Individual differences in spatial working memory strategies differentially reflected in the engagement of control and default brain networks

Nina Purg Suljič^{a*}, Aleksij Kraljič^a, Masih Rahmati^b, Youngsun T. Cho^b, Anka Slana Ozimič^a, John D. Murray^{b,c,d}, Alan **Anticevic**b,c+**, and Grega Repovš**a+

> ^aDepartment of Psychology, Faculty of Arts, University of Ljubljana, Ljubljana, Slovenia **b** Department of Psychiatry, Yale University School of Medicine, New Haven, CT, USA ^cDepartment of Psychology, Yale University School of Medicine, New Haven, CT, USA dDepartment of Physics, Yale University, New Haven, CT, USA Corresponding author

Co-senior authors

Spatial locations can be encoded and maintained in working 40 2 memory using different representations and strategies. Fine-3 grained representations provide detailed stimulus information, $_{42}$ but are cognitively demanding and prone to inexactness. The $_{43}$ ⁵ uncertainty in fine-grained representations can be compensated by the use of coarse, but robust categorical representations. In this study, we employed an individual differences approach to identify brain activity correlates of the use of fine-grained and categorical representations in spatial working memory. We ¹⁰ combined data from six fMRI studies, resulting in a sample of $11 - 155$ (77 women, 25 ± 5 years) healthy participants performing a ¹² spatial working memory task. Our results showed that individ-¹³ ual differences in the use of spatial representations in working 14 memory were associated with distinct patterns of brain activ- $_{52}$ ¹⁵ ity. Higher precision of fine-grained representations was related to greater engagement of attentional and control brain systems $_{54}$ ¹⁷ throughout the task trial, and the stronger deactivation of the ¹⁸ default network at the time of stimulus encoding. In contrast, ¹⁹ the use of categorical representations was associated with lower ²⁰ default network activity during encoding and higher frontopari-²¹ etal network activation during maintenance. These results may 22 indicate a greater need for attentional resources and protection 59 ²³ against interference for fine-grained compared to categorical ²⁴ representations.

25 **spatial cognition, working memory, fMRI, fine-grained representation,** 26 **categorical representation**

27 **Correspondence:** *nina.purg@ff.uni-lj.si*

²⁸ **Introduction**

 Research on working memory has shown that individuals use a variety of different representations and strategies to encode and maintain information over short periods of time in sup- 70 32 port of an ongoing task (e.g., [Curtis,](#page-15-0) [2004;](#page-15-0) [Oblak et al.,](#page-16-0) [2024,](#page-16-0) 71 [2022;](#page-16-2) [Purg et al.,](#page-16-2) 2022; Slana Ozimič et al., [2023;](#page-16-3) [Starc et al.,](#page-16-4) 72 $34 \quad 2017$). While mental representations describe the content of 73 information encoded in working memory, cognitive strate- gies refer to the selection of mental representations and pro-37 cesses that are either explicitly or implicitly used by an in- 76 38 dividual to perform a working memory task [\(Miller et al.,](#page-16-5) 77 [2012;](#page-16-5) [Oblak et al.,](#page-16-0) [2024,](#page-16-0) [2022;](#page-16-1) Slana Ozimič et al., [2023\)](#page-16-3). 78

The specific representations and strategies used in working memory depend on several factors, such as the type of in[f](#page-16-3)ormation to be retained [\(Oblak et al.,](#page-16-1) [2022;](#page-16-1) Slana Ozimič [et al.,](#page-16-3) [2023\)](#page-16-3), the type and predictability of a response to be generated [\(Curtis,](#page-15-0) [2004;](#page-15-0) [Purg et al.,](#page-16-2) [2022\)](#page-16-2), the availability of attentional resources [\(Adam et al.,](#page-14-0) [2015;](#page-14-0) [Starc et al.,](#page-16-4) [2017\)](#page-16-4), and behavioral relevance [\(Klyszejko et al.,](#page-15-1) [2014;](#page-15-1) [Yoo et al.,](#page-16-6) 47 [2022\)](#page-16-6). Increasingly, research also shows that even when faced with the same task requirements, individuals may use different representations and strategies to perform the task 50 [\(Oblak et al.,](#page-16-0) [2024,](#page-16-0) [2022;](#page-16-1) Slana Ozimič et al., [2023;](#page-16-3) [Starc](#page-16-4) ⁵¹ [et al.,](#page-16-4) [2017\)](#page-16-4). Here, we investigate the neural correlates of individual differences in the use of working memory strategies in a multi-study, multi-site dataset of spatial working memory ⁵⁴ performance during functional magnetic resonance imaging ⁵⁵ (fMRI).

Spatial working memory enables the short-term storage ⁵⁷ of spatial information, such as the location of a stimulus. Extensive research has shown that memory for a stimulus lo[c](#page-15-2)ation is affected by systematic distortions (e.g., [Crawford](#page-15-2) ϵ_0 [et al.,](#page-15-2) [2016;](#page-15-2) [Huttenlocher et al.,](#page-15-3) [2004,](#page-15-3) [1991\)](#page-15-4). In particular, it ⁶¹ has been observed that when individuals are asked to repro-⁶² duce a stimulus location stored in working memory within ⁶³ an empty circle, they exhibit systematic shifts in their re-⁶⁴ sponses towards the diagonals of the four quadrants, formed ⁶⁵ by dividing the circle using the horizontal and vertical axes ⁶⁶ of symmetry [\(Huttenlocher et al.,](#page-15-3) [2004,](#page-15-3) [1991\)](#page-15-4). These sys-⁶⁷ tematic biases in spatial working memory performance have been suggested to reveal a hierarchical organization of spatial representations [\(Huttenlocher et al.,](#page-15-4) [1991\)](#page-15-4).

According to the category adjustment model [\(Hutten](#page-15-4)[locher et al.,](#page-15-4) [1991,](#page-15-4) [2000\)](#page-15-5), a stimulus location is encoded and maintained at two levels of representation – first, as a precise, fine-grained representation that stores the information of the actual location in memory, and second, as a categori-⁷⁵ cal representation that assigns the stimulus location to one of a limited number of spatial categories (e.g., quadrants). The model predicts that the estimation of a stimulus location results from the combination of information at both levels, with

 79 the use of a categorical representation helping to compensate 136 80 for the loss of precision in a fine-grained representation. Even 137 81 though this process introduces a systematic bias in individual 138 82 responses away from the correct position toward the proto-139 ⁸³ typical location of the spatial category, it is assumed to in-84 crease the overall response accuracy by decreasing the vari-141 85 ability of responses. At the neural level, the dynamic field 142 86 theory [\(Schutte et al.,](#page-16-8) [2003;](#page-16-8) [Simmering et al.,](#page-16-9) [2006\)](#page-16-9) suggests 143 87 that spatial boundaries, such as perceivable edges or sponta-144 88 neously imposed axes of symmetry in task space, have a de-145 89 flecting effect on memory-guided behavioral responses due 146 ⁹⁰ to their lateral inhibitory effects, causing the activation pro-91 duced by the target stimulus stored in working memory to 148 92 drift in the opposite direction.

93 The degree of reliance on fine-grained and categorical ₁₅₀ 94 coding of spatial locations has been related to variability 151 95 in cognitive resources. In our previous work [\(Starc et al.,](#page-16-4) 152 $96\quad 2017$), we separately estimated the use of fine-grained and $_{153}$ 97 categorical representations during the performance of a spa-98 tial working memory task, while measuring pupil responses. 155 99 We assumed that increased pupil dilation would reflect in- $_{156}$ 100 creased cognitive effort exerted toward the formation and 157 101 maintenance of either fine-grained or categorical representa-102 tions. Our results were consistent with a compensatory use of 159 103 fine-grained and categorical representations within individu-104 als, where a drop in attentional resources directed towards 161 105 the formation of fine-grained representations during stimulus 162 106 encoding resulted in increased reliance on categorical repre- 107 sentations during late maintenance and response phases of 164 108 the task. Additionally, we observed that individuals who 165 ¹⁰⁹ showed on average worse fine-grained precision also exhib-¹¹⁰ ited greater overall use of categorical representations, sug- 111 gesting stable individual differences in the use of specific rep- $_{168}$ ¹¹² resentations and strategies.

113 Similarly, [Crawford et al.](#page-15-2) (2016) found individual dif- $_{170}$ ferences in fine-grained and categorical spatial coding that $_{171}$ 115 were correlated with individual spatial working memory ca- $_{172}$ pacity. Specifically, individuals with better spatial working 173 memory capacity showed higher fine-grained memory preci- 174 sion and lower reliance on categorical representations. Since 175 119 working memory capacity describes the limited cognitive re- $_{176}$ 120 sources that can be directed towards storage of information $_{177}$ in working memory, either at the level of attentional alloca- 178 tion or representational capacities (Slana Ozimič and Repovš, 122 [2020\)](#page-16-10), these results suggest that individual differences in the $_{180}$ use of fine-grained and categorical representations might be $_{181}$ explained by the availability of cognitive resources with fine- 182 grained representations requiring more resources than cate- $_{183}$ 127 gorical representations.

128 Despite the extensive behavioral and computational char-185 acterization of fine-grained and categorical spatial coding, 130 not much is known about the underlying neurobiological 187 131 mechanisms. Spatial working memory is consistently charac-188 terized by sustained activation in frontal and parietal brain ar- eas as measured with electrophysiological recordings in non[h](#page-15-7)uman primates (e.g., [Chafee and Goldman-Rakic,](#page-15-6) [1998;](#page-15-6) Fu-191 [nahashi et al.,](#page-15-7) [1989;](#page-15-7) [Fuster,](#page-15-8) [1973;](#page-15-8) [Fuster and Alexander,](#page-15-9)

[1971;](#page-15-9) [Kubota and Niki,](#page-15-10) [1971\)](#page-15-10) and fMRI in humans (e.g., [Brown et al.,](#page-15-11) [2004;](#page-15-0) [Courtney et al.,](#page-15-12) [1998;](#page-15-12) [Curtis,](#page-15-0) 2004; [Sri](#page-16-11)[mal and Curtis,](#page-16-11) [2008;](#page-16-11) [Zarahn et al.,](#page-16-12) [1999\)](#page-16-12). This activity is thought to reflect active engagement of these areas in working memory processes, however, the specific function of this activity has been more difficult to identify. Relating brain activity with behavioral performance of working memory tasks during fMRI has shown that brain activity varies with the level of response precision [\(Curtis,](#page-15-0) [2004;](#page-15-0) [Hallenbeck et al.,](#page-15-13) [2021\)](#page-15-13), specific strategy use [\(Curtis,](#page-15-0) [2004;](#page-15-0) [Purg et al.,](#page-16-2) [2022\)](#page-16-2), general memory load [\(Adam et al.,](#page-15-14) [2018;](#page-15-14) [Glahn et al.,](#page-15-15) [2002;](#page-15-15) [Leung et al.,](#page-15-16) [2004;](#page-15-16) [Linden et al.,](#page-15-17) [2003;](#page-15-17) [Proskovec et al.,](#page-16-13) [2019\)](#page-16-13) and behavioral prioritization [\(Klyszejko et al.,](#page-15-1) [2014;](#page-15-1) ¹⁴⁹ [Yoo et al.,](#page-16-6) [2022\)](#page-16-6).

In a previous fMRI study [\(Anticevic et al.,](#page-15-18) [2010\)](#page-15-18), we investigated the relationship between response accuracy in a visual working memory task and brain activity during the task. Our results showed that stronger deactivation in the temporo-parietal junction (TPJ) and the default network during stimulus encoding predicted higher accuracy of working memory performance. Since TPJ and the default network have been associated with stronger deactivation during increased cognitive effort and inhibition of distractors [\(Raichle,](#page-16-14) [2015a;](#page-16-14) [Shulman et al.,](#page-16-15) [2003;](#page-16-15) [Todd et al.,](#page-16-16) [2005\)](#page-16-16), these results suggest that their suppression may be related ¹⁶¹ to increased cognitive effort that is required to ensure good memory accuracy and protection from interference. However, the study used non-spatial visual stimuli and match-tosample responses that do not allow the estimation of separate contribution of fine-grained and categorical representations ¹⁶⁶ to behavioral responses. Therefore, the brain systems and related mechanisms underlying fine-grained and categorical spatial coding have yet to be determined.

In the present study, we were interested in brain activity correlates of individual differences in the use of fine-grained and categorical representations in spatial working memory. Due to the hypothesized relationship between the use of these working memory representations and the level of cognitive resources required, we focused on brain systems that have been previously associated with general engagement of attention and cognitive control, specifically the cinguloopercular, dorsal-attention, and frontoparietal networks (e.g., [Barch et al.,](#page-15-19) [2013;](#page-15-19) [Cole et al.,](#page-15-20) [2014;](#page-15-20) [Ji et al.,](#page-15-21) [2019;](#page-15-21) [Raichle,](#page-16-14) [2015a;](#page-16-14) [Smith et al.,](#page-16-17) [2009\)](#page-16-17). In addition, we investigated the role of the default network in the use of fine-grained and categorical representations, which has been associated with stronger inhibition during high attentional demands and the function of providing protection from distractors in working ¹⁸⁴ [m](#page-15-21)emory tasks (e.g., [Barch et al.,](#page-15-19) [2013;](#page-15-19) [Cole et al.,](#page-15-20) [2014;](#page-15-20) [Ji](#page-15-21) [et al.,](#page-15-21) [2019;](#page-15-21) [Raichle,](#page-16-14) [2015a;](#page-16-14) [Smith et al.,](#page-16-17) [2009\)](#page-16-17). We hypothesized that a greater reliance on precise, fine-grained representations would be supported by increased activation of attentional and control brain systems, and a stronger inhibition of the default network. On the other hand, we assumed that uncertainty in fine-grained representations, such as due to a loss of precision or task interference, would be accompanied by a greater reliance on categorical representations that require

¹⁹³ fewer attentional and control resources.

¹⁹⁴ To test these hypotheses, we investigated brain activ-¹⁹⁵ ity measured with fMRI during the performance of a spa-¹⁹⁶ tial working memory task. A methodological challenge in ¹⁹⁷ the investigation of individual differences in brain-behavior ¹⁹⁸ relationships are low effect sizes that require large sample 199 sizes to be detected [\(Elliott et al.,](#page-15-22) [2020;](#page-15-22) [Grady et al.,](#page-15-23) [2021;](#page-15-23) 255 200 [Marek et al.,](#page-15-24) [2022\)](#page-15-24). To overcome this challenge, we com-256 201 bined six fMRI studies conducted at two different recording 257 ²⁰² sites. Together, we used data from 155 (77 women, 25 ± 5 ²⁵⁸ ²⁰³ years) healthy individuals, which largely exceeded the aver-²⁰⁴ age sample sizes of similar studies (e.g., around 25 partici-205 pants, [Marek et al.,](#page-15-24) [2022\)](#page-15-24). Based on behavioral performance 261 ²⁰⁶ on the task, we estimated the overall reliance on fine-grained ²⁰⁷ and categorical representations of each participant by decom-²⁰⁸ posing their contributions to task response errors. Individ-²⁰⁹ ual use of fine-grained and categorical representations was ²¹⁰ then related to differences in levels of brain activity. Our results revealed individual differences in the use of spatial 267 212 representations in working memory that were related to dis-268 ²¹³ tinct patterns of brain activity. Ongoing engagement of atten-²¹⁴ tional and control brain networks throughout the entire task ²¹⁵ trial, and stronger deactivation of the default network at the ²¹⁶ time of encoding a stimulus location were found to predict²⁷⁰ 217 higher fine-grained precision in spatial working memory per-²⁷¹ 218 formance. In contrast, the use of a categorical representa-²⁷² 219 tion was associated with lower default network activity in the ²⁷³ 220 encoding period and higher frontoparietal network activation^{274} $_{221}$ in the delay period. These results suggest that the forma- 275 222 tion, maintenance and recall of fine-grained representations is 276 223 supported by an increased allocation of attentional resources 277 224 provided by attentional and control brain networks, whereas²⁷⁸ ²²⁵ the categorical representations do not seem to impose such²⁷⁹ attentional demands and may be associated with an inability²⁸⁰ ₂₂₇ to protect the fine-grained representation from interference, ²⁸¹ ²²⁸ resulting in higher reliance on the categorical representation

²²⁹ when providing the response.

