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Flexible coding of object motion in multiple reference frames by parietal cortex neurons

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22 **ABSTRACT**

23 Neurons represent spatial information in diverse reference frames, but it remains unclear
24 whether neural reference frames change with task demands and whether these changes can
25 account for behavior. We examined how neurons represent the direction of a moving object
26 during self-motion, while monkeys switched, from trial to trial, between reporting object
27 direction in head- and world-centered reference frames. Self-motion information is needed to
28 compute object motion in world coordinates, but should be ignored when judging object motion
29 in head coordinates. Neural responses in the ventral intraparietal area are modulated by the task
30 reference frame, such that population activity represents object direction in either reference
31 frame. In contrast, responses in the lateral portion of the medial superior temporal area primarily
32 represent object motion in head coordinates. Our findings demonstrate a neural representation of
33 object motion that changes with task requirements.

34

35

36 **INTRODUCTION**

37 Sensory signals are encoded in modality-specific reference frames at the sensory periphery,
38 such as an eye-centered reference frame for visual signals and a head-centered reference frame
39 for vestibular signals. In downstream brain areas, signals are often transformed into non-native
40 reference frames, including intermediate or mixed reference frames¹⁻¹³, and it is generally
41 presumed that different reference frames are useful for guiding different behaviors.

42 Natural behavior is flexible, however, and may require the observer to interpret the same
43 sensory signals in different reference frames, depending on task context (e.g., a soccer player
44 might judge motion of the ball relative to their head or relative to the goal). Having neural
45 representations that flexibly adapt to task demands may thereby simplify sensorimotor
46 transformations. While task-switching has been studied extensively with behavior and
47 neuroimaging^{14,15}, and single-neuron correlates of switching task sets have been reported^{16,17},
48 little is known about whether the reference frame of neural representations changes dynamically
49 when judgements are made in different reference frames. When the reference frame required for
50 a task changes, do neural representations change accordingly or are neural reference frames
51 fixed?

52 Perception of object motion during self-motion provides an attractive model system for
53 studying this issue. When an observer is stationary, there is a unique mapping of object motion in
54 the world to image motion on the retina. However, when a translating observer views the same
55 moving object, image motion also depends on self-motion (Fig. 1). To judge object motion
56 relative to their head (e.g., a soccer player who wants to head the ball), one can rely on retinal
57 image motion, which is equivalent to motion in a head-centered reference frame if the eyes do
58 not move. However, to judge object motion relative to the world (e.g., a soccer goalie trying to

59 judge whether a shot will be on goal), one needs to compensate for the visual consequences of
60 self-motion. How neural circuits incorporate information about self-motion to represent object
61 motion in the world is not well understood^{18, 19}. Moreover, it is not known whether neural
62 representations are dynamically updated to represent object motion in head or world coordinates
63 based on task instructions.

64 Numerous psychophysical studies have examined how perception of object motion discounts
65 image motion caused by self-motion, and have identified visual mechanisms that attempt to
66 isolate image motion produced by independent object motion²⁰⁻²⁴. Vestibular signals also aid in
67 dissociating the components of retinal image motion due to object motion and self-motion²⁴⁻²⁶.
68 Thus, we hypothesize that vestibular input may contribute to generating flexible representations
69 of object motion that update with task reference frame.

70 We trained macaque monkeys to discriminate object direction in either a world-centered or a
71 head-centered reference frame. Monkeys successfully switched between reference frames,
72 randomly across trials, and their performance was enhanced when both visual and vestibular self-
73 motion signals were available. We recorded neural activity from the lateral subdivision of the
74 medial superior temporal (MSTl) area, which has been suggested to play a role in coding object
75 motion^{27, 28}. MSTl neurons combine retinal image motion with extra-retinal signals related to
76 eye and head rotation²⁹, suggesting that MSTl might be a viable candidate for representing
77 direction of object motion in different reference frames. We also recorded neural activity from
78 the ventral intraparietal (VIP) area, which is well known for its roles in representing visual
79 motion, as well as for carrying multisensory representations of visual, vestibular, tactile, and
80 auditory signals in diverse reference frames^{2, 4, 11, 30, 31}. Thus, VIP is a good candidate to flexibly
81 represent object motion in a head- or world-centered reference frame.

82 We find that responses of individual VIP neurons are modulated by the task reference frame,
83 such that their tuning shifts toward world coordinates when the task requires a world-centered
84 reference frame. In contrast, MSTl neurons do not show this effect. At the population level,
85 linear decoding of VIP activity accounts nicely for behavioral effects, whereas decoding of MSTl
86 activity does not. Strikingly, a single set of decoding weights can accurately classify object
87 direction in either head or world coordinates based on VIP activity, but not MSTl activity. The
88 time course of the reference frame transformation in VIP is delayed relative to onset of visual
89 responses. Together, our findings demonstrate that VIP flexibly represents object motion in
90 different reference frames depending on task instructions, with self-motion signals being
91 incorporated into the computation when needed. More generally, our results provide striking
92 evidence that the reference frames of neural representations can be highly dynamic, and that the
93 same neural populations can carry information in different reference frames from moment to
94 moment.

95

96 **RESULTS**

97 We trained two macaque monkeys to report whether an object moves upward and rightward
98 or upward and leftward during lateral self-motion (Fig. 2a). In the world coordinate task,
99 monkeys judged object motion relative to vertical in a world-centered reference frame; in the
100 head coordinate task, animals reported object motion relative to a head-centered vertical
101 reference. Thus, for some stimulus conditions, the same patterns of random-dot motion on the
102 retina could give rise to opposite perceptual reports in the two different reference frames. In each
103 trial, the color and shape of the fixation point instructed the reference frame to be used (Fig. 2b),
104 and the two reference frame conditions were randomly interleaved. Self-motion information was

105 provided by optic flow of background dots or by a congruent combination of optic flow and
106 physical translation of the animal on a motion platform. A partial cube frame that faded out
107 during object motion helped to instruct the task reference frame (Fig. 2c). If monkeys could
108 switch perfectly between task reference frames, psychometric functions for the two directions of
109 self-motion should overlap in the world coordinate task (Fig. 2d). In contrast, for the head
110 coordinate task, psychometric functions for the two opposite directions of self-motion should be
111 shifted by a predictable amount ($\Delta\text{PSE} = 35.5$ deg, see Methods).

112

113 **Monkeys can switch between world- and head-centered reference frames while judging** 114 **object motion**

115 To summarize behavior, we computed average psychometric functions across all recording
116 sessions from the two monkeys (Fig. 3a). In the absence of self-motion (Object Only condition),
117 monkeys reported object direction with very little bias (Fig. 3a, black curve). During self-motion,
118 monkeys also performed quite accurately in the head coordinate task, as evidenced by
119 psychometric functions that were shifted by approximately the predicted amount (Fig. 3a, blue
120 and cyan curves). In the world coordinate task, psychometric functions for the two directions of
121 self-motion showed much smaller shifts, indicating that animals largely compensated for the
122 effect of their self-motion on object motion (Fig. 3a, magenta and brown curves). In the
123 Object+Combined condition, this compensation was nearly complete (magenta), whereas
124 compensation was substantially less complete in the Object+Visual condition (brown). This
125 finding demonstrates that vestibular signals enhance the monkeys' ability to judge object
126 direction in a world-centered reference frame. This effect presumably arises because adding
127 vestibular signals provides a more accurate and precise estimate of self-motion velocity, rather

128 than facilitating switching between reference frames per se. However, we cannot differentiate
129 these possibilities.

130 To quantify these effects, we fit psychometric functions with cumulative Gaussian curves
131 and measured the point of subjective equality (PSE) for each self-motion direction, reference
132 frame, and recording session. Then, we computed the difference in PSE (Δ PSE) between
133 leftward and rightward self-motion directions and compared the results between Object+Visual
134 and Object+Combined conditions (Fig. 3b). For the head coordinate task, mean Δ PSE values for
135 both conditions (Fig. 3b, black triangle) are close to predicted values, although both are 10-15%
136 greater than expected (two-tailed t-test, Object+Visual: $t(184)=2.03$, $p=0.044$;
137 Object+Combined: $t(184)=6.05$, $p=8.0\times 10^{-9}$). Moreover, there is no substantive difference
138 between mean Δ PSE values for Object+Visual and Object+Combined conditions in the head
139 coordinate task (two-tailed paired t-test, $t(368)=-0.71$, $p=0.47$). This is expected because self-
140 motion information is not needed to perform the head coordinate task.

141 The pattern of results is strikingly different for the world coordinate task, where Δ PSE
142 values are much smaller. The mean Δ PSE value in the Object+Visual condition is small (9.4 deg)
143 but consistently greater than zero ($t(184)=13.9$, $p=1.3\times 10^{-50}$), whereas the corresponding value
144 (0.75 deg) for the Object+Combined condition is not substantially different from zero (two-tailed
145 t-test, $t(184)=1.34$, $p=0.054$). The difference in Δ PSE values between Object+Visual and
146 Object+Combined conditions is quite robust across sessions in the world coordinate task (two-
147 tailed paired t-test, $t(368)=11.9$, $p=1.7\times 10^{-35}$), unlike the head coordinate task. These behavioral
148 results demonstrate that monkeys successfully switch between world and head reference frames
149 from trial to trial, and that vestibular signals facilitate this transformation.

150 Note that the slope of psychometric functions in Fig. 3a depends on the presence of self-
151 motion. Additional analysis showed that object motion discrimination thresholds are
152 significantly lower in the Object Only condition than in the conditions with self-motion, with
153 only modest differences between Object+Visual and Object+Combined conditions (Extended
154 Data Fig. 1).

155

156 **Effects of task reference frame on single-unit responses in VIP and MSTl**

157 We next investigated whether the activity of VIP and MSTl neurons was modulated by task
158 reference frame. We recorded from 223 VIP neurons and 177 MSTl neurons that met basic
159 inclusion criteria (see Methods). In general, responses of neurons in both areas are influenced by
160 both object motion and self-motion directions (Fig. 4, filled vs. open symbols). This is not
161 surprising since motion stimuli were transparent such that both object and self-motion vectors
162 impinged upon the receptive fields of the recorded populations (Extended Data Fig. 2). The key
163 question is whether responses depend on the instructed reference frame for the same object and
164 background motions, which can be assessed by comparing responses in the world coordinate task
165 with responses in the head coordinate task. Differences would suggest a representation of object
166 motion that changes with task demands. Indeed, the two example VIP neurons in Fig. 4a show
167 clear response differences between task reference frames in the Object+Combined condition
168 (magenta vs. cyan). In contrast, for the two example MSTl neurons in Fig. 4b, responses are
169 much more similar between the two task reference frames. These examples suggest that VIP
170 responses are more strongly modulated by task reference frame than MSTl responses. Data from
171 the Object+Visual condition for these same example neurons are shown in Fig. 4c,d; data from
172 additional example neurons are shown in Extended Data Fig. 3.

173 To quantify response modulations related to task reference frame (without assuming a
174 functional form of those modulations), we first computed a modulation index (MI) that captures
175 the net response difference between head and world coordinate tasks (see Methods): larger
176 values of MI indicate greater response differences between task reference frames. We find that
177 MI values are significantly greater for VIP than MSTl neurons (Fig. 5a; Wilcoxon signed rank
178 test, Object+Visual condition: $Z=4.10$, $p=4.1\times 10^{-5}$; Object+Combined: $Z=4.46$, $p=8.2\times 10^{-6}$).
179 This suggests that the representation of object motion in VIP is more dependent on task
180 requirements than that in MSTl.

181 To characterize the temporal dynamics of response modulations related to task reference
182 frame, we computed MI values within a 300 ms sliding window that was shifted in increments of
183 50 ms. While MI values are greater in VIP than MSTl at almost all time points (Fig. 5b), they
184 grow substantially over time in both areas. Comparison of the time course of MI (Fig. 5b) with
185 the time course of population responses to the most effective object direction for each neuron
186 (Extended Data Fig. 4a) reveals that response modulations related to task reference frame arise
187 later than stimulus-driven modulations in both MSTl and VIP.

188 Differences in receptive field sizes and/or locations between brain areas could potentially
189 confound interpretation of the MI data. To evaluate this possibility, we performed an analysis of
190 covariance (ANCOVA) to test whether the difference in MI between MSTl and VIP was robust
191 to including receptive field size and eccentricity as covariates. We found a significant main
192 effect of brain area ($F(1, 201)= 5.0$, $p = 0.026$), with no significant dependence on receptive field
193 size ($F(3, 201)= 0.7$, $p = 0.55$) or eccentricity ($F(3, 201)= 1.85$, $p = 0.14$).

