Induced Numerical Instabilities in Analytical Pipelines Lead to Impactful Variability in Brain Networks

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Abstract

The analysis of brain-imaging data requires complex processing pipelines to support findings on brain function or pathologies. Recent work has shown that variability in analytical decisions, small amounts of noise, or computational environments can lead to substantial differences in the results, endangering the trust in conclusions¹⁻⁷. We explored the instability of results by instrumenting a connectome estimation pipeline with Monte Carlo Arithmetic^{8,9} to introduce random noise throughout. We evaluated the reliability of the connectomes, their features 10,11, and the impact on analysis 12,13. The stability of results was found to range from perfectly stable to highly unstable. This paper highlights the potential of leveraging induced variance in estimates of brain connectivity to reduce the bias in networks alongside increasing the robustness of their applications in the classification of individual differences. We demonstrate that stability evaluations are necessary for understanding error inherent to brain imaging experiments, and how numerical analysis can be applied to typical analytical workflows both in brain imaging and other domains of computational science. Overall, while the extreme variability in results due to analytical instabilities could severely hamper our understanding of brain organization, it also leads to an increase in the reliability of datasets.

Keywords

Stability — Reproducibility — Network Neuroscience — Neuroimaging

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The modelling of brain networks, called connectomics, 9 but potentially pave the way for therapeutics 19-23. ² has shaped our understanding of the structure and function 3 of the brain across a variety of organisms and scales over 4 the last decade 11, 14-18. In humans, these wiring diagrams are 6 and show promise towards identifying biomarkers of disease. 7 This can not only improve understanding of so-called "connec-

However, the analysis of brain imaging data relies on complex computational methods and software. Tools are trusted to ₁₂ perform everything from pre-processing tasks to downstream 5 obtained in vivo through Magnetic Resonance Imaging (MRI), 13 statistical evaluation. While these tools undoubtedly undergo 14 rigorous evaluation on bespoke datasets, in the absence of 15 ground-truth this is often evaluated through measures of re-8 topathies", such as Alzheimer's Disease and Schizophrenia, 16 liability^{24–27}, proxy outcome statistics, or agreement with

 $_{17}$ existing theory. Importantly, this means that tools are not $_{53}$ individual and 100 samples total ($25 \times 2 \times 2$ samples). Struc-24 and it is likely that software instabilities played a role.

29 cations of the observed instabilities on downstream analyses 30 were quantified. We accomplished this through the use of 31 Monte Carlo Arithmetic (MCA)⁸, a technique which enables 32 characterization of the sensitivity of a system to small nu-33 merical perturbations. This is importantly distinct from data 34 perturbation experiments where the underlying datasets are 35 manipulated or pathologies may be simulated, and allows 36 for the evaluation of experimental uncertainty in real-world 37 settings. We explored the impact of numerical perturbations 38 through the direct comparision of structural connectomes, the 39 consistency of their features, and their eventual application in 40 a neuroscience study. We also characterized the consequences 41 of instability in these pipelines on the reliability of derived 42 datasets, and discuss how the induced variability may be har-43 nessed to increase the discriminability of datasets. Finally, 44 we make recommendations for the roles perturbation analyses 45 may play in brain imaging research and beyond.

46 Graphs Vary Widely With Perturbations

48 understanding of the induced variability was required. A sub- 85 tinct, connectomes generated with sparse perturbations show 49 set of the Nathan Kline Institute Rockland Sample (NKIRS) 86 considerable variability, often reaching deviations equal to 50 dataset²⁹ was randomly selected to contain 25 individuals 87 or greater than those observed across individuals or sessions 51 with two sessions of imaging data, each of which was sub- 88 (Figure 1A; right). Interpretting these results with respect to

18 necessarily of known or consistent quality, and it is not un- 54 tural connectomes were generated with canonical determinis-19 common that equivalent experiments may lead to diverging 55 tic and probabilistic pipelines 30,31 which were instrumented 20 conclusions^{1,5–7}. While many scientific disciplines suffer 56 with MCA, replicating computational noise either sparsely 21 from a lack of reproducibility²⁸, this was recently explored 57 or densely throughout the pipelines^{4,9}. In the sparse case, a 22 in brain imaging by a 70 team consortium which performed 58 small subset of the libraries were instrumented with MCA, al-23 equivalent analyses and found widely inconsistent results¹, 59 lowing for the evaluation of the cascading effects of numerical 60 instabilities that may arise. In the dense case, operations are The present study approached evaluating reproducibility 61 more uniformly perturbed and thus the law of large numbers 26 from a computational perspective in which a series of brain 62 suggests that perturbations will quickly offset one-another and 27 imaging studies were numerically perturbed in such a way 63 only dramatic local instabilities will have propagating effects. 28 that the plausibility of results was not affected, and the impli- 64 Importantly, the perturbations resulting from the sparse setting 65 represent a strict subset of the possible outcomes of the dense 66 implementation. The random perturbations are statistically 67 independent from one another across both settings and sim-68 ulations. Instrumenting pipelines with MCA increases their 69 computation time, in this case by multiplication factors of ₇₀ 1.2× and $7\times$ for the sparse and dense settings, respectively⁴. 71 The results obtained were compared to unperturbed (e.g. ref-72 erence) connectomes in both cases. The connectomes were 73 sampled 10 times per sample and once without perturbations, 74 resulting in a total of 4,200 connectomes. Two versions of 75 the unperturbed connectomes were generated and compared 76 such that the absence of variability aside from that induced 77 via MCA could be confirmed.