²³⁰ **Materials and Methods**

²³¹ **Participants**

232 We combined data from six studies (Figure [1A](#page-4-0)). Three stud-288 ies (Studies I-III; Table S1) were conducted at the University of Ljubljana, Slovenia, and three studies (Studies IV-VI; Ta- ble S1) at Yale University, USA. Between 11 and 37 partici- pants took part in each study, for a total of 166 participants. 237 All participants were healthy adults with no current or previ-293 ous neurological, psychiatric, or substance-use disorders. Ex- clusion criteria also included contraindications to MR, such as the presence of metal implants or any other metal parti-241 cles in the body, history of epileptic seizures, tremor or other 297 motor disorders, and pregnancy. All participants had nor- mal or corrected-to-normal vision. Several participants were excluded from further data analysis due to incomplete data 245 collection ($N = 5$), failure to follow instructions ($N = 1$), 301 poor data quality, or excessive movement during data collec- $_{247}$ tion ($N = 2$). We also excluded participants who deviated $_{303}$ 248 greatly from the group mean age (i.e., greater than $3 \times SD$) 304

249 to ensure a more homogeneous sample $(N = 2)$. Furthermore, we excluded an outlier in neuroimaging data $(N = 1)$, as explained in detail in the section *fMRI acquisition, pre-*²⁵² *processing and analysis*. Data from the remaining 155 (77 women, $25±5$ years) participants were used for further analysis. Most participants were right-handed (90.9%), while the rest of the participants were left-handed (11 participants, ²⁵⁶ 7*.*14%) or ambidextrous (3 participants, 1*.*95%). All participants performed the behavioral task with their dominant hand. Detailed demographic information of the participants included in the data analysis are presented in Table S1. The studies carried out at the University of Ljubljana were approved by the Ethics Committee of the Faculty of Arts, Uni-²⁶² versity of Ljubljana, and the National Medical Ethics Committee, Ministry of Health of the Republic of Slovenia. The studies conducted at Yale University were approved by the Yale Institutional Review Board. Participants gave written informed consent before participating in the study. In all studies, participants had to perform a spatial working memory task while their brain activity was measured with fMRI.

Spatial working memory task

Individual studies were primarily conducted to address different research questions related to spatial working memory. Some of the studies are described elsewhere [\(Moujaes et al.,](#page-16-18) [2024;](#page-16-18) [Purg et al.,](#page-16-2) [2022\)](#page-16-2), while others are yet unpublished. The studies also differed slightly in the exact details of the spatial working memory task, which included different task conditions in each study. For the purposes of this paper, we only analyzed the task conditions that were most comparable across the studies. In particular, we focused our investigation on the task condition in which participants were asked to remember the position of a briefly presented target stimulus and, after a short delay period, to move a probe using a joystick to the position of the remembered target (Figure ²⁸³ [1B](#page-4-0)). Despite minor differences in task design across studies, 284 the goal of the task was always the same – a single stimulus ²⁸⁵ location had to be remembered, maintained, and then recre-²⁸⁶ ated with a joystick on each trial. The task was displayed ²⁸⁷ on an MR-compatible screen that was visible to participants from the MR scanner via a head mirror. The specific screen sizes and resolutions varied depending on the recording site and study as described in detail in Table S2. The tasks were ²⁹¹ prepared using custom scripts and run in PsychoPy (Studies I–III; Table S2; [Peirce et al.,](#page-16-19) [2019\)](#page-16-19) or E-Prime 2.0 (Stud-ies IV-VI; Table S2; [Schneider et al.,](#page-16-20) [2012\)](#page-16-20). Participants responded with an MR-compatible joystick (Hybridmojo LLC, Washington, USA).

The spatial working memory task differed in the time course of task events and the exact range of target locations across studies (for details see Table S2). In three studies (Studies I-III; Table S2), the trial started with the presentation of a fixation point (2.5 s) in the center of the screen, followed by a brief presentation of a target disk stimulus. In the remaining three studies (Studies IV-VI; Table S2), the trial started immediately with the presentation of a target disk stimulus. Target stimulus presentation lasted between 0.1 s

305 and 2 s, depending on the study. The target stimuli were 361 306 presented at variable locations that were pseudorandomly se- 362 307 lected from 20 to 36 different possible locations, depending 363 308 on the study. Target locations were chosen such that the target 364 309 amplitude (i.e., radial distance) from the center of the screen 365 310 was constant for each participant, whereas target angles from 366 311 the center of the screen varied between trials for the same 367 312 participant (see Table S2 for details on the exact target am-368) 313 plitudes and angles for each study). The target stimuli were 369 314 never presented on the cardinal axes to prevent verbalization 370 315 of precise locations [\(Srimal and Curtis,](#page-16-11) [2008\)](#page-16-11). Participants 371 316 were instructed to memorize the exact position of the target 372 317 stimulus. In one study (Study I; Table S2), the presentation of 373 ³¹⁸ the target stimulus was followed by a masking pattern (0.05 374) 319 s) with the aim of disrupting iconic visual memory [\(Curtis,](#page-15-0) 375 $320 \ 2004$). In all studies, the target presentation was followed by 376 321 a delay period (8 s to 10.4 s, depending on the study) dur- 377 322 ing which a fixation point was presented in the center of the 378 323 screen to which participants were asked to direct their gaze. 379 324 In three studies (Studies IV-VI; Table S2), gaze fixation was 380 325 additionally enforced by instructing participants to press a 381 326 button upon a change of color of the fixation cross, which 382 327 occurred randomly in 50% of trials. After the delay, a probe 383 ³²⁸ (i.e., a disk stimulus of the same size as the target stimulus, 329 but a different color) appeared in the center of the screen, and 385 330 participants were instructed to move the probe using a joy-386 331 stick to the location of the previously presented target stimu-387 ³³² lus, as precisely as possible. The time of their response was ³³³ limited due to the concurrent fMRI recording between 2.3 s³⁸⁹ 334 and 3 s, depending on the study. Individual trials were sep-390 335 arated by an inter-trial interval (ITI) that was either fixed in 391 336 duration (Studies IV-VI; Table S2) or randomly varied to al-392 337 low for better task-related fMRI signal decomposition (Stud-393 338 ies I-III; Table S2). Participants performed between 20 and 394 339 80 trials of the task, divided into 1 to 4 blocks, depending on 395 ³⁴⁰ the study.

³⁴¹ **Behavioral data analysis**

 342 In behavioral data analysis, we first converted all behavioral $_{400}$ 343 data from pixel-based measurements into degrees of visual $_{401}$ $_{344}$ angle (\degree va) to provide standardization across different screen $_{402}$ ³⁴⁵ resolutions and viewing distances. At the level of individ-³⁴⁶ ual participants, we calculated trial-to-trial response errors as 347 the difference between the final location of the response in 348 relation to the target location, which are thought to reflect 403 349 the precision of spatial working memory. Since the find-404 ³⁵⁰ ings of single-neuron recordings suggest that spatial repre-351 sentations are encoded at the neural level in terms of angle 406 352 [a](#page-15-6)nd amplitude in the polar coordinate system (e.g., [Chafee](#page-15-6) 407 ³⁵³ [and Goldman-Rakic,](#page-15-6) [1998;](#page-15-6) [Funahashi et al.,](#page-15-7) [1989;](#page-15-7) [Rainer](#page-16-21) 354 [et al.,](#page-16-21) [1998\)](#page-16-21), we decomposed the response error on each trial 409 355 into angular and amplitude differences between target and 410 356 response locations measured from the center of the screen. 411 357 Next, we excluded all invalid or outlier responses to ensure 412 ³⁵⁸ that the results reflected the engagement of spatial working 359 memory and not any technical errors or inattention to the 414 ³⁶⁰ task. We defined outliers as any response that was located

During the performance of the task, only the stimulus angle was varied, while the stimulus amplitude remained constant for each participant. Thus, we assumed that memory processes would be more strongly reflected in angular response errors than in amplitude response errors, and focused our further analyses on angular response errors only. To delineate the individual effects of fine-grained and categorical representations on response errors we relied on the as-sumptions of the category adjustment model [\(Crawford et al.,](#page-15-2) ³⁷⁸ [2016;](#page-15-2) [Duffy et al.,](#page-15-25) [2010;](#page-15-25) [Huttenlocher et al.,](#page-15-3) [2004,](#page-15-3) [1991,](#page-15-4) [2000\)](#page-15-5). The model proposes that the estimation of stimulus location retained in working memory results from the combined use of fine-grained and categorical representations, each prone to decay and associated inexactness. Addition-ally, studies [\(Haun et al.,](#page-15-26) [2005;](#page-15-26) [Huttenlocher et al.,](#page-15-3) [2004,](#page-15-3) ³⁸⁴ [1991;](#page-15-4) [Purg et al.,](#page-16-2) [2022;](#page-16-2) [Starc et al.,](#page-16-4) [2017\)](#page-16-4) have shown that when participants are asked to recall the position of a stimulus in a blank space, such as in the case of our study, they use four quadrants, delineated by the horizontal and vertical axes, as spatial categories, with the central value located at their corresponding diagonals, acting as the category prototype. Hence, behavioral responses collected during the spatial working memory task are assumed to be composed of a systematic shift toward the categorical center (i.e., the prototype) with the associated inexactness of this information, in addition to variability around the shifted representation due to a loss of fine-grained precision.

³⁹⁶ Computationally, we used a Bayesian model (Figure [1C](#page-4-0)), ³⁹⁷ [p](#page-15-2)reviously explained in detail in several publications [\(Craw-](#page-15-2)³⁹⁸ [ford et al.,](#page-15-2) [2016;](#page-15-2) [Duffy et al.,](#page-15-25) [2010;](#page-15-25) [Huttenlocher et al.,](#page-15-3) 399 [2004,](#page-15-3) [1991,](#page-15-4) [2000\)](#page-15-5), where the response (R) was modeled as a weighted sum of the fine-grained memory location (M) and the location of the categorical prototype (P) , while the contribution of each component was defined by λ :

$R = \lambda M + (1 - \lambda)P$

The memory location (M) was defined as the true target location (i.e., the target angle from the center of the screen; μ) with the associated standard deviation (σ_M) reflecting memory inexactness. Similarly, the prototype location (P) was ⁴⁰⁷ centered on the diagonal of the quadrant in which the target stimulus was presented (i.e., the angle of the corresponding diagonal; ρ), its inexactness reflected by the standard deviation around the prototype location (σ *P*). λ reflected confidence in the memory representation, while $1 - \lambda$ defined the degree of bias toward the use of prototype information. Mathematically, λ was defined as the ratio between the inexactness of the prototype compared to the combined inexact[n](#page-15-2)ess of the prototype and memory representations [\(Crawford](#page-15-2)

Site: University of Ljubljana Vale University of Ljubljana Vale University Study: | Study I *n* = 27 Study II *n* = 26 Study III *n* = 30 Study IV *n* = 37 Study V *n* = 25 Study VI Participants: | n = 27 || n = 26 || n = 30 || n = 37 || n = 25 || n = 10 Total: $n = 155$

A. Multi-site and multi-study fMRI dataset

B. Spatial working memory task

C. Estimation of behavioral measures

Fig. 1. Overview of the dataset structure and behavioral methods. A. [The dataset included six fMRI studies of spatial working memory, conducted at two different sites.](#page-15-2) In total, 155 participants were included in the data analysis. **B.** [Common elements of a spatial working memory task across all studies. Each task trial consisted of a brief](#page-15-2) [presentation of a target stimulus at different angles and a constant amplitude from the center of the screen, followed by a hand response to the target location using a joystick](#page-15-2) [after a short delay. ITI refers to the inter-trial interval.](#page-15-2) **C.** An illustration of how the memory inexactness (*σM*) and the prototype bias (1−*λ*) were calculated based on angular [response errors as measures of the use of fine-grained and categorical representations, respectively.](#page-15-2)

⁴¹⁶ [et al.,](#page-15-2) [2016;](#page-15-2) [Duffy et al.,](#page-15-25) [2010;](#page-15-25) [Huttenlocher et al.,](#page-15-4) [1991\)](#page-15-4):

$$
\lambda = \frac{\sigma_P^2}{\sigma_P^2 + \sigma_M^2}
$$

⁴¹⁷ In this way, we modeled the assumption that the more⁴³⁵ 418 inexact the memory representation is compared to the proto-436 419 type, the lower the reliance on the fine-grained memory of the 437 420 target and the higher the contribution of the prototype when 438 ⁴²¹ estimating stimulus location.

⁴²² The parameters of the Bayesian model were estimated ⁴⁴⁰ 423 using the probabilistic programming language Stan [\(Team,](#page-16-22) 441) 424 [2022b\)](#page-16-22) in R [\(Team,](#page-16-23) [2022a\)](#page-16-23). We estimated the posterior prob-442 425 abilities of λ , σ_M and σ_P for each participant using a two-⁴⁴³ 426 level linear model by fitting the Student's t-distribution to 444 427 the data. Estimates were obtained based on multiple task 445 428 trials per each participant, thus participants were used as a 446 429 grouping variable at the first level to model varying intercepts 447 430 across participants. The model was run separately for each 448

⁴³¹ study to prevent the potential influence of different study de-⁴³² signs and protocols on behavioral performance. Weakly in-⁴³³ formative prior distributions were used for all model param-⁴³⁴ eters, ensuring that the standard deviation of the prior distribution was at least 10 times larger than that of the posterior distribution. Specifically, we used normal prior distribution for regression parameters and half-normal distributions for standard deviations. The prior distributions were ⁴³⁹ centered at mean values of the posterior parameter estimates computed with a preliminary one-level regression model to ensure stable sampling convergence. The prior distribution for the degrees of freedom parameter was set to $\Gamma(2,0.1)$ as recommended by [Juárez and Steel](#page-15-27) [\(2010\)](#page-15-27). The stability of the Hamiltonian Monte Carlo (HMC) sampling algorithm was analyzed by verifying that all estimated parameters had estimated effective sample sizes in the bulk of the distributions and in the tails of the distributions larger than 400 sam-ples [\(Vehtari et al.,](#page-16-24) [2021\)](#page-16-24), and that the potential scale reduc-

tion statistics (R) did not deviate from 1.0. To ensure stable 505 450 convergence of our models, we visually inspected the trace 506 451 plots of the posterior parameters and performed prior and 507 452 posterior predictive checks. We verified that the maximum 508 453 tree depth was not saturated. Strong degeneracies inherent ₅₀₉ 454 to multilevel models were addressed by reparametrizing the $_{510}$ 455 [m](#page-15-28)odels to a non-centered parameterization [\(Betancourt and](#page-15-28) $_{511}$) ⁴⁵⁶ [Girolami,](#page-15-28) [2013\)](#page-15-28).