194 While the MI data of Fig. 5a,b suggest greater task dependency of responses to object
195 motion in VIP, MI is sensitive to any differences in response between tasks, and does not

196 necessarily reflect a shift in the neural representation toward world coordinates in the world task
197 condition. To assess this, we computed a (Pearson) correlation coefficient between tuning in the
198 Object Only condition and tuning in the conditions with self-motion. These correlations were
199 computed among tuning curves expressed in world coordinates (e.g., bottom row of Fig. 4), such
200 that alignment of the tuning curves in world coordinates would yield a large positive correlation
201 coefficient. This analysis was performed by pooling data across the Object+Visual and
202 Object+Combined conditions to gain statistical power, but results were very similar when
203 computed for the Object+Visual and Object+Combined conditions separately (not shown).
204 Across the populations of VIP and MSTd neurons, tuning correlations cover a broad range of
205 values (Fig. 5c), indicating that object tuning is not generally expressed in world coordinates.
206 Our data did not allow us to compute the correlation between tuning curves in head coordinates,
207 since there was insufficient overlap of object directions across self-motion directions when
208 expressed in head coordinates (e.g., top row of Fig. 4).

209 Critically, our design allowed us to test the hypothesis that correlation among tuning curves
210 in world coordinates becomes stronger in the world coordinate task as compared to the head
211 coordinate task. Indeed, for VIP, we find a robust increase in this correlation for the world
212 coordinate task (Fig. 5c, orange), consistent with a shift in neural tuning toward world
213 coordinates (Wilcoxon signed-rank test, $Z=-4.45$, $p=8.4\times 10^{-6}$). The time course of this
214 correlation reveals that the shift toward world coordinates begins around 1000ms (Fig. 5d), just
215 as stimulus-driven responses in VIP are rising rapidly (Extended Data Fig. 4a). In contrast, we
216 find no significant dependence of tuning correlation on task reference frame for MSTl (Fig. 5c,
217 green, $Z=0.175$, $p=0.86$), with only a small shift toward world coordinates occurring late in the

218 trial (Fig. 5d). These data suggest that VIP activity might account for the reference frame shifts
219 seen in behavior, whereas MSTl activity cannot.

220

221 **Linear decoding of VIP population activity predicts behavioral performance across** 222 **reference frame conditions**

223 To test whether the observed changes in single-unit responses with task reference frame
224 could account for behavioral performance, we used linear decoders (Fisher Linear Discriminant)
225 to classify object motion as rightward or leftward relative to vertical, based on responses of
226 pseudo-populations of 223 VIP neurons or 177 MSTl neurons (see Methods). We first
227 considered whether neural activity could account for behavioral performance if we trained
228 separate decoders to classify object motion direction in world or head coordinates (Fig. 6a). For
229 the head coordinate task, decoding either MSTl or VIP activity produced a pattern of results (Fig.
230 6b, blue and cyan curves) similar to the measured behavior (Fig. 3a, blue and cyan curves). This
231 implies that head-centered motion signals can be extracted from VIP and MSTl in the presence
232 of self-motion. Critically, the world coordinate task revealed clear differences in decoding
233 performance between brain areas. For MSTl, decoder performance curves (Fig. 6b right, brown
234 and magenta curves) shifted with self-motion direction to a much greater extent than seen in
235 behavior (Fig. 3a). In contrast, decoding of VIP responses (Fig. 6b left, brown and magenta
236 curves) reveals a pattern of results quite similar to behavior in the world coordinate task,
237 including substantially smaller shifts in the Object+Combined condition than the Object+Visual
238 condition. Because decoder weights were common across Object+Combined and Object+Visual
239 conditions, this improvement in decoder performance with inclusion of vestibular signals is not
240 guaranteed.

241 To summarize decoder performance, we computed Δ PSE values from the decoder-predicted
242 psychometric functions exactly as done for behavioral data. Results for the VIP decoder lie fairly
243 close to behavioral performance for both world and head coordinate tasks (Fig. 6c, orange vs.
244 black symbols). In contrast, Δ PSE values for the MSTl decoder differ greatly from behavioral
245 metrics in the world coordinate task (Fig. 6c, green vs. black symbols). These findings
246 demonstrate that MSTl responses only partially integrate self-motion signals and cannot be
247 effectively decoded in world coordinates, whereas the representation of object motion in VIP can
248 be decoded linearly to estimate object motion in either reference frame. Very similar results
249 (Extended Data Fig. 5) were obtained using a logistic regression decoder³², and results were
250 consistent across animals (Extended Data Fig. 6a-d).

251 In the analysis described above, we computed separate decoders for head and world
252 coordinate tasks, effectively assuming that the brain can apply different read-out weights to the
253 two task contexts. We can further ask whether it is possible to find a single set of decoding
254 weights that allows object direction to be read out in either reference frame. For this purpose, we
255 computed a single decoder that classifies object direction across both task reference frames (Fig.
256 6d). Strikingly, we found that a single decoder of VIP activity predicts performance in both
257 reference frames that is similar to behavior, whereas a single decoder of MSTl activity fails
258 almost completely for the world coordinate task (Fig. 6e). This finding, which was also quite
259 consistent across animals (Extended Data Fig. 6e-h), suggests that self-motion signals are
260 incorporated into VIP activity when animals are instructed to perform the task in a world-
261 centered reference frame (see Discussion). These results provide strong evidence for a novel role
262 of VIP in constructing a flexible representation of object motion.

263 A potential confound is that differential responses between the world and head coordinate
264 tasks might be driven by retinal motion of the partial cube frame, which is different between task
265 reference frames. The partial cube frame was generally kept outside of the receptive fields and
266 was faded out during motion of the object and background dots (Fig. 2c) to avoid this confound;
267 nevertheless, it is possible that this could account for some of the task-related response
268 modulations. To assess this possibility, we computed a measure of direction-selective response to
269 the partial cube frame (cube effect index, CEI, see Methods) for both world and head coordinate
270 tasks, focusing on the first 500ms of the trial during which the partial cube is visible and moving
271 but background dots are still largely invisible. For both brain areas, we found very few cells that
272 showed significant directionally-selective responses to the partial cube frame (Extended Data Fig.
273 7 a,b,d,e), with little difference in median CEI between world and head task conditions
274 (Wilcoxon signed rank test, VIP: $Z=1.62$, $p=0.10$; MSTl: $Z=-0.83$, $p=0.41$). Nevertheless, we
275 divided the neural populations in half based on the absolute difference in CEI, $|\Delta\text{CEI}|$, between
276 world and head tasks (Extended Data Fig. 7c,f), and we performed population decoding for these
277 two subsets of neurons. We found no reliable differences in decoder accuracy between subset of
278 neurons with relatively small and large values of $|\Delta\text{CEI}|$ (Extended Data Fig. 7g,h for separate
279 decoders, Extended Data Fig. 7i,j for single decoder), suggesting that the partial cube frame was
280 not responsible for differences in VIP responses between world and head task conditions.

281

282 **Temporal dynamics of reference frame transformations**

283 Since monkeys were trained to switch reference frames based on a visual cue, there might be
284 some delay in gating self-motion signals into the computation of object direction, especially if
285 monkeys rely on judging motion of the partial cube frame, in addition to the color of the fixation

286 point. Thus, we examined the time course of decoder performance using a 300 ms sliding
287 window that was shifted in increments of 50 ms. For this analysis, we used separate decoders for
288 head and world coordinate tasks to give each area the best chance of success. We computed the
289 time course of decoder performance separately for stimulus conditions in which the correct
290 answer is the same for both tasks (“response matched”) and conditions in which the correct
291 answers are different for the two task reference frames (“response conflict”, Fig. 7a). Decoder
292 performance on response conflict trials is of special interest, as it should reveal the clearest
293 differences in neural representations across task reference frames.

294 Indeed, time courses of decoder accuracy for response conflict trials revealed striking
295 differences between task reference frames and brain areas (Fig. 7b). For VIP on response
296 conflict trials, decoder accuracy rose ~500ms later for the world coordinate task than the head
297 coordinate task (Fig. 7b, left), roughly consistent with the time course of average MI values (Fig.
298 5b). This late rise in VIP decoder performance for the world task, starting around 1000ms, is too
299 early to be attributed to re-appearance of the partial cube frame, which reaches zero luminance at
300 ~1120ms and does not become clearly visible for another few hundred ms. In contrast, for MSTl,
301 decoder accuracy never reached much above chance for response conflict trials in the world
302 coordinate task, whereas accuracy rose quickly to high levels in the head coordinate task (Fig. 7b,
303 right). Thus, VIP responses undergo a delayed transformation that represents object motion in
304 world coordinates, whereas MSTl responses do not reliably represent motion in world
305 coordinates at any time. Note that decoder performance for VIP in the world task initially dips
306 below chance (0.5) levels, before rising precipitously (Fig. 7b, left). This below-chance
307 performance on response conflict trials is consistent with an early representation of object
308 direction in head coordinates in VIP, which later transitions to world coordinates. Below chance

309 performance for response conflict trials is also seen for MSTl during much of the stimulus period,
310 again reflecting a representation of object direction in head coordinates (Fig. 7b, right).

311 In response matched conditions, the time course of decoder performance is largely similar
312 for MSTl and VIP in both head and world coordinate tasks (Fig. 7c). The only notable difference
313 is that MSTl decoder accuracy drops off somewhat toward the end of the stimulus period,
314 whereas accuracy of the VIP decoder is more sustained. This difference was not attributable to
315 receptive field coverage (Extended Data Fig. 2) or to the temporal profiles of neural responses
316 (Extended Data Fig. 4). We also examined decoder performance separately for trials that
317 followed a switch between task reference frames, as compared to trials for which the reference
318 frame did not change, and we found no clear differences (data not shown).

319 Our findings suggest that self-motion signals are incorporated into the computation of object
320 motion in VIP during the world coordinate task. To probe this idea further, we took advantage of
321 the fact that monkeys judged object direction in both reference frames for each unique random-
322 dot stimulus. This allowed us to perform cross-task decoding by training a classifier to perform
323 the world coordinate task using neural responses from the head coordinate task, or vice-versa.
324 For this analysis, we focused on the response conflict conditions, for which differences between
325 areas and task reference frames are most clear.

326 For VIP, the decoder could perform the head coordinate task using responses from the world
327 coordinate task (Fig. 8b, gray/black), but could not perform the world coordinate task using
328 responses from the head coordinate task (Fig. 8a, gray/black). This suggests that VIP activity
329 contains information about object motion in head coordinates in both task conditions, but only
330 represents object motion in world coordinates when the animal is performing the world
331 coordinate task. By comparison, a decoder of MSTl activity fails to perform the world

332 coordinate task at all times using responses from either task condition (Fig. 8c), whereas it
333 accurately reports object direction in head coordinates when trained on responses from either
334 task condition (Fig. 8d). These observations are consistent with the hypothesis that both areas
335 carry robust information about object direction in a head-centered reference frame under all
336 conditions, whereas self-motion signals are incorporated into the computation of object motion
337 in VIP (but not MSTl) when the task requires a world-centered reference frame.

338

339 **Dissociation of choice signals from task reference frame signals**

340 Given that monkeys' choices are clearly different between the world and head task
341 conditions (Fig. 3), can the effects of task instruction on VIP responses be simply accounted for
342 by choice-related modulations? This question is especially relevant given that VIP neurons often
343 have strong choice-related activity that is not predictable from their stimulus tuning³³. We
344 performed analyses, at both the single neuron and population levels, which demonstrate that
345 response modulations related to task instruction are distinct from choice-related activity.

346 At the single-unit level, we separately quantified choice- and task-related activity (see
347 Methods). For choice-related activity, we computed the familiar choice probability (CP) metric³⁴,
348 which involves sorting responses into choice groups separately for each distinct stimulus and
349 task condition. Analogously, we quantified task-related modulations by computing a 'task
350 probability (TP)' metric, which involves sorting responses into groups based on task reference
351 frame. Critically, this is done separately for each choice and stimulus condition before z-scoring
352 and pooling, to ensure that TP is not influenced by choice. Many neurons in VIP and MSTl show
353 significant CP and/or TP values (Extended Data Fig. 8a,b); however, we find no correlation
354 between CP and TP across the population (VIP: $r=-0.062$, $p=0.36$; MSTl: $r = -0.041$, $p=0.59$,

355 Pearson correlation). To probe this dissociation further, we computed TP separately for left and
356 right choices (Extended Data Fig. 8c,d), and we find these values to be strongly correlated (VIP:
357 $r=0.76$, $p=9.9 \times 10^{-42}$; MSTl: $r = 0.76$, $p=1.6 \times 10^{-32}$, Pearson correlation), indicating that task-
358 related modulations for individual neurons are consistent across left and right choices. Similarly,
359 we find that CP values are correlated across task reference frames (VIP: $r=0.50$, $p=2.3 \times 10^{-15}$;
360 MSTl: $r = 0.33$, $p=1.7 \times 10^{-5}$, Extended Data Fig. 8e,f, Pearson correlation). Together, these
361 results show that choice and task reference frame have separable effects on responses of
362 individual neurons.

