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35 Highlights

- 36 • The perirhinal cortex (PrC) is damaged early in Alzheimer's disease (AD)
- 37 • It allows for conceptually and perceptually confusable objects discrimination
- 38 • We investigated perceptual and conceptual discrimination in different tasks
- 39 • Conceptual confusability disambiguation was linked to the left PrC
- 40 • Conceptual discrimination of confusable objects can be a marker of PrC damage

41

42 Abstract

43 The perirhinal cortex (PrC) stands among the first brain areas to deteriorate in Alzheimer's
44 disease (AD). Many efforts have been made to understand its role in different cognitive
45 functions, to eventually identify early cognitive markers of AD. Based on recent research, the
46 present paper tests to what extent the PrC is involved in representing and discriminating
47 confusable objects based on the conjunction of their perceptual and conceptual features. To this
48 aim, AD patients and control counterparts performed three tasks: a naming task, a recognition
49 memory task, and a conceptual matching task, where we manipulated conceptual confusability
50 and, in the recognition memory task, perceptual confusability. In addition, a high-resolution
51 structural MRI of the antero-lateral parahippocampal subregions was obtained for each
52 participant. The main results were that the sensibility to conceptual confusability was associated
53 with the left PrC volume in both AD patients and control participants for the recognition
54 memory task, while it was specifically associated with the volume of the left PrC in AD patients
55 for the conceptual matching task. This suggests that a decreased volume of the PrC is related to
56 the ability to disambiguate conceptually confusable items. Therefore, testing recognition
57 memory or conceptual matching of easily conceptually confusable items can provide a potential
58 cognitive marker of PrC atrophy.

59 Key words: perirhinal cortex, Alzheimer's disease, episodic memory, semantic memory,
60 conceptual distance, perceptual distance

61

62 **Introduction**

63 It is well-established that the medial temporal lobe (MTL), comprising the
64 hippocampus, entorhinal (ErC), perirhinal (PrC) and parahippocampal cortices, is critical for
65 declarative long-term memory (Squire et al., 2004). In this system, the PrC, composed of
66 Brodmann areas (BA) 35 and 36, has received increasing interest in the last decades with a
67 particular focus on its role in cognition (for a review, see Suzuki & Naya, 2014). The PrC plays
68 a key role in memory-related behaviors such as the ability to recognize a previously experienced
69 stimulus (Squire et al., 2007) and especially using familiarity, as opposed to recollection (e.g.,
70 Brown & Aggleton, 2001. For instance, a rare lesion-based human neuropsychological case
71 (NB), who had a surgical resection of a large portion of the PrC sparing the hippocampus,
72 presents impaired familiarity with preserved recollection (Bowles et al., 2007). This line of
73 evidence has also received strong support by a wide variety of neuroimaging studies (see Bastin
74 et al., 2019, for review).

75 The MTL is widely connected with the neocortex, and the PrC receives much of its
76 neocortical input from the ventral visual stream (VVS; Suzuki & Amaral, 1994), which is
77 critical to build perceptual representations of objects (Lee et al., 2012). This has led some
78 researchers to interpret the PrC as an extension of the representational hierarchy within the VVS
79 for object identification (Murray & Bussey, 1999), making this cortex responsible for
80 processing and storing representations of complex feature conjunctions (Bussey & Saksida,
81 2007). Based on this idea, Bussey et al. (2002) introduced the perceptual–mnemonic/feature
82 conjunction (PMFC) model that emphasizes the conjunctive processing function of the PrC,
83 required to perform fine-grained perceptual discrimination between highly similar objects
84 composed of overlapping features. This model has been supported by multiple subsequent
85 imaging and lesion studies, which have linked the PrC with fine-grained visual discrimination
86 of perceptually ambiguous objects (e.g., Barense et al., 2007; Buckley & Gaffan, 2006; Bussey

87 et al., 2005; Inhoff et al., 2019). In addition, some recent lesion and neuroimaging studies have
88 suggested that this role of fine-grained object discrimination could be extended to the antero-
89 lateral ErC (alErC) onto which the PrC projects directly, through the integration of the object
90 with additional spatial features (Connor & Knierim, 2017; Ferko et al., 2022).

91 Nevertheless, not all inputs received by the PrC come from the VVS. Indeed, the PrC
92 also receives multi-modal projections, such as from the insular cortex and area 13 of the
93 orbitofrontal cortex (Suzuki & Amaral, 1994), and is also part of the anterior temporal lobe,
94 that is considered by predominant semantic memory models as a semantic hub responsible for
95 the amodal integration of conceptual information (Patterson et al., 2007; Ralph et al., 2016). As
96 such, the left PrC has recently been identified as key for the integration of the meaning of object
97 representations derived from visual information (Clarke & Tyler, 2015; Martin et al., 2018;
98 Price et al., 2017; see Bastin et al., 2019) and is recruited when conceptually confusable objects
99 must be differentiated (Bruffaerts et al., 2019; Clarke & Tyler, 2015; Kivisaari et al., 2012;
100 Wright et al., 2015). Therefore, the cognitive role of the PrC is not only dealing with the
101 conjunction of perceptual features, but also with the conjunction of conceptual features (Martin
102 et al., 2018), even though it is sometimes difficult to distinguish between a perceptual and
103 conceptual feature (e.g., “has four legs”). This also implies that the PrC should not be
104 confined to a role in mnemonic similarity tasks (Kent et al., 2016) developed under Yassa and
105 Stark’s model of pattern separation (Yassa & Stark, 2011). Indeed, under the assumption of a
106 role in conceptual or perceptual disambiguation, its function is thus not restricted to memory,
107 nor to discrimination between similar exemplars of the same concept but also to discrimination
108 between exemplars representing different concepts or percepts, as long as they overlap. Taken
109 together, these theories seem to point to the idea that the role of the PrC could be better
110 understood by considering the type of representation it underlies (i.e., fine-grained
111 representation of fully specified object concepts), rather than the type of process it supports

112 (episodic memory *vs.* semantic memory *vs.* perception) (Sheldon et al., 2019; Bastin et al.,
113 2019). However, as to what type of representation it precisely supports, the question remains
114 elusive to date.

115 Critically, the study of the cognitive function of the PrC is particularly relevant in the
116 frame of Alzheimer’s disease (AD) research. AD-related neuropathology starts years before the
117 onset of behavioral symptoms leading to AD diagnosis, and occurs as accumulation of tau
118 neurofibrillary tangles propagates in the brain (see Sexton et al., 2022). More precisely, Stage
119 1 of AD concerns the transentorhinal cortex, corresponding to BA 35 within the PrC and to the
120 alErC (see Braak & Braak, 1991, Braak & Braak, 1997). As such, one may predict that patients
121 diagnosed with AD should display impairments when it comes to recruiting the cognitive
122 functions supported by this region. We recently proposed that the very first impairment
123 occurring early in the course of AD would thereby affect representations at the level of entity
124 by binding its perceptuo-conceptual features enabling to differentiate it from similar but distinct
125 entities (Bastin & Delhaye, 2023).