 457 The resulting mean estimate of $σ_M$ for each participant 513 ⁴⁵⁸ was used as a measure of memory inexactness, since it re-459 flected the variability around the true target location thought 515 ⁴⁶⁰ to result from the loss of precision in a fine-grained repre-461 sentation. To estimate the degree of reliance on a categorical 517 462 representation, we used the measure of prototype bias defined 518 463 as $1-\lambda$, which reflected the relative contribution of the pro- 519 464 totypical location to behavioral responses, while mean λ was 520 ⁴⁶⁵ computed from posterior probabilities for individual partici-⁴⁶⁶ pants.

⁴⁶⁷ **fMRI acquisition, preprocessing and analysis**

⁴⁶⁸ fMRI data were collected with Philips Achieva 3TX (Studies 469 I-III; Table S3), Siemens Tim Trio (Study IV; Table S3), and ⁵²⁶ ⁴⁷⁰ Prisma (Studies IV-VI; Table S3) scanners. We acquired T1- 471 and T2-weighted structural images and several BOLD images 472 using T2*-weighted echo-planar imaging sequences. We also 529 ⁴⁷³ collected pairs of spin-echo images with opposite phase en-474 coding to estimate field maps for the purpose of distortion⁵³¹ ⁴⁷⁵ correction during data preprocessing. Acquisition parame-476 ters for specific images varied between different studies, as ⁵³³ ⁴⁷⁷ described in Table S3.

 The preprocessing and analysis of the MRI data was performed with the Quantitative Neuroimaging Environment⁵³⁶ 480 and Toolbox (QuNex; [Ji et al.,](#page-15-29) [2023\)](#page-15-29). Several steps of analy-481 sis and visualizations were prepared using R [\(Team,](#page-16-23) [2022a\)](#page-16-23), Matlab (R2021a, Natick, Massachusetts, USA), and Connec- tome Workbench (Human Connectome Project, Washington University, St. Louis, Missouri, USA).

 MR images were preprocessed using Human Connec[t](#page-15-30)ome Project (HCP) minimal preprocessing pipeline [\(Glasser](#page-15-30)⁵⁴² [et al.,](#page-15-30) [2013\)](#page-15-30). Specifically, structural images were corrected 543 for magnetic field distortions and registered to the MNI at- las, brain tissue was segmented into white and gray mat- ter, and the cortical surface was reconstructed. Functional BOLD images were sliced-time aligned, corrected for spa- tial distortions, motion-corrected, registered to the MNI atlas, 493 and the BOLD signal was mapped to the joint surface vol-549 494 ume representation (CIFTI) and spatially smoothed ($\sigma = 4$ 550 495 mm). Further analyses were performed on "dense" whole-551 brain data (i.e., each grayordinate independently). To ob-497 serve general patterns across functional brain systems and to 553 increase statistical power, we also performed analyses on par-499 cellated whole-brain data. Parcellated data were obtained by 555 extracting the mean signal of 360 cortical brain regions iden-501 tified based on the HCP-MMP1.0 parcellation [\(Glasser et al.,](#page-15-31) 557 [2016\)](#page-15-31) and, additionally, for 358 subcortical regions and 12 558 brain networks based on the Cole-Anticevic Network Parti-504 tion [\(Ji et al.,](#page-15-21) [2019\)](#page-15-21). Although the exploratory analyses were 560 performed for all brain areas and networks, we were primarily interested in the cingulo-opercular, dorsal-attention, frontoparietal, and default networks as defined in the Cole-Anticevic Network Partition [\(Ji et al.,](#page-15-21) [2019\)](#page-15-21).

We performed the activation analysis using a general linear modeling (GLM) approach in which event regressors were convolved with the assumed double-gamma haemody-512 namic response function (HRF; [Friston et al.,](#page-15-32) [1998\)](#page-15-32). For each participant, we modeled each phase of a task trial separately. Specifically, we estimated the β coefficients for the encoding, delay, and response phases (Figure S1). For three studies (Studies IV-VI; Table S2), we also separately modeled the attention cue in the middle of the delay period when present (Figure S1). Trials with outlier responses based on the behavioral data analysis were modeled as separate events using unassumed modeling and excluded from the group-level statistical analyses of the fMRI data. We additionally mod-⁵²² eled motion parameters, their first derivatives, and squared ⁵²³ motion parameters to account for any signal artifacts due to ⁵²⁴ movement. To identify outlier participants based on brain activity, we computed Pearson correlation coefficients between the β maps for encoding, delay and response activity of each participant with a corresponding group average β map. We identified one participant who deviated more than $3 \times SD$ from the group average β map and excluded this participant from further analysis.

To identify significant activation and deactivation during the task, we next analyzed the β estimates at the group level using permutation analysis (500 permutations, tail acceler-ation) in PALM [\(Winkler et al.,](#page-16-25) [2014\)](#page-16-25). To test the significance of the β estimates based on the "dense" grayordinate data, we conducted two-tailed one-sample t-tests with TFCE $(H = 2, E = 0.5, C = 26)$ FWE correction. To test the significance of the β estimates based on the parcellated data, ⁵³⁹ we conducted two-tailed one-sample t-tests with FDR correction. The resulting corrected *p*-value maps were thresholded at the whole-brain corrected significance level of $\alpha < 0.05$.

⁵⁴² **The estimation of brain-behavior relationship**

To estimate the relationship between brain activity in specific networks and behavioral measures, we performed Bayesian two-level linear modeling with factors memory inexactness and prototype bias. The models were numerically estimated using the probabilistic programming language Stan [\(Team,](#page-16-22) ⁵⁴⁸ [2022b\)](#page-16-22) in R [\(Team,](#page-16-23) [2022a\)](#page-16-23). To obtain standardized *β* coefficients and provide easier comparison of results across both behavioral factors, brain activity estimates, memory inexactness, and prototype bias were standardized to $\mu = 0$, $\sigma = 1$, across all participants. We used study as the grouping variable at the first level to model varying intercepts across studies and Student's t-distribution to describe the data. Weakly informative prior distributions were used for all model parameters, ensuring that the standard deviation of the prior dis-⁵⁵⁷ tribution was at least 10 times larger than that of the posterior distribution. Specifically, we used normal prior distributions $(\mu = 0, \sigma = 10)$ for regression parameters and half-Cauchy prior distributions ($\mu = 0$, $\lambda = 2.5$) for standard deviations,

561 as recommended by [Gelman](#page-15-33) [\(2006\)](#page-15-33). The prior distribution 616 562 for the degree of freedom parameter was set to $\Gamma(2,0.1)$ as 617 563 recommended by [Juárez and Steel](#page-15-27) [\(2010\)](#page-15-27). The stability of 618 the HMC sampling algorithm was analyzed by verifying that 619 ⁵⁶⁵ all estimated parameters had estimated effective sample sizes

 $_{566}$ in the bulk of the distributions and in the tails of the distri- $_{520}$ $_{567}$ butions larger than 400 samples [\(Vehtari et al.,](#page-16-24) [2021\)](#page-16-24), and $_{621}$ that the potential scale reduction statistics (R) did not deviate ₅₆₉ from 1.0. To ensure a stable convergence of our models, we ⁶²² 570 visually inspected the trace plots of the posterior parameters⁶²³ 571 and performed prior and posterior predictive checks. We ver- 624 ⁵⁷² ified that the maximum tree depth was not saturated. Strong ⁵⁷³ degeneracies inherent to multilevel models were addressed 574 by reparametrizing the models to a non-centered parameteri-⁶²⁷ 575 zation [\(Betancourt and Girolami,](#page-15-28) [2013\)](#page-15-28).

 To examine the effect of sample size on the detection of brain-behavior relationships, we conducted Bayesian linear 578 modeling for sample sizes ranging from 15 to 155 partici-⁶³¹ pants. At each sample size, 1000 samples were created by sampling with replacement from the set of all participants. We then performed Bayesian two-level normal linear model 534 582 with factors memory inexactness and prototype bias with ⁶³⁵ study as a random effect for each separate sample. The mod- els were computed in the same manner as described in the 637 previous paragraph.

⁵⁸⁶ **Results**

⁵⁸⁷ **Individual differences in the use of spatial coding** ⁵⁸⁸ **strategies**

 We first examined the pattern of response errors at different target angles in order to identify any behavioral indicators of the use of categorical representations during spatial working memory performance. We observed that participants system- atically shifted their responses toward the nearest diagonals, with a greater bias occurring at target angles further away from the diagonals (Figures [2A](#page-7-0)-B and S2A). This finding indicates the use of categorical representations, where par- ticipants formed spatial categories defined by the four quad- rants of the screen, delineated by the vertical and horizon[t](#page-15-3)al axes, each best represented by its diagonal [\(Huttenlocher](#page-15-3) 655 [et al.,](#page-15-3) [2004,](#page-15-3) [1991;](#page-15-4) [Starc et al.,](#page-16-4) [2017\)](#page-16-4).

⁶⁰¹ To separately measure the contribution of fine-grained ⁶⁵⁷ and categorical representations in spatial working memory, we next derived two behavioral measures based on the mod-604 eling of the responses – memory inexactness and prototype 660 bias. We used memory inexactness as a measure of the preci- sion of fine-grained representations, and prototype bias as a measure of the extent to which participants relied on categorical representations. For each behavioral measure, we calcu-664 lated a mean estimate for each participant, reflecting their use of fine-grained and categorical representations (for distribu- tions across participants see Figure [2C](#page-7-0) and for study differ- 667 ences in both measures see Figures S2B-C). We then com- puted the Pearson correlation coefficient to examine the re-614 lationship between both measures across studies. Our results 670 F_{615} revealed a positive correlation, $r = 0.660$, $p < 0.001$, between F_{671} memory inexactness and prototype bias (Figure [2D](#page-7-0)), suggesting that participants who relied more heavily on categorical representations showed poorer precision of fine-grained representations and vice versa.

⁶²⁰ **Task-related brain activity across different levels of** ⁶²¹ **parcellation**

In the analysis of the fMRI data, we first examined the areas of the brain that were activated or deactivated during different phases of a task trial, namely the encoding, delay, ⁶²⁵ and response phases (Figure S3A). During all phases of the trial, significant activation (i.e., $p < 0.05$ corrected for multiple comparisons) was observed in a number of brain regions, spanning the frontal, parietal, and occipital cortices. Subcor-⁶²⁹ tical activation was consistently observed in the cerebellum, thalamus, putamen, caudate, and brainstem. Phase-specific activations differed mainly in the early and ventral stream ⁶³² visual areas, where extensive activation was observed only ⁶³³ during the encoding and response phases. Significant deactivation was observed in all phases of the trial in the posterior cingulate cortex, and in areas of the medial prefrontal cor-⁶³⁶ tex, and inferior frontal cortex. Additional deactivation was observed in the lateral temporal cortex for the delay and response, and in the inferior parietal cortex, early and ventral ⁶³⁹ stream visual areas for the delay phase only. Subcortical de-⁶⁴⁰ activation was mainly observed during the delay and response ⁶⁴¹ phases in the cerebellum, hippocampus, and amygdala.

⁶⁴² To investigate the integration of activity within functional brain regions and networks, and their average responses to ⁶⁴⁴ the task, we also performed the activation analysis of the fMRI data averaged within cortical regions of the HCP-MMP1.0 parcellation [\(Glasser et al.,](#page-15-31) [2016\)](#page-15-31), and within subcortical regions and networks based on the Cole-Anticevic Network Partition [\(Ji et al.,](#page-15-21) [2019\)](#page-15-21). The results based on parcellated data showed additional significant task-related acti-vations and deactivations (Figures [3A](#page-9-0), S3B, and S3C). When looking at more general networks, increased activity was observed during encoding in the primary and secondary visual networks, somatomotor, cingulo-opercular, dorsal attention, frontoparietal, and language networks, in addition to the posterior and ventral multimodal networks. Deactivation was ob-656 served only in the default network. The delay phase showed significant activation in the secondary visual, somatomotor, cingulo-opercular, dorsal-attention, and posterior multimodal networks. In contrast, decreased activity was observed in the default, ventral multimodal, and orbito-affective networks during the delay. Finally, the response phase was characterized by activation in the primary and secondary visual networks, somatomotor, cingulo-opercular, dorsal attention, frontoparietal, auditory, posterior multimodal, and ventral multimodal networks. Significant deactivation was again observed only in the default network.

Lastly, we examined whether the analysis on parcellated fMRI data improved effect sizes or, alternatively, diluted effects due to inhomogeneous activity within individual brain regions and networks. Similar to the analysis described in [Glasser et al.](#page-15-31) (2016) and [Ji et al.](#page-15-21) (2019) , we compared the

C. Distributions of behavioral measures across individuals D. Correlation between measures

Fig. 2. Systematic biases and individual differences in spatial working memory performance. A. An example pattern of the systematic bias in spatial working memory performance computed as average angular response errors at different target angles across participants in Studies IV, V and VI, which used the same stimulus angles in relation to the center of the screen. The start of the arrow denotes the target position, while the head of the arrow points to the average response position. **B.** Angular response errors at different target angles across all participants. Red lines represent diagonals of each quadrant, delineated by the horizontal and vertical axes shown as gray lines. C. The distribution of memory inexactness and prototype bias across participants. The points present the mean of each measure, with the range indicating the standard deviation of the measure. **D.** Relationship between memory inexactness and prototype bias across all participants estimated using Pearson correlation coefficient.

672 unthresholded Z-values for delay-related activity between the 691 ⁶⁷³ "dense" grayordinate data and brain regions, and additionally ⁶⁹² 674 between the brain regions and network data (Figure S3D). 693 675 Our results showed that although the Z-values of individual 694 676 grayordinates exceeded the Z-values obtained for the brain 695 677 regions and networks to which they belonged, the analysis of 696 678 the parcellated data resulted in higher overall effect size es-697 ₆₇₉ timates than the analysis of the grayordinate data. Similarly, 698 ⁶⁸⁰ analysis of network average data resulted in higher effect size ⁶⁸¹ estimates than analysis of the brain regions. Although work-682 ing with grayordinate data provides better spatial precision of 701 683 results and is preferable when precise localization is of inter-702 ⁶⁸⁴ est, these results suggest that working with parcellated data ⁶⁸⁵ is preferable when testing hypotheses related to functional ⁶⁸⁶ regions or networks, as was the case in our study.

⁶⁸⁷ **Individual differences in spatial coding strategies re-**⁶⁸⁸ **flected in brain activity**

689 Next, we were interested in whether individual differences in 709 690 the use of fine-grained and categorical representations are re- 710

flected in brain activity. To this end, we used Bayesian linear modeling to predict the activity of brain networks of interest based on measures of memory inexactness and prototype bias (Figure [3B](#page-9-0)). Specifically, we used hierarchical linear modeling with behavioral measures as fixed factors and study as a random effect. We focused on the average activity within networks (i) to identify the engagement of broad brain systems during the use of different spatial coding strategies and (ii) to increase the effect sizes and statistical power of the analysis. Specifically, we examined brain-behavior relationships for the cingulo-opercular, dorsal-attention, frontoparietal, and default networks, separately for different task phases $(Figure 3C)$ $(Figure 3C)$ $(Figure 3C)$.