363 To assess whether performance of our decoders could be confounded with choice-related
364 activity, we devised an approach (see Methods) to largely remove either the choice-related or
365 task-related response modulations for each neuron. Our approach for removing choice-related
366 activity virtually eliminated significant CP values while leaving TP values largely unchanged
367 (compare Extended Data Fig. 9a,b with Extended Data Fig. 8a,b). Similarly, our method for
368 removing task-related modulations largely eliminated significant TP values while leaving CP
369 values largely unchanged (Extended Data Fig. 9c,d vs. Extended Data Fig. 8a,b). Critically,
370 decoder performance on response conflict trials was greatly impaired when task-related
371 modulations were removed (compare Extended Data Fig. 9e to Fig. 7b), whereas there was little
372 effect on decoder performance of removing choice-related modulations in response (Extended
373 Data Fig. 9f). These findings demonstrate that the flexible reference frame exhibited by VIP
374 activity cannot simply be attributed to choice-related activity.

375

376 **DISCUSSION**

377 By training monkeys to report object motion in either head or world coordinates, we
378 examined whether neurons represent object direction in a fixed reference frame or one that
379 changes with task requirements. Our findings demonstrate that VIP, but not MSTl, contains a
380 flexible representation of object motion that dynamically changes from moment to moment to
381 represent object direction relative to the head or world. The dynamics of these effects suggest
382 that self-motion signals are incorporated into the representation of object motion in VIP when the
383 task requires a world-centered representation.

384 Our findings provide an important advance in understanding how the brain represents
385 object motion during self-motion, providing the first evidence for a flexible multi-sensory
386 representation that can signal object motion relative to the head or world. The fact that addition
387 of vestibular stimulation facilitates reference frame transitions is consistent with previous
388 psychophysical and neurophysiological studies showing that vestibular input helps to dissociate
389 object motion and self-motion^{19, 24, 25, 35, 36}. More generally, our findings provide compelling
390 evidence that the reference frame of neural representations is not static, and can be powerfully
391 modulated by task instructions.

392

393 **Caveats and limitations**

394 It was difficult for animals to switch reference frames from trial to trial without the
395 partial cube frame, potentially because the screen edges provide a strong head-centered frame in
396 the absence of the partial cube (see Methods). Although the partial cube frame gradually became
397 invisible while the moving object became visible (Fig. 2c), it is possible that animals might have
398 tried to judge the horizontal velocity of object motion relative to specific edges or corners of the
399 partial cube, rather than judging object velocity relative to the world. Two factors argue strongly

400 against this interpretation: 1) a visual strategy of judging object velocity relative to specific
401 features of the partial cube frame would not explain the behavioral and neural effects of physical
402 motion of the platform, which provided vestibular signals. 2) We performed control experiments
403 in which we varied the location of the partial cube frame in depth from trial to trial. If animals
404 were reporting object motion relative to the near or far edges of the cube frame, their Δ PSE
405 values would be expected to depend systematically on cube depth, since the retinal velocity of
406 cube features depends on their distance from the observer. In contrast, we found no significant
407 dependence on depth of the cube (Extended Data Fig. 10), suggesting that the animals did not
408 employ this strategy.

409 Because we were not able to record from large ensembles of neurons simultaneously, our
410 decoding analyses were based on pseudo-populations of neurons for which the noise correlations
411 were largely unknown (see Methods). Thus, our analyses effectively assumed that neurons had
412 independent noise, which is not accurate^{37,38}. It is well established that correlated noise among
413 neurons can influence the information content (or sensitivity) of a population code³⁹⁻⁴¹.
414 Importantly, all of our main conclusions are based upon estimates of biases in decoding
415 performance, not on sensitivity measures. While we cannot rule out the possibility that
416 correlated noise would influence biases in decoder performance, it seems unlikely that the
417 pattern of results would change qualitatively.

418 Our analyses assume that the monkeys always identified the object as moving relative to
419 the background field of dots, which is a safe assumption for two reasons. First, object motion
420 always contained a substantial vertical component that was not compatible with the horizontal
421 self-motion of the animal. Second, the moving object was visually distinct from background
422 dots (see Methods), such that it was easily segmented from the background. More generally, the

423 brain has to solve a causal inference problem to discern whether the retinal motion of an object is
424 produced by self-motion or also reflects independent movement of the object relative to the
425 scene⁴². The neural basis of this causal inference process will be the topic of future studies.

426

427 **Relationship to previous studies**

428 A previous study²⁹ reported that visual tracking neurons in area MSTl represent visual
429 target motion in world coordinates while macaques tracked a target using voluntary eye and head
430 rotations. It was suggested that MSTl neurons represent world-centered target motion by
431 combining retinal motion signals, efference copy signals related to smooth eye movement, and
432 vestibular signals related to head rotation. On the surface, the findings of Ilg et al.²⁹ appear to
433 conflict with our finding that MSTl does not represent object motion in world coordinates.
434 However, our subjects performed the discrimination task while their eyes and head remained
435 oriented straight ahead. Whereas the tracking task used by Ilg et al.²⁹ elicited extra-retinal
436 signals related to eye and head rotation, our stimuli involved real or simulated head translations.
437 The findings of Ilg et al.²⁹ are therefore not incompatible with ours, and collectively they
438 suggest that some MSTl neurons may account for eye and head rotations, but that MSTl does not
439 contain a generalized representation of world-referenced object motion.

440 Reference frames of different sensory signals in area VIP have been the focus of several
441 previous studies. Facial tactile receptive fields (RFs) are coded in a head-centered reference
442 frame², whereas auditory RFs are organized in a continuum between eye- and head-centered
443 coordinates^{10,11}. Visual RFs and heading tuning (optic flow) are represented mainly in an eye-
444 centered reference frame^{11,31,43}, although some studies have described head-centered visual RFs
445 in VIP as well⁴. In contrast, vestibular heading signals in VIP are coded in body- or world-

446 centered reference frames^{30, 44}. One recent study showed that the reference frame of vestibular
447 heading tuning in VIP depended on whether gaze was focused on a head- or world-fixed target⁴⁴,
448 consistent with the idea that VIP has flexible reference frames. However, our findings show that
449 VIP reference frames can change just by task instructions, and do not require different motor
450 actions⁴⁴.

451 Our findings substantially extend previous work on the context-dependence and
452 dynamics of spatial reference frames. Human neuroimaging studies have reported that visual
453 motion signals can be represented in retinal or head coordinates depending on the spatial
454 allocation of attention⁴⁵, although this finding has been refuted by other studies⁴⁶. Human fMRI
455 studies also demonstrated that activity in parietal and premotor cortex reflected different spatial
456 reference frames depending on the sensory modality used to specify target location⁴⁷. Previous
457 studies have also demonstrated that the reference frame of neural activity in monkeys is dynamic,
458 changing over time relative to task events^{8, 48}. Our findings extend this work in two important
459 ways. First, we demonstrate that neural activity is modified by task instructions to represent
460 object motion in the reference frame required for each task condition. Second, we directly
461 compare neural and behavioral correlates of dynamically changing task reference frames,
462 allowing for a more direct assessment of whether changes in neural activity with task reference
463 frame can explain behavior. In contrast, previous neurophysiological studies of reference frames
464 have generally just varied the position of an effector without requiring animals to make a
465 perceptual report.

466 The Duncker illusion^{49, 50} describes biases in the perceived trajectory of an object when it
467 moves relative to a moving background. The perceptual biases exhibited by our monkeys cannot
468 simply be accounted for by the Duncker illusion because the image motion of the target object

469 and background dots are identical in the world and head coordinate tasks, yet the perceptual
470 reports are strikingly different (Fig. 3a).

471

472 **Implications of flexible reference frames**

473 The primary contribution of our study is to demonstrate that neural reference frames can
474 change dramatically based on task instructions. Secondly, these results have implications for
475 understanding the variability of outcomes across previous studies of neural reference frames.

476 Two examples of such variability, as noted above, include the incidence of head-centered visual
477 receptive fields in VIP^{4,43} and the existence of spatiotopic representations in human visual
478 cortex^{45,46}. Given that the vast majority of neurophysiological studies of reference frames have
479 not used a behavioral task that enforces a specific task reference frame, findings could vary with
480 the intrinsic (and uncontrolled) reference frame that the animal employs, which in turn may
481 depend on the animal's previous experience or training history. Findings could also vary with the
482 stimuli used, which might bias the animal toward adopting a specific task reference frame.

483 We found that a single set of decoding weights could be used to classify object direction
484 in either head or world coordinates, based on VIP activity. This result could arise because task
485 instructions simply shift the 'population hill' of neural activity along the stimulus axis, similar to
486 changing object direction itself. A pure horizontal shift of the population hill, in which the
487 pattern of population activity is simply translated along the object direction axis, would occur if
488 all tuning curves for object direction simply shifted with self-motion in the world coordinate task.
489 This was clearly not the case based on inspection of tuning curves from individual neurons (Fig.
490 4 and Extended Data Fig. 3), as well as the broad distribution of tuning correlation values in Fig.
491 5c. We found that self-motion has diverse effects on object motion tuning in the world

492 coordinate task, including shifts, gain changes, and changes in shape of tuning. Thus, it remains
493 an interesting topic for future studies to determine how a single decoder can estimate object
494 direction in head or world coordinates based on such diverse modulations at the single-unit level.

495

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503 interpreted results of experiments; R.S. prepared figures; R.S. and G.C.D. drafted manuscript;
504 R.S., A.A., D.E.A., and G.C.D. edited and revised manuscript; R.S., A.A., D.E.A., and G.C.D.
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506

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508

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- 624

625 **FIGURE LEGENDS**

626 **Figure 1. Schematic illustration of interactions between object motion and self-motion.** (a)

627 An object (gray sphere) moves upward in the world while an observer is translated rightward or
628 leftward at two speeds. (b) Resultant image motion vectors. Without self-motion, image motion
629 is upward (white). During self-motion, image motion is biased according to the direction and
630 speed of self-motion. For simplicity, the image view in panel b does not reflect image reversals
631 that would be caused by projection onto the retina.

632

633 **Figure 2. Behavioral task design and predicted psychometric functions.** (a) A sphere of dots

634 moves up-right (+ θ) or up-left (- θ) in the world. Rightward or leftward self-motion occurs while
635 the animal views the moving object. (b) In the world coordinate task, a world-fixed partial cube
636 indicates that the monkey should report object motion relative to the world. In the head
637 coordinate task, the partial cube remains fixed relative to the head, and cues a report in head
638 coordinates. Dashed vertical lines indicate fixed world-centered locations as a reference.

639 Background dots were presented at 40% coherence, but background motion here is depicted with
640 100% coherence for visual clarity. (c) Time course of the luminance of visual stimulus

641 components. The luminance of the object and partial cube were changed dynamically such that
642 the partial cube faded out during the portion of the trial when the object faded in. (d)

643 Hypothetical psychometric functions that plot the proportion of 'rightward' choices as a function
644 of object direction in world coordinates. If the animal compensates fully for self-motion in the
645 world coordinate task, psychometric functions for rightward and leftward self-motion should
646 overlap (magenta). On the other hand, those functions should shift with self-motion by a specific

647 amount (horizontal bar) in the head coordinate task (cyan). Dashed/solid curves:
648 leftward/rightward self-motion.

649

650 **Figure 3. Summary of behavioral performance for each task reference frame. (a)**

651 Psychometric functions showing the proportion of rightward choices as a function of object
652 direction in world coordinates (positive = rightward). Data are shown for trials in which there is
653 no self-motion (Object Only, black), for trials with self-motion in which the animal performs the
654 head coordinate task (cyan/navy), and for trials in which the animal performs the world-
655 coordinate task (magenta/brown). Darker colors (navy/brown) represent the Object+Visual
656 condition and lighter colors (cyan/magenta) represent the Object+Combined condition.

657 Filled/open symbols: rightward/leftward self-motion. Smooth curves are cumulative Gaussian
658 fits to data pooled across 128 sessions for Monkey N and 57 sessions for monkey K. (b)

659 Summary of behavioral biases, quantified as the difference in point of subjective equality
660 (Δ PSE) between psychometric functions for rightward and leftward self-motion. Δ PSE values
661 are compared between the Object+Combined and Object+Visual conditions for each recording
662 session and each monkey (squares: monkey N; diamonds: monkey K). Data are shown separately
663 for the world coordinate (red) and head coordinate (blue) task conditions. Error bars represent
664 95% confidence intervals around the mean values (black symbols) across 185 sessions.

665

666 **Figure 4. Data from example neurons recorded from areas VIP and MSTl. (a)** Data from

667 two example VIP neurons (one neuron per column) recorded in the Object+Combined condition.