The stability of connectomes was evaluated through the 79 normalized percent deviation from reference⁴ and the num-80 ber of significant digits (Figure 1). The comparisons were 81 grouped according to differences across simulations, subsam-82 pling of data, sessions of acquisition, or subjects, and accord-83 ingly sorted from most to least similar. While the similarity 47 Prior to exploring the analytic impact of instabilities, a direct 84 of connectomes decreases as the collections become more dis-52 sampled into two components, resulting in four samples per 89 the distinct MCA environments used suggests that the tested

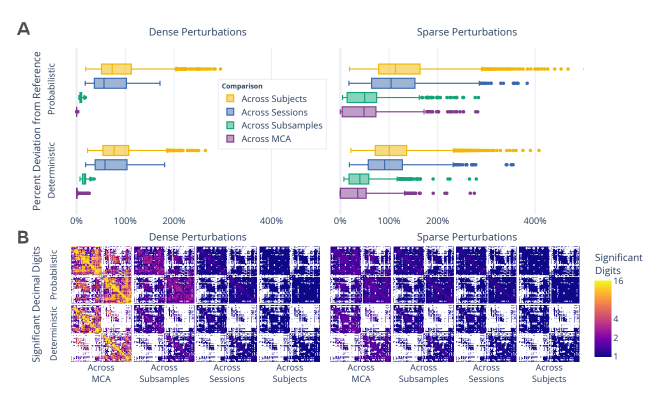


Figure 1. Exploration of perturbation-induced deviations from reference connectomes. (**A**) The absolute deviations between connectomes, in the form of normalized percent deviation from reference. The difference in MCA-perturbed connectomes is shown as the across MCA series, and is presented relative to the variability observed across subsamples, sessions, and subjects. (**B**) The number of significant decimal digits in each set of connectomes as obtained by evaluating the complete distribution of networks. In the case of 16, values can be fully relied upon, whereas in the case of 1 only the first digit of a value can be trusted. Dense and sparse perturbations are shown on the left and right, respectively.

90 pipelines may not suffer from single dominant sources of 104 creasing similarity between comparison groups. While the 91 instability, but that nevertheless there exist minor local in- 105 cross-MCA comparison of connectomes generated with dense 92 stabilities which may the propagate throughout the pipeline. 106 perturbations show nearly perfect precision for many edges 93 Furthermore, this finding suggests that instabilities inherent 107 (approaching the maximum of 15.7 digits for 64-bit data), 94 to these pipelines may mask session or individual differences, 108 this evaluation uniquely shows considerable drop off in per-95 limiting the trustworthiness of derived connectomes. While 109 formance when comparing networks across subsamplings ₉₆ both pipelines show similar performance, the probabilistic ₁₁₀ (average of < 4 digits). In addition, sparsely perturbed con-97 pipeline was more stable in the face of dense perturbations 111 nectomes show no more than an average of 3 significant digits 98 whereas the deterministic was more stable to sparse perturba- 112 across all comparison groups, demonstrating a significant lim-₉₉ tions (p < 0.0001 for all; exploratory). As an alternative to 113 itation in the reliability of independent edge weights. The 100 the normalized percent deviation, the stability of correlations 114 number of significant digits across individuals did not exceed between networks can be found in Supplemental Section S1. 115 a single digit per edge in any case, indicating that only the order of magnitude of edges in naively computed groupwise The number of significant digits per edge across conaverage connectomes can be trusted. The combination of 103 nectomes (Figure 1B) similarly decreases alongside the de118 these results with those presented in Figure 1A suggests that 154 settings perfectly preserved differences between sessions with while specific edge weights are largely affected by instabili- 155 a score of 1.0 (p < 0.005; optimal score: 0.5; chance: 0.5), 120 ties, macro-scale network organization is stable.

122 inability

124 ing and extending a typical test-retest experiment²⁶ in which 125 the similarity of samples across sessions were compared to distinct samples in the dataset (Table 1, with additional experiments and explanation in Supplemental Section S2). The abil-128 ity to discriminate connectomes across subjects (Hypothesis 1) 165 129 is an essential prerequisite for the application of brain imaging 166 uate the interaction between data acquisition and tool, the 130 towards identifying individual differences 18. In testing hypoth- 167 use of subsampling allowed for characterizing the discrimesis 1, we observe that the dataset is discriminable with a score inability of networks sampled from within a single acquisition of 0.64 and 0.65 (p < 0.001; optimal score: 1.0; chance: 0.04) 169 (Hypothesis 3). While this experiment could not be evaluated as p = 0.064 and 0.65 (p < 0.001; optimal score: 1.0; chance: 0.04) 169 (Hypothesis 3). 133 for the Deterministic and Probabilistic pipelines, respectively, 170 ated using reference executions, the networks generated with in the absence of MCA. However, we can see that inducing 171 dense perturbations showed near perfect discrimination beinstabilities through MCA improves the discriminability of tween subsamples, with scores of 0.99 and 1.0 (p < 0.005; the dataset to over 0.75 in each case (p < 0.001 for all), sig- 173 optimal: 0.5; chance: 0.5). Given that there is no variability in nificantly higher than without instrumentation (p < 0.005 for 174 data acquisition, due to undesired effects such as participant 138 all). The definition of the discriminability statistic is such that 175 motion, or preprocessing, the ability to discriminate between 139 if all samples derived from the a given session were near repli- 176 equivalent subsamples in this experiment may only be due 140 cates of one another the score would be unchanging despite 177 to instability or bias inherent to the pipelines. The high varithe increased sample size²⁶. The resulting increase in discrim- 178 ability introduced through sparse perturbations considerably 142 inability therefore suggests the utility of perturbation methods 179 lowered the discriminability towards chance (score: 0.71 and for synthesizing robust and reliable individual estimates of $180 \ 0.61$; p < 0.005 for all), further supporting this as an effec-144 connectivity, serving as a cost effective and context-agnostic 181 tive method for obtaining lower-bias estimates of individual 145 method for dataset augmentation.

While the discriminability of individuals is essential for 183 147 the identification of individual brain networks, it is similarly 184 creased ability to discriminate networks on the basis of mean-148 reliant on network similarity – or lack of discriminability – 185 ingful biological signal alongside a reduction in discriminabil-149 across equivalent acquisitions (Hypothesis 2). In this case, 186 ity due to of off-target signal. This result appears strikingly 150 connectomes were grouped based upon session, rather than 187 like a manifestation of the well-known bias-variance trade-151 subject, and the ability to distinguish one session from an- 188 off³² in machine learning, a concept which observes a de-152 other based on subsamples was computed within-individual 189 crease in bias as variance is favoured by a model. In particular, 153 and aggregated. Both the unperturbed and dense perturbation 190 this highlights that numerical perturbations can be used to not

156 indicating a dominant session-dependent signal for all indi-157 viduals despite no intended biological differences. However, Perturbations Increase Biologically-Driven Discrim- 158 while still significant relative to chance (score: 0.85 and 0.88; p < 0.005 for both), sparse perturbations lead to significantly We assessed the reproducibility of the dataset through mimick- 160 lower discriminability of the dataset (p < 0.005 for all). This 161 reduction of the difference between sessions suggests that 162 the added variance due to perturbations reduces the relative 164 in the networks.