126 While studies exploring familiarity processing in patients with AD or with Mild
127 Cognitive Impairment (MCI) at risk of AD led to inconsistent results (see Koen & Yonelinas,
128 2014; Schoemaker et al., 2014), some studies showed evidence for impairments in MCI patients
129 to perform tasks requiring the fine perceptual discrimination of objects (Newsome et al., 2012;
130 Yeung et al., 2013). Moreover, recent work showed that the volume of the alErC significantly
131 and selectively predicts the processing of the spatial arrangement of conjunctive objects features
132 in healthy aging (Yeung et al., 2017) as well as the ability to discriminate in memory between
133 similar objects despite being differently presented at recognition using familiarity in amnesic
134 MCI patients (Besson et al., 2020). In addition, studies have also reported a greater vulnerability
135 to distinguish between distinctive features (e.g., “has stripes”) than shared ones (e.g., “has four
136 legs”) early in the course of AD, causing close concepts to become gradually supported solely

137 by shared features, and eventually merge these concepts together into a single unit (Laisney et
138 al., 2011). This degradation was shown across a variety of tasks where distinctive features were
139 manipulated such as in naming (Garrard et al., 2005), semantic priming (Laisney et al., 2011),
140 or recognition memory (Flanagan et al., 2013). Finally, two studies have demonstrated an
141 impaired capacity in AD patients to name (Kivisaari et al., 2012) and to discriminate in memory
142 (Kivisaari et al., 2013) what the authors considered as “confusable” concepts, and this
143 impairment was related to the atrophy of the left medial PrC.

144 Yet, these aforementioned studies considered “conceptual confusability” between object
145 concepts as the distinction between belonging (confusable) or not (not confusable) to the same
146 category-domain (living vs. non-living). In other words, in these studies, all living things were
147 considered more “confusable” than non-living things. This is based on the idea put forward by
148 some models of semantic memory that living things are inherently more confusable than non-
149 living things due to their conceptual structure (see the Conceptual Structure Account (CSA);
150 see Clarke & Tyler, 2015, for review). According to these models, concepts confusability could
151 be precisely characterized by computing their feature-based statistics using feature-based
152 matrices, where conceptual confusability would be defined by the number of conceptual
153 features that they share, their tendency to co-occur, as well as the number of distinctive features
154 that a particular concept has as compared to other concepts from the same category. Depending
155 on these measures, a concept might be more or less confusable with other concepts. Yet, despite
156 the fact that living concepts are thought to be inherently more confusable in nature, because
157 they share a greater number of features that tend to co-occur more often, and tend to have less
158 distinctive features, there should still be more and less confusable concepts in both living and
159 non-living domains. On this basis, Wright et al. (2015) developed a quantitative measure of the
160 sensitivity to conceptual confusability, which relates performance to a quantitative distance
161 between objects, based on their internal conceptual structure defined by their features. Thereby,

162 this method goes beyond the approximative distinction between living vs. non-living used in
163 previous studies (Kivisaari et al., 2012, 2013). Here we suggest that this method can then also
164 be extended to perceptual distances between objects. Yet, to date, no study has ever used it to
165 assess sensibility to perceptual confusability.

166 In this study, with the aim to better characterize the cognitive role of the MTL regions,
167 and more specifically of the PrC region, we tested the hypothesis according to which this region
168 is involved in conceptual and/or perceptual fine-grained discrimination, regardless of the type
169 of memory involved, be it semantic or episodic, or the type of task. To do so, we sought to
170 better quantify and characterize conceptual confusability among living and non-living things.
171 We implemented several tasks assessing a variety of cognitive functions all requiring fine
172 conceptual discrimination (naming task, subsequent recognition memory and conceptual
173 matching task). Across these tasks, we manipulated conceptual similarity across living and non-
174 living things, using the quantification metrics of conceptual similarity developed by feature-
175 based models. In addition, for the recognition memory task only, we accounted for perceptual
176 similarity between to-be-discriminated items (see Wright et al., 2015 and Naspi et al., 2022 for
177 a similar method, and the Methods section for details). Our main hypothesis was that in both
178 the recognition memory and conceptual matching tasks, higher conceptual similarity would be
179 related to the integrity of the left PrC in patients with AD but not in control participants. Indeed,
180 we expected patients' variability in volumes and cognitive scores to come predominantly from
181 a similar factor (the AD pathology within the transentorhinal region) while controls' variability
182 - in the absence of such a common factor - to come from a more diverse set of factors not
183 necessarily affecting simultaneously volumes and cognitive scores.

184 **Methods**

185 *Participants*

186 A total of 24 patients diagnosed with mild probable AD (clinical criteria from McKhann
 187 et al., 2011) and 23 control participants took part in the study. They were matched in terms of
 188 age and education level. All participants were community-dwelling individuals; they were all
 189 French-speaking, had normal or corrected-to-normal vision, and reported no neurological or
 190 psychiatric history (except for the disease in the case of AD patients). They all underwent a
 191 short neuropsychological evaluation. Demographics and neuropsychological data are shown in
 192 Table 1. The study was approved by the Ethics committee of the Medicine Faculty of the
 193 University of Liège. Participants signed an informed consent form prior to taking part in the
 194 experiment.

195 **Table 1. Demographic information and comparison of the neuropsychological evaluation**
 196 **between AD patients and control participants (t-tests).**

| | AD patients Mean (SD) | Control Participants Mean (SD) | <i>p</i> -value |
|---|--------------------------|-----------------------------------|-----------------|
| Female/Male | 10/14 | 11/12 | |
| Age | 74.79 (6.30) | 72.26 (4.18) | .113 |
| Education | 12.46 (3.09) | 13.09 (3.34) | .506 |
| MoCA | 21.00 (3.16) | 27.43 (1.27) | < .001 |
| Letter fluency | 16.29 (7.36) | 18.96 (4.47) | .142 |
| Category fluency | 18.62 (6.83) | 26.26 (5.32) | < .001 |
| WAIS-3 digit symbol substitution | 16.37 (14.50) | 18.91 (18.72) | .605 |
| WAIS-3 vocabulary | 31.85 (9.09) | 37.21 (7.82) | .056 |
| WMS-3 Logical Memory immediate recall | 7.96 (3.42) | 13.13 (3.44) | < .001 |

197

198 *Materials*

199 276 object concepts were selected from 16 categories of living and non-living things
200 from existing feature norms (CSLB property norms, Devereux et al., 2014) and were translated
201 into French. Each selected concept was associated with a picture representing the object it
202 defines. In a pilot test, an independent group of 10 young participants evaluated the exemplarity
203 of the picture for the concept-label on a scale from 1 (not representative at all) to 7 (very
204 representative). Pictures received mean exemplarity judgements of 6.18 (range: 4.5 - 7).
205 Concepts pairs (e.g., target-distractor pairs in the memory task, or test-pairs in the conceptual
206 matching task) were characterized by an index of conceptual similarity as the cosine between
207 their production frequency vectors within the feature norms database (McRae et al., 2005). We
208 also extracted measures from the CSA representing the interaction between feature sharedness
209 and their correlational strength ('correlation x distinctiveness', CxD), which represents the
210 extent to which concepts are distinctive (with more or less distinctive features, that more or less
211 co-occur with one another) and thus, whether they will be more easily identified or will require
212 additional differentiation processes.