668 The top and bottom rows plot firing rates as a function of object direction in head and world

669 coordinates, respectively. (b) Data from two example MSTl neurons in the Object+Combined

670 condition. Note the greater differences in response between the head (cyan) and world (magenta)
671 coordinate task conditions for the example VIP neurons, as compared to the example MSTl
672 neurons. Error bars denote SEM (across $n=10$ stimulus repetitions). (c, d) Data for the same 4
673 example neurons from the Object+Visual condition.

674

675 **Figure 5. Summary of single-unit results for VIP and MSTl.** (a) Summary of modulation
676 index (MI) values for populations of neurons recorded from VIP (orange, $N=223$) and MSTl
677 (green, $N=177$) in the Object+Visual (top) and Object+Combined (bottom) conditions. MI
678 measures the response difference between a pair of object tuning curves in the head- and world-
679 coordinate tasks. Filled bars represent MI values significantly greater than zero (permutation test,
680 $p < 0.05$). Numbers in the legend indicate the total number of neurons, as well as the number
681 with MI values significantly greater than zero, for each brain area. Arrowheads and numbers
682 indicate the median values for each brain area and self-motion condition. (b) Time course of
683 average MI values for VIP (orange, $N=223$) and MSTl (green, $N=177$) neurons in the
684 Object+Visual (lighter hues) and Object+Combined (darker hues) conditions. Error bars
685 represent 95% confidence intervals. Gray curve shows the Gaussian temporal profile of object
686 speed. (c) The correlation between object direction tuning (computed in world coordinates) is
687 compared for the world and head coordinate tasks. Data from VIP and MSTl are shown in
688 orange and green, respectively. Data are included in this panel only for neurons (VIP: $N=57$;
689 MSTl: $N=44$) that had significant tuning (ANOVA, $p < 0.05$) in the Object Only condition. Star
690 symbols denote the three neurons in Figure 4 that met this criterion. (d) Time course of the
691 difference in tuning correlation between world and head coordinate tasks for the same

692 populations of VIP (orange, N=223) and MSTl (green, N=177) neurons described in panel c.
693 Error bars represent 95% confidence intervals.

694

695 **Figure 6. Summary of population decoding results.** Panels a-c correspond to results from
696 training separate decoders to perform the world and head coordinate tasks; panels d-f correspond
697 to results from a single decoder trained to perform in both task reference frames. (a) Schematic
698 diagram of separate decoders for the head and world coordinate task conditions. (b) Results for
699 separate world/head task decoders, plotted in the same format as the behavioral data of Fig. 3a.
700 Decoding VIP activity produces a pattern of results very similar to behavior, whereas decoding
701 MSTl produces large biases in the world coordinate task. (c) Summary comparison of monkey
702 behavior and performance of the separate decoders. Δ PSE for the Object+Combined condition is
703 plotted against Δ PSE for the Object+Visual condition. Results from the VIP decoder (orange) are
704 largely similar to behavior (black, same data from Fig. 3b), whereas results from the MSTl
705 decoder (green) depart sharply from behavioral performance for the world coordinate task. Error
706 bars on decoder performance values represent 95% confidence intervals obtained by
707 bootstrapping (n=1000 bootstraps, see Methods). Pink and cyan dashed lines: expected Δ PSE for
708 perfect performance in the world and head coordinate tasks, respectively. (d) Schematic diagram
709 for the single decoder. (e) Analogous results to panel b, but from a single decoder trained to
710 classify object direction in both task reference frames. (f) Summary comparison of single
711 decoder results with behavior. Format as in panel c; error bars represent 95% confidence
712 intervals across n=1000 bootstraps.

713

714 **Figure 7. Time course of decoder performance.** (a) Schematic illustration of examples of
715 “response matched” and “response conflict” conditions. In a response matched condition (left),
716 correct answers are the same for both task reference frames; in a response conflict condition
717 (right), correct reports are opposite for the two reference frames. Magenta and cyan vectors
718 indicate object direction in world and head coordinates, respectively. (b) Time course of decoder
719 classification accuracy for populations of VIP (left, n=223) and MSTl (right, n=177) neurons,
720 evaluated in the subset of response conflict conditions. Separate decoders were trained to classify
721 object direction in the world (magenta/brown) and head (cyan/navy) reference frames at each
722 time point. Error bars represent 95% confidence intervals (across n=100 bootstraps). (c) Time
723 course of decoder classification accuracy in the subset of response matched conditions, format as
724 in b. Time courses were obtained by computing each variable within a 300 ms sliding time
725 window that was advanced across the trial epoch in steps of 50 ms.

726

727 **Figure 8. Time courses of classification accuracy using within-task vs. cross-task decoding.**

728 (a, b) Results for decoding VIP activity. In panel a, the decoder is trained to classify object
729 direction in world coordinates using responses from the world task condition (magenta/brown,
730 within-task) or using responses from the head task condition (light/dark gray, cross-task). In
731 panel b, the decoder is trained to classify object direction in head coordinates using responses
732 from the head task condition (cyan/navy, within-task) or the world task condition (light/dark gray,
733 cross-task). (c,d) Analogous results for within-task (colors) and cross-task (gray) decoders of
734 MSTl activity. In all panels, error bars denote 95% confidence intervals (across n=100
735 bootstraps).

736

738 **METHODS**

739 **General**

740 Two male rhesus monkeys (*Macaca mulatta*) participated in this study. During this study,
741 monkey K ranged in age from 4 to 6 years and ranged in weight from 5.8 to 8.5 kg. Monkey N
742 ranged in age from 5 to 7 years and weight from 7.2 to 9.7 kg. General procedures have been
743 described previously^{19,51}. All experimental procedures conformed to National Institutes of
744 Health guidelines and were approved by the University Committee on Animal Resources at the
745 University of Rochester. Additional information can be found in the Life Sciences Reporting
746 Summary.

747

748 **Vestibular and visual stimuli**

749 A 6 degree-of-freedom motion platform (MOOG 6DOF2000E; Moog) was used to passively
750 translate animals leftward or rightward along the interaural axis. Visual stimuli were projected
751 onto a tangent screen by a three-chip digital light projector (Mirage S⁺3K ; Christie Digital
752 Systems, Cypress, CA). The display screen measured 60 x 60 cm and was mounted ~30 cm in
753 front of the monkey, thus subtending ~90 x 90° of visual angle. Visual stimuli simulated
754 translational self-motion through a three-dimensional field of stars. Each star was a triangle that
755 measured 0.15 cm x 0.15 cm, and the field of stars measured 100 cm wide by 100 cm tall by 40
756 cm deep, with a star density of 0.01 stars per cm³. To provide stereoscopic cues, the star field
757 was rendered as a red-green anaglyph and viewed through custom red-green goggles, consisting
758 of Kodak Wratten2 filters (#29 and #61). The entire display was visible through the colored
759 filters.

760 The optic flow field contained naturalistic cues simulating lateral translation of the observer
761 in the horizontal plane; these included motion parallax, size, and binocular disparity cues. While
762 the monkey was translated leftward or rightward, an object also moved upward in the world with
763 a small leftward or rightward component (Fig. 1). The moving object was a transparent sphere
764 (diameter 10°) composed of random dots, with a density (0.25 dots/cm^3) that was higher than
765 that of the star-field background, such that the object was easily segmented from the background.
766 The moving object's center was located in depth within the plane of the visual display, such that
767 it consisted of dots with a mixture of crossed and uncrossed disparities. At the start of each trial,
768 the object appeared with its center located 5 deg left of fixation and 10 deg below fixation. The
769 object moved in one of 7 directions relative to upward in the world: -21, -14, -7, 0, 7, 14 and 21
770 deg, where negative angles represent upward/left motion, positive angles represent upward/right
771 motion, and 0 means straight upward (in world coordinates). All self-motion and object motion
772 trajectories were straight translational movements with a duration of 2 sec, and having a
773 Gaussian velocity profile with a SD of $1/3 \text{ sec}^{51}$. Peak stimulus velocity occurred at 1120ms
774 after stimulus onset, due to delays and dynamics of the motion platform; visual stimuli were
775 synchronized to platform motion. The total excursion (0.25 m) and peak velocity (0.75 m/s) of
776 object motion were greater than those for self-motion (0.08 m and 0.24 m/s, respectively).
777 Because the head-centered velocity of the object is determined by both self-motion velocity and
778 object velocity relative to the world, the object could move up/left in world coordinates and
779 up/right in head coordinates, or vice-versa. While we shall distinguish between world- and head-
780 centered reference frames in this study, we cannot distinguish head-centered and retinal
781 reference frames because the fixation target was always head-fixed.

782 Two different versions of the task were interleaved that required the animal to report object
783 direction in either head or world coordinates (Fig. 2). In the world coordinate task (Fig. 2b), a
784 partially-visible cube defined a world-fixed reference frame that was updated every video frame.
785 The cube dimensions were 76 cm wide by 76 cm tall by 40 cm deep and the center of the cube
786 was located in depth at the fixation point. Thus, the cube moved relative to the head during self-
787 motion in the world coordinate task (Fig. 2b) but remained head-fixed in the head coordinate task
788 (Fig. 2b).

789 We attempted to train monkeys to switch between the world and head coordinate tasks
790 based solely on the color of the fixation point. While one animal could partially achieve this, the
791 other animal could not. Both animals were much better able to switch between tasks when the
792 partially-visible cube was presented. We think that the partial cube was particularly important
793 because it was not possible to eliminate luminance boundaries at the edge of the display. Since
794 the display screen translated with the animal, the luminance boundaries always provided a head-
795 centered reference frame, and thus the partially visible cube was important to help define a world
796 reference frame (anecdotally, this was the case for human observers also).

797 A potential concern about use of the partially visible cube is that animals might learn to
798 report object direction relative to the moving visual elements of the cube. Two measures helped
799 to prevent this possibility. First, the luminance of the moving object was dynamically changed
800 according to the same Gaussian envelope that governed the speed of object and self-motion, such
801 that the moving object was initially invisible, reached maximum brightness in the middle of the
802 presentation when it also reached maximum speed, and then decayed again to become invisible
803 at the end of the trial (Fig. 2b,c). Simultaneously, the luminance of the partially visible cube
804 followed an inverted Gaussian velocity profile, such that the cube was maximally visible at the

805 beginning and end of each trial and disappeared in the middle of the trial (Fig. 2b,c). This
806 allowed the partial cube to define the reference frame while having little overlap with the
807 visibility of the moving object.

808 Second, to assess whether animals might have still judged object motion relative to the
809 partially-visible cube, we performed behavioral control experiments in which we randomly
810 varied the location of the partial cube in depth from trial to trial. If animals reported object
811 direction relative to the cube, then their performance should depend systematically on the depth
812 of the cube. We found no such dependence (Extended Data Fig. 10), indicating that animals
813 were successfully prevented from adopting this strategy. Thus, we believe that both animals
814 successfully learned to report object direction in world or head coordinates.

815

816 **Behavioral task**

817 Monkeys were trained to report whether the object moved up-left or up-right in either head-
818 or world-coordinates (Fig. 2). This was a very challenging task for animals to learn, and required
819 1.5-2 years of training for each animal. We initially trained the animals to perform the head and
820 world tasks in separate blocks of trials. We then gradually reduced the length of these blocks,
821 and then transitioned animals to trial-by-trial interleaving of the two tasks.

822 In each trial, a fixation point initially appeared. Once the monkey looked at the fixation
823 point, the object appeared and moved upward in the virtual environment, with a small rightward
824 or leftward component. The monkey reported whether the object moved upward/rightward or
825 upward/leftward by making a saccade to one of two targets that appeared (10 degrees to the right
826 and left of the fixation target) after a 500 ms delay period following the end of the visual
827 stimulus. In the world and head coordinate tasks (Fig. 2b), the monkey reported whether object

828 motion moved leftward or rightward relative to vertical in world or head coordinates,
829 respectively. The two versions of the task were cued by the shape and color of the fixation point
830 (Fig. 2b), such that they could be randomly interleaved. Animals were rewarded for reporting
831 the correct direction of object motion in each reference frame condition. When object direction
832 was exactly vertical (in the relevant reference frame), monkeys were rewarded randomly on 50%
833 of trials.

834 Crucially, for each particular combination of a self-motion direction and an object motion
835 direction in the world, the motion trajectory of the object in head (or retinal) coordinates was
836 identical across the two task conditions. Thus, for the same exact motion of all of the dots on the
837 screen, the animal might be required to make a rightward choice in the world coordinate task and
838 a leftward choice in the head coordinate task, or vice-versa. For other stimulus conditions, the
839 correct choice would be the same in both reference frames. This allowed us to compare decoder
840 performance for subsets of trials in which the correct answers were the same or different for the
841 two tasks.