> Though the previous sets of experiments inextricably eval-182 connectivity.

Across all cases, the induced perturbations led to an in-

Table 1. The impact of instabilities as evaluated through the discriminability of the dataset based on individual (or subject) differences, session, and subsample. The performance is reported as mean discriminability. While a perfectly discriminable dataset would be represented by a score of 1.0, the chance performance, indicating minimal discriminability, is 1/the number of classes. H_3 could not be tested using the reference executions due to too few possible comparisons. The alternative hypothesis, indicating significant discrimination, was accepted for all experiments, with p < 0.005.

			Reference Execution		Dense Perturbations		Sparse Perturbations	
Comparison	Chance	Target	Det.	Prob.	Det.	Prob.	Det.	Prob.
<i>H</i> ₁ : Across Subjects	0.04	1.0	0.64	0.65	0.82	0.82	0.77	0.75
H_2 : Across Sessions	0.5	0.5	1.00	1.00	1.00	1.00	0.88	0.85
<i>H</i> ₃ : Across Subsamples	0.5	0.5			0.99	1.00	0.71	0.61

only evaluate the stability of pipelines, but that the induced 216 tion modes.

192 variance may be leveraged for the interpretation as a robust 217

193 distribution of possible results.

Distributions of Graph Statistics Were Reliable, But 220 feature-moments were stable with more than 10 significant 195 Individual Statistics Were Not

196 Exploring the stability of topological features of connectomes 222 ministic pipeline, though the probabilistic pipeline was more 197 is relevant for typical analyses, as low dimensional features are 223 stable for all comparisons (p < 0.0001; exploratory). In stark often more suitable than full connectomes for many analytical 224 contrast, sparse perturbations led to highly unstable featuremethods in practice¹¹. A separate subset of the NKIRS dataset 225 moments (Figure 2D), such that none contained more than 200 was randomly selected to contain a single non-subsampled ses- 226 5 significant digits of information and several contained less 201 sion for 100 individuals (100 × 1 × 1) using the pipelines and 227 than a single significant digit, indicating a complete lack of re-202 instrumentation methods to generate connectomes as above. 228 liability. This dramatic degradation in stability for individual 203 Connectomes were generated 20 times each, resulting in a 229 measures strongly suggests that these features may be unre-204 dataset which also contained 4,200 connectomes with the 230 liable as individual biomarkers when derived from a single 205 MCA simulations serving as the only source of repeated mea- 231 pipeline evaluation, though their reliability may be increased

208 features 10 were explored and are presented in Figure 2. The 204 obtained similar findings and can be found in Supplemental 209 cumulative density of the features was computed within in- 235 Section S3. 210 dividuals and the mean cumulative density and associated 211 standard error were computed for across individuals (Fig- 236 Uncertainty in Brain-Phenotype Relationships

212 ures 2A and 2B). There was no significant difference between 237 While the variability of connectomes and their features was 213 the distributions for each feature across the two perturbation 238 summarized above, networks are commonly used as inputs to 214 settings, suggesting that the topological features summarized 239 machine learning models tasked with learning brain-phenotype by these multivariate features are robust across both perturba- 240 relationships 18. To explore the stability of these analyses, we

In addition to the comparison of distributions, the stabil-

218 ity of the first 5 moments of these features was evaluated 219 (Figures 2C and 2D). In the face of dense perturbations, the

221 digits with the exception of edge weight when using the deter-

232 when studying their distributions across perturbations. A sim-

The stability of several commonly-used multivariate graph 233 ilar analysis was performed for univariate statistics which

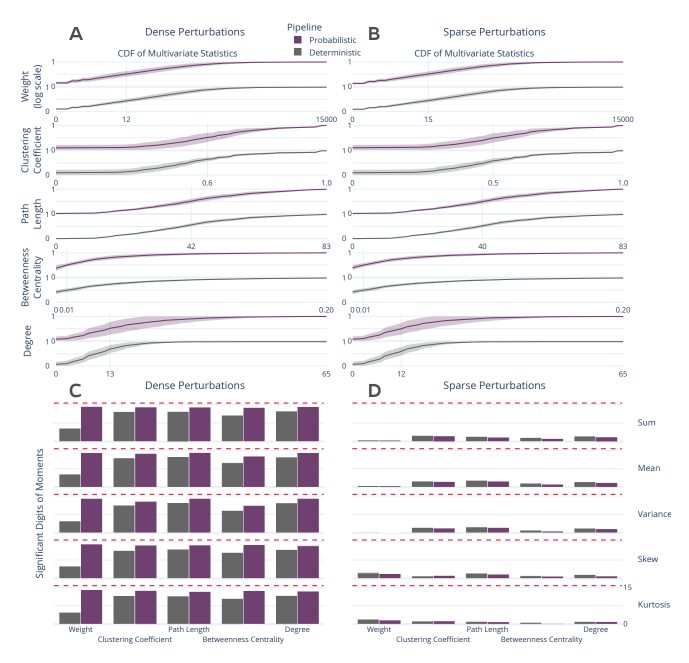


Figure 2. Distribution and stability assessment of multivariate graph statistics. (**A**, **B**) The cumulative distribution functions of multivariate statistics across all subjects and perturbation settings. There was no significant difference between the distributions in A and B. (**C**, **D**) The number of significant digits in the first 5 five moments of each statistic across perturbations. The dashed red line refers to the maximum possible number of significant digits.

modelled the relationship between high- or low- Body Mass 246 the dataset across both pipelines and perturbation methods. Index (BMI) groups and brain connectivity using standard di- 247 The accuracy and F1 score for the perturbed models varied mensionality reduction and classification tools 12,13, and com- 248 from 0.520 – 0.716 and 0.510 – 0.725, respectively, rang- pared this to reference and random performance (Figure 3). 249 ing from at or below random performance to outperforming 250 performance on the reference dataset. This large variability

The analysis was perturbed through distinct samplings of

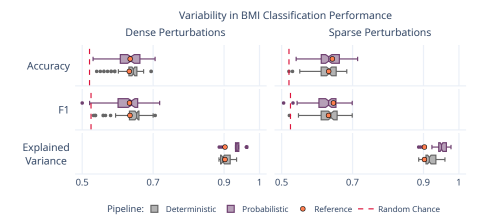


Figure 3. Variability in BMI classification across the sampling of an MCA-perturbed dataset. The dashed red lines indicate random-chance performance, and the orange dots show the performance using the reference executions.