213 In addition, perceptual distances between stimuli pairs in the episodic recognition task
214 were computed using the HMax computational model of vision (available at
215 <http://cbcl.mit.edu/software-datasets/standardmodel>), following Clarke et al.'s study (2015).
216 HMax models different hierarchical stages of the ventral processing stream in different layers,
217 progressing from early visual cortex (V1/V2) to posterior inferior temporal cortex (IT). The C1
218 layers correspond to increasingly position- and scale-invariant early visual cortex (V1/V2) that
219 maintain feature specificity, while C2 layers simulate the extrastriate visual area cells (V4/IT)
220 that integrate visual features from previous layers to represent object shape. Measures based on
221 these two layers have been validated in studies of visual object recognition that have
222 distinguished the time courses and neural correlates of semantic versus visual processing
223 (Clarke & Tyler, 2014; Clarke et al., 2015). Here, we captured the C1 and C2 responses of

224 HMax IT (hence respectively capturing low- and mid-level visual object information
225 (Riesenhuber & Poggio, 1999; Serre et al., 2007), on our images resized to 92 x 92, using the
226 same setting (i.e., Serre et al., 2005) and precomputed S2 features from natural image
227 fragments). Principal components analyses were then performed on each matrix, concatenating
228 respectively C1 and C2 features across all stimuli, and only the respectively 12 and 6 best
229 components were kept and concatenated in a single matrix of 18 visual features per stimuli. The
230 perceptual distance between two stimuli was then computed as the euclidean distance between
231 the 18-values vectors of each stimulus.

232 *Procedure*

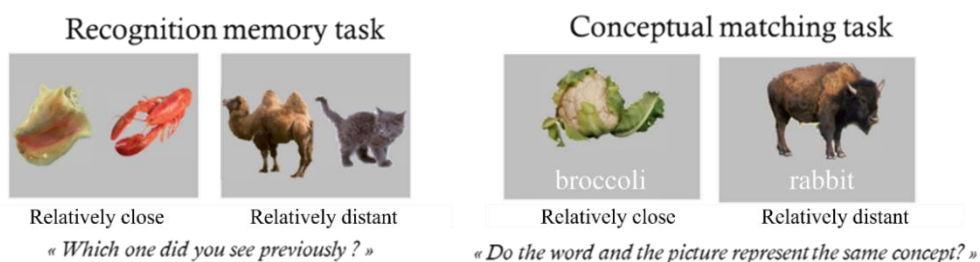
233 Participants completed three tasks: a naming task, a recognition memory task and a
234 conceptual matching task. These tasks are described in the following.

235 Naming task. 96 pictures of object concepts were presented one by one on a monitor for 3
236 seconds, preceded by a fixation cross (500 ms) and followed by a blank screen (500 ms). Half
237 of the pictures represented living things and the other half represented non-living things.
238 Participants were asked to name the object represented in the picture at a basic-level within the
239 3 seconds presentation to avoid any ceiling effects, especially in control participants.
240 Participants' verbal answers given within the 3-seconds presentation of the stimuli were
241 collected by the experimenter using a dedicated answer-sheet. The naming task served as
242 incidental encoding for the subsequent forced-choice recognition memory task that directly
243 followed.

244 Recognition memory task. 60 unique pictures (30 living & 30 non-living) from the naming task
245 were randomly selected and matched with a distractor from the same subordinate categories
246 (e.g., birds, mammals, etc.) (Figure 1). Each target-distractor pair was associated with an index
247 of their conceptual distance extracted from the feature-norms database (cosine), as well as with

248 an index of their perceptual distance. There was no difference of conceptual distance between
 249 living and non-living pairs ($M_{\text{Living Pairs}} = 0.74$ vs. $M_{\text{Non-Living Pairs}} = 0.72$, $t = 0.30$, $df = 58$, $p =$
 250 $.765$) but this was not true for perceptual distance, with non-living pairs being more perceptually
 251 distant than living pairs ($M_{\text{Living Pairs}} = 3.75$ vs. $M_{\text{Non-Living Pairs}} = 4.53$, $t = -3.24$, $df = 58$, $p = .002$).
 252 Each trial began with a fixation cross (500 ms) followed by the target-distractor pairs. Each pair
 253 was presented on the monitor for 3 seconds and ended with a 500 ms blank screen. Within the
 254 3-seconds presentation, participants were asked to indicate which of the two presented pictures
 255 was presented in the previous naming task using the right and left arrows of the keyboard.

256 Conceptual matching task. Trials for the conceptual matching task began with a fixation cross
 257 (500 ms) followed by a word-picture pair (120 trials in total) presented for 3 seconds and ended
 258 by a 500 ms blank screen. In half of the pairs, the picture matched the concept label (filler
 259 trials), while in the other half, the word and the picture did not correspond to the same concept,
 260 although the two concepts represented by the word and by the picture belonged to the same
 261 superordinate category. Cosine similarity was computed between each word-picture non-
 262 matching pair, half of the non-matching trials being livings, and the other half, non-livings.
 263 There was no difference of conceptual distance between living and non-living pairs ($M_{\text{Living Pairs}}$
 264 $= 0.34$ vs. $M_{\text{Non-Living Pairs}} = 0.35$, $t = -0.25$, $df = 118$ $p = .892$). Participants were instructed to
 265 determine whether the word referred to the same concept as the one represented in the picture
 266 or not (Figure 1). Participants answered verbally, and their answers were encoded by the
 267 experimenter using the response-keys “1” or “2”.



269 Figure 1. Example of trials from the recognition memory task (left) and of the conceptual
270 matching task (right) for living object concepts, illustrating concepts from higher vs. lower
271 conceptual distance values from the distribution of our sampled materials.

272 *MRI acquisition*

273 Images were acquired on a 3T Siemens Prisma scanner with a 64-channel head coil.
274 Two anatomical images were acquired: a T1-weighted structural MRI (acquisition matrix = 240
275 x 256 x 224, voxel size = 1 x 1 x 1 mm³) and a high-resolution T2-weighted structural MRI
276 (acquisition matrix = 448 x 448 x 60, voxel size = .4 x .4 x 1.2 mm³) with a partial field of view
277 covering the entire MTL with an oblique coronal orientation perpendicular to the long axis of
278 the hippocampus. The quality of each image was systemically visually checked, especially the
279 T2-MRI that is highly sensitive to movement (after reminding the participant to stay still during
280 the entire following 8 minutes of acquisition). When T2-MRI were acquired twice (N=14), we
281 chose the image with the best quality for further processing.