842 Three self-motion conditions were interleaved for each task reference frame. 1) In the Object
843 Only condition, there was no self-motion such that world- and head-centered reference frames
844 are aligned. Background dots were stationary on the display in the Object Only condition, since
845 the only source of image motion for the background dots is self-motion. 2) In the Object+Visual
846 condition, the motion platform remained stationary while a background of random dots provided
847 optic flow that simulated leftward or rightward self-motion. Background optic flow had a motion
848 coherence of 40% such that the object was easy to segment from the background. 3) In the
849 Object+Combined condition, self-motion was indicated by both optic flow and physical
850 translation of the motion platform (which provided vestibular cues). Since cue combination is

851 known to enhance heading perception^{52,53}, we expected that the monkeys would be best able to
852 compensate for self-motion in this condition.

853

854 **Behavioral data analysis**

855 Psychometric functions were constructed by plotting the proportion of ‘rightward’ choices
856 as a function of object direction in world coordinates. Plotted in this fashion, psychometric
857 functions for rightward and leftward self-motion should overlap in the world coordinate task if
858 the animal compensates fully for self-motion (Fig. 2d). If the animal does not account for self-
859 motion and reports object direction in head coordinates, there will be a large horizontal shift
860 between psychometric functions corresponding to leftward and rightward self-motion (Fig. 2d).
861 To quantify these shifts, we fit each psychometric curve with a cumulative Gaussian function
862 and used the mean parameter of the fit to estimate the point of subjective equality (PSE) for each
863 direction of self-motion. The difference in PSE (Δ PSE) between rightward and leftward self-
864 motion directions was then taken as an index of the reference frame used by the monkeys to
865 judge object direction (Fig. 3b). If the monkey correctly estimates object direction in head
866 coordinates, then we expect Δ PSE=35.5 deg (horizontal bar, Figs. 2d, 3a).

867

868 **Physiological recording procedures**

869 Neural recordings were obtained from the right hemisphere of two monkeys while the
870 animals performed the behavioral task. We recorded 223 VIP neurons (monkey N, N=93;
871 monkey K, N=130) and 177 MSTl neurons (monkey N, N=94; monkey K, N=83), with most
872 neurons recorded in separate sessions. We attempted to record from any VIP and MSTl neuron
873 that could be isolated; there were no selection criteria based on response properties other than

874 receptive field location. Recordings were included if we obtained data for at least 3 repetitions
875 for each stimulus condition. For Monkey N, 68 VIP and 50 MSTl neurons were recorded with
876 single tungsten microelectrodes (FHC, Bowdoinham, ME; 0.5 – 1 M Ω impedance). Single-unit
877 action potentials were sorted on-line using a hardware window discriminator (Bak Electronics).
878 The remaining 25 VIP and 44 MSTl neurons from Monkey N were recorded with linear
879 electrode arrays that were inserted into either VIP or MSTl daily (Plexon U-probes with two
880 rows of 12 channels spaced 100 μ m vertically and 50 μ m horizontally or Plexon V-probes with
881 24 channels spaced 50 μ m vertically). For array recordings, single-unit action potentials were
882 isolated using Plexon Offline Sorter. For Monkey K, all 130 VIP and 83 MSTl neurons were
883 recorded with linear arrays. There were a total of 104 recording sessions for VIP (monkey N,
884 N=69; monkey K, N=35) and 81 recording sessions for MSTl (monkey N, N=59; monkey K,
885 N=22). In experiments using linear arrays, a mean of 3.8 and 3.7 neurons were recorded
886 simultaneously for areas MSTl and VIP, respectively.

887 Both VIP and MSTl were initially localized via structural MRI scans as described previously
888 for VIP⁵⁴ and MST⁵⁵. We were careful to distinguish MSTl from the dorsal subdivision of area
889 MST (MSTd) and MT³⁶. To do this, we carefully mapped the portions of area MT that were
890 found beneath MSTd, in the posterior bank of the superior temporal sulcus. We located the
891 foveal representation of area MT, which is generally located at the anterior-lateral extent of MT.
892 We then carefully mapped regions around that area, and MSTl was localized primarily in regions
893 anterior to the foveal representation of MT.

894

895 **Experimental protocol**

896 We first performed standard tests to map receptive fields and assess response properties
897 qualitatively. These tests, along with mapping recording sites onto structural MRI images⁵⁴,
898 allowed us to confidently assign recording sites to MSTl or VIP. Neurons were isolated while
899 presenting a large field of flickering dots that could be varied in position, size, and velocity. For
900 some neurons, we used a reverse-correlation technique to measure the spatial and directional
901 receptive field structure of each neuron⁵⁶. From these maps, we fit the receptive field with a
902 two-dimensional Gaussian, and used the contour of the Gaussian at half-maximal response to
903 define the receptive field contours shown in Extended Data Fig. 2 (17% of VIP neurons and 13%
904 of MSTl neurons). Due to a technical difficulty, reverse correlation maps were not available for a
905 substantial fraction of neurons. In other recordings, receptive fields were mapped by hand, and
906 receptive field location and size was estimated when the map was clearly noted. We also
907 performed a standard measurement of directional tuning within the fronto-parallel plane by
908 presenting 8 directions of motion, 45 deg apart. These preliminary tests typically required 150-
909 200 trials of fixation behavior.

910 We recorded from all neurons regardless of their direction and speed preferences. To
911 facilitate population decoding, we used exactly the same stimulus set for all recorded neurons.
912 This allowed us to construct pseudo-population responses for decoding, although these pseudo-
913 population responses do not contain accurate correlated noise since the vast majority of neurons
914 were recorded separately. After extensively mapping out the receptive field coverage of neurons
915 in areas VIP and MSTl, we focused our recordings on a set of penetrations for which receptive
916 fields were concentrated on the same region of space for both brain areas (Extended Data Fig. 2).
917 We carefully selected the starting location and trajectory of object motion based on the
918 distributions of receptive fields of MSTl and VIP neurons in our selected penetrations, such that

919 object motion was centered on the receptive field locations for the populations of neurons in both
920 VIP and MSTl (Extended Data Fig. 2). The main experimental protocol involved 7 directions of
921 object motion, 2 reference frame conditions (head or world), 2 self-motion directions, and 10
922 stimulus repetitions for each of the Object+Visual and Object+Combined conditions (560 trials),
923 as well as 7 directions and 20 stimulus repetitions for the Object Only condition (140 trials), for a
924 total of 700 trials.

925

926 **Neural data analyses**

927 Neural responses were computed as firing rates over a time window from 500-2500 ms
928 following stimulus onset. Since the stimulus duration was 2000 ms, this window included most
929 of the stimulus period during which neurons were active, as well as the 500ms delay period after
930 stimulus offset. This analysis window was based on inspection of population response profiles
931 (Extended Data Fig. 4a). The initial 500ms of the stimulus period was not included in our main
932 analysis window because there is an early response to luminance onset of dots during this time
933 (see Extended Data Fig. 4a) and because object and background motion is small during the first
934 500ms (due to the Gaussian velocity profile used). Our main analyses were also conducted as a
935 function of time, using a moving window of 300 ms that was slid across the data in steps of 50
936 ms. All analyses are performed on all trials, including both correct and incorrect trials, unless
937 indicated otherwise.

938 *Modulation Index for the effect of reference frame on neural responses:* To quantify how
939 neural responses are modulated by the task reference frame, we computed a modulation index
940 (MI) as follows:

941
$$MI = \frac{1}{N} \left(\frac{|\sum_{\theta} (R(\theta)_{W,L} - R(\theta)_{H,L})| + |\sum_{\theta} (R(\theta)_{W,R} - R(\theta)_{H,R})|}{|\sum_{\theta} (R(\theta)_{W,L} + R(\theta)_{H,L})| + |\sum_{\theta} (R(\theta)_{W,R} + R(\theta)_{H,R})|} \right) \quad (1)$$

942 In this formulation, R_W and R_H , denote the mean responses of a neuron in the world and head-
 943 coordinate tasks, respectively, whereas additional subscripts L and R denote leftward and
 944 rightward self-motion directions. θ represents object direction, and N denotes the number of
 945 object directions. MI ranges from 0 (no difference between responses in the two reference
 946 frames) to 1 (if, for example, responses to one reference frame condition are completely
 947 suppressed).

948 In formulating MI, we sought a simple metric to quantify response modulations related to
 949 the task reference frame across all object and self-motion directions. If neural responses were
 950 identical in the world and head coordinate tasks, MI would be zero; however, in practice, MI
 951 values are unlikely to be very close to zero due to response variability. If world and head
 952 coordinate tasks produce different average neural responses, then MI values will become
 953 substantially greater than zero. Note that MI is not sensitive to the nature of response
 954 modulations (e.g., peak shifts vs. gain modulations vs. tuning shape changes). Given that our
 955 direction discrimination task covered a relatively narrow range of directions relative to the full
 956 tuning curves, it is difficult to examine the exact nature of tuning changes from our data.

957 *Direction discrimination index:* To quantify the strength of tuning for object direction, we
 958 used a direction discrimination index (DDI) that was defined as follows:

959
$$DDI = \frac{R_{\max} - R_{\min}}{R_{\max} - R_{\min} + 2\sqrt{SSE/(N-M)}} \quad (2)$$

960 where R_{\max} and R_{\min} represent the maximum and minimum responses from the measured
 961 direction tuning function, respectively. SSE is the sum squared error around the mean responses,

962 N is the total number of observations (trials), and M is the number of tested object directions (M
 963 $= 7$). DDI is a signal-to-noise metric, conceptually similar to d' , that is normalized to range from
 964 0 to 1. Neurons with stronger response modulations relative to their variability will take on
 965 values closer to 1.

966 *Effect of partial cube frame on responses:* To quantify effects of the partial cube frame on
 967 neural responses, we computed a cube effect index (CEI). This index measures neural responses
 968 over the initial 500ms of each trial, when the partial cube is visible while background dots are
 969 largely invisible (Fig. 2c). For each object direction, θ , CEI takes the absolute difference in
 970 response to the cube frame between rightward and leftward self-motion directions, and divides
 971 by the sum of those responses. The resultant is then averaged across the N object directions. For
 972 the world coordinate task, the calculation of CEI is as follows:

$$973 \quad CEI_W = \frac{1}{N} \left(\frac{|\sum_{\theta} (R(\theta)_{W,R} - R(\theta)_{W,L})|}{|\sum_{\theta} (R(\theta)_{W,R} + R(\theta)_{W,L})|} \right)_{t \in [0-500 \text{ ms}]} \quad (3)$$

974 where R_W denotes the mean responses of a neuron in the world task, and subscripts L and R
 975 denote leftward and rightward self-motion directions. The calculation of CEI for the head
 976 coordinate task is identical, with replacing R_W by R_H .

977 *Metrics of choice-related and task-related activity:* To quantify choice-related activity in
 978 single neurons, we computed the well-established choice probability (CP) metric³⁴. For each
 979 unique object direction, self-motion direction, self-motion modality (visual, combined), and task
 980 reference frame condition (world vs. head), the distribution of responses was z-scored and then
 981 divided into two groups based on whether the animal made a leftward or rightward saccade. Z-
 982 scored responses were then pooled across unique stimulus/task conditions as long as there were
 983 at least 3 choices made in each direction. ROC analysis was then applied to the pooled z-scores

984 for the two choice groups, and CP was defined as the area under the ROC curve. For our
985 purposes, CP was not referenced to each neuron's preferred direction; rather $CP > 0.5$
986 corresponds to a preference for rightward choices and $CP < 0.5$ corresponds to a preference for
987 leftward choices. This avoids potential issues with defining the "preferred" stimulus when choice
988 effects are large³³.

989 We devised an analogous ROC-based metric to quantify single-unit activity related to task
990 reference frame. This 'task probability' (TP) metric is computed just like CP, but swapping the
991 roles of variables that represent choice (left vs. right) and task (head vs. world). For each distinct
992 combination of object direction, self-motion direction, self-motion modality, and choice,
993 responses were z-scored and sorted into two groups based on task reference frame. If there were
994 at least 3 trials for world and head reference frames, normalized responses from that condition
995 were pooled with other conditions that met the same criteria. ROC analysis was applied to the
996 pooled z-scores that were sorted into world and head task groups. $TP > 0.5$ corresponds to
997 greater responses in the world coordinate task, and $TP < 0.5$ corresponds to greater responses in
998 the head coordinate task.