During the encoding phase of the task, our results re-⁷⁰⁵ vealed 98*.*2% and 100% posterior probabilities for a nega-⁷⁰⁶ tive relationship between memory inexactness and activity ⁷⁰⁷ in the cingulo-opercular and dorsal-attention networks, re-⁷⁰⁸ spectively. These results suggest that increased encodingrelated activity in the cingulo-opercular and dorsal-attention networks was related to decreased memory inexactness, or

 in other words, increased memory precision. We also ob- 768 served a 96*.*2% posterior probability of a positive relation- ship between memory inexactness and activity in the default $_{770}$ network, indicating that decreased activity in the default net- $_{771}$ work was associated with increased memory precision. Re- $_{772}$ lating encoding-related activity with prototype bias revealed $_{773}$ a negative relationship between prototype bias and the de- $_{774}$ fault network activity with a posterior probability of 98.5%, 775 showing that decreased activity in the default network was 776 associated with increased prototype bias.

 For the delay phase, the results indicated 98*.*8%, 99*.*9%, 722 and 95.1% posterior probabilities for a negative relationship 779 between memory inexactness and activity in the cingulo- opercular, dorsal-attention, and frontoparietal networks, re-725 spectively. This result again suggests that increased memory 782 precision was related to increased activity in these networks 727 during spatial working memory performance. On the other 784 hand, the results showed 96*.*4% and 90*.*9% posterior proba- bilities of a positive relationship between prototype bias and activity in the frontoparietal and default networks, respec- tively. These relationships suggest that both increased fron- 788 toparietal activation and weaker deactivation of the default network are associated with increased prototype bias during the spatial working memory task.

 Relating response-related activity with memory inexact- ness revealed a negative relationship between memory inex- actness and activity in the cingulo-opercular and dorsal at- 794 tention networks with posterior probabilities of 98*.*5% and 98*.*8%, respectively. The results also showed a 92*.*6% poste- rior probability of a positive relationship between prototype 797 bias and activity in the cingulo-opercular network. These re- 798 ⁷⁴² sults suggest that increased response-related activity in these ⁷⁹⁹ networks was related with increased memory precision, as well as increased prototype bias. We also observed a negative 745 relationship between prototype bias and the default network 802 activity with a posterior probability of 92*.*7%, suggesting that 747 decreased activity in this network was associated with in-804 creased prototype bias.

749 The general whole-brain patterns of the relationship be-₈₀₅ tween brain activity and behavioral measures of memory in- exactness and prototype bias for brain regions and networks ⁷⁵² can be observed in Figures S4-5. These analyses revealed ⁸⁰⁷ several additional relationships with both behavioral mea- sures and activity in other brain networks. For memory in-⁷⁵⁵ exactness, a negative relationship with activity in the primary ⁸¹⁰ visual, secondary visual, and posterior-multimodal networks during the encoding was found with posterior probabilities of 812 98*.*9%, 100%, and 99*.*0%, respectively (Figure S5). We also observed a negative relationship between memory inexact-760 ness and response-related activity in the primary visual, sec-⁸¹⁵ 761 ondary visual, somatomotor, and posterior-multimodal net-⁸¹⁶ works with posterior probabilities of 96*.*1%, 99*.*2%, 97*.*5%, and 98*.*7%, respectively (Figure S5). For the prototype bias, a positive relationship with activity in the language and orbito- 818 affective networks during the delay was observed with poste- rior probabilities of 97*.*7%, and 98*.*8%, respectively (Figure ⁷⁶⁷ S5).

The effect of sample size on the detection of brain**behavior relationships**

A comparatively large multi-study sample provided us with an increased power to detect brain-behavior relationships with relatively small effect sizes. To further validate the stability of the results and assess statistical power in evaluating brain-behavior relationships, we conducted a comprehensive resampling analysis. Specifically, for each sample size from 15 to 155, we randomly selected a set of participants from our 777 original sample with replacement 1000 times and repeated the Bayesian hierarchical linear regression for the four networks of interest, i.e. the cingulo-opercular, dorsal-attention, frontoparietal, and default networks, for the delay period for each sample. This allowed us to evaluate the effects of sample size on model estimates, their confidence intervals, and statistical power.

While mean β coefficients estimated in the linear model were generally stable across different sample sizes (Figure 56), our results indicated that the variability of β estimates within each sample size changed significantly with sample size. Zero was robustly excluded from the 95% confidence interval computed across 1000 resamplings for the relationships between memory inexactness and activity in the ⁷⁹¹ cingulo-opercular, dorsal-attention, and frontoparietal networks only after sample sizes of 93, 73, and 151, respectively. Zero was also consistently excluded from the 95% confidence interval across 1000 resamplings for the relationship between prototype bias and the frontoparietal network activity after a sample size of 149. Statistical power, computed as the proportion of samples in which 95% of posterior distribution was above or below 0, linearly increased with in-⁷⁹⁹ creasing sample size and reached 61*.*3%, 87*.*8%, and 26*.*0% for the relationships between memory inexactness and activity in the cingulo-opercular, dorsal-attention, and frontoparietal networks, respectively, and 31.2% for the relationship between prototype bias and the frontoparietal network activity (Figure S6).

⁸⁰⁵ **Discussion**

A spatial location can be encoded and maintained in working memory using different representations and strategies. Finegrained representations provide detailed stimulus informa-809 tion, but are cognitively demanding and prone to inexactness. On the other hand, categorical representations may provide a more robust and less demanding strategy, but at the cost of loss of fine-grained precision. In our study, we were interested in the extent to which individuals rely on fine-grained and categorical representations to encode and maintain spatial information in working memory, and how these individual differences in spatial working memory strategies are reflected in brain activity.

⁸¹⁸ **Individual differences in spatial coding strategies**

The investigation of behavioral performance in the spatial working memory task revealed the presence of a systematic 821 bias in behavioral responses. Specifically, we observed that

B. Estimation of brain-behavior relationships

C. Brain-behavior relationship for different brain networks and task phases

Fig. 3. Average network activity in relation to individual spatial working memory performance. A. The average activity in the cingulo-opercular, frontoparietal, dorsalattention, and default networks during different task phases. **B.** Steps in the analysis of the relationship between brain activity in specific networks and behavioral measures of memory inexactness and prototype bias. For each participant, we computed average brain activity within networks of interest defined by Cole-Anticevic Network Partition [\(Ji et al.,](#page-15-21) [2019\)](#page-15-21), and individual measures of memory inexactness and prototype bias. Next, we ran a Bayesian hierarchical linear model across participants predicting brain network activity with memory inexactness and prototype bias, and controlling for study as a random effect. **C.** Posterior distributions of the relationship between the activity of specific networks and behavioral measures of memory inexactness (red) and prototype bias (blue). Points indicate mean *β*-estimates, and lines 95% confidence intervals.

822 participants tended to shift their responses closer to the near-879 823 est diagonals of the four quadrants, formed by dividing the 880 824 screen at the vertical and horizontal axes of symmetry. Sev-881 eral previous studies [\(Haun et al.,](#page-15-26) [2005;](#page-15-26) [Huttenlocher et al.,](#page-15-3) ssa 826 [2004,](#page-15-3) [1991;](#page-15-4) [Purg et al.,](#page-16-2) [2022;](#page-16-2) [Starc et al.,](#page-16-4) [2017\)](#page-16-4) have sug-883 827 gested that such a bias reflects the use of categorical repre-884 828 sentations, where participants spontaneously impose spatial 885 829 categories in coding stimulus position. [Huttenlocher et al.](#page-15-3) 886 830 (2004) have shown that this bias is replicated even when dif-887 831 ferent spatial categories are imposed by the task by clustering 888 832 stimuli around the horizontal and vertical axes, as well as en-889 833 couraging participants to use categories centered on the car-890 834 dinal axes and bounded by the diagonals. This suggests that 891 835 the horizontal and vertical axes represent the most robust cat-892 836 egory boundaries, resulting in the lowest misclassification of 893 837 spatial information [\(Huttenlocher et al.,](#page-15-3) [2004\)](#page-15-3). Nevertheless, 894 838 the use of different reference points [\(Holyoak and Mah,](#page-15-34) [1982;](#page-15-34) 895 839 [Sadalla et al.,](#page-16-26) [1980\)](#page-16-26) or spatial borders [\(Nelson and Chaiklin,](#page-16-27) 896 840 [1980;](#page-16-27) [Newcombe and Liben,](#page-16-28) [1982\)](#page-16-28), and specific instructions 897 841 on the context of the space [\(Tversky and Schiano,](#page-16-29) [1989\)](#page-16-29) have 898 842 been shown to affect the type of categories constructed in spa-899 843 tial estimation tasks, suggesting that the categories formed 900 844 [a](#page-15-4)re, at least to some extent, context-dependent [\(Huttenlocher](#page-15-4) 901 ⁸⁴⁵ [et al.,](#page-15-4) [1991\)](#page-15-4).

846 Our results are in line with the category adjustment model 903 847 [Huttenlocher et al.](#page-15-4) [\(1991,](#page-15-4) [2000\)](#page-15-5), which proposes that a spa-⁹⁰⁴ tial location in working memory is simultaneously repre-⁹⁰⁵ 849 sented as a fine-grained and categorical representation. The 906 850 model predicts that the uncertainty in remembered fine- 907 851 grained information is compensated for by using information 908 852 of a broader stimulus category, which introduces a system-⁹⁰⁹ 853 atic bias in responses towards a prototypical value, but in-910 854 creases an overall response accuracy by decreasing response 911 855 variability. We used the assumptions of the category adjust-912 856 ment model to mathematically describe behavioral responses 913 857 during spatial working memory performance and to identify 914 858 individual contributions of fine-grained and categorical rep-915 resentations to response errors. Specifically, we estimated the 916 860 inexactness in fine-grained memory as a spread of responses 917 861 around the true target value, while the effect of a categorical 918 862 representation on the estimation of stimulus location was de-919 863 scribed in terms of a degree of systematic bias towards the 920 864 prototype. We were particularly interested in the relation-921 865 ship between the use of both representations across individ- 922 866 uals. Our results replicated previous observation of a posi- 923 867 tive correlation between a loss of fine-grained memory preci-924 868 sion and the use of a categorical representation [\(Starc et al.,](#page-16-4) 925) 869 [2017\)](#page-16-4). Specifically, our results suggest that there are individ- 926 870 ual differences in the balance between the use of fine-grained $\frac{1}{2}$ 871 and categorical spatial coding – individuals with higher fine-872 grained precision of spatial representations relied less on cat- 873 egorical information, whereas individuals who showed lower 874 precision in fine-grained representations seemed to rely more $\frac{1}{931}$ 875 strongly on categorical representations.

876 At the interindividual level, the degree of reliance on cate-933 877 gorical versus fine-grained representations has been related to 934 878 individual working memory capacity [\(Crawford et al.,](#page-15-2) [2016;](#page-15-2) 935

[Stukken et al.,](#page-16-30) [2016\)](#page-16-30). Studies on working memory capacity have traditionally focused on estimating the number of items a participant can maintain over short periods of time by comparing task performance under different working mem-ory loads (for a review see [Luck and Vogel,](#page-15-35) [2013\)](#page-15-35). However, recent studies [\(Bays and Husain,](#page-15-36) [2008;](#page-15-36) [Spencer and Hund,](#page-16-31) [2002;](#page-16-31) [Zhang and Luck,](#page-16-32) [2008\)](#page-16-32) suggest that increasing the detail or precision of these objects requires additional working memory resources at the cost of reducing the number of objects that can be remembered simultaneously. Therefore, the formation of high-precision representations might be easier for individuals with a high working memory capacity, whereas a low working memory capacity would require a reduction in stimulus complexity, such as by using coarse categorical coding. [Crawford et al.](#page-15-2) [\(2016\)](#page-15-2) estimated the relationship between spatial working memory capacity and the use of fine-grained or categorical representations during spatial working memory performance based on a sample of 778 adults. Their results showed a correlation between spatial working memory capacity and different spatial coding strategies, with higher capacity predicting higher spatial precision and lower categorical bias. Moreover, consistent with these results is also the observation that introducing dis-⁹⁰² tractor stimuli that need to be retained during spatial working memory performance or an interference task which put additional strain on working memory resources results in an increased use of categorical representations [Crawford et al.](#page-15-2) (2016) ; [Huttenlocher et al.](#page-15-4) (1991) . To sum, the use of different spatial coding strategies might be related to the availability of cognitive resources, which could explain interindividual differences in the preference for a specific strategy.

It is important to note that due to the complex hierarchical structure of our model of the effect of fine-grained and categorical representations on behavioral responses, the assumptions of the model were to some extent simplified, which could potentially affect our estimates. For example, in our model we assumed the same prototype location for all participants, which we centered on the diagonal of each quadrant. Some studies have shown that the prototype might not be located exactly on the diagonal, and might even differ between different quadrants of the task display or between participants [\(Huttenlocher et al.,](#page-15-3) [2004,](#page-15-3) [1991\)](#page-15-4). This variability in the prototype location was captured to an extent by the measure of prototype inexactness in our model, although larger incosistencies in the assumed and actual prototype location could increase the estimation of prototype inexactness and, in turn, underestimate the degree of reliance on categorical representations in spatial working memory performance.

Furthermore, our model did not account for the potential influence of inexact boundaries in the estimation of stimulus locations near boundaries. In the case of our study, participants gave their responses on a blank screen, which meant that no spatial boundary was explicitly presented, but 932 participants spontaneously imposed boundaries in the form of horizontal and vertical symmetry axes. Their estimation of boundaries could therefore be uncertain or inexact, which could lead to misclassification of stimuli near bound-

936 aries. For instance, responses within the boundary inexact-992 937 ness could fall into any of the two categories delineated by 993 938 the boundary and adjusted towards its center. When aver-994 939 aged, these responses with opposing directions of prototype 995 940 bias would cancel each other out, resulting in an overall de-996 941 creased effect of prototype bias around the inexact boundary 997 942 [\(Huttenlocher et al.,](#page-15-3) [2004,](#page-15-3) [1991\)](#page-15-4). Our results showed that av- 998 943 erage response errors increased with target angle further from 999 944 the diagonal, with a slight decrease near the boundaries, es-1000 945 pecially in the study in which the stimuli were presented the 946 closest to the cardinal axes. For this reason, we excluded all1002 947 misclassified stimuli from our data analysis to prevent dilu-1003 948 tion of the effects of categorical representations. We identi-1004 949 fied only 1.86% of misclassified stimuli per participant, with 1005 950 all misclassification occuring up to the target angle 15° from1006 951 any boundary. The dynamic field theory [\(Schutte et al.,](#page-16-8) [2003;](#page-16-8)1007) 952 [Simmering et al.,](#page-16-9) [2006\)](#page-16-9) assumes that boundaries, perceived 953 or spontaneously imposed, have a deflecting effect on behav-1009 $_{954}$ ioral responses during the estimation of a stimulus location in₁₀₁₀ ⁹⁵⁵ working memory due to their lateral inhibitory effects at the 956 neural level, which results in a drift of the activation produced $\frac{958}{957}$ by the remembered target stimulus away from the boundary. 958 Despite the overall decreased response errors near boundaries $\frac{959}{959}$ in our study, our results might still be in line with the assump- $\frac{1}{1015}$ $_{960}$ tions of the dynamic field theory when looking at individual $\frac{961}{961}$ responses – i.e. the boundary might still have a deflecting $\frac{962}{962}$ effect, but in different directions for the correctly and incor- $\frac{100}{1018}$ ⁹⁶³ rectly classified stimuli.