282 *Volume segmentation*

283 High-resolution T2-MRI was labeled using the Automatic Segmentation of
284 Hippocampal Subfields software package using an atlas package available online
285 ('ashs_atlas_upennpmc_20161128', from the NITRC repository made available on ASHS
286 website) (Yushkevich, Amaral, et al., 2015). This atlas package was generated from images
287 manually segmented following classical documentation for the hippocampus segmentation
288 (Adler et al., 2014; Duvernoy, 1988) and Ding and van Hoesen (2010) procedure for the ErC
289 and PrC segmentation (the landmark used by this manual protocol for the anterior extent of the
290 PrC - i.e., 2 mm anterior to the first slice of the hippocampal head – cuts off an anterior portion
291 of the PrC (see also Yushkevich et al., 2015). The hippocampus, the ErC, BA35 and BA36 in
292 the left and right hemisphere were thereby labeled in each participant. Each ASHS output was

293 visually checked for quality control. For the ErC and PrC subregions, volumes were normalized
294 by the extent of their segmentation in the slice direction (hippocampal axis), dividing their
295 volume by the product of the number of slices and the slice thickness (Yushkevich et al., 2015).
296 In addition, regional volumes were adjusted before analyses to account for total estimated
297 intracranial volume (ICV) for each participant using the formula $\text{Volume}_{\text{adjusted}} = \text{Volume}_{\text{raw}} -$
298 $\beta_{\text{ICV}}(\text{ICV}_{\text{indiv}} - \text{ICV}_{\text{mean}})$, where β refers to the regression coefficient of the model on a given
299 regional volume of interest while using ICV as predictor, based on extensive prior work (e.g.,
300 Delhaye et al., 2019, Gellersen et al., 2022, Yeung et al., 2017).

301 *Data analyses*

302 The data were analyzed with R version 4.1.2 (Team R Core, 2021). We analyzed
303 accuracy on the three behavioral tasks on a trial-by-trial basis with binomial Generalized Linear
304 Mixed Models (GLMMs) to account for the binary outcome (0, 1) of the dependent variable.
305 These models were fit with the package *lme4* (Bates et al., 2015). This model was run on
306 accuracy of the naming task with group (AD, control) as a between-subjects factor, domain
307 (living, non-living) as a within-subject factor, and CxD (to account for conceptual
308 confusability) as a continuous factor (centered scale). Additionally, participant's ID and trial
309 number were set as random factors. Another binomial GLMM was run for the recognition
310 memory task with conceptual distance and perceptual distance based on the indices of
311 perceptual and conceptual distance (both centered scale) as continuous factors. A third GLMM
312 as that for the recognition memory task was run for the matching task with the same factors,
313 with the exclusion of perceptual distance (because the pairs consisted of a word and a picture).
314 Following these GLMMs, pairwise comparisons were used with Tukey's adjustments when
315 there were multiplicity issues using the *emmeans* package (Lenth, 2020) and the function
316 *lstrends* from *lsmeans* package to deal with continuous factors; estimated marginal means

317 (EMMs) from the models are reported. Plots of the results were obtained using the *ggplot2*
318 package (Wickham, 2016) and error bars represent standard errors.

319 To investigate the recognition memory and matching tasks in relation with the integrity
320 of the sub-hippocampal regions, we used a measure of the ‘accuracy sensitivity to
321 conceptual/perceptual similarity’. To compute this measure, we adapted Wright et al. (2015)’s
322 method by correlating each participant’s accuracy to the conceptual distance value on a trial-
323 by-trial level using Pearson correlations and then transforming each Rho-value obtained for
324 each participant by a Fisher transformation to give a Z-score. We also implemented the same
325 method to compute a measure of ‘accuracy sensitivity to perceptual similarity’ using the
326 perceptual distance in the recognition memory task only. We then examined how these scores
327 were associated with eight volumes of brain regions of interest: left and right ErC, left and right
328 BA35, left and right BA36, and left and right hippocampus score (average of the volumes CA
329 fields, dentate gyrus, and subiculum) separately for AD patients and control participants using
330 multiple regressions. Importantly, these relations were controlled by the cognitive level as
331 evidenced by the score at the MoCA, which was added as a control variable in our regression
332 analyses¹. Finally, we conducted these analyses separately between living and non-living
333 stimuli for conceptual distance in AD patients only (we did not run these in control participants
334 due to ceiling effects, nor for the perceptual distance, because of the observed significant
335 difference between living and non-living stimuli in terms of this distance).

336 The data used for these analyses as well as the analytic codes are publicly available on
337 the Open Science Framework repository: <https://osf.io/r4gfy/>

¹ We also explored how the performance at the naming task could explain the relation between the brain volumes and conceptual and perceptual distance in the recognition and matching tasks. Yet, as these analyses did not reveal any significant effect of the naming performance, see Analytic Code “Supp_Analyses_NamingVar_AD.R” on <https://osf.io/r4gfy/>.

338 Results

339 Naming task

340 Regarding the accuracy on the naming task, there were main effects of group, $\chi^2 = 22.46$,
 341 $df = 1, p < .001$, and domain, $\chi^2 = 8.45, df = 1, p = .004$, but not of CxD, $p = .290$. This indicated
 342 that, overall, control participants were more accurate than AD participants ($M_{\text{control}} = .91$ vs.
 343 $M_{\text{AD}} = .78$) and accuracy was higher for non-living stimuli than for living stimuli ($M_{\text{living}} = .80$
 344 vs. $M_{\text{non-living}} = .90$). Interestingly, group and domain significantly interacted, $\chi^2 = 24.52, df = 1$,
 345 $p < .001$, revealing that whereas accuracy was similar between living and non-living stimuli for
 346 control participants ($M_{\text{living}} = .90$ vs. $M_{\text{non-living}} = .92, p = .250$), it was higher for non-living
 347 stimuli than for living stimuli in AD participants ($M_{\text{living}} = .64$ vs. $M_{\text{non-living}} = .87, p < .001$;
 348 Figure 2). Control participants showed higher accuracy than AD participants for both living and
 349 non-living stimuli, $ps < .026$. Naming was not influenced by conceptual confusability.