999 *Removal of choice- and task-related response modulations:* To test whether choice- or task-
1000 related signals make specific contributions to decoder performance, we devised a method to
1001 remove either choice- or task-related response modulations from neural activity. First, we
1002 identified a set of trials corresponding to each unique combination of object direction, self-
1003 motion direction, and self-motion modality. If this set of trials included at least 3 trials each for
1004 left and right choices and 3 trials each for world and head task conditions, then we proceeded to
1005 remove either the choice- or task-related response component. To remove the choice-related
1006 response component, we shifted the mean responses for right and left choices toward each other

1007 to equate the mean responses. Comparison of Extended Data Fig. 9a,b to Extended Data Fig.
1008 8a,b indicates that this manipulation eliminated most of the choice-related modulations, while
1009 preserving task-related modulations. Similarly, to remove the task-related component, we shifted
1010 the mean responses for world and head task conditions to equate the means. Extended Data Fig.
1011 9c,d indicates that this manipulation was successful in eliminating most of the task-related
1012 modulations, while preserving choice effects. These manipulations cannot completely remove all
1013 choice- or task-related activity because they can only be performed when there are at least a few
1014 trials in each choice and task group, and estimates of mean responses based on a few trials are
1015 noisy.

1016 *Population decoding by a linear classifier:* Linear decoding was performed to classify object
1017 direction as rightward or leftward of vertical in each reference frame. Pseudo-population
1018 responses of 223 neurons for VIP and 177 neurons for MSTl were used for this purpose. We
1019 used a linear classifier to categorize object motion as rightward or leftward relative to vertical in
1020 either world or head coordinates:

$$1021 \quad f = \sum_{i=1}^N w_i \cdot r_i + k \quad (4)$$

1022 Here, N is the number of neurons in the pseudo-population for either VIP or MTL, r_i is the
1023 response of the i^{th} neuron, w_i is the decoding weight for the i^{th} neuron, and k is constant scalar.

1024 The decoder's choice is determined by the sign of the output variable, f . We used a Fisher linear
1025 discriminant (FLD) to compute the parameters (w , k) as follows:

$$1026 \quad w = \Sigma^{-1} \cdot (\mu_R - \mu_L) \quad (5)$$

$$1027 \quad k = \frac{1}{2} \cdot [(\mu_L^T \cdot \Sigma^{-1} \cdot \mu_L) - (\mu_R^T \cdot \Sigma^{-1} \cdot \mu_R)] \quad (6)$$

1028 where μ_L and μ_R indicate the mean population response vectors for rightward and leftward object
1029 directions relative to the world (for the world task decoder) or head (for the head task decoder),
1030 and Σ is the response covariance matrix.

1031 Since most of our neurons were not recorded simultaneously, all neurons did not see the
1032 same number of repetitions of each unique stimulus. Thus, we constructed a population response
1033 matrix in which each neuron had responses corresponding to 10 stimulus repetitions. For neurons
1034 recorded for >10 repetitions (199/223 for VIP, 152/177 for MSTl), we randomly removed some
1035 repetitions; for neurons recorded for <10 repetitions (23/223 for VIP, 24/177 for MSTl), we
1036 filled in data by sampling with replacement. Once this was done, we computed the covariance
1037 matrix using the ‘cov()’ function in Matlab, as though all neurons had been recorded
1038 simultaneously. Since most pairs of neurons were not recorded simultaneously (simultaneous
1039 pairs: 226/24753 for VIP, 187/15576 for MSTl), the off-diagonal elements of the resulting
1040 covariance matrix do not reflect correlated noise for the vast majority of neuron pairs. However,
1041 the off-diagonal elements are generally non-zero since they reflect covariance that is driven by
1042 stimulus variations, and which is also dependent on the similarity of tuning properties of a pair.
1043 Separate covariance matrices were computed for leftward and rightward object direction classes
1044 and were averaged to get the covariance matrix used in Eqn. 5, $\Sigma = \frac{1}{2} \cdot (\Sigma_R + \Sigma_L)$. However,
1045 results were very similar if a single covariance matrix was used for both object direction classes.
1046 We also compared our results to performance of a standard decoder based on logistic regression
1047 ^{32, 57}, which was trained on data and does not require explicit computation of a covariance matrix.
1048 Cross-validated output of the logistic regression decoder produced nearly identical results
1049 (Extended Data Fig. 5).

1050 We took multiple approaches to decoding object direction from VIP and MSTl responses.

1051 1) *Separate decoders for each task reference frame.* In this approach, we trained separate
1052 decoders to classify object direction in world or head coordinates for each brain area. This
1053 approach assumes that the animal could have learned to read out VIP or MSTl activity in
1054 different ways for each task reference frame. For each task condition (head vs. world), FLD
1055 parameters (w_{world} , k_{world} and w_{head} , k_{head}) were computed separately from neural responses
1056 recorded in the corresponding task condition. Otherwise, each decoder was trained to report
1057 object direction across all stimulus conditions, including both self-motion directions and both
1058 Object+Visual and Object+Combined conditions. For each decoder, we randomly sampled 20
1059 trials (with replacement) from each neuron. 80% of these trials were used for computing the
1060 classifier parameters as described above, and the remaining 20% were used for computing
1061 classifier performance (fivefold cross-validation approach). This was repeated 1000 times and
1062 overall performance was found by averaging the results. For sliding window analyses, this
1063 resampling approach was repeated 100 times for each time bin.

1064 2) *Common decoder for both reference frame conditions.* We also investigated whether a
1065 single decoder with one set of common weights could correctly classify object direction in both
1066 head and world coordinates. This decoder examines the hypothesis that VIP or MSTl responses
1067 are modulated by self-motion signals in a task-dependent manner that allows for the same
1068 readout weights to be used for computing object direction in either head or world coordinates.
1069 For this analysis, FLD parameters were computed from neural responses that were recorded in
1070 both the head- and world-coordinate task conditions, as well as across both self-motion
1071 directions and both Object+Visual and Object+Combined conditions. All other aspects of the
1072 computation (e.g., cross-validation) were as described above for the separate decoders.

1073 3) *Cross-task decoders*. Because neural responses were obtained for identical conditions of
1074 object and background-dot motion (in screen coordinates) under both task reference frames, we
1075 could test how well a decoder trained to perform the task in a particular reference frame would
1076 perform when supplied with neural responses from the other task reference frame condition.
1077 Specifically, for the cross-task decoders, we trained a decoder to perform the world coordinate
1078 task based on neural responses from the head coordinate task, and we trained a decoder to
1079 perform the head coordinate task using responses from the world task. All other aspects of the
1080 decoding procedure were as described above. This approach allowed us to test how well the
1081 neural representations in VIP or MSTl could generalize across tasks.

1082

1083 **Statistics and reproducibility**

1084 In cases where the data met assumptions of normality, as assessed by Lilliefors test,
1085 parametric statistical tests were used, including t-tests, paired t-tests, and Pearson correlations.
1086 When data were not normally distributed, we used non-parametric tests, including the Wilcoxon
1087 rank sum test and the Wilcoxon signed rank test (for paired data).

1088 No statistical methods were used to pre-determine sample sizes but our sample sizes are
1089 comparable to, if not greater than, those reported in previous publications of a similar nature^{37, 52,}
1090⁵³. Data collection and analysis were not performed blind to the conditions of the experiments.
1091 However, all experimental conditions followed a standard protocol for each recording site and
1092 were entirely under computer control. Within each recording session, all stimulus conditions
1093 were block-randomized, such that the distinct stimuli were presented in a random order for each
1094 repetition. No animals were excluded from the analysis. Neurons were selected for analysis only

1095 based on their receptive field location (as described above), and if they could be recorded for at
1096 least 3 stimulus repetitions in the main experiment.

1097

1098

1099 Code availability. Custom analysis code was written using MATLAB (v. 2018a). Matlab scripts
1100 employed are available from the corresponding author upon reasonable request.

1101

1102 Data availability: The data that support the findings of this study are available from the
1103 corresponding author upon reasonable request.

1104

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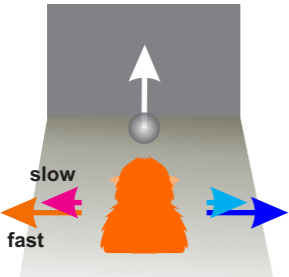
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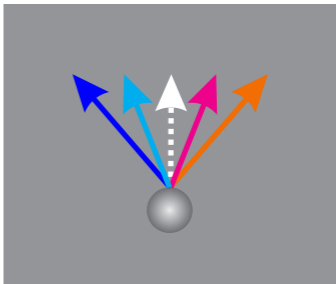
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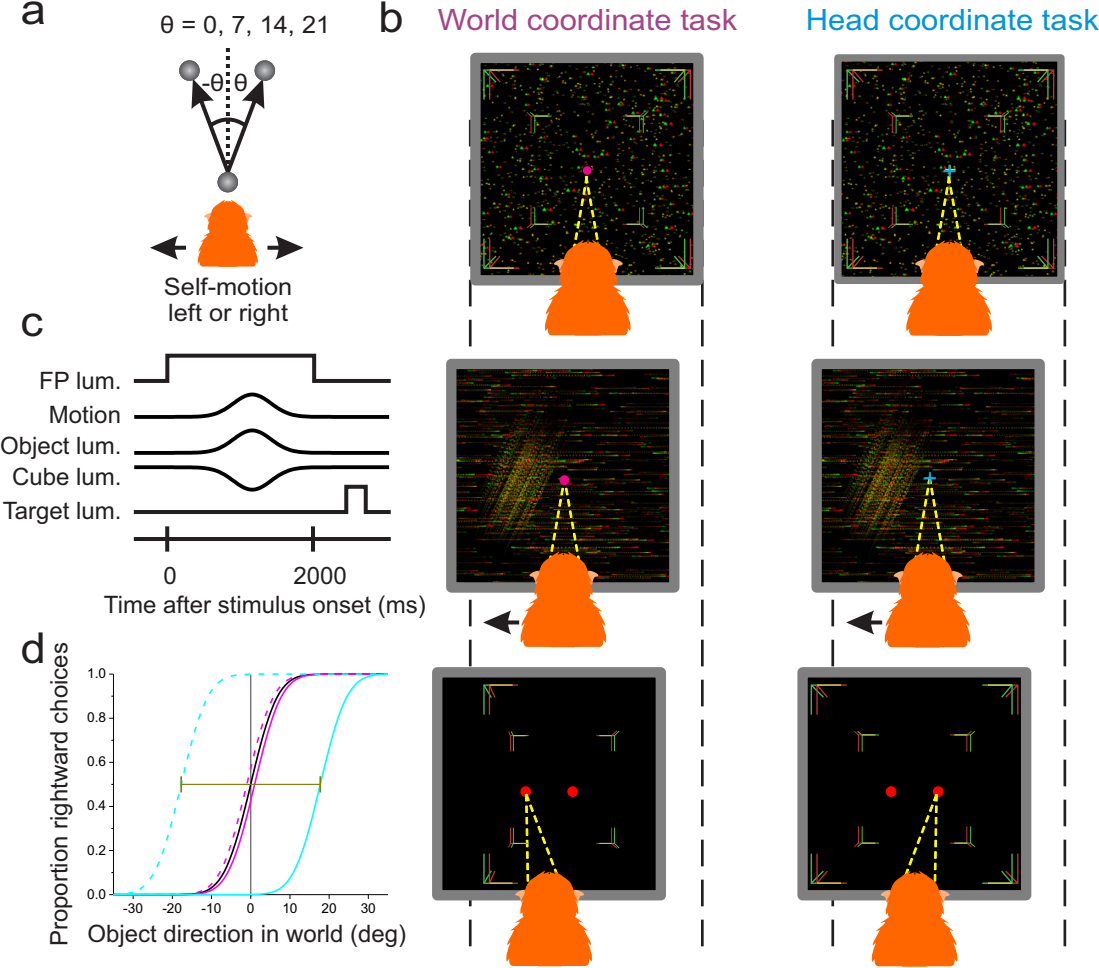
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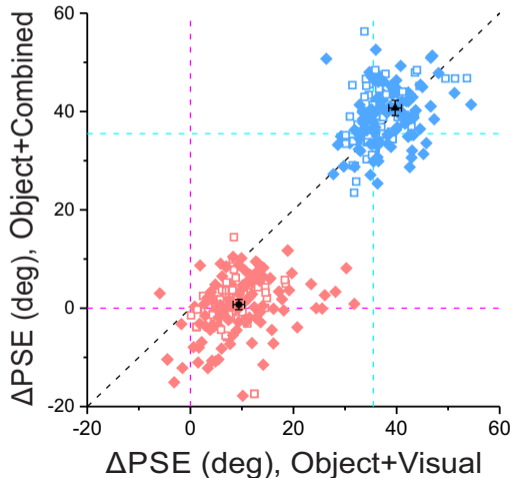
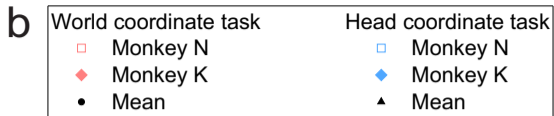
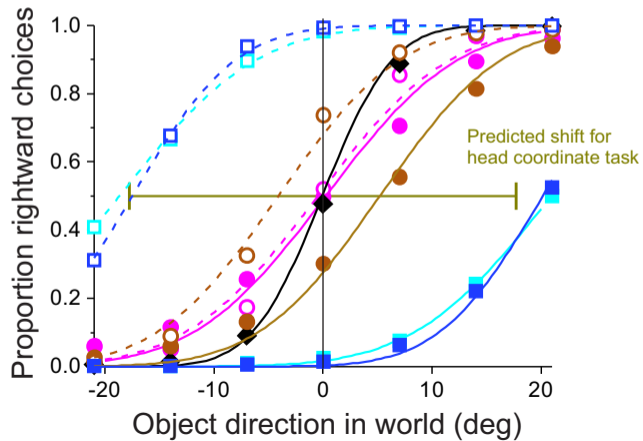
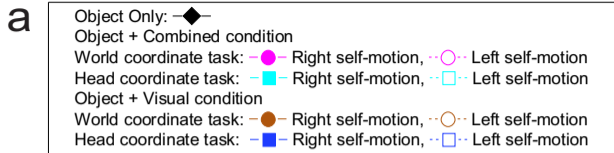
a World coordinate view