251 illustrates a previously uncharacterized margin of uncertainty 275 modelling brain-phenotype relationships. 252 in the modelling of this relationship, and limits confidence in 253 reported accuracy scores on singly processed datasets. The 254 portion of explained variance in these samples ranged from 255 88.6% — 97.8%, similar to the reference of 90.3%, suggest-256 ing that the range in performance was not due to a gain or 257 loss of meaningful signal, but rather the reduction of bias 258 towards specific outcome. Importantly, this finding does not 259 suggest that modelling brain-phenotype relationships is not 260 possible, but rather it sheds light on impactful uncertainty that 261 must be accounted for in this process, and supports the use of 262 ensemble modeling techniques.

276 Discussion

279 led to considerable variability in derived brain graphs. Across 280 all analyses the stability of results ranged from nearly per-281 fectly trustworthy (i.e. no variation) to completely unreliable 282 (i.e. containing no trustworthy information). Given that the 283 magnitude of introduced numerical noise is to be expected 284 in computational workflows, this finding has potentially sig-285 nificant implications for inferences in brain imaging as it is One distinction between the results presented here and 286 currently performed. In particular, this bounds the success of 264 the previous is that while networks derived from dense pertur- 287 studying individual differences, a central objective in brain 265 bations had been shown to exhibit less dramatic instabilities 266 imaging 18, given that the quality of relationships between 266 in general, the results here show similar variability in clas- 269 phenotypic data and brain networks will be limited by the 267 sification performance across the two methods. This consis- 290 stability of the connectomes themselves. This issue is accen-268 tency suggests that the desired method of instrumentation may 291 tuated through the crucial finding that individually derived 269 vary across experiments. While sparse perturbations result 292 network features were unreliable despite there being no signif-270 in considerably more variability in networks directly, the two 293 icant difference in their aggregated distributions. This finding 271 techniques capture similar variability when relating networks 294 is not damning for the study of brain networks as a whole, but 272 to this phenotypic variable. Given the dramatic reduction 295 rather is strong support for the aggregation of networks, either 273 in computational overhead, a sparse instrumentation may be 296 across perturbations for an individual or across groups, over 274 preferred when processing datasets for eventual application in 297 the use of individual estimates.

277 The perturbation of structural connectome estimation pipelines

278 with small amounts of noise, on the order of machine error,

298 Underestimated False Positive Rates While the instabil- 395 the discriminability due to off-target differences across ac-299 ity of brain networks was used here to demonstrate the lim- 396 quisitions and subsamples opens the door for a promising 300 itations of modelling brain-phenotype relationships in the 337 paradigm shift. Given that MCA is data-agnostic, this tech-301 context of machine learning, this limitation extends to classi- 338 nique could be used effectively in conjunction with, or in lieu 302 cal hypothesis testing, as well. Though performing individual 339 of, realistic noise models to augment existing datasets. While 303 comparisons in a hypothesis testing framework will be accom- 340 this of course would not replace the need for repeated measuregod panied by reported false positive rates, the accuracy of these 341 ments when exploring the effect of data collection paradigm 305 rates is critically dependent upon the reliability of the samples 342 or study longitudinal progressions of development or disease, 306 used. In reality, the true false positive rate for a test would be 343 it could be used in conjunction with these efforts to increase 307 a combination of the reported confidence and the underlying 344 the reliability of each distinct sample within a dataset. In 308 variability in the results, a typically unknown quantity.

313 unknown, regardless of the reported false positive rate. In 350 collection. 314 fact, it is a virtual certainty that the true false positive rate 315 for a given hypothesis exceeds the reported value simply as 316 a result of numerical instabilities. This uncertainty inherent 317 to derived data is compounded with traditional arguments 318 limiting the trustworthiness of claims³³, and hampers the 319 ability of researchers to evaluate the quality of results. The accompaniment of brain imaging experiments with direct 321 evaluations of their stability, as was done here, would allow 322 researchers to simultaneously improve the numerical stability 323 of their analyses and accurately gauge confidence in them. 324 The induced variability in derived brain networks may be 325 leveraged to estimate aggregate connectomes with lower bias 326 than any single independent observation, leading to learned 327 relationships that are more generalizable and ultimately more 328 useful.

329 Cost-Effective Data Augmentation The evaluation of reli- 366 and ability in brain imaging has historically relied upon the expen- 367 compare this to numerical instability. Recently, the nearly 331 sive collection of repeated measurements choreographed by 368 boundless space of analysis pipelines and their impact on outmassive cross-institutional consortia^{34,35}. The finding that per- 369 comes in brain imaging has been clearly demonstrated¹. The 333 turbing experiments using MCA both increased the discrim- 370 approach taken in these studies complement one another and 334 inability of the dataset due to biological signal and decreased 371 explore instability at the opposite ends of the spectrum, with

345 contexts where repeated measurements are typically collected When performing these experiments outside of a repeated-346 to increase the fidelity of the dataset, MCA could potentially measure context, such as that afforded here through MCA, it 347 serve as an alternative solution to to increase the reliability of 311 is impossible to empirically estimate the reliability of samples. 348 the dataset and capture more biological variability, with the This means that the reliability of accepted hypotheses is also 349 added result being the savings of millions of dollars on data

> 351 Shortcomings and Future Questions Given the complex-352 ity of recompiling complex software libraries, pre-processing was not perturbed in these experiments as the instrumentation 354 of the canonical workflow used in diffusion image process-355 ing would have added considerable technical complexity and 356 computational overhead to the large set of experiments per-357 formed here. Other work has shown that linear registration, a 358 core piece of many elements of pre-processing such as motion ³⁵⁹ correction and alignment, is sensitive to minor perturbations. 360 It is likely that the instabilities across the entire processing 361 workflow would be compounded with one another, resulting in 362 even greater variability. While the analyses performed in this paper evaluated a single dataset and set of pipelines, extending 364 this work to other modalities and analyses is of interest for 365 future projects.