350 Recognition memory task

351 Regarding accuracy on the recognition memory task, the analysis revealed main effects
 352 of group, $\chi^2 = 37.35, df = 1, p < .001$, domain, $\chi^2 = 8.75, df = 1, p = .003$, and conceptual distance,
 353 $\chi^2 = 7.66, df = 1, p = .006$, but not of perceptual distance, $p = .751$. Overall, control participants
 354 were significantly more accurate than AD patients ($M_{\text{control}} = .99$ vs. $M_{\text{AD}} = .88$), accuracy for
 355 non-living stimuli was significantly higher than for living stimuli ($M_{\text{living}} = .94$ vs. $M_{\text{non-living}} =$
 356 $.97$) and the estimated marginal mean of linear trend for conceptual distance was positive (1.98)
 357 meaning that accuracy significantly increased as conceptual distance increased (Figure 2). No
 358 interactions were significant, $ps > .168$.

359 Table 2 shows the results of the multiple regression assessing the relation between
 360 accuracy sensitivity to conceptual similarity and the volume of the different brain regions within
 361 the MTL. As indicated in Table 2, only the left BA36 was significantly associated with accuracy

362 sensitivity to conceptual similarity in both AD patients, $t = 2.20$, $p = .045$, and control
363 participants, $t = 2.39$, $p = .038$. No other regions turned out to be significant in both populations,
364 $ps > .088$.

365 Regarding accuracy sensitivity to perceptual similarity, none of the regions of the MTL
366 was associated with this measure for both AD patients and control participants, $ps > .099$ (Table
367 2).

Table 2. Multiple regressions between accuracy sensitivity to conceptual and perceptual similarity and the volumes of the different brain regions of the MTL for the recognition memory task.

| Variables | Conceptual Similarity | | | | | | Perceptual Similarity | | | | | |
|----------------------------|-----------------------|--------------|--------------|--------------|--------------|--------------|-----------------------|--------|------------|-------------|--------|------------|
| | Control participants | | | AD patients | | | Control participants | | | AD patients | | |
| | β | t | p -value | β | t | p -value | B | t | p -value | β | t | p -value |
| Left BA35 | -0.010 | -0.574 | 0.579 | -0.017 | -1.506 | 0.154 | -0.034 | -1.818 | 0.099 | -0.0003 | -0.032 | 0.975 |
| Right BA35 | -0.008 | -0.916 | 0.381 | 0.005 | 0.492 | 0.630 | -0.006 | -0.624 | 0.546 | -0.007 | -0.762 | 0.459 |
| Left BA36 | 0.008 | 2.392 | 0.038 | 0.006 | 2.204 | 0.045 | 0.001 | 0.415 | 0.687 | 0.0002 | 0.082 | 0.936 |
| Right BA36 | -0.002 | -0.586 | 0.571 | -0.001 | -0.404 | 0.693 | 0.003 | 1.218 | 0.251 | 0.003 | 1.140 | 0.272 |
| Left ERC | 0.008 | 0.743 | 0.474 | -0.005 | -0.379 | 0.7103 | -0.0003 | -0.025 | 0.980 | 0.013 | 1.366 | 0.194 |
| Right ERC | -0.030 | -1.888 | 0.088 | 0.008 | 0.738 | 0.473 | 0.010 | 0.603 | 0.560 | 0.005 | 0.560 | 0.585 |
| Left Hippocampus Score | 0.010 | 1.225 | 0.249 | -0.008 | -1.638 | 0.124 | 0.007 | 0.673 | 0.516 | -0.002 | -0.551 | 0.590 |
| Right Hippocampus Score | -0.002 | -0.313 | 0.761 | 0.006 | 1.257 | 0.229 | -0.002 | -0.363 | 0.724 | 0.002 | 0.408 | 0.689 |
| MoCA | -0.0003 | -0.019 | 0.985 | 0.002 | 0.146 | 0.886 | 0.024 | 1.172 | 0.268 | -0.003 | -0.306 | 0.764 |

Conceptual matching task

This analysis revealed a main effect of group, $\chi^2 = 29.84$, $df = 1$, $p < .001$, domain, $\chi^2 = 5.45$, $df = 1$, $p = .020$, and conceptual distance, $\chi^2 = 34.69$, $df = 1$, $p < .001$. Overall, control participants showed significantly higher accuracy than AD participants ($M_{\text{control}} = .97$ vs. $M_{\text{AD}} = .87$), accuracy for non-living stimuli was significantly higher than for living stimuli ($M_{\text{living}} = .90$ vs. $M_{\text{non-living}} = .96$) and the direction for conceptual distance was positive (estimated marginal mean of linear trend = 4.32), revealing that accuracy significantly increased as conceptual distance increased (Figure 2). No interactions were significant, $ps > .083$.

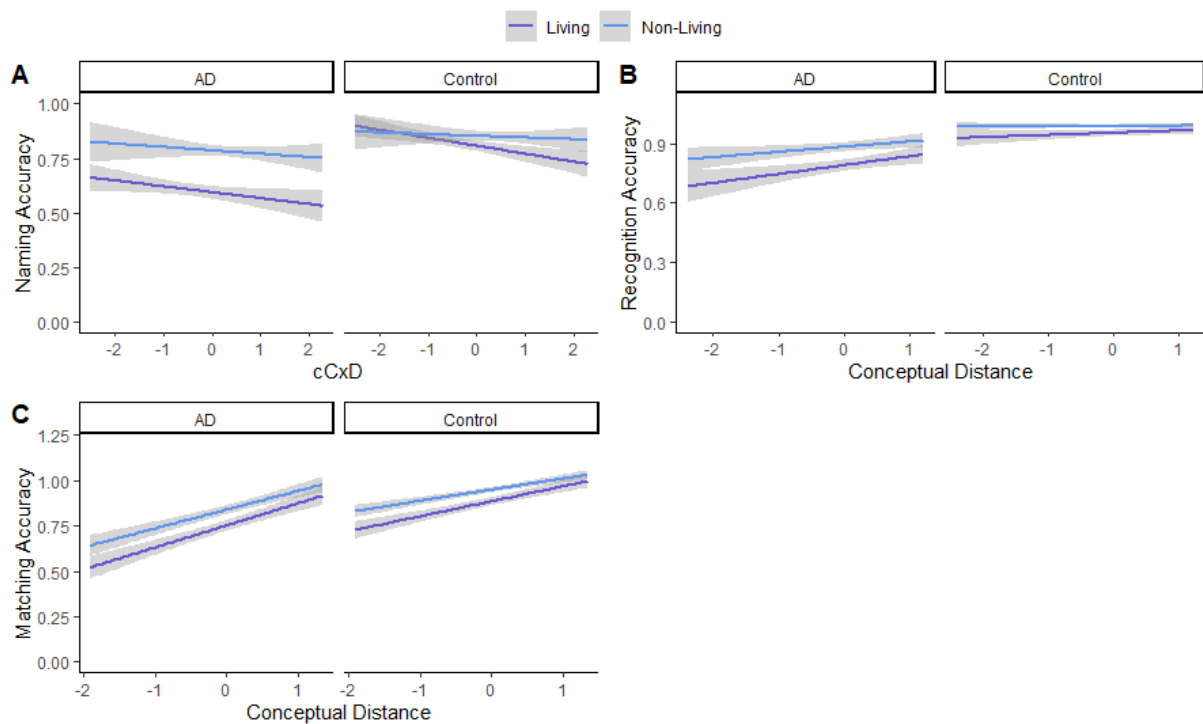


Figure 2. Naming accuracy as function of group, domain and CxD (A), recognition memory accuracy (B) and conceptual matching accuracy (C) as a function of group, domain and conceptual distance.