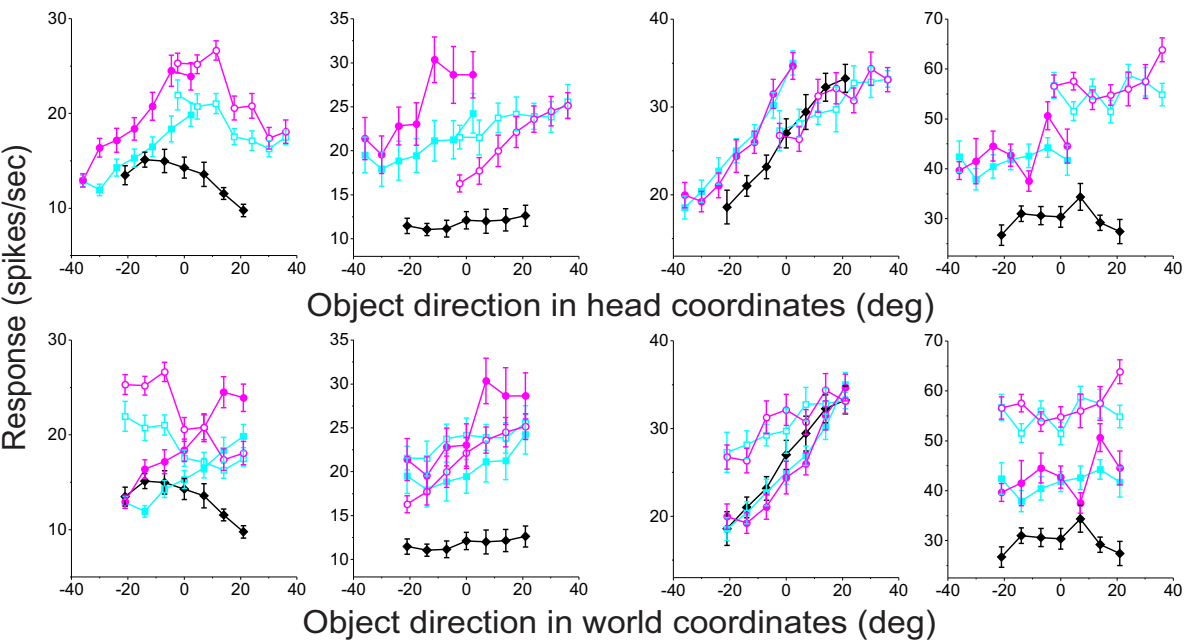
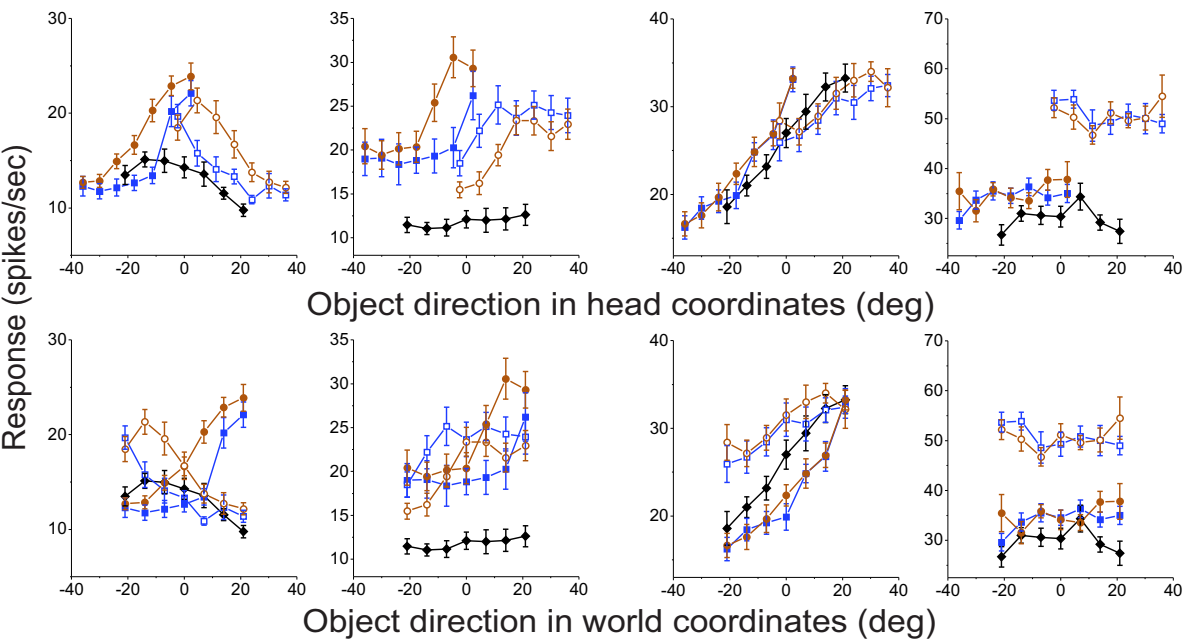


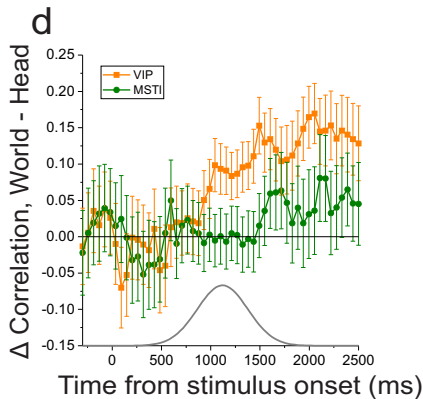
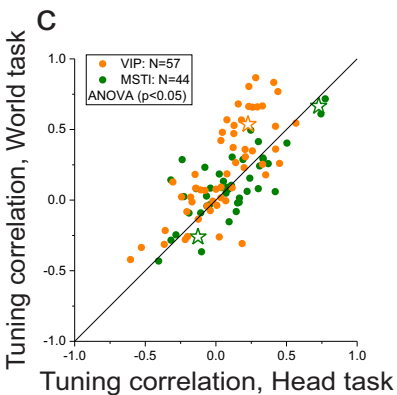
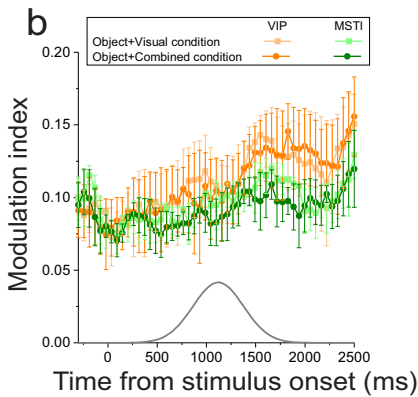
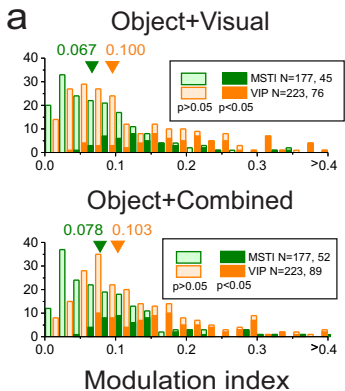
b Image motion in head coordinates



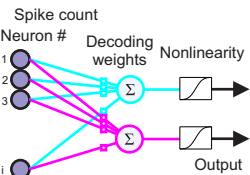




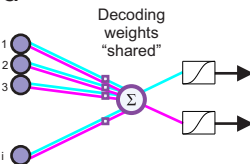
Object + Combined conditionObject Only: \blacklozenge World-coordinate task: \bullet Right self-motion, \circ Left self-motionHead-coordinate task: \square Right self-motion, \square Left self-motion**a****VIP****b****MSTI****Object + Visual condition**Object Only: \blacklozenge World-coordinate task: \bullet Right self-motion, \circ Left self-motionHead-coordinate task: \square Right self-motion, \square Left self-motion**c****VIP****d****MSTI**



a



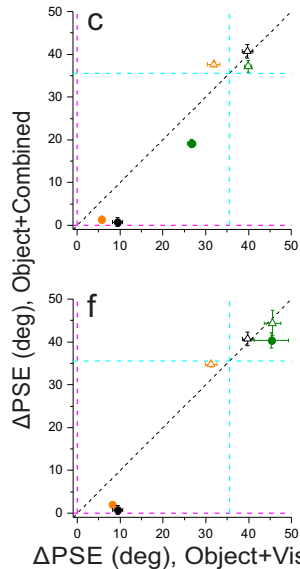
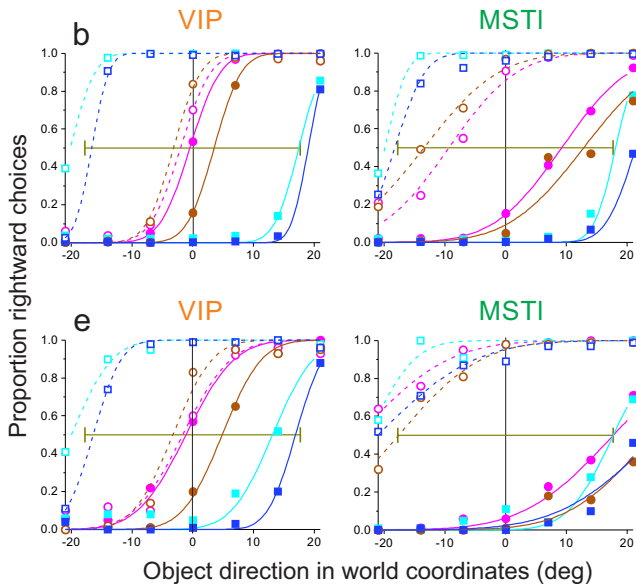
d

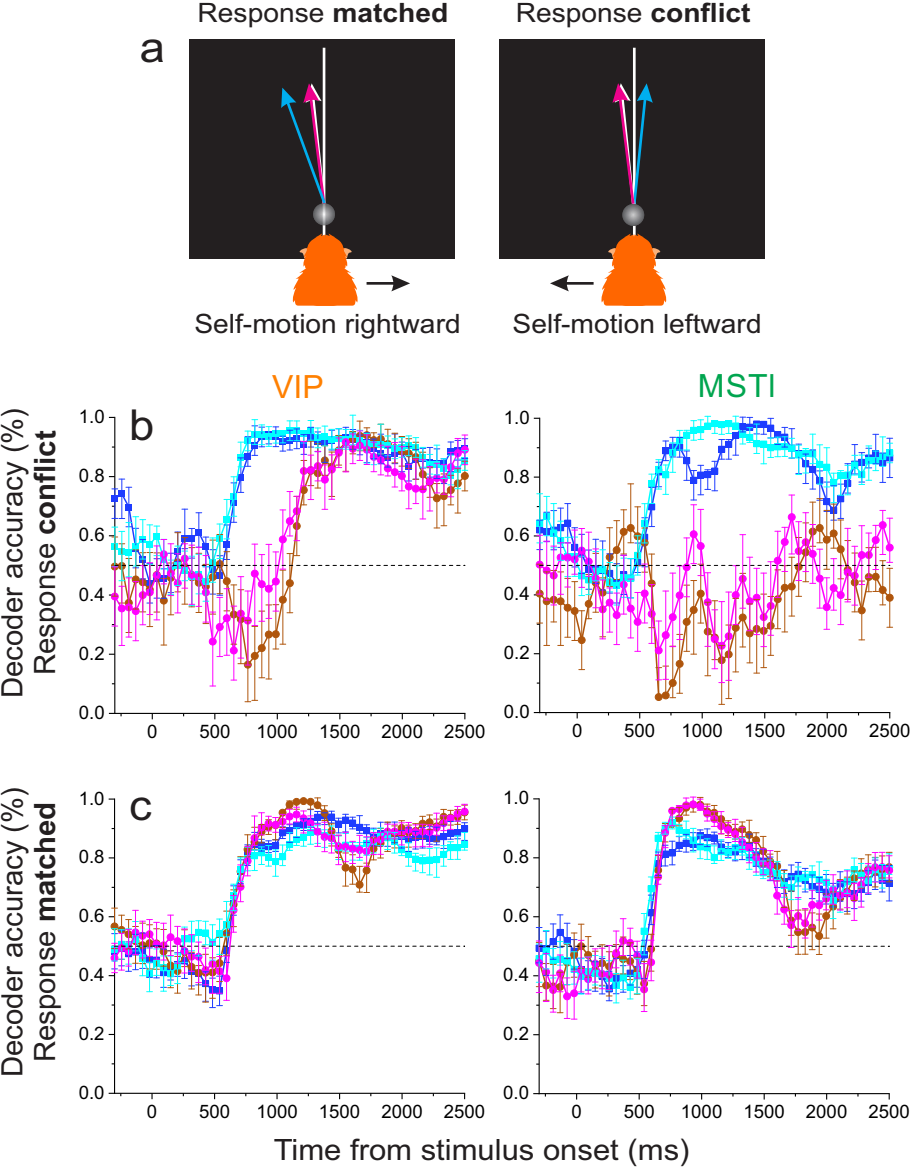


Object + Combined condition
 World coordinate task: ● Right self-motion, ○ Left self-motion
 Head coordinate task: ■ Right self-motion, □ Left self-motion