964 Different coding strategies related to the engagement ⁹⁶⁵ **of separable brain systems**

966 The assumed advantage of categorical spatial coding is that¹⁰²³ 967 it is less demanding on cognitive resources without compro-1024 968 mising the overall accuracy of responses. In contrast, encod-1025 ⁹⁶⁹ ing fine-grained information yields precise responses, but re-¹⁰²⁶ 970 quires greater engagement of attention and cognitive control.¹⁰²⁷ 971 Therefore, we hypothesized that the use of specific spatial¹⁰²⁸ 972 representations would be related to the level of engagement¹⁰²⁹ 973 of the attentional and control brain systems. Specifically, we¹⁰³⁰ 974 expected that a stronger reliance on precise, fine-grained rep-1031 975 resentations would be supported by increased activation of at-1032 976 tentional and control brain systems, and stronger inhibition of¹⁰³³ 977 the default network. On the other hand, we assumed that un-1034 978 certainty in fine-grained representations, such as due to a loss¹⁰³⁵ 979 of precision or task interference, would be related to an in-1036 980 creased use of categorical representations that would require¹⁰³⁷ 981 fewer attentional and control resources.

982 In the investigation of the relationship between brain¹⁰³⁹ 983 activity with behavioral measures of the precision of fine-1040 984 grained representations and the use of categorical represen-¹⁰⁴¹ 985 tations, we observed a strong positive relationship between 986 fine-grained memory precision and activity in the cingulo-1043 987 opercular and dorsal-attention networks during all phases of 1044 the task, the encoding, delay, and response. We also ob-1045 989 served a slightly weaker positive relationship between mem-1046 990 ory precision and the frontoparietal network activity during 1047 991 the delay. These results suggest that increased memory pre-1048 cision is indeed accompanied by an increased enagagement of these networks. The cingulo-opercular, dorsal-attention, and frontoparietal networks are consistently activated during different working memory tasks and have been widely recognized to play an important role in active maintenance of [i](#page-15-37)nformation in working memory [\(Brown et al.,](#page-15-11) [2004;](#page-15-11) [Cur](#page-15-37)[tis,](#page-15-37) [2006,](#page-15-37) [2004;](#page-15-0) [D'Esposito and Postle,](#page-15-38) [2015;](#page-15-38) [Eriksson et al.,](#page-15-39) [2015;](#page-15-39) [Liu et al.,](#page-15-40) [2017;](#page-15-40) [Purg et al.,](#page-16-2) [2022;](#page-16-2) [Zarahn et al.,](#page-16-12) [1999\)](#page-16-12). In addition, increases in the level of activity and functional connectivity within these networks have been found to scale with increased attentional demands, working memory load, and memory accuracy (e.g., [Assem et al.,](#page-15-41) [2020;](#page-15-41) [Barch et al.,](#page-15-19) [2013;](#page-15-19) [Bray et al.,](#page-15-42) [2015;](#page-15-42) [Cole et al.,](#page-15-20) [2014;](#page-15-20) [Fox et al.,](#page-15-43) [2005;](#page-15-43) [Liu et al.,](#page-15-40) [2017,](#page-15-40) [2018;](#page-15-44) [Magnuson et al.,](#page-15-45) [2015;](#page-15-45) [Smith et al.,](#page-16-17) [2009\)](#page-16-17). Therefore, our findings support the notion that the formation and active maintenance of fine-grained representations presents a cognitive load and engages attentional and cognitive control systems.

Our results also revealed a negative relationship between fine-grained memory precision and the default network activity during encoding only, showing that decreased activity in this network was related to increased memory precision. Traditionally, fMRI studies investigating functional connectivity at rest have identified the role of the default network in spontaneous intrinsic activity in the absence of cognitive [l](#page-15-43)oad (e.g., [Cole et al.,](#page-15-20) [2014;](#page-15-20) [Damoiseaux et al.,](#page-15-46) [2006;](#page-15-46) [Fox](#page-15-43) [et al.,](#page-15-43) [2005;](#page-15-43) [Greicius et al.,](#page-15-47) [2003;](#page-15-47) [Moussa et al.,](#page-16-33) [2012;](#page-16-33) [Smith](#page-16-17) 1019 [et al.,](#page-16-17) [2009\)](#page-16-17). Moreover, the default network shows robust de-¹⁰²⁰ activation during the performance of various cognitive tasks, including during working memory performance, which be-¹⁰²² [c](#page-15-18)omes stronger with increasing cognitive load (e.g., [Antice](#page-15-18)[vic et al.,](#page-15-18) [2010;](#page-15-18) [Cole et al.,](#page-15-20) [2014;](#page-15-20) [Fox et al.,](#page-15-43) [2005;](#page-15-43) [Liu et al.,](#page-15-44) [2018;](#page-15-44) [Raichle,](#page-16-14) [2015a](#page-16-14)[,b;](#page-16-34) [Smith et al.,](#page-16-17) [2009\)](#page-16-17). Such decreases in the activity of the default network are thought to reflect the allocation of cognitive resources to task-relevant information and protection from distraction [\(Liu et al.,](#page-15-40) [2017\)](#page-15-40). For exam-ple, [Anticevic et al.](#page-15-18) [\(2010\)](#page-15-18) showed that stronger suppression of the default network during the encoding of target stimuli, prior to the presentation of distractors, predicted higher response accuracy in a working memory task. These results are consistent with our observations, although the study included non-spatial visual stimuli and match-to-sample responses that do not allow the uncoupling of separate contributions of fine-grained and categorical representations to response accuracy, which makes it difficult to directly relate the two studies. In summary, our results suggest that stronger inhibition of the default network is required to ensure good fine-grained memory precision, likely as a result of allocating attentional and control resources toward task-relevant stimuli and protection from interference.

Conversely, the relationship between the use of categorical representations and brain activity was somewhat less clear. Our results revealed opposing relationships with the default network activity and the prototype bias during the encoding and response phases of the task compared to the delay period. Specifically, we observed that increased use of prototype bias was related to stronger deactivation during

 the encoding and response, and weaker deactivation during the delay phase of the task. These results suggest temporal differences in the engagement of the default network in re- lation to the use of categorical representations. Stronger de- activation during the encoding and response might reflect in- creased attentional engagement and inhibition of distractors directed toward the formation and recall of categorical repre- sentations, respectively. On the other hand, weaker deactiva-1057 tion during the delay might suggest decreased attentional and 1114 control demands when individuals rely on categorical repre- sentations, supporting the hypothesis that categorical coding of spatial positions provides a less demanding spatial work-ing memory strategy.

1062 The investigation of the relationship between the use of¹¹⁹ 1063 categorical representations and the activity in attentional and¹¹²⁰ ¹⁰⁶⁴ control brain networks revealed a positive relationship be-¹⁰⁶⁵ tween the prototype bias and activity in the frontoparietal 1066 network during the delay, as well as a slightly weaker pos-¹¹²³ 1067 itive relationship between the prototype bias and the activ-¹¹²⁴ $_{1068}$ ity in the cingulo-opercular network during the response. In 1069 other words, increased engagement in these networks pre- 1070 dicted a higher use of categorical representations. While the 127 ¹⁰⁷¹ formation, maintenance and recall of fine-grained representa-1072 tions required constant engagement of attentional and control 1073 systems, the results on the use of categorical representations $\frac{1}{1130}$ ¹⁰⁷⁴ suggest that these brain systems were engaged only later in 1075 a task trial during stimulus maintenance and recall. Simi- 132 1076 larly, [Starc et al.](#page-16-4) [\(2017\)](#page-16-4) reported a compensatory use of fine- 1133 1077 grained and categorical representations during an individual 1078 task trial, where the failure to encode fine-grained informa- 1079 tion with high precision at the time of encoding of spatial in- $\frac{1}{1136}$ 1080 formation could then be compensated for by the reconstruc- $\frac{1}{1137}$ 1081 tion of target location based on categorical information in the 1138 1082 late delay and response periods of the trial. These results are 139 1083 somewhat inconsistent with the hypothesis of reduced $\cos_{\frac{1}{1400}}$ 1084 nitive load and reliance on cognitive resources when using $_{1141}$ ¹⁰⁸⁵ categorical representations, but may indicate a need for at- 1086 tentional and cognitive control during the recall of categorical 1143 ¹⁰⁸⁷ representations just before the response has to be given.

1088 However, our assumption that the observed relationship₁₁₄₅ with the delay-related frontoparietal activity reflects the use 1090 of a categorical representation may be wrong. The category1147 1091 adjustment model [\(Huttenlocher et al.,](#page-15-4) [1991,](#page-15-4) [2000\)](#page-15-5) proposes₁₁₄₈ that participants resort to the use of a categorical represen- tation when their confidence in a memory representation is 1094 low, which would arguably be assessed just before or at the₁₁₅₁ time of the response. Taking this into account, we can hy- pothesize that the increased frontoparietal activity does not reflect the cognitive processes engaged in the maintenance of a categorical representation, but rather the processes that predict a loss of confidence in the fine-grained memory rep- resentation and subsequent increased reliance on the cate- gorical representation. Even though the frontoparietal net- work has been strongly implicated in allocation of attention and active maintenance of task-relevant information in work- ing memory, studies have also shown its role in protection 1105 from task-irrelevant information (e.g., [Jerde and Curtis,](#page-15-48) [2013;](#page-15-48)1162)

[Ptak,](#page-16-35) [2012;](#page-16-35) [Zhang et al.,](#page-16-36) [2017\)](#page-16-36). The increase in frontoparietal activity may reflect an increased effort in protecting the memory from task interference and suppression of distractors due to lower ability or confidence in the precision of the finegrained memory representation, leading to larger reliance on categorical representation when providing the response. This is consistent with the finding that introducing an interference during the delay of spatial estimation tasks increased the re-liance on categorical information [\(Huttenlocher et al.,](#page-15-3) [2004,](#page-15-3) [1991\)](#page-15-4). When assessing the role of the frontoparietal network, it is also prudent to consider the functional heterogeneity of the network. Specifically, our results of task-related activity ¹¹¹⁸ based on voxel-wise fMRI data showed activation in some, and deactivation in other areas within the frontoparietal network, suggesting that the role of the frontoparietal network in spatial working memory processes might be more complex than initially thought, and the averaging of the activity within the network might mask diverging functions within the network.

Together, the observed patterns of associations between brain and behavior reflect important relationships between the two strategies of encoding, maintenance and recall of spatial information. While the negative relationship between fine-grained memory precision and the use of categorical representations suggests a complementary use of categorical and fine-grained representations with the goal to increase the overall response accuracy, the two strategies relate to the engagement of separable brain systems. In particular, the precision of fine-grained representations is related to the level of attentional engagement, which is reflected in the activation of the attentional and control brain networks. Additionally, greater deactivation of the default network during the formation of fine-grained representations appears to predict higher memory precision, perhaps by providing suppression of distractors and the allocation of resources toward task-relevant information. In contrast, the extent of reliance on categorical representations does not seem to impose such attentional demands. Compared with the ongoing engagement of atten-¹¹⁴⁴ tional and control systems necessary to ensure high precision of fine-grained representations, some evidence was found for the activation of these systems in relation to the use of categorical representations later in the task trial during maintenance and response. Interestingly, the relationship between categorical representations and the activity in the default network appears to change over the course of the trial, where stronger inhibition of the default network is required during stimulus encoding and recall, whereas decreased inhibition is observed during the maintenance of spatial information. Since the use of a categorical representation is predicted by the uncertainty or loss of confidence in a fine-grained representation the increased deactivation of the default network during the stimulus presentation and response may reflect an increased effort in protecting the memory from task interference and suppression of distractors. On the other hand, the relaxation of the default network during the delay possibly reflects a decrease in cognitive demands in the maintenance of categorical representations.

¹¹⁶³ By exploring the relationship between the use of fine-¹¹⁶⁴ grained or categorical representations with the activity of ¹¹⁶⁵ other brain networks, we identified several relationships that ¹¹⁶⁶ suggest that the use of two strategies is related to different ¹¹⁶⁷ modalities. The precision of fine-grained memory was asso-¹¹⁶⁸ ciated with the activity in primary visual, secondary visual, 1169 and posterior multimodal networks during the encoding and 1226 1170 response, with increased activity in these networks predicting-1227 1171 higher memory precision. Additionally, higher memory pre-1228 ¹¹⁷² cision was also associated with increased somatomotor net-1173 work activity during the response. These results suggest that 1174 fine-grained representations might be encoded as a visual in- 1230 ¹¹⁷⁵ formation which is reactivated during the response, when it is 1176 converted into a motor plan used to execute the task response¹²³¹ 1177 [\(Purg et al.,](#page-16-2) [2022\)](#page-16-2). In contrast, we observed that the use of²³² 1178 categorical representations was predicted by the delay-related¹²³³ 1179 activity in the language network, with an increased categor-¹²³⁴ 1180 ical bias related to increased activity in this network. The¹²³⁵ 1181 engagement of the language network in the maintenance of²³⁶ 1182 categorical representations might indicate the transformation¹²³⁷ ¹¹⁸³ of spatial information into verbal codes during spatial work-1184 ing memory. For example, spatial categories defined as the¹²³⁹ 1185 four quadrants of the screen, delineated by the horizontal and¹²⁴⁰ 1186 vertical axes, could be remembered in terms of verbal codes¹²⁴¹ ¹¹⁸⁷ "up-right", "up-left", "down-left", and "down-right". Simi-1188 larly, studies that collected subjective reports on the strate-¹²⁴³ 1189 gies used during the performance of visuospatial working¹²⁴⁴ 1190 memory tasks have found that both, visualization and verbal-¹²⁴⁵ 1191 ization, are common strategies used to encode and maintain¹²⁴⁶ 1192 information in working memory [\(Brown and Wesley,](#page-15-49) 2013 ;¹²⁴⁷ ¹¹⁹³ [Oblak et al.,](#page-16-0) [2024,](#page-16-0) [2022;](#page-16-1) [Sanfratello et al.,](#page-16-37) [2014;](#page-16-37) [Slana Oz-](#page-16-3)1194 imič et al., [2023\)](#page-16-3). In addition, several studies have related¹²⁴⁹ 1195 individual differences in the use of these strategies to dis-1250 1196 tinct patterns of brain activity [\(Kirchhoff and Buckner,](#page-15-50) 2006 ;¹²⁵¹
Miller et al. 2012; Septembelle at al. 2014) 1197 [Miller et al.,](#page-16-5) [2012;](#page-16-5) [Sanfratello et al.,](#page-16-37) [2014\)](#page-16-37).