Table 3 indicates the results of the multiple regression analyses regarding the relation between accuracy sensitivity to conceptual similarity and the volumes of the different brain regions within the MTL. For AD patients, the analysis yielded significant effects on the left BA36

(PrC), $t = -2.55$, $p = .023$, but not on other regions, $ps > .059$. For control participants, none of the brain volumes were significantly associated with this measure, $ps > .056$.

Table 3. Multiple regressions between accuracy sensitivity to conceptual similarity and the volume of the different brain regions of the MTL in the conceptual matching task

| Variables | Control participants | | | AD patients | | |
|-------------------------|----------------------|--------|------------|---------------|---------------|--------------|
| | β | t | p -value | β | t | p -value |
| Left BA35 | 0.007 | 0.375 | 0.714 | 0.024 | 2.052 | 0.059 |
| Right BA35 | -0.016 | -1.206 | 0.249 | 0.021 | 1.896 | 0.079 |
| Left BA36 | -0.0002 | -0.041 | 0.968 | -0.007 | -2.554 | 0.023 |
| Right BA36 | 0.005 | 1.292 | 0.219 | -0.005 | -1.435 | 0.173 |
| Left ERC | -0.003 | -0.210 | 0.837 | -0.015 | -1.297 | 0.215 |
| Right ERC | 0.018 | 1.024 | 0.324 | -0.014 | -1.301 | 0.214 |
| Left Hippocampus Score | 0.019 | 1.558 | 0.143 | 0.003 | 0.552 | 0.590 |
| Right Hippocampus Score | -0.020 | -2.102 | 0.056 | 0.002 | 0.498 | 0.626 |
| MoCA | 0.013 | 0.478 | 0.640 | 0.005 | 0.351 | 0.731 |

Living vs. non-living items in the recognition and matching tasks

We applied the same regression analyses to investigate the relation between accuracy sensitivity to conceptual similarity and the brain volumes of the MTL and doing so by considering living and non-living items separately for AD patients in both the recognition memory and conceptual matching tasks (Table 4). For the recognition task, these analyses revealed that the right PrC was significantly associated with the accuracy sensitivity measure for non-living items (right BA35: $t = 3.85$, $p = .006$; right BA36: $t = -2.65$, $p = .033$) but not other regions, $ps > .063$. For living items, no significant associations between the accuracy sensitivity measure and any brain regions were significant, $ps > .082$.

Regarding the conceptual matching task, no significant associations were found for living items, $ps > .067$, whereas for non-living items, both the left and right PrC and ErC were associated with the measure of accuracy sensitivity to conceptual similarity (right BA35: $t = 3.65$, $p = .003$; left BA35: $t = 2.30$, $p = .037$; right BA36: $t = -2.24$, $p = .042$; left BA36: $t = -2.38$, $p = .032$; right ERC: $t = -2.26$, $p = .040$; left ERC: $t = 2.17$, $p = .047$). The right and left hippocampus regions were not significantly associated with the accuracy sensitivity measure, $ps > .319$.

Table 4. Multiple regressions between accuracy sensitivity to conceptual similarity and the volumes of the different brain regions of the MTL for living and non-living stimuli in AD patients in the recognition memory task

| Variables | Recognition Task | | | | | | Matching Task | | | | | |
|----------------------------|------------------|--------|------------|---------------|---------------|--------------|---------------|--------|------------|---------------|---------------|--------------|
| | Living | | | Non-Living | | | Living | | | Non-Living | | |
| | β | t | p -value | β | t | p -value | B | t | p -value | β | t | p -value |
| Left BA35 | -0.020 | -1.147 | 0.271 | -0.027 | -2.206 | 0.063 | 0.023 | 1.333 | 0.204 | 0.026 | 2.299 | 0.037 |
| Right BA35 | -0.009 | -0.589 | 0.565 | 0.051 | 3.849 | 0.006 | 0.007 | 0.451 | 0.659 | 0.040 | 3.650 | 0.003 |
| Left BA36 | 0.007 | 1.875 | 0.082 | 0.007 | 2.125 | 0.071 | -0.008 | -1.982 | 0.067 | -0.006 | -2.377 | 0.032 |
| Right BA36 | 0.002 | 0.417 | 0.683 | -0.010 | -2.650 | 0.033 | -0.003 | -0.521 | 0.610 | -0.007 | -2.243 | 0.042 |
| Left ERC | -0.005 | -0.287 | 0.778 | -0.06 | -0.445 | 0.670 | -0.008 | -0.430 | 0.673 | -0.025 | -2.173 | 0.047 |
| Right ERC | 0.013 | 0.882 | 0.419 | 0.014 | 1.227 | 0.259 | 0.005 | -0.314 | 0.758 | -0.024 | -2.259 | 0.040 |
| Left Hippocampus Score | -0.007 | -0.947 | 0.360 | -0.007 | -0.987 | 0.356 | 0.003 | 0.377 | 0.712 | 0.003 | 0.573 | 0.576 |
| Right Hippocampus Score | 0.008 | 1.090 | 0.294 | -0.004 | -0.519 | 0.620 | -0.0001 | -0.017 | 0.986 | 0.005 | 1.033 | 0.319 |
| MoCA | -0.001 | -0.056 | 0.956 | 0.021 | 1.440 | 0.1930 | 0.016 | 0.795 | 0.440 | -0.008 | -0.570 | 0.578 |

Comparison between mild and moderate AD patients

In the light of the results obtained in the previous sections, we conducted exploratory analyses to examine the idea that the matching task might be particularly relevant to track PrC atrophy due to the AD neuropathology, for instance in individuals at risk to develop AD. We ran further analyses on conceptual confusability for the recognition memory and matching tasks by splitting the AD patients tested in our study into two groups based on the median of their scores at the MoCA. These low MoCA patients and high MoCA patients were matched in terms of age and education (for demographic information about these two groups, see Table S1 in the Supplementary Material). These analyses revealed that the only significant association was between the conceptual matching task and the volume of left BA36 of the AD patients with a low MoCA score ($t = -2.60$, $p = .048$). Other associations failed to reach significance, $ps > .069$ (Table 5).