This paper does not explore methodological flexibility or

human variability in the construction of an analysis workflow on one end and the unavoidable error implicit in the digital representation of data on the other. It is of extreme interest to combine these approaches and explore the interaction of these scientific degrees of freedom with effects from software implementations, libraries, and parametric choices.

Finally, it is important to state explicitly that the work presented here does not invalidate analytical pipelines used in brain imaging, but merely sheds light on the fact that many studies are accompanied by an unknown degree of uncertainty due to machine-introduced errors. The presence of unknown error-bars associated with experimental findings limits the impact of results due to increased uncertainty. The desired outcome of this paper is to motivate a shift in scientific computing – both in neuroimaging and more broadly – towards a paradigm that favours the explicit evaluation of the trustworsthiness of claims alongside the claims themselves.

Methods

390 Dataset

The Nathan Kline Institute Rockland Sample (NKI-RS)²⁹ dataset contains high-fidelity imaging and phenotypic data from over 1,000 individuals spread across the lifespan. A subset of this dataset was chosen for each experiment to both match sample sizes presented in the original analyses and to minimize the computational burden of performing MCA. The selected subset comprises 100 individuals ranging in age from selected subset comprises 100 individuals ranging in age from 6 – 79 with a mean of 36.8 (original: 6 – 81, mean 37.8), 60% female (original: 60%), with 52% having a BMI over 25 (original: 54%).

Each selected individual had at least a single session of both structural T1-weighted (MPRAGE) and diffusion-weighted (DWI) MR imaging data. DWI data was acquired with 137 diffusion directions in a single shell; more information regarding the acquisition of this dataset can be found in the NKI-RS data release²⁹.

In addition to the 100 sessions mentioned above, 25 individuals had a second session to be used in a test-retest analysis.
Two additional copies of the data for these individuals were
generated, including only the odd or even diffusion directions
(64 + 9 B0 volumes = 73 in either case) such that the acquired
data was evenly represented across both portions. This allowed for an extra level of stability evaluation to be performed
between the levels of MCA and session-level variation.

In total, the dataset is composed of 100 subsampled ses-416 sions of data originating from 50 acquisitions and 25 indi-417 viduals for in depth stability analysis, and an additional 100 418 sessions of full-resolution data from 100 individuals for sub-419 sequent analyses.

420 Processing

The dataset was preprocessed using a standard FSL³⁶ work-flow consisting of eddy-current correction and alignment. The MNI152 atlas³⁷ was aligned to each session of data via the structural images, and the resulting transformation was ap-

426 sion data took place after preprocessing was performed on 460 while performing MCA on the output of an operation high-427 full-resolution sessions, ensuring that an additional confound 461 lights round-off errors that may be introduced. The former is 428 was not introduced in this process when comparing between 462 referred to as Precision Bounding (PB) and the latter is called 429 downsampled sessions. The preprocessing described here was 463 Random Rounding (RR). 430 performed once without MCA, and thus is not being evaluated. 464 $_{432}$ data using two canonical pipelines from Dipy 30 : deterministic $_{466}$ distribution of these results can then lead to insights on the and probabilistic. In the deterministic pipeline, a constant 467 stability of the instrumented tools or functions. To this end, 434 solid angle model was used to estimate tensors at each voxel 468 a complete software stack was instrumented with MCA and and streamlines were then generated using the EuDX algo- 469 is made available on GitHub at https://github.com/ 436 rithm³¹. In the probabilistic pipeline, a constrained spherical 437 deconvolution model was fit at each voxel and streamlines 438 were generated by iteratively sampling the resulting fiber ori-439 entation distributions. In both cases tracking occurred with 8 440 seeds per 3D voxel and edges were added to the graph based 441 on the location of terminal nodes with weight determined by 442 fiber count.

The random state of both pipelines was fixed for all anal-444 yses. Fixing this random state led to entirely deterministic 445 repeated-evaluations of the tools, and allowed for explicit at-446 tribution of observed variability to Monte Carlo simulations 447 rather than internal state of the algorithm.

448 Perturbations

449 All connectomes were generated with one reference execu-450 tion where no perturbation was introduced in the processing. 451 For all other executions, all floating point operations were ⁴⁵³ Verificarlo⁹. MCA simulates the distribution of errors im-454 plicit to all instrumented floating point operations (flop). This 488 455 rounding is performed on a value x at precision t by:

$$inexact(x) = x + 2^{e_x - t}\xi \tag{1}$$

dom variable in the range $(-\frac{1}{2}, \frac{1}{2})$. MCA can be introduced in description 494 the 100 sample dataset and 10 times for the repeated measures

425 plied to the DKT parcellation³⁸. Downsampling the diffu- 459 ing MCA on the inputs of an operation limits its precision,

Using MCA, the execution of a pipeline may be performed Structural connectomes were generated from preprocessed 465 many times to produce a distribution of results. Studying the 470 verificarlo/fuzzy.

> Both the RR and PB variants of MCA were used indepen-472 dently for all experiments. As was presented in⁴, both the 473 degree of instrumentation (i.e. number of affected libraries) and the perturbation mode have an effect on the distribution 475 of observed results. For this work, the RR-MCA was applied 476 across the bulk of the relevant operations (those occurring 477 in BLAS, LAPACK, Python, Cython, and Numpy) and is 478 referred to as dense perturbation. In this case the bulk of and numerical operations were affected by MCA.

Conversely, the case in which PB-MCA was applied across 481 the operations in a small subset of operations (those ocurring 482 in Python and Cython) is here referred to as sparse perturba-483 tion. In this case, the inputs to operations within the instru-484 mented libraries were perturbed, resulting in less frequent, 485 data-centric perturbations. Alongside the stated theoretical 452 instrumented with Monte Carlo Arithmetic (MCA)⁸ through 486 differences, sparse perturbation is considerably less computa-487 tionally expensive than dense perturbation.