 In this study, we focused on general behavioral and neu- ral strategies used in spatial working memory rather than spe- cific mechanisms. Our results provide insight into the level of 1201 general cognitive demand involved in the use of fine-grained¹²⁵⁷ versus categorical representations. However, they do not in- dicate the specific brain regions in which the different types of information are represented. fMRI studies that used multi- variate pattern analysis (MVPA) have shown that fine-grained 1206 stimulus-specific information can be decoded from early sen-1262 [s](#page-15-51)ory areas that initially processed the stimulus [\(Harrison and](#page-15-51) 1263 [Tong,](#page-15-51) [2009;](#page-15-51) [Serences et al.,](#page-16-38) [2009\)](#page-16-38). In contrast, other stud- ies have shown that the prefrontal and parietal areas can store 1210 more abstract representations, such as goals, task rules, and 1266 categories [\(Christophel et al.,](#page-15-52) [2017;](#page-15-52) [D'Esposito and Postle,](#page-15-38) [2015;](#page-15-38) [Meyers et al.,](#page-16-39) [2008;](#page-16-39) [Riggall and Postle,](#page-16-40) [2012\)](#page-16-40). Addi- tionally, single-neuron recordings in the prefrontal cortex of 1214 monkeys during the performance of a spatial working mem-1270 ory task have shown that neurons, exhibiting directional se- lectivity for presented target angles, differed in the width of their tuning curves, suggesting that certain neurons respond to more specific directions and others to a broader range of 1219 directions [\(Funahashi et al.,](#page-15-7) [1989\)](#page-15-7). Based on these findings, 1275 it has been proposed that brain areas in the posterior-anterior axis respond to different levels of abstraction, with low-level posterior areas responding to fine-grained information and high-level anterior areas to more abstract information and [r](#page-15-38)egulatory signals [\(Christophel et al.,](#page-15-52) [2017;](#page-15-52) [D'Esposito and](#page-15-38) [Postle,](#page-15-38) [2015;](#page-15-38) [Rahmati et al.,](#page-16-41) [2018\)](#page-16-41). However, further studies are needed to identify areas of the brain that are involved in the storage of fine-grained and categorical representations used in spatial working memory.

The ability to detect significant brain-behavior rela**tionships**

Several recent studies [\(Elliott et al.,](#page-15-22) [2020;](#page-15-22) [Grady et al.,](#page-15-23) [2021;](#page-15-23) [Marek et al.,](#page-15-24) [2022;](#page-15-24) [Poldrack et al.,](#page-16-42) [2017\)](#page-16-42) have discussed the problem of highly variable brain-behavior relationships that require large sample sizes to obtain stable and reliable re-sults. For example, [Marek et al.](#page-15-24) [\(2022\)](#page-15-24) have shown that brain-wide association studies with typical sample sizes (i.e., around 25 participants) resulted in low statistical power, inflated effect sizes, and a failure to replicate results. We addressed this challenge by using a multi-site and multi-study fMRI dataset, which afforded us with a relatively large sample size $(n = 155)$ compared to other task-related fMRI stud-ies [\(Elliott et al.,](#page-15-22) [2020;](#page-15-22) [Marek et al.,](#page-15-24) [2022\)](#page-15-24). To the best of our knowledge, this is the largest fMRI dataset on spatial working memory to date. An additional advantage of a larger sample size was that it allowed us to explore the effect of the sample size on the findings of interest.

The investigation of the effect of sample size on β estimates as a measure of brain-behavior relationships revealed that β estimates can vary substantially from sample to sample when employing relatively small sample sizes. As indicated by the confidence intervals, the variability of the estimates decreased steeply at first and then slowly approached ¹²⁵³ the population mean. These results are consistent with the observation that the sampling variability is large for small [s](#page-15-24)ample sizes and stabilizes at larger sample sizes [\(Marek](#page-15-24) [et al.,](#page-15-24) 2022). In the case of our study, brain-behavior associations appeared to stabilize roughly between 73 and 151 [o](#page-15-23)bservations, consistent with the result obtained by [Grady](#page-15-23) [et al.](#page-15-23) [\(2021\)](#page-15-23) and [Schönbrodt and Perugini](#page-16-43) [\(2013\)](#page-16-43). Moreover, similar to other fMRI studies on brain-behavior associations [\(Marek et al.,](#page-15-24) [2022;](#page-15-24) [Poldrack et al.,](#page-16-42) [2017\)](#page-16-42), statistical power, i.e. the ability to detect a significant effect, increased monotonically with increasing sample size, and remained fairly low even at larger sample sizes. The maximum statistical power we observed was 88.4% at $n = 153$ for the relationship between memory inexactness and activity in the dorsalattention network.

In order to maximize sample size and statistical power, we combined data from multiple sites and studies, which presented additional challenges and limitations. Notably, there were minor differences in task designs and data collection protocols between studies, potentially contributing to the observed variability across participants. We addressed this issue using a multilevel approach. First, we analyzed the task condition that was directly comparable across studies and always

 had the same goal, i.e. to remember a single random target location for a few seconds on any individual trial. However, even though the task was essentially the same across stud- ies, there could potentially be some differences in task dif- ficulty, as a result of a different number of possible target locations, different duration of target stimulus presentation, or different length of the delay period. While we observed 1283 some differences in behavioral performance across studies, any differences in task difficulty were hard to delineate from 1285 the effects of different strategies on responses. Second, we¹³³⁹ 1286 used a hierarchical model with study as a random effect to¹³⁴⁰ 1287 account for any variability due to systematic differences be-¹³⁴¹ tween studies. Nevertheless, the final sample size was still 1289 relatively small compared to the recommendation of recent¹³⁴³ 1290 studies [\(Elliott et al.,](#page-15-22) [2020;](#page-15-22) [Marek et al.,](#page-15-24) [2022\)](#page-15-24) indicating that¹³⁴⁴ 1291 thousands of participants are required to prevent the inflation¹³⁴⁵ 1292 of effect sizes and replication failure in brain-behavior asso-1346 ciation analyses. In addition, the hierarchical structure of our data increased the complexity of the linear model used and might require even larger sample sizes to obtain reliable esti- 1348 1296 mates [\(Kerkhoff and Nussbeck,](#page-15-53) [2019;](#page-15-53) [Maas and Hox,](#page-15-54) [2004,](#page-15-54) [2005\)](#page-15-55). To further increase statistical power we performed analyses on brain networks rather than grayordinates with should provide a better signal-to-noise ratio due to averag-1300 ing data and ensure fewer statistical comparisons. We also¹³⁵¹ used Bayesian statistical methods which have been found to [g](#page-16-44)ive more robust results even at low sample sizes (e.g., [Van](#page-16-44)₁₃₅₃ [De Schoot et al.,](#page-16-44) [2021\)](#page-16-44). Additionally, we have provided de-1354 tailed power analysis to allow better insight into the stability of brain-behavior relationships in our study. However, more data or replication on an independent dataset would be wel- come to further ensure the validity and generalizability of the relationships observed in our analyses.

¹³⁰⁹ **Conclusion**

¹³¹⁰ In this multi-site, multi-study analysis, we found that indi-¹³¹¹ viduals differ in the extent to which they rely on fine-grained ¹³¹² versus categorical representations to encode and maintain a ¹³¹³ spatial location in working memory, and that these differ-¹³¹⁴ ences correlate with the engagement of brain networks dur-¹³¹⁵ ing the encoding, delay, and response phases of the task trial. ¹³¹⁶ Behaviorally, individuals with lower fine-grained precision ¹³¹⁷ relied more on categorical representations, which led to a 1318 higher categorical bias. Increased activation of attentional₁₃₆₈ ¹³¹⁹ and control brain networks throughout the entire task trial, 1320 and stronger deactivation of the default network in the en-¹³⁶⁹ 1321 coding period were found to predict higher precision in spa-¹³⁷⁰ ¹³²² tial working memory performance, possibly reflecting the 1323 importance of attentional resources for successful encoding¹³⁷¹ $_{1324}$ and maintenance of the fine-grained representation. In con- $_{1372}$ trast, the use of a categorical representation was associated₁₃₇₃ $_{1326}$ with lower default network activity in the encoding period_{$_{1374}$} ¹³²⁷ and higher frontoparietal network engagement in the delay ¹³²⁸ period, the latter possibly reflecting an inability to protect the ¹³²⁹ fine-grained representation from interference, which led to $_{1330}$ higher reliance on the categorical representation when pro $_{1377}^{1376}$ 1331 viding the response. The results stress the need to consider₁₃₇₈

individual differences in the use of specific representations and strategies when studying complex cognitive functions, such as working memory. They also illustrate the insights that the individual differences approach can provide in the study of brain-behavior relationships when a sufficient number of participants is ensured.

Author Contributions

Conceptualization: N.P.S., J.D.M., A.A., and G.R.; Project administration: N.P.S.; Data curation: N.P.S., Y.T.C., and A.S.O.; Methodology: N.P.S., A.K., M.R., J.D.M., A.A., and G.R.; Formal analysis: N.P.S., A.K., and G.R.; Visualization: N.P.S.; Writing – original draft: N.P.S.; Writing – review & editing: N.P.S., A.K., M.R., Y.T.C., A.S.O., J.D.M., A.A., and G.R.; Supervision: J.D.M., A.A., and G.R.; Funding acquisition: A.A. and G.R.; Resources: A.A. and G.R.

Conflict of Interest

J.D.M., A.A., and G.R. consult for and hold equity in Neumora Therapeutics and Manifest Technologies. Other authors declare that they have no conflict of interest.

Funding

This work was supported by the Slovenian Research and Innovation Agency $(Z5-50177)$ to N.P.S., J7-5553 and J3-¹³⁵⁴ 9264 to G.R., P3-0338 to A.S.O., and G.R., P5-0110 to A.K.), the National Institutes of Health (DP5OD012109-01, ¹³⁵⁶ 1U01MH121766, 5R01MH112189, and 5R01MH108590 to A.A.), the National Institute on Alcohol Abuse and Alcoholism (2P50AA012870-11 to A.A.), the Brain and Be-¹³⁵⁹ havior Research Foundation Young Investigator Award (to ¹³⁶⁰ A.A.), and Simons Foundation Autism Research Initiative Pi- $_{1361}$ lot Award (to A.A.).

¹³⁶² **Acknowledgments**

The authors would like to thank colleagues and students who helped with data collection, as well as all participants in the studies for their time and cooperation. We also thank the reviewers and editors for their constructive comments and suggestions.

Supplementary Material

Supplementary tables and figures are available at [the link to the supplementary material].

Data Availability

Data and analysis scripts for this paper can be found in the Open Science Framework (OSF) repository available at [https://osf.io/k8mvb/.](https://osf.io/k8mvb/)

¹³⁷⁵ **References**

Adam KCS, Mance I, Fukuda K, Vogel EK. The Contribution of Attentional Lapses to Individual Differences in Visual Working Memory Capacity. Journal of Cognitive Neuroscience. 2015 1378 27:1601–1616. https://doi.org/10.1162/jocn_a_00811.