Table 5. Multiple regressions between accuracy sensitivity to conceptual similarity and the volume of the different brain regions of the MTL for AD patients with low scores at the MoCA (≤ 22 ; Low MoCA Group) and with high scores at the MoCA (> 22 ; High MoCA Group) in the recognition memory task and the conceptual matching task.

| Variables | Recognition Memory | | | | | | Conceptual Matching | | | | | |
|----------------------------|--------------------|--------|------------|--------------------|--------|------------|---------------------|---------------|--------------|--------------------|--------|------------|
| | Low MoCA patients | | | High MoCA patients | | | Low MoCA patients | | | High MoCA patients | | |
| | β | t | p -value | β | t | p -value | B | t | p -value | β | t | p -value |
| Left BA35 | -0.042 | -2.308 | 0.069 | -0.042 | -2.308 | 0.069 | 0.031 | 1.539 | 0.184 | -0.003 | -0.179 | 0.888 |
| Right BA35 | 0.0002 | 0.012 | 0.991 | 0.0002 | 0.012 | 0.991 | 0.011 | 0.717 | 0.506 | -0.063 | -3.094 | 0.199 |
| Left BA36 | 0.006 | 1.659 | 0.158 | 0.006 | 1.659 | 0.158 | -0.011 | -2.597 | 0.048 | 0.015 | 3.503 | 0.972 |
| Right BA36 | -0.001 | -0.353 | 0.738 | -0.001 | -0.353 | 0.738 | -0.004 | -0.829 | 0.445 | 0.005 | 0.558 | 0.676 |
| Left ERC | 0.006 | 0.447 | 0.673 | 0.006 | 0.447 | 0.673 | -0.025 | -1.665 | 0.156 | 0.066 | 2.814 | 0.217 |
| Right ERC | -0.003 | -0.182 | 0.863 | -0.003 | -0.182 | 0.863 | -0.033 | -1.888 | 0.117 | -0.010 | -0.984 | 0.505 |
| Left Hippocampus Score | -0.014 | -1.686 | 0.153 | -0.014 | -1.686 | 0.153 | -0.0001 | -0.015 | 0.989 | -0.059 | -4.699 | 0.133 |
| Right Hippocampus Score | 0.019 | 1.733 | 0.144 | 0.019 | 1.733 | 0.144 | 0.015 | 1.204 | 0.282 | 0.014 | 1.489 | 0.376 |

Discussion

In the present study, we investigated fine-grained episodic and semantic discriminations of both perceptually (for episodic memory only) and conceptually confusable objects and their associations with the integrity of the brain structures of the MTL in AD patients and control counterparts, with the aim to improve our understanding of the role of the PrC region in cognition. More specifically, we used a quantitative measure to capture the structural conceptual and perceptual confusability of objects and their relation to performance to provide refined examinations of how volumes from the MTL structures are associated with finer-grained discrimination in AD (we called this score accuracy sensitivity to conceptual similarity).

First, behavioral results from the naming task showed that AD patients named less object concepts than control participants, but more particularly, that patients had difficulties for naming living stimuli as compared to non-living stimuli, regardless of their confusability, whereas no such difference was found for healthy volunteers. This result is in line with previous reports showing that AD patients experience word-finding difficulties and produce naming errors (e.g., ‘hippopotamus’ for ‘rhinoceros’) to a larger extent than healthy individuals, and that these difficulties seem to be especially important for living stimuli (see Laws et al., 2007). Interestingly, when subsequently asked to recognize the previously seen objects on a forced-choice recognition memory task, although AD patients showed poorer recognition memory performance than control participants (e.g., Goldstein et al., 2019), both populations showed lower recognition accuracy for living stimuli than for non-living stimuli, and for both groups and across domains, accuracy was particularly lower when the pairs of items were relatively conceptually close (e.g., seashell and crayfish) than when the pairs showed items that are conceptually distant (e.g., cat and camel). These results are partially consistent with the previous findings showing that AD patients’ recognition memory is poorer for living than for

non-living stimuli (Kivisaari et al., 2013). Yet, contrary to this study, we found that poorer recognition of living stimuli as compared to non-living stimuli also holds for control participants, and that poorer memory in case of high conceptual confusability occurred regardless of the domain. One possible explanation for this difference could be related to the fact that there were 96 stimuli in our naming (and encoding) task whereas in Kivisaari et al. (2013), there were only 60 stimuli at encoding, therefore potentially making our encoding task costlier. Moreover, in our task, participants had to give their responses rapidly (3 seconds, see Methods), which measures a rapid access to concept, contrary to Kivisaari et al. (2013) who did not use such timing constraints in their design. In addition, this effect of conceptual confusability on memory in healthy subjects is relatively consistent with a study by Montefinese et al. (2015) showing that conceptual proximity between memory targets and lures, calculated using feature norms, induces an increase in false alarm rates, even in young subjects. However, controls' performance on this task is close to ceiling, limiting the variability in the dataset and potentially hindering statistical effects, so we are cautious about any further interpretations. In addition, a similar pattern was observed for the conceptual matching task.

At the brain level, our analyses focused on the accuracy sensitivity to similarity score (i.e., correlating accuracy with the conceptual or perceptual distance at the trial level for each subject) adapted from Wright et al. (2015). This method allows for finer conceptualization of conceptual confusability (see also Taylor et al., 2012) than considering objects' confusability as reflected by their domain, either living or non-living, hence using two discrete categories. Using this measure, we found that the left PrC (BA 36) was associated with greater accuracy sensitivity to conceptual similarity in both AD patients and control participants in the recognition memory task, and in AD patients only in the conceptual matching task, with a trend of an association with left BA 35 as well. Altogether, these results support the idea of a hemispheric specialization of the PrC, in accordance with previous findings on the importance

of the left PrC in fine-grained disambiguation for highly confusable objects specifically, relative to less confusable ones (see Bruffaerts et al. 2019 for review; Bruffaerts et al., 2013; Clarke & Tyler, 2014; Duke et al., 2017).

In addition, when disentangling stimuli as livings vs non-livings (Table 4) and investigating the association between our measure of accuracy sensitivity to conceptual similarity and the different brain regions in AD patients, we observed strong relations between the volume of the perirhinal and entorhinal cortices, with some interesting lateralization differences depending on the task. For the recognition task, the right PrC (BA 35 and BA 36) was associated with the measure of accuracy sensitivity for non-living items while, for living items, only the left PrC (BA 36) was marginally associated with this measure. In the matching task, on the other hand, both the left and right PrC (BA 35 and BA 36) and ErC were associated with accuracy sensitivity for non-living items whereas only the left PrC was marginally associated with performance for living items. The association between the ability to disambiguate non-living close concepts and the integrity of the right PrC might be explained by the coarse activation hypothesis (Jung-Beeman, 2005), which argues that semantic processing is coarser in the right than in the left hemisphere. Indeed, non-living stimuli are generally less complex than living stimuli in terms of their conceptual structure, and this might explain why distinguishing highly conceptually similar non-living concepts was more associated with the right PrC in AD patients. As for the left PrC, while it was also involved in the disambiguation of non-living items for the matching task (see also Liuzzi et al., 2019), it was specifically marginally associated with conceptual discrimination of more confusable items (i.e., living) in both tasks, fitting with the idea of a specific left-lateralized involvement of the PrC in fine-grained conceptual disambiguation (see Bruffaerts et al., 2019). Yet, more research is needed at this stage to unravel any potential lateralization of the PrC based on the type of items that have to be disambiguated and the type of the task.