All perturbations targeted the least-significant-bit for all 489 data (t = 24 and t = 53 in float 32 and float 64, respectively⁹). 490 Perturbing the least significant bit importantly serves as a 491 perturbation of machine error, and thus is the appropriate 492 precision to be applied globally in complex pipelines. Simulawhere e_x is the exponent value of x and ξ is a uniform ran-493 tions were performed 20 times for each pipeline execution for 458 two places for each flop: before or after evaluation. Perform- 495 dataset. A detailed motivation for the number of simulations

496 can be found in³⁹.

497 Evaluation

498 The magnitude and importance of instabilities in pipelines 499 can be considered at a number of analytical levels, namely: 500 the induced variability of derivatives directly, the resulting 501 downstream impact on summary statistics or features, or the 502 ultimate change in analyses or findings. We explore the na-503 ture and severity of instabilities through each of these lenses. 504 Unless otherwise stated, all p-values were computed using 505 Wilcoxon signed-rank tests. To ensure avoid biasing these 506 statistics in this unique repeated-measures context, tests were 507 performed across sets of independent obversations and then 508 the results were aggregated in all cases.

509 Direct Evaluation of the Graphs

measured directly through both a direct variance quantifica- 542 ing a direct measure of the tool-introduced variability across 512 tion and a comparison to other sources of variance such as 543 perturbations. A distribution was formed by aggregating these 513 individual- and session-level differences.

521 the reference graph, and is defined as⁴:

$$\%Dev(A,B) = \sqrt{\sum_{i=1}^{m} \sum_{j=1}^{n} |a_{ij} - b_{ij}|^2} / \sqrt{\sum_{i=1}^{m} \sum_{j=1}^{n} |a_{ij}|^2}, (2)$$

where A and B each represent a graph, and \square_{ii} are el-523 ements therein corresponding to row and column i and j, 554 see respectively. For these experiments, the A graph always refers see at observation j, where $i \neq i'$ and $j \neq j'$. 525 to the reference, where B represents a perturbed value. The 556 526 purpose of this comparison is to provide insight on the scale 557 an observation belonging to a given class will be more simi-527 of differences in observed graphs relative to the original signal 558 lar to other observations within that class than observations

529 in complement to normalized percent deviation to identify 530 the consistency of structure and not just intensity between observed graphs, though the result of this experiment is shown 532 only in Supplemental Section S1.

Finally, the estimated number of significant digits, s', for 534 each edge in the graph is calculated as:

$$s' = -log_{10} \frac{\sigma}{|\mu|} \tag{3}$$

where μ and σ are the mean and unbiased estimator of 536 standard deviation across graphs, respectively. The upper 537 bound on significant digits is 15.7 for 64-bit floating point 538 data.

The percent deviation, correlation, and number of signifi-540 cant digits were each calculated within a single session of data, The differences between perturbation-generated graphs was 541 thereby removing any subject- and session-effects and provid-544 individual results.

Graphs, in the form of adja- 545 Class-based Variability Evaluation To gain a concrete un-515 cency matrices, were compared to one another using three 546 derstanding of the significance of observed variations we exmetrics: normalized percent deviation, Pearson correlation, plore the separability of our results with respect to understood and edgewise significant digits. The normalized percent devi- 548 sources of variability, such as subject-, session-, and pipelineation measure, defined in⁴, scales the norm of the difference 549 level effects. This can be probed through Discriminability²⁶, between a simulated graph and the reference execution (that 550 a technique similar to ICC²⁴ which relies on the mean of a 520 without intentional perturbation) with respect to the norm of 551 ranked distribution of distances between observations belong-552 ing to a defined set of classes. The discriminability statistic is 553 formalized as follows:

$$Disc. = Pr(\|g_{ij} - g_{ij'}\| \le \|g_{ij} - g_{i'j'}\|)$$
 (4)

where g_{ij} is a graph belonging to class i that was measured

Discriminability can then be read as the probability that 528 intensity. A Pearson correlation coefficient 40 was computed 559 of a different class. It is a measure of reproducibility, and ₅₆₀ is discussed in detail in²⁶. This definition allows for the ex-₅₉₄ Evaluating Graph-Theoretical Metrics ploration of deviations across arbitrarily defined classes that 562 in practice can be any of those listed above. We combine 563 this statistic with permutation testing to test hypotheses on 564 whether differences between classes are statistically signifises cant in each of these settings. This statistic is similar to ICC^{24} 566 in a two-measurement setting, however, given the dependence 567 on a rank distribution from all measurements, discriminabil-568 ity scores are not artificially inflated by the addition of more 569 samples which are highly similar to the originals, making it 570 appropriate in this context.

With this in mind, three hypotheses were defined. For 576 dently for each pipeline and perturbation mode.

 H_{A1} : Individuals are distinct from one another Class definition: Subject ID session), MCA (1 subsample, 1 session)

581 H_{A2} : Sessions within an individual are distinct Class definition: Session ID | Subject ID Comparisons: Subsample, MCA (1 subsample) 583

Comparisons: MCA

584 H_{A3} : Subsamples are distinct Class definition: Subsample | Subject ID, Session ID

₅₈₈ periments and 3 reference experiments on 2 pipelines and 2 ₆₂₂ tures, the cumulative density functions of their distributions 589 perturbation modes, resulting in a total of 30 distinct tests. 623 were evaluated over a fixed range and subsequently aggre-590 While results from all tests can be found within Supplemental 624 gated across individuals. The number of significant digits 591 Section S2, only the bolded comparisons in the list above have 625 for each moment of these distributions (sum, mean, variance, 592 been presented in the main body of this article. Correction for 626 skew, and kurtosis) were calculated across observations within 593 repeated testing was performed.