- 1379 Adam KCS, Robison MK, Vogel EK. Contralateral Delay Activity Tracks Fluctuations in Working1465 1380 Memory Performance. Journal of Cognitive Neuroscience. 2018. 30:1229-1240. [https:](https://doi.org/10.1162/jocn_a_01233)1466 1381 [//doi.org/10.1162/jocn_a_01233](https://doi.org/10.1162/jocn_a_01233).
- 1382 Anticevic A, Repovs G, Shulman GL, Barch DM. When less is more: TPJ and default network 1383 deactivation during encoding predicts working memory performance. NeuroImage. 2010.1469 1384 49:2638–2648. <https://doi.org/10.1016/j.neuroimage.2009.11.008>.
- 1385 Assem M. Blank IA, Mineroff Z, Ademoğlu A, Fedorenko E. Activity in the fronto-parietal multiple-1471 1386 demand network is robustly associated with individual differences in working memory and 472 1387 fluid intelligence. Cortex. 2020. 131:1–16. [https://doi.org/10.1016/j.cortex.](https://doi.org/10.1016/j.cortex.2020.06.013) 1388 [2020.06.013](https://doi.org/10.1016/j.cortex.2020.06.013).
- 1389 Barch DM, Burgess GC, Harms MP, Petersen SE, Schlaggar BL, Corbetta M, Glasser MF, Curtiss
- 1390 S, Dixit S, Feldt C, Nolan D, Bryant E, Hartley T, Footer O, Bjork JM, Poldrack R, Smith S, 1476 [25217](https://doi.org/10.1002/hbm.25217). 1391 Johansen-Berg H, Snyder AZ, Van Essen DC. Function in the human connectome: Task-1477 1392 fMRI and individual differences in behavior. NeuroImage. 2013. 80:169-189. [https://](https://doi.org/10.1016/j.neuroimage.2013.05.033)1478 1393 doi.org/10.1016/j.neuroimage.2013.05.033.
- 1394 Bays PM, Husain M. Dynamic Shifts of Limited Working Memory Resources in Human Vision. 1395 Science. 2008. 321:851–854. <https://doi.org/10.1126/science.1158023>.
- 1396 Betancourt MJ, Girolami M. Hamiltonian Monte Carlo for Hierarchical Models. 2013. Number: 1397 arXiv:1312.0906 arXiv:1312.0906 [stat].
- 1398 Bray S, Almas R, Arnold AEGF, Iaria G, MacQueen G. Intraparietal Sulcus Activity and Func-1399 tional Connectivity Supporting Spatial Working Memory Manipulation. Cerebral Cortex. 2015.1485 1400 25:1252–1264. <https://doi.org/10.1093/cercor/bht320>.
- 1401 Brown LA, Wesley RW. Visual working memory is enhanced by mixed strategy use and seman-1487 1402 tic coding. Journal of Cognitive Psychology. 2013. 25:328-338. [https://doi.org/10.](https://doi.org/10.1080/20445911.2013.773004)1488 1403 [1080/20445911.2013.773004](https://doi.org/10.1080/20445911.2013.773004).
- 1404 Brown M, DeSouza J, Goltz HC, Ford K, Menon RS, Goodale MA, Everling S. Comparison of 1490 1405 Memory- and Visually Guided Saccades Using Event-Related fMRI. Journal of Neurophysi-1406 ology. 2004. 91:873–889. <https://doi.org/10/d4rkwq>.
- 1407 Chafee MV, Goldman-Rakic PS. Matching Patterns of Activity in Primate Prefrontal Area 8a and 1408 Parietal Area 7ip Neurons During a Spatial Working MemoryTask. Journal of Neurophysiol-1494 1409 ogy. 1998. 79:2919–2940. <https://doi.org/10.1152/jn.1998.79.6.2919>.
- Christophel TB, Klink PC, Spitzer B, Roelfsema PR, Haynes JD. The Distributed Nature of Work-1496 1411 ing Memory. Trends in Cognitive Sciences. 2017. 21:111-124. [https://doi.org/10.](https://doi.org/10.1016/j.tics.2016.12.007)1497 1412 [1016/j.tics.2016.12.007](https://doi.org/10.1016/j.tics.2016.12.007).
- 1413 Cole M, Bassett D, Power J, Braver T, Petersen S. Intrinsic and Task-Evoked Network Architec1414 tures of the Human Brain. Neuron. 2014. 83:238–251. [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.neuron.2014.05.014)1500
1501 1501 1501 1501 1501 1501 [neuron.2014.05.014](https://doi.org/10.1016/j.neuron.2014.05.014).
- 1416 Courtney SM, Petit L, Maisog JM, Ungerleider LG, Haxby JV. An Area Specialized for Spatialisoz
1417 Working Memory in Human Frontal Cortex. Science, 1998, 279:1347-1351. https://1503 Working Memory in Human Frontal Cortex. Science. 1998. 279:1347-1351. [https://](https://doi.org/10.1126/science.279.5355.1347)1503 1418 doi.org/10.1126/science.279.5355.1347.
- 1419 Crawford LE, Landy D, Salthouse TA. Spatial working memory capacity predicts bias in estimates 1420 of location. Journal of Experimental Psychology: Learning, Memory, and Cognition. 2016.1506 1421 42:1434–1447. <https://doi.org/10.1037/xlm0000228>.
- 1422 Curtis C. Prefrontal and parietal contributions to spatial working memory. Neuroscience. 2006.1508 1423 139:173–180. <https://doi.org/10.1016/j.neuroscience.2005.04.070>.
- 1424 Curtis CE. Maintenance of Spatial and Motor Codes during Oculomotor Delayed Response Tasks. 1425 Journal of Neuroscience. 2004. 24:3944–3952. <https://doi.org/10/fs74wg>.
- 1426 Damoiseaux JS, Rombouts SARB, Barkhof F, Scheltens P, Stam CJ, Smith SM, Beckmann 1427 CF. Consistent resting-state networks across healthy subjects. Proceedings of the National 1428 Academy of Sciences. 2006. 103:13848–13853. [https://doi.org/10.1073/pnas.](https://doi.org/10.1073/pnas.0601417103) 1429 [0601417103](https://doi.org/10.1073/pnas.0601417103).
- 1430 D'Esposito M, Postle BR. The Cognitive Neuroscience of Working Memory. An1431 nual Review of Psychology. 2015. 66:115-142. [https://doi.org/10.1146/](https://doi.org/10.1146/annurev-psych-010814-015031)1517 1432 [annurev-psych-010814-015031](https://doi.org/10.1146/annurev-psych-010814-015031).
- 1433 Duffy S, Huttenlocher J, Hedges LV, Crawford LE. Category effects on stimulus estimation: Shift-1434 ing and skewed frequency distributions. Psychonomic Bulletin & Review. 2010. 17:224-230.1520 1435 <https://doi.org/10.3758/PBR.17.2.224>.
- 1436 Elliott ML, Knodt AR, Ireland D, Morris ML, Poulton R, Ramrakha S, Sison ML, Moffitt TE, Caspi 1437 A, Hariri AR, What Is the Test-Retest Reliability of Common Task-Functional MRI Measures?1523 1438 New Empirical Evidence and a Meta-Analysis. Psychological Science. 2020. 31:792–806. 1439 <https://doi.org/10.1177/0956797620916786>.
- 1440 Eriksson J, Vogel E, Lansner A, Bergström F, Nyberg L. Neurocognitive Architecture of Working 1441 Memory. Neuron. 2015. 88:33-46. [https://doi.org/10.1016/j.neuron.2015.](https://doi.org/10.1016/j.neuron.2015.09.020)1527 1442 [09.020](https://doi.org/10.1016/j.neuron.2015.09.020).
- 1443 Fox MD, Snyder AZ, Vincent JL, Corbetta M, Van Essen DC, Raichle ME. The human brain ist529 1444 intrinsically organized into dynamic, anticorrelated functional networks. Proceedings of the 1445 National Academy of Sciences. 2005. 102:9673–9678. [https://doi.org/10.1073/](https://doi.org/10.1073/pnas.0504136102) 1446 [pnas.0504136102](https://doi.org/10.1073/pnas.0504136102).
- 1447 Friston K, Fletcher P, Josephs O, Holmes A, Rugg M, Turner R. Event-Related fMRI: Characteriz1448 ing Differential Responses. NeuroImage. 1998. 7:30-40. [https://doi.org/10.1006/](https://doi.org/10.1006/nimg.1997.0306)1534 1449 [nimg.1997.0306](https://doi.org/10.1006/nimg.1997.0306).
- 1450 Funahashi S, Bruce CJ, Goldman-Rakic PS. Mnemonic coding of visual space in the monkey's 1451 dorsolateral prefrontal cortex. Journal of Neurophysiology. 1989. 61:331-349. [https://](https://doi.org/10/ggcg98)1537 1452 doi.org/10/ggcg98.
- 1453 Fuster JM. Unit activity in prefrontal cortex during delayed-response performance: neuronal 1454 correlates of transient memory. Journal of Neurophysiology. 1973. 36:61-78. [https:](https://doi.org/10.1152/jn.1973.36.1.61)1540 1455 [//doi.org/10.1152/jn.1973.36.1.61](https://doi.org/10.1152/jn.1973.36.1.61).
- 1456 Fuster JM, Alexander GE. Neuron Activity Related to Short-Term Memory. Science. 1971.1542
1457 173:652-654 https://doi.org/10.1126/science. 173.3997.652. 1543. 1457 173:652–654. <https://doi.org/10.1126/science.173.3997.652>.
- 1458 Gelman A. Prior distributions for variance parameters in hierarchical models (comment on arti-1544 1459 cle by Browne and Draper). Bayesian Analysis. 2006. 1. [https://doi.org/10.1214/](https://doi.org/10.1214/06-BA117A)1545 1460
- 1461 Glahn D, Kim J, Cohen M, Poutanen VP, Therman S, Bava S, Van Erp T, Manninen M, Huttunen 1462 M, Lönnqvist J. Maintenance and Manipulation in Spatial Working Memory: Dissociations im 548 1463 the Prefrontal Cortex. NeuroImage. 2002. 17:201–213. [https://doi.org/10.1006/](https://doi.org/10.1006/nimg.2002.1161) 1464 [nimg.2002.1161](https://doi.org/10.1006/nimg.2002.1161).
- 1465 Glasser MF, Coalson TS, Robinson EC, Hacker CD, Harwell J, Yacoub E, Ugurbil K, Andersson J. Beckmann CF, Jenkinson M, Smith SM, Van Essen DC, A multi-modal parcellation 1467 of human cerebral cortex. Nature. 2016. 536:171–178. [https://doi.org/10.1038/](https://doi.org/10.1038/nature18933) [nature18933](https://doi.org/10.1038/nature18933).
- Glasser MF, Sotiropoulos SN, Wilson JA, Coalson TS, Fischl B, Andersson JL, Xu J, Jbabdi S, 1470 Webster M, Polimeni JR, Van Essen DC, Jenkinson M. The minimal preprocessing pipelines for the Human Connectome Project. NeuroImage. 2013. 80:105–124. [https://doi.org/](https://doi.org/10.1016/j.neuroimage.2013.04.127) 1472 [10.1016/j.neuroimage.2013.04.127](https://doi.org/10.1016/j.neuroimage.2013.04.127).
- Grady CL, Rieck JR, Nichol D, Rodrigue KM, Kennedy KM. Influence of sample size and analytic 1474 approach on stability and interpretation of brain-behavior correlations in task-related fMRI data. Human Brain Mapping. 2021. 42:204-219. [https://doi.org/10.1002/hbm.](https://doi.org/10.1002/hbm.25217)
- Greicius MD, Krasnow B, Reiss AL, Menon V, Functional connectivity in the resting brain: A network analysis of the default mode hypothesis. Proceedings of the National Academy of 1479 Sciences. 2003. 100:253–258. <https://doi.org/10.1073/pnas.0135058100>.
- Hallenbeck GE, Sprague TC, Rahmati M, Sreenivasan KK, Curtis CE. Working memory represen-1481 tations in visual cortex mediate distraction effects. Nature Communications. 2021. 12:4714. <https://doi.org/10.1038/s41467-021-24973-1>.
- 1483 Harrison SA, Tong F. Decoding reveals the contents of visual working memory in early visual areas. Nature. 2009. 458:632-635. https://doi.org/10.1038/nature078
	- Haun DB, Allen GL, Wedell DH. Bias in spatial memory: a categorical endorsement. Acta Psychologica. 2005. 118:149-170. https://doi.org/10.1016/j.actpsy.2004.10.
- Holyoak KJ, Mah WA. Cognitive reference points in judgments of symbolic magnitude. Cognitive Psychology. 1982. 14:328-352. [https://doi.org/10.1016/0010-0285\(82\)](https://doi.org/10.1016/0010-0285(82)90013-5) 1489 [90013-5](https://doi.org/10.1016/0010-0285(82)90013-5).
- Huttenlocher J, Hedges LV, Corrigan B, Crawford L. Spatial categories and the estimation of location. Cognition. 2004. 93:75-97. [https://doi.org/10.1016/j.cognition.2003.](https://doi.org/10.1016/j.cognition.2003.10.006) 1492 [10.006](https://doi.org/10.1016/j.cognition.2003.10.006)
- Huttenlocher J, Hedges LV, Duncan S. Categories and Particulars: Prototype Effects in Estimating Spatial Location. Psychological Review. 1991. 98:352–376. [https://doi.org/10.](https://doi.org/10.1037/0033-295X.98.3.352) 1495 1037/0033-295X.98.3.35
	- Huttenlocher J, Hedges LV, Vevea JL. Why Do Categories Affect Stimulus Judgment? Journal of Experimental Psychology: General. 2000. 129:220-241. [https://doi.org/10.1Q37/](https://doi.org/10.1Q37//0096-3445.129.2320) $96 - 3445.$
	- Jerde TA, Curtis CE. Maps of space in human frontoparietal cortex. Journal of Physiology-Paris 2013. 107:510-516. https://doi.org/10/ggrf9
- 1501 Ji JL, Demšar J, Fonteneau C, Tamayo Z, Pan L, Kraljič A, Matkovič A, Purg N, Helmer M, Warrington S, Winkler A, Zerbi V, Coalson TS, Glasser MF, Harms MP, Sotiropoulos SN, Murray JD, Anticevic A, Repovš G. QuNex—An integrative platform for reproducible neuroimaging an-1504 alytics. Frontiers in Neuroinformatics. 2023. 17:1104508. [https://doi.org/10.3389/](https://doi.org/10.3389/fninf.2023.1104508) 1505 [fninf.2023.1104508](https://doi.org/10.3389/fninf.2023.1104508).
- Ji JL, Spronk M, Kulkarni K, Repovš G, Anticevic A, Cole MW. Mapping the human brain's cortical1507 subcortical functional network organization. NeuroImage. 2019. 185:35-57. [https://doi.](https://doi.org/10.1016/j.neuroimage.2018.10.006) 1508 [org/10.1016/j.neuroimage.2018.10.006](https://doi.org/10.1016/j.neuroimage.2018.10.006).
- 1509 Juárez MA, Steel MFJ. Model-Based Clustering of Non-Gaussian Panel Data Based on Skew- *t* Distributions. Journal of Business & Economic Statistics. 2010. 28:52–66. [https://doi.](https://doi.org/10.1198/jbes.2009.07145) 1511 [org/10.1198/jbes.2009.07145](https://doi.org/10.1198/jbes.2009.07145).
	- Kerkhoff D, Nussbeck FW. The Influence of Sample Size on Parameter Estimates in Three-Level Random-Effects Models. Frontiers in Psychology. 2019. 10:1067. [https://doi.org/10.](https://doi.org/10.3389/fpsyg.2019.01067) 1514 [3389/fpsyg.2019.01067](https://doi.org/10.3389/fpsyg.2019.01067).
- 1515 Kirchhoff BA, Buckner RL. Functional-Anatomic Correlates of Individual Differences in Memory. 1516 Neuron. 2006. 51:263–274. <https://doi.org/10.1016/j.neuron.2006.06.006>.
- Klyszejko Z, Rahmati M, Curtis CE. Attentional priority determines working memory precision. 1518 Vision Research. 2014. 105:70–76. [https://doi.org/10.1016/j.visres.2014.](https://doi.org/10.1016/j.visres.2014.09.002) [09.002](https://doi.org/10.1016/j.visres.2014.09.002)
- Kubota K, Niki H. Prefrontal cortical unit activity and delayed alternation performance in mon-1521 keys. Journal of Neurophysiology. 1971. 34:337–347. [https://doi.org/10.1152/jn.](https://doi.org/10.1152/jn.1971.34.3.337) 1522 [1971.34.3.337](https://doi.org/10.1152/jn.1971.34.3.337).
- Leung HC, Seelig D, Gore JC. The effect of memory load on cortical activity in the spatial working memory circuit. Cognitive, Affective, & Behavioral Neuroscience. 2004. 4:553-563. [https:](https://doi.org/10.3758/CABN.4.4.553) 1525 [//doi.org/10.3758/CABN.4.4.553](https://doi.org/10.3758/CABN.4.4.553).
- Linden DE, Bittner RA, Muckli L, Waltz JA, Kriegeskorte N, Goebel R, Singer W, Munk MH. Cortical capacity constraints for visual working memory: dissociation of fMRI load effects 1528 in a fronto-parietal network. NeuroImage. 2003. 20:1518–1530. [https://doi.org/10.](https://doi.org/10.1016/j.neuroimage.2003.07.021) 1529 [1016/j.neuroimage.2003.07.021](https://doi.org/10.1016/j.neuroimage.2003.07.021).
- 1530 Liu J, Xia M, Dai Z, Wang X, Liao X, Bi Y, He Y. Intrinsic Brain Hub Connectivity Underlies Individual Differences in Spatial Working Memory. Cerebral Cortex. 2017. 27:5496-5508. 1532 https://doi.org/10.1093/cercor/bhw31
- Liu S, Poh JH, Koh HL, Ng KK, Loke YM, Lim JKW, Chong JSX, Zhou J. Carrying the past to the future: Distinct brain networks underlie individual differences in human spatial work1535 ing memory capacity. NeuroImage. 2018. 176:1-10. [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.neuroimage.2018.04.014) [neuroimage.2018.04.014](https://doi.org/10.1016/j.neuroimage.2018.04.014).
- Luck SJ, Vogel EK. Visual working memory capacity: from psychophysics and neurobiology to 1538 individual differences. Trends in Cognitive Sciences. 2013. 17:391-400. [https://doi.](https://doi.org/10.1016/j.tics.2013.06.006) 1539 [org/10.1016/j.tics.2013.06.006](https://doi.org/10.1016/j.tics.2013.06.006).
- Maas CJM, Hox JJ. Robustness issues in multilevel regression analysis. Statistica Neerlandica 1541 2004. 58:127–137. <https://doi.org/10.1046/j.0039-0402.2003.00252.x>.
- 1542 Maas CJM, Hox JJ. Sufficient Sample Sizes for Multilevel Modeling. Methodology. 2005. 1:86–92. 1543 <https://doi.org/10.1027/1614-2241.1.3.86>.
	- Magnuson ME, Thompson GJ, Schwarb H, Pan WJ, McKinley A, Schumacher EH, Keilholz SD. Errors on interrupter tasks presented during spatial and verbal working memory performance are linearly linked to large-scale functional network connectivity in high temporal resolution resting state fMRI. Brain Imaging and Behavior. 2015. 9:854-867. [https://doi.org/](https://doi.org/10.1007/s11682-014-9347-3) 10.1007/s11682-014-9347-
- Marek S, Tervo-Clemmens B, Calabro FJ, Montez DF, Kay BP, Hatoum AS, Donohue MR, Foran 1550 W, Miller RL, Hendrickson TJ, Malone SM, Kandala S, Feczko E, Miranda-Dominguez O, Gra-

- 1551 ham AM, Earl EA, Perrone AJ, Cordova M, Doyle O, Moore LA, Conan GM, Uriarte J, Snideris37 1552 K, Lynch BJ, Wilgenbusch JC, Pengo T, Tam A, Chen J, Newbold DJ, Zheng A, Seider NA,1638
- 1553 Van AN, Metoki A, Chauvin RJ, Laumann TO, Greene DJ, Petersen SE, Garavan H, Thomp-1554 son WK, Nichols TE, Yeo BTT, Barch DM, Luna B, Fair DA, Dosenbach NUF. Reproducible
- 1555 brain-wide association studies require thousands of individuals. Nature. 2022. 603:654–660.