Interestingly, our results showed correlations in both patients and controls in the recognition memory task, while only in patients in the conceptual matching task. We did not have expectations as to observing correlations in our control group, with the reasoning that controls' variability in PrC integrity would come from a diverse set of factors not necessarily affecting simultaneously volumes and cognitive scores. These correlations in controls suggest that our measure of sensitivity to conceptual confusability in recognition memory is highly sensitive to variations in PrC volume, and not only to variations due to AD neuropathology, probably due to the episodic nature of the task, known to favor the use of familiarity (Bastin & Vander Linden, 2003), that is also highly reliant on the PrC. As for the matching task, the association between the PrC and accuracy sensitivity to conceptual similarity did not hold for control participants. We interpret this discrepancy in control participants in terms of the nature of the tasks. Indeed, the recognition memory task is tapping at the interface of episodic and semantic processes, leaving room for some variability, especially coming from episodic memory. Conversely, the matching task is purely semantic, where it is not expected that healthy controls would show variability as semantic memory, if anything, improves in aging (Lalla et al., 2022). [In line with this idea, we observed that, when splitting the sample according to the MoCA scores, there were non-significant associations between the brain volumes and the accuracy sensitivity to conceptual similarity for the recognition memory task](#) whereas in the matching task, the left PrC (BA 36) was associated with this measure only for patients having lower scores at the MoCA. Therefore, it seems that the matching task might not be used as an early marker of PrC atrophy in the course of AD, although future studies using a sample of MCI patients would best allow to answer this question given the statistical limits of median split analyses (DeCoster et al., 2011), and the resulting small sample size for group comparisons and correlational analyses, that call for caution in their interpretation.

Concerning the recognition memory task, the absence of relation between our measure of sensitivity to perceptual similarity and the PrC was surprising in the light of the extensive literature showing PrC involvement in fine-grained perceptual discrimination (e.g., Inhoff et al., 2019; but see Gellersen et al., 2022). A possible explanation for this result is that the HMax model we used captures visual features from low- and mid-level visual information, which might not finely reflect human perceived similarity. Previous studies have indeed shown that models based on objects conceptual structure are better predictors of neural activity patterns associated with individual objects than the HMax model (Clarke et al., 2015), as the HMax model does not reflect abstract object information that is not directly related to the visual input, such as semantic domain (livings vs. non-livings) that has been used previously to characterize similarity in association with PrC integrity in healthy and pathological aging (Kivisaari et al., 2012, 2013).

Despite the important new findings evidenced in this study, it has nevertheless some limitations. First, it is now clear that brain volumes are not the most sensitive measure of the presence of AD in the brain, and more refined biomarkers exist to track the presence of AD neuropathology (Jack et al., 2018). Second, we used brain volumes as a proxy of brain function, although we reckon that functional alterations are not linearly linked with structural integrity changes (Jack et al., 2013). Illustrating these two limitations, it was convincingly shown that changes to the functional connectivity between MTL sub-regions related to tau pathology are associated with cognitive behavioral measures similar to ours in the absence of structural damage in cognitively unimpaired older adults (Berron et al., 2019). This limitation could thus explain why here, the brain volumes are not associated with cognitive performance for AD patients with higher scores at the MoCA as compared to those with lower scores (Table 5). Yet, our study aligns with a now extensive body of research using this correlational approach, all pointing to a clear association between regional volumes in the transentorhinal region and key

cognitive functions such as conceptual and perceptual discriminations, not only in AD, but also in MCI (Delhaye et al., 2019) and in at-risk older adults without complaints (Gellersen et al., 2022, Olsen et al., 2017, Yeung et al., 2017, 2019). In addition, the cross-sectional design employed might not be best to deal with the high variability of the neuro-cognitive profiles of AD patients, which could weaken the associations we investigated here (for a similar point albeit in healthy aging, see Armstrong et al., 2020). Future studies should use a longitudinal design to confirm cross-sectional findings that start to accumulate on the role of perirhinal shrinkage in the ability to disambiguate highly confusable objects across different tasks. Another limitation concerns the fact that our observed group differences for the associations between brain volumes and cognitive measures are based on significant and non-significant effects resulting from different statistical models. We acknowledge that conclusions could be stronger if the statistical design allows for the examination of group interactions in a single model, and if behavioral effects were characterized by significant group interactions, which was not the case.

To conclude, the present study reports that across different tasks, namely a recognition memory task and an item-matching task, the volume of the left PrC accounts for difficulties in distinction between highly confusable objects, supporting existing evidence on the role of the left PrC in fine-grained conceptual disambiguation of confusable objects, here using refined measures to quantify conceptual confusability.

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Supplemental Material

Table S1. Demographic and brain volumes comparisons (t-tests) between AD patients with Lower MoCA scores (n = 14) and AD patients with Higher MoCA scores (n = 10).

| | Low MoCA Mean (SD) | High MoCA Mean (SD) | <i>p</i> -value |
|---|-----------------------|------------------------|-----------------|
| Female/Male | 7/7 | 3/7 | |
| Age | 75.50 (6.41) | 73.80 (6.34) | .526 |
| Education | 12.14 (3.63) | 12.90 (2.18) | .564 |
| MoCA | 19.00 (2.63) | 23.80 (0.79) | < .001 |
| Letter fluency | 15.00 (7.30) | 18.10 (7.43) | .320 |
| Category fluency | 15.93 (6.04) | 22.40 (6.26) | .018 |
| WAIS-3 digit symbol substitution | 17.29 (15.56) | 15.10 (13.58) | .724 |
| WAIS-3 vocabulary | 30.45 (7.84) | 33.56 (10.65) | .463 |
| WMS-3 Logical Memory immediate recall | 6.71 (2.64) | 9.70 (3.74) | .032 |
| Left ERC | 20.85 (3.94) | 22.79 (3.18) | .214 |
| Right ERC | 18.81 (4.93) | 20.00 (4.39) | .549 |
| Left BA 35 | 15.41 (3.87) | 19.28 (4.81) | .040 |
| Right BA 35 | 17.48 (4.79) | 16.51 (4.42) | .620 |
| Left BA 36 | 59.93 (14.98) | 65.42 (16.11) | .395 |
| Right BA 36 | 60.09 (17.12) | 55.29 (13.02) | .464 |
| Left Hippocampus Score | 67.22 (10.26) | 71.96 (7.63) | .231 |
| Right Hippocampus Score | 68.66 (12.12) | 71.90 (10.12) | .498 |
| | | | |