Object + Visual condition
 World coordinate task: ● Right self-motion, ○ Left self-motion
 Head coordinate task: ■ Right self-motion, □ Left self-motion

World Head
 ● △ behavior
 ● △ VIP decoder
 ● △ MSTI decoder

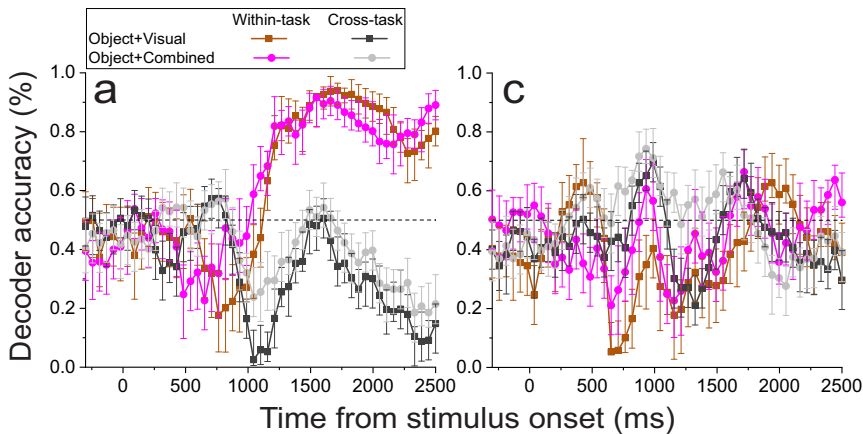




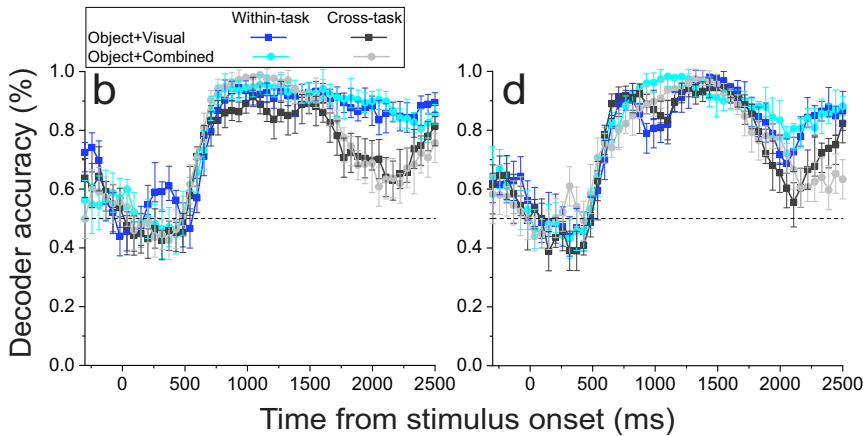
VIP

MSTl

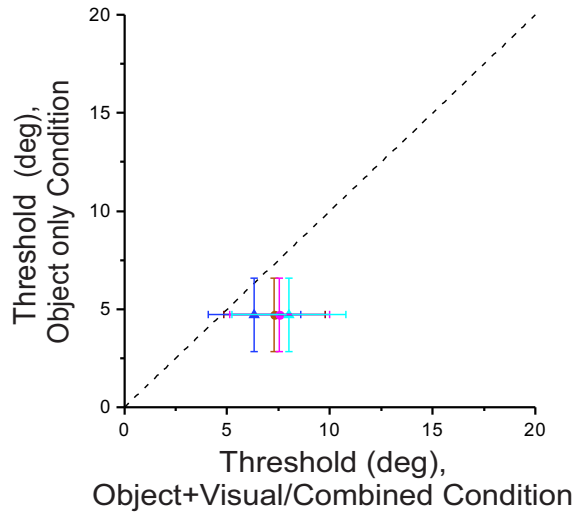
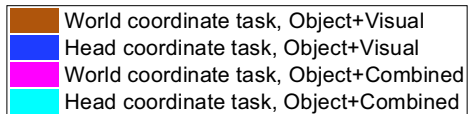
World coordinate task



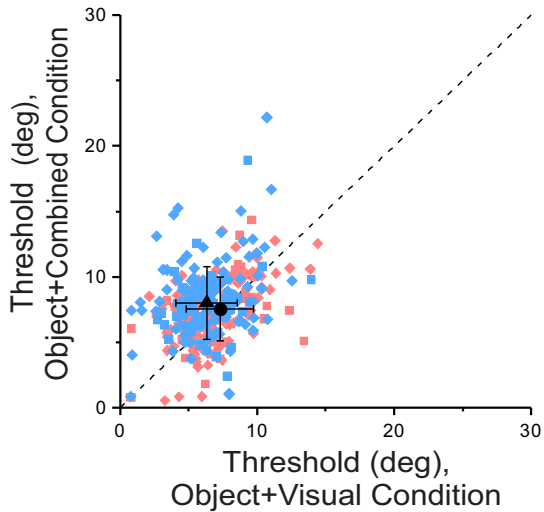
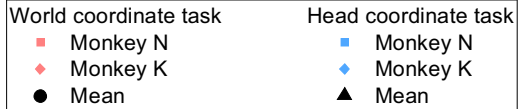
Head coordinate task

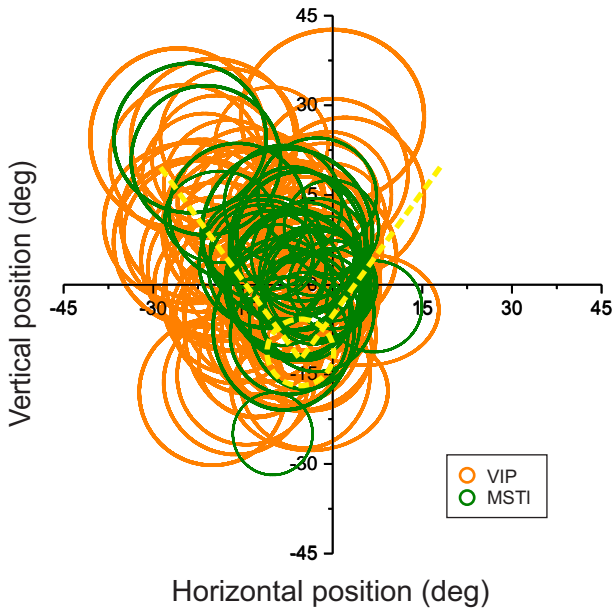


a



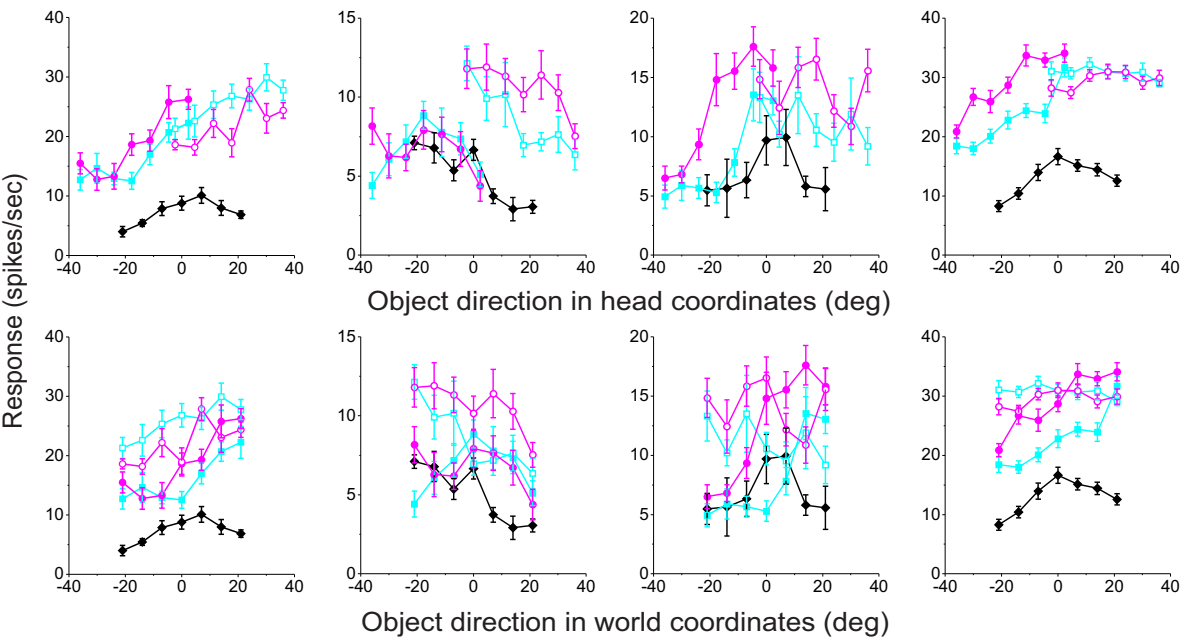
b



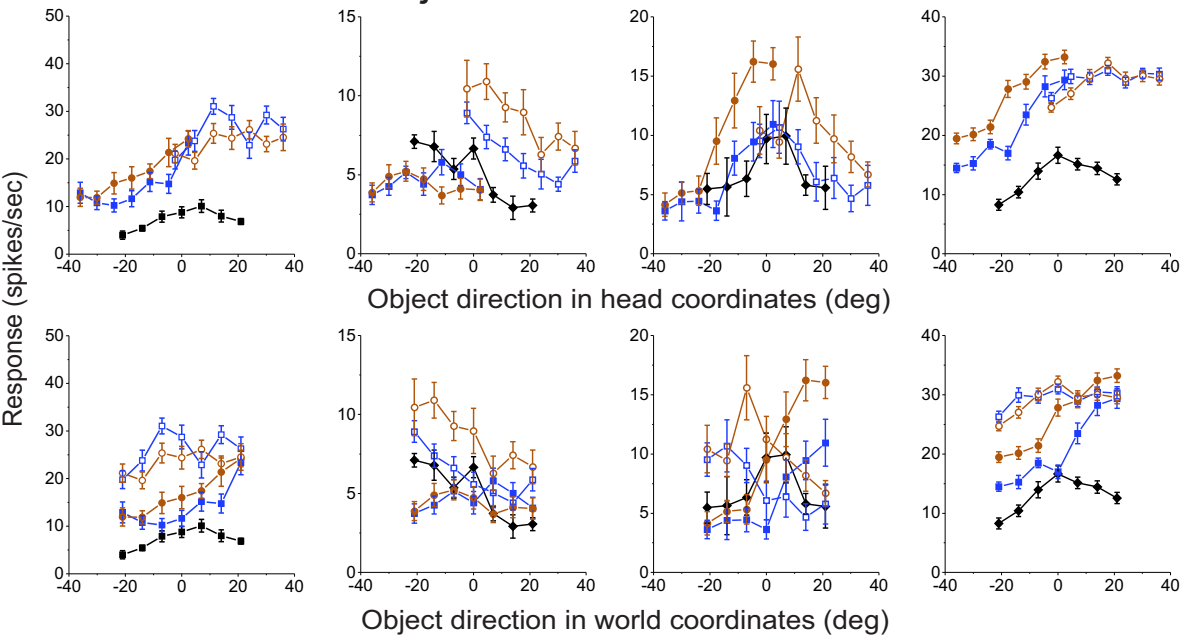


Object Only: ◆—
 Object + Combined condition
 World coordinate task: ●— Right self-motion, ○— