope using vector sum calculation. 653 C. Parameters for heritability analysis. Both PSD characteristics (beta peak power and frequency, total 654 beta power at 14-30 Hz (periodic part), 1/f exponent; upper panel) and time-resolved beta oscillatory 655 characteristics (beta events; lower panel) were used in the heritability analysis.

656

### 657 Figure 2: Beta phenotypes

A. Phenotypic spectrum of beta activity. Examples of typical beta range PSD patterns: (a) narrow
 beta peak, (b) broad range, 'beta brush' like activity, (c) double peaks of comparable strength, one in
 the lower, one in the higher beta range. B. Beta PSD patterns in siblings. Examples of siblings' beta
 PSD patterns (two families with two siblings, one family with three siblings, one family with four sib lings).

663

#### Figure 3. Sources of MEG signals and their putative relationship to the MEG parameters examined in the present study.

The schematic figure's upper panel summarizes the main anatomical and morphological factors, as well as factors determining timing of events, that contribute to the generation of MEG signals. The lower part indicates the putative relationship of those factors to the MEG parameters examined here. We postulate that the 1/f signal and beta event amplitude parameters are more heavily dependent on fixed, anatomical parameters, whereas beta event duration and its modulation are more dynamic characteristics, yet keeping in mind that timing is very much constrained by network anatomy. Brain slice modified from

https://commons.wikimedia.org/wiki/File:Human\_basal\_ganglia\_nuclei\_as\_shown\_in\_two\_coro-nal\_slices\_and\_with\_reference\_to\_an\_illustration\_in\_the\_sagital\_plane.svg 

			Left he	nere	Right hemisphere				
PSD characteristics		mea n	me- dian	std	range	mea n	me- dian	std	range
peak beta free	quency (Hz)	19.7	19.3	3.0	14.1-25.8	19.8	19.3	3.1	14.1-29.3
peak beta pov	ver (fT/cm) <sup>2</sup>	276	144	374	15-3231	133	70	173	5-1251
total beta ban odic)	d power (peri-	297 9	1686	331 8	164- 16284	109 3	619	132 2	26-9652
1/f component exponent		1.03	1.01	0.1 9	0.34-1.76	1.07	1.05	0.1 8	0.67-1.66
1/f component offset		۔ 22.9 5	- 22.99	0.3 7	-23.75 21.75	- 23.2 8	23.32	0.3 4	-24.19 22.25
Beta event ch	naracteristics	mea n	me- dian	std	range	mea n	me- dian	std	range
duration (ms)	mean	256. 9	248.0	49. 4	181.7- 498.2	265. 2	253.7	50. 0	182.0- 454.0
	median	199. 0	195.0	32. 3	152.5- 420.0	201. 4	195.0	34. 9	150.0- 355.0
	standard de- viation	198. 4	182.4	63. 2	99.8- 487.5	213. 0	198.3	67. 4	67.3- 531.7
	robust maxi- mum	858. 0	788.6	257 .4	439.5- 2145.0	912. 1	852.2	263 .9	382.7- 2135.0
amplitude (fT/cm)	mean	325	285	165	104-994	221	188	113	73-654
	median	301	260	155	93-984	203	173	105	70-618
	standard de- viation	86	71	48	21-264	62	52	34	14-202
6	robust maxi- mum	564	486	286	174-1467	390	335	196	114-1128
event rate (1/s)		1.00	1.00	0.1 6	0.50-1.36	0.97	0.98	0.1 6	0.55-1.37
dispersion		1.14	1.05	0.4 4	0.65-5.59	1.16	1.06	0.4 4	0.41-5.58

### left hemi-sphere

ċ	Left he sphere	emi- e		Right he sphere	emi-	
PSD characteristics	h²	р	n sig. (/6000)	h²	р	n sig. (/6000)
peak beta frequency	0.45	0.0047	28	0.41	0.0103	62
peak beta power	0.28	0.0648	389	0.58	0.0072	43
total beta band power (peri- odic)	0.49	0.0068	41	0.44	0.0157	94
1/f component exponent *	0.47	0.0035	21	0.87	0.0000	0
1/f component offset *	0.35	0.0258	155	0.69	0.0000	0
Beta event characteristics	h²	р	n sig. (/6000)	h²	р	n sig. (/6000)
duration mean	0.45	0.1350	810	0.36	0.0222	133
median	0.28	0.1338	803	0.40	0.0172	103

	standard deviation	0.49	0.2495	1497	0.32	0.0412	247
	robust maximum	0.47	0.2383	1430	0.33	0.0372	223
amplitude	mean *	0.35	0.0060	36	0.75	0.0002	1
	median *	0.45	0.0110	66	0.72	0.0002	1
tion *	standard devia-	0.28	0.0005	3	0.81	0.0000	
	robust maximum	0.20	0.0005	5	0.01	0.0000	
*		0.49	0.0007	4	0.79	0.0000	0
event rate		0.47	0.0543	326	0.38	0.0137	82
dispersion		0.35	0.2850	1710	0.00	1.0000	6000
	çi	C		ed			







1/f signal component
tissue/apical dendrite
filtering properties
→ highly heritable

beta event amplitude & its dynamic range size, morphology and biophysical properties of cortical cells, cortical microstructure → heritable beta event duration & dynamics interplay of inputs from different sources and their interaction, network resonance properties → not significantly heritable