595 While connectomes may be used directly for some analyses, 596 it is common practice to summarize them with structural mea-597 sures, that can then be used as lower-dimensional proxies 598 of connectivity in so-called graph-theoretical studies¹¹. We 599 explored the stability of several commonly-used univariate 600 (graphwise) and multivariate (nodewise or edgewise) features. 601 The features computed and subsequent methods for comparison in this section were selected to closely match those com-603 puted in 10.

each setting, we state the alternate hypotheses, the variable(s) 604 Univariate Differences For each univariate statistic (edge 573 which were used to determine class membership, and the 605 count, mean clustering coefficient, global efficiency, modu-574 remaining variables which may be sampled when obtaining 606 larity of the largest connected component, assortativity, and multiple observations. Each hypothesis was tested indepen- 607 mean path length) a distribution of values across all perturba-608 tions within subjects was observed. A Z-score was computed 609 for each sample with respect to the distribution of feature 610 values within an individual, and the proportion of "classically significant" Z-scores, i.e. corresponding to p < 0.05, was Comparisons: Session (1 subsample), Subsample (1 612 reported and aggregated across all subjects. There was no 613 correction for multiple comparisons in these statistics, as they 614 were not used to interpret a hypothesis but demonstrate the 615 false-positive rate due to perturbations. The number of signifi-616 cant digits contained within an estimate derived from a single 617 subject were calculated and aggregated. The results of this 618 analysis can be found in Supplemental Section S3.

619 Multivariate Differences In the case of both nodewise (de-620 gree distribution, clustering coefficient, betweenness central-As a result, we tested 3 hypotheses across 6 MCA ex- 621 ity) and edgewise (weight distribution, connection length) fea-627 a sample and aggregated.

628 Evaluating A Brain-Phenotype Analysis

629 Though each of the above approaches explores the instabil- 665 630 ity of derived connectomes and their features, many modern 666 publicly available on GitHub at https://github.com/ 632 instance to learn brain-phenotype relationships or identify dif-668 were launched using Boutiques⁴² and Clowdr⁴³ in Compute 633 ferences across groups. We carried out one such study and ex- 669 Canada's HPC cluster environment. MCA instrumentation plored the instability of its results with respect to the upstream 670 was achieved through Verificarlo available on Github at 636 tions. We performed the modeling task with a single sampled 672 A set of MCA instrumented software containers is available connectome per individual and repeated this sampling and 673 on Github at https://github.com/gkiar/fuzzy. 638 modelling 20 times. We report the model performance for 639 each sampling of the dataset and summarize its variance.

 $_{649}$ which explained > 90% of the variance when averaged across 650 the training set for each fold within the cross validation of 686 Concordia.ca. 651 the original graphs; this resulted in a feature of 20 compo-652 nents. We trained the model using k-fold cross validation, with k = 2, 5, 10, and N (equivalent to leave-one-out; LOO).

654 Data & Code Provenance

655 The unprocessed dataset is available through The Consortium 656 of Reliability and Reproducibility (http://fcon_ 1000. 657 projects.nitrc.org/indi/enhanced/), including 658 both the imaging data as well as phenotypic data which may 694 References 659 be obtained upon submission and compliance with a Data Us- 695 [1] age Agreement. The connectomes generated through simula-661 tions have been bundled and stored permanently (https:// $_{662}$ doi.org/10.5281/zenodo.4041549), and are made $_{699}$ [2] C. M. Bennett, M. B. Miller, and G. L. Wolford, "Neural correlates of available through The Canadian Open Neuroscience Platform 700

664 (https://portal.comp.ca/search, search term "Kiar").

All software developed for processing or evaluation is studies employ modeling or machine-learning approaches, for 667 gkpapers/2020ImpactOfInstability. Experiments 835 variability of connectomes characterized in the previous sec-671 https://github.com/verificarlo/verificarlo.

674 Author Contributions

675 GK was responsible for the experimental design, data pro-BMI Classification Structural changes have been linked to 676 cessing, analysis, interpretation, and the majority of writing. obesity in adolescents and adults⁴¹. We classified normal- 677 All authors contributed to the revision of the manuscript. YC, weight and overweight individuals from their structural net- 678 POC, and EP were responsible for MCA tool development and $_{643}$ works (using for overweight a cutoff of BMI $> 25^{13}$). We $_{679}$ software testing. AR, GV, and BM contributed to experimenreduced the dimensionality of the connectomes through prin- 680 tal design and interpretation. TG contributed to experimental 645 cipal component analysis (PCA), and provided the first N-681 design, analysis, and interpretation. TG and ACE were re-646 components to a logistic regression classifier for predicting 682 sponsible for supervising and supporting all contributions BMI class membership, similar to methods shown in 12,13. 683 made by GK. The authors declare no competing interests for The number of components was selected as the minimum set 684 this work. Correspondence and requests for materials should 685 be addressed to Tristan Glatard at tristan.glatard@

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S1. Graph Correlation

The correlations between observed graphs (Figure S1) across each grouping follow the same trend to as percent deviation, as shown in Figure 1. However, notably different from percent deviation, there is no significant difference in the correlations between dense or sparse instrumentations. By this measure, the probabilistic pipeline is more stable in all cross-MCA and cross-directions except for the combination of sparse perturbation and cross-MCA (p < 0.0001 for all; exploratory).

The marked lack in drop-off of performance across these settings, inconsistent with the measures show in Figure 1 is likely due to the nature of the measure and the structure of graphs being compared. Given that structural graphs are sparse and contain considerable numbers of zero-weighted edges, the presence or absense of edges dominated the correlation measure where it was less impactful for the others. For this reason and others 44, correlation is not a commonly used measure in the context of structural connectivity, and thus this analysis was demoted to the supplement material.

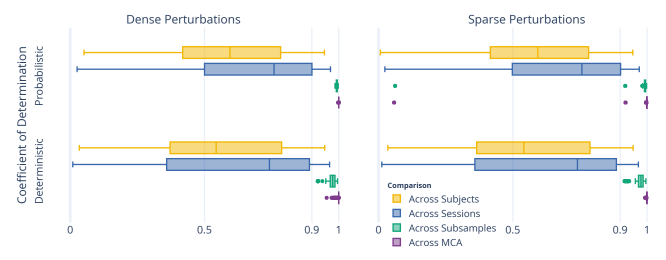


Figure S1. The correlation between perturbed connectomes and their reference.