1556 <https://doi.org/10.1038/s41586-022-04492-9>. 1557 Meyers EM, Freedman DJ, Kreiman G, Miller EK, Poggio T. Dynamic Population Coding of Cat-

- 1558 egory Information in Inferior Temporal and Prefrontal Cortex. Journal of Neurophysiology. 1559 2008. 100:1407–1419. <https://doi.org/10.1152/jn.90248.2008>.
- 1560 Miller MB, Donovan CL, Bennett CM, Aminoff EM, Mayer RE. Individual differences in cognitive 1561 style and strategy predict similarities in the patterns of brain activity between individuals.1647 1562 NeuroImage. 2012. 59:83–93. [https://doi.org/10.1016/j.neuroimage.2011.](https://doi.org/10.1016/j.neuroimage.2011.05.060) 1563 [05.060](https://doi.org/10.1016/j.neuroimage.2011.05.060).
- 1564 Moujaes F, Ji JL, Rahmati M, Burt JB, Schleifer C, Adkinson BD, Savic A, Santamauro N, Tamayo 1565 Z, Diehl C, Kolobaric A, Flynn M, Rieser N, Fonteneau C, Camarro T, Xu J, Cho Y, Repovs 1566 G, Fineberg SK, Morgan PT, Seifritz E, Vollenweider FX, Krystal JH, Murray JD, Preller KH, 1567 Anticevic A. Ketamine induces multiple individually distinct whole-brain functional connectivity 1653
1568 Signatures, eLife, 2024, 13:e84173, https://doi.org/10.7554/eLife.84173. signatures. eLife. 2024. 13:e84173. <https://doi.org/10.7554/eLife.84173>.
- 1569 Moussa MN, Steen MR, Laurienti PJ, Hayasaka S. Consistency of Network Modules in Resting1570 State fMRI Connectome Data. PLoS ONE. 2012. 7:e44428. [https://doi.org/10.](https://doi.org/10.1371/journal.pone.0044428)1656 1571 [1371/journal.pone.0044428](https://doi.org/10.1371/journal.pone.0044428).
- 1572 Nelson TO, Chaiklin S. Immediate memory for spatial location. Journal of Experimental Psychol-1658 1573 ogy: Human Learning and Memory. 1980. 6:529–545. [https://doi.org/10.1037/](https://doi.org/10.1037/0278-7393.6.5.529) 1574 [0278-7393.6.5.529](https://doi.org/10.1037/0278-7393.6.5.529).
- 1575 Newcombe N, Liben LS. Barrier effects in the cognitive maps of children and adults. Jour-1576 nal of Experimental Child Psychology. 1982. 34:46–58. [https://doi.org/10.1016/](https://doi.org/10.1016/0022-0965(82)90030-3) 1577 0022-0965 (82) 90030
- 1578 Oblak A, Dragan O, Slana Ozimič A, Kordeš U, Purg N, Bon J, Repovš G. What is it like to do at664 1579 visuo-spatial working memory task: A qualitative phenomenological study of the visual span 1580 task. Consciousness and Cognition. 2024. 118:103628. [https://doi.org/10.1016/](https://doi.org/10.1016/j.concog.2023.103628)1666 1581 j.concog. 2023. 103628
1582 Oblak A. Slana Ozimič A. Repov
- Oblak A, Slana Ozimič A, Repovš G, Kordeš U. What Individuals Experience During Visuo-Spatial 668 1583 Working Memory Task Performance: An Exploratory Phenomenological Study. Frontiers in
- 1584 Psychology. 2022. 13:811712. <https://doi.org/10.3389/fpsyg.2022.811712>. 1585 Peirce J, Gray JR, Simpson S, MacAskill M, Höchenberger R, Sogo H, Kastman E, Lindeløv 1586 JK. PsychoPy2: Experiments in behavior made easy. Behavior Research Methods. 2019.1672
1587 51:195–203. https://doi.org/10.3758/s13428-018-01193-v.
- 1587 51:195–203. <https://doi.org/10.3758/s13428-018-01193-y>. 1588 Poldrack RA, Baker CI, Durnez J, Gorgolewski KJ, Matthews PM, Munafò MR, Nichols TE, Poline
- JB, Vul E, Yarkoni T. Scanning the horizon: towards transparent and reproducible neuroimag-1675 1590 ing research. Nature Reviews Neuroscience. 2017. 18:115-126. [https://doi.org/10.](https://doi.org/10.1038/nrn.2016.167)1676 1591 [1038/nrn.2016.167](https://doi.org/10.1038/nrn.2016.167).
- 1592 Proskovec AL, Wiesman AI, Heinrichs-Graham E, Wilson TW. Load effects on spatial working 1593 memory performance are linked to distributed alpha and beta oscillations. Human Brain Map-1594 ping. 2019. 40:3682–3689. <https://doi.org/10.1002/hbm.24625>.
- 1595 Ptak R. The Frontoparietal Attention Network of the Human Brain: Action, Saliency, and a Priority 1596 Map of the Environment. The Neuroscientist. 2012. 18:502-515. [https://doi.org/10.](https://doi.org/10.1177/1073858411409051)1682 1597 [1177/1073858411409051](https://doi.org/10.1177/1073858411409051).
- 1598 Purg N, Starc M, Slana Ozimič A, Kraljič A, Matkovič A, Repovš G, Neural Evidence for Different 684 1599 Types of Position Coding Strategies in Spatial Working Memory. Frontiers in Human Neuro-1600 science. 2022. 16:821545. <https://doi.org/10.3389/fnhum.2022.821545>.
- 1601 Rahmati M, Saber GT, Curtis CE. Population Dynamics of Early Visual Cortex during Working 1602 Memory. Journal of Cognitive Neuroscience. 2018. 30:219–233. [https://doi.org/10.](https://doi.org/10.1162/jocn_a_01196) 1603 [1162/jocn_a_01196](https://doi.org/10.1162/jocn_a_01196).
- 1604 Raichle ME. The Brain's Default Mode Network. Annual Review of Neuroscience. 2015a. 38:433– 1605 447. <https://doi.org/10.1146/annurev-neuro-071013-014030>.
- 1606 Raichle ME. The restless brain: how intrinsic activity organizes brain function. Philosophical 1607 Transactions of the Royal Society B: Biological Sciences. 2015b. 370:20140172. [https:](https://doi.org/10.1098/rstb.2014.0172) 1608 [//doi.org/10.1098/rstb.2014.0172](https://doi.org/10.1098/rstb.2014.0172).
- 1609 Rainer G, Asaad WF, Miller EK. Selective representation of relevant information by neurons in the 1610 primate prefrontal cortex. Nature. 1998. 393:577–579. <https://doi.org/10/df47cn>.
- 1611 Riggall AC, Postle BR. The Relationship between Working Memory Storage and Elevated Activity 1612 as Measured with Functional Magnetic Resonance Imaging. Journal of Neuroscience. 2012.
- 1613 32:12990–12998. <https://doi.org/10.1523/JNEUROSCI.1892-12.2012>. 1614 Sadalla EK, Burroughs WJ, Staplin LJ. Reference points in spatial cognition. Journal of Experi-
- 1615 mental Psychology: Human Learning and Memory. 1980. 6:516–528. [https://doi.org/](https://doi.org/10.1037/0278-7393.6.5.516) 1616 [10.1037/0278-7393.6.5.516](https://doi.org/10.1037/0278-7393.6.5.516).
- 1617 Sanfratello L, Caprihan A, Stephen JM, Knoefel JE, Adair JC, Qualls C, Lundy SL, Aine CJ. Same 1618 task, different strategies: How brain networks can be influenced by memory strategy. Human 1619 Brain Mapping. 2014. 35:5127-5140. https://doi.org/10.1002/hb
- 1620 Schneider W, Eschman A, Zuccolotto A. E-Prime Reference Guide. 2012.
- 1621 Schutte AR, Spencer JP, Schöner G. Testing the Dynamic Field Theory: Working Memory for 1622 Locations Becomes More Spatially Precise Over Development. Child Development. 2003. 1623 74:1393–1417. <https://doi.org/10.1111/1467-8624.00614>.
- 1624 Schönbrodt FD, Perugini M. At what sample size do correlations stabilize? Journal of Research in 1625 Personality. 2013. 47:609–612. <https://doi.org/10.1016/j.jrp.2013.05.009>.
- 1626 Serences JT, Ester EF, Vogel EK, Awh E. Stimulus-Specific Delay Activity in Human Primary 1627 Visual Cortex. Psychological Science. 2009. 20:207–214. [https://doi.org/10.1111/](https://doi.org/10.1111/j.1467-9280.2009.02276.x) 1628 **[j.1467-9280.2009.02276.x](https://doi.org/10.1111/j.1467-9280.2009.02276.x)**
- 1629 Shulman GL, McAvoy MP, Cowan MC, Astafiev SV, Tansy AP, d'Avossa G, Corbetta M. Quantita-1630 tive Analysis of Attention and Detection Signals During Visual Search. Journal of Neurophys-1631 iology. 2003. 90:3384-3397. https://doi.org/10.1152/jn.00343.200
- 1632 Simmering VR, Spencer JP, Schöner G. Reference-related inhibition produces enhanced position 1633 discrimination and fast repulsion near axes of symmetry. Perception & Psychophysics. 2006. 1634 68:1027–1046. <https://doi.org/10.3758/BF03193363>.
- 1635 Slana Ozimič A, Oblak A, Kordeš U, Purg N, Bon J, Repovš G. The diversity of strategies used 1636 in working memory for colors, orientations, and positions: A quantitative approach to a first-

person inquiry. BioRxiv. 2023.

- Slana Ozimič A, Repovš G. Visual working memory capacity is limited by two systems that change across lifespan. Journal of Memory and Language. 2020. 112:104090. [https://doi.](https://doi.org/10.1016/j.jml.2020.104090) [org/10.1016/j.jml.2020.104090](https://doi.org/10.1016/j.jml.2020.104090)
- Smith SM. Fox PT. Miller KL. Glahn DC, Fox PM, Mackay CE, Filippini N, Watkins KE, Toro R, Laird 1642 AR, Beckmann CF. Correspondence of the brain's functional architecture during activation and rest. Proceedings of the National Academy of Sciences. 2009. 106:13040-13045. [https:](https://doi.org/10.1073/pnas.0905267106) 1644 [//doi.org/10.1073/pnas.0905267106](https://doi.org/10.1073/pnas.0905267106).
- 1645 Spencer JP, Hund AM. Prototypes and particulars: Geometric and experience-dependent spatial categories. Journal of Experimental Psychology: General. 2002. 131:16-37. [https://](https://doi.org/10.1037/0096-3445.131.1.16) 1647 doi.org/10.1037/0096-3445.131.1.16.
- Srimal R, Curtis CE. Persistent neural activity during the maintenance of spatial position in 1649 working memory. NeuroImage. 2008. 39:455–468. [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.neuroimage.2007.08.040) 1650 [neuroimage.2007.08.040](https://doi.org/10.1016/j.neuroimage.2007.08.040).
- Starc M, Anticevic A, Repovš G. Fine-grained versus categorical: Pupil size differentiates between strategies for spatial working memory performance: Pupil size predicts working memory strategies. Psychophysiology. 2017. 54:724–735. [https://doi.org/10.1111/psyp.](https://doi.org/10.1111/psyp.12828) 1654 [12828](https://doi.org/10.1111/psyp.12828).
- Stukken L, Van Rensbergen B, Vanpaemel W, Storms G. Understanding individual differences in representational abstraction: The role of working memory capacity. Acta Psychologica. 2016. 1657 170:94–102. <https://doi.org/10.1016/j.actpsy.2016.06.002>.
	- Team RC. R: A Language and Environment for Statistical Computing. 2022a.
	- Team SD. Stan Modeling Language Users Guide and Reference Manual. 2022b.
- 1660 Todd JJ, Fougnie D, Marois R. Visual Short-Term Memory Load Suppresses Temporo-Parietal Junction Activity and Induces Inattentional Blindness. Psychological Science. 2005. 16:965–972. https://doi.org/10.1111/j.1467-9280.2005.01645.x. /doi.org/10.1111/j.1467-9280.2005.01645.x.
- 1663 Tversky B, Schiano DJ. Perceptual and conceptual factors in distortions in memory for graphs and maps. Journal of Experimental Psychology: General. 1989. 118:387-398. [https:](https://doi.org/10.1037/0096-3445.118.4.387) 1665 [//doi.org/10.1037/0096-3445.118.4.387](https://doi.org/10.1037/0096-3445.118.4.387).
- 1666 Van De Schoot R, Depaoli S, King R, Kramer B, Märtens K, Tadesse MG, Vannucci M, Gelman 1667 A, Veen D, Willemsen J, Yau C. Bayesian statistics and modelling. Nature Reviews Methods Primers. 2021. 1:1. https://doi.org/10.1038/s43586-020-00001-
	- Vehtari A, Gelman A, Simpson D, Carpenter B, Bürkner PC. Rank-normalization, folding, and localization: An improved \$\widehat{R}\$ for assessing convergence of MCMC. Bayesian Analysis. 2021. 16. <https://doi.org/10.1214/20-BA1221>. ArXiv:1903.08008 [stat].
- Winkler AM, Ridgway GR, Webster MA, Smith SM, Nichols TE. Permutation inference for the 1673 general linear model. NeuroImage. 2014. 92:381–397. [https://doi.org/10.1016/j.](https://doi.org/10.1016/j.neuroimage.2014.01.060) iroimage.2014.01.060
- Yoo AH, Bolaños A, Hallenbeck GE, Rahmati M, Sprague TC, Curtis CE. Behavioral Prioritization Enhances Working Memory Precision and Neural Population Gain. Journal of Cognitive 1677 Neuroscience. 2022. 34:365–379. https://doi.org/10.1162/jocn_a_01804.
- Zarahn E, Aguirre GK, D'Esposito M. Temporal isolation of the neural correlates of spatial mnemonic processing with fMRI. Cognitive Brain Research. 1999. 7:255–268. [https:](https://doi.org/10.1016/S0926-6410(98)00029-9) 1680 [//doi.org/10.1016/S0926-6410\(98\)00029-9](https://doi.org/10.1016/S0926-6410(98)00029-9).
- Zhang R, Geng X, Lee TMC. Large-scale functional neural network correlates of response inhibition: an fMRI meta-analysis. Brain Structure and Function. 2017. 222:3973-3990. 1683 <https://doi.org/10.1007/s00429-017-1443-x>.
	- Zhang W, Luck SJ. Discrete fixed-resolution representations in visual working memory. Nature. 1685 2008. 453:233–235. <https://doi.org/10.1038/nature06860>.