S2. Complete Discriminability Analysis

Table S1. The complete results from the Discriminability analysis, with results reported as mean \pm standard deviation Discriminability. As was the case in the condensed table, the alternative hypothesis, indicating significant separation across groups, was accepted for all experiments, with p < 0.005.

				Reference Execution		Dense Pertu	rbations	Sparse Perturbations	
Exp.	Subj.	Sess.	Samp.	Det.	Prob.	Det.	Prob.	Det.	Prob.
1.1	All	All	1	0.64 ± 0.00	0.65 ± 0.00	0.82 ± 0.00	0.82 ± 0.00	0.77 ± 0.00	0.75 ± 0.00
1.2	All	1	All	1.00 ± 0.00	1.00 ± 0.00	1.00 ± 0.00	1.00 ± 0.00	0.93 ± 0.02	0.90 ± 0.02
1.3	All	1	1			1.00 ± 0.00	1.00 ± 0.00	0.94 ± 0.02	0.90 ± 0.02
2.4	1	All	All	1.00 ± 0.00	1.00 ± 0.00	1.00 ± 0.00	1.00 ± 0.00	0.88 ± 0.12	0.85 ± 0.12
2.5	1	All	1			1.00 ± 0.00	1.00 ± 0.00	0.89 ± 0.11	0.84 ± 0.12
3.6	1	1	All			0.99 ± 0.03	1.00 ± 0.00	0.71 ± 0.07	0.61 ± 0.05

The complete discriminability analysis includes comparisons across more axes of variability than the condensed version.

The reduction in the main body was such that only axes which would be relevant for a typical analysis were presented. Here,
eso each of Hypothesis 1, testing the difference across subjects, and 2, testing the difference across sessions, were accompanied
with additional comparisons to those shown in the main body.

Subject Variation Alongside experiment 1.1, that which mimicked a typical test-retest scenario, experiments 1.2 and 1.3 could be considered a test-retest with a handicap, given a single acquisition per individual was compared either across subsamples or simulations, respectively. For this reason, it is unsurprising that the dataset achieved considerably higher discriminability scores.

Session Variation Similar to subject variation, the session variation was also modelled across either both or a single subsample in experiments 2.4 and 2.5. In both of these cases the performance was similar, and the finding that sparse perturbations reduced the off-target signal was consistent.

S3. Univariate Graph Statistics

Figure S2 explores the stability of univariate graph-theoretical metrics computed from the perturbed graphs, including modularity, global efficiency, assortativity, average path length, and edge count. When aggregated across individuals and perturbations, the distributions of these statistics (Figures S2A and S22B) showed no significant differences between perturbation methods for either deterministic or probabilistic pipelines, consistent with the comparison of the cumulative density of the multivariate statistics compared in 2.

However, when quantifying the stability of these measures across connectomes derived from a single session of data, the two perturbation methods show considerable differences. The number of significant digits in univariate statistics for dense perturbation instrumented connectome generation exceeded 11 digits for all measures except modularity, which contained more than 4 significant digits of information (Figure S2C). When detecting false-positives from the distributions of observed statistics for a given session, the rate (using a threshold of p = 0.05) was approximately 2% for all statistics with the exception of modularity which again was less stable with an approximately 10% false positive rate. The probabilistic pipeline is significantly more stable than the deterministic pipeline (p < 0.0001; exploratory) for all features except modularity. When similarly evaluating these features from connectomes generated in the sparse perturbation setting, no statistic was stable with more than 3 significant digits or a false positive rate lower than nearly 6% (Figure S2D). The deterministic pipeline was more stable than the probabilistic pipeline in this setting (p < 0.0001; exploratory).

Two notable differences between the two perturbation methods are, first, the uniformity in the stability of the statistics, and second, the dramatic decline in stability of individual statistics in the sparse perturbation setting despite the consistency in the overall distribution of values. This result is consistent with that obtained from the multivariate exploration performed in the body of this article. It is unclear at present if the discrepancy between the stability of modularity in the pipeline perturbation context versus the other statistics suggests the implementation of this measure is the source of instability or if it is implicit to the measure itself. The dramatic decline in the stability of features derived from sparse perturbed graphs despite no difference in their overall distribution both shows that while individual estimates may be unstable the comparison between aggregates or groups may be considered much more reliable; this finding is consistent with that presented for multivariate statistics.

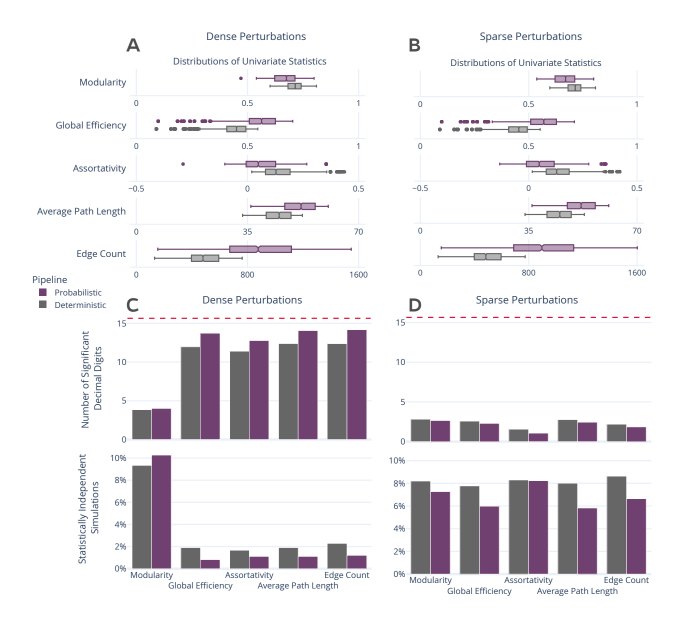


Figure S2. Distribution and stability assessment of univariate graph statistics. (**A**, **B**) The distributions of each computed univariate statistic across all subjects and perturbations for dense and sparse settings, respectively. There was no significant difference between the distributions in A and B. (**C**, **D**; top) The number of significant decimal digits in each statistic across perturbations, averaged across individuals. The dashed red line refers to the maximum possible number of significant digits. (**C**, **D**; bottom) The percentage of connectomes which were deemed significantly different (p < 0.05) from the others obtained for an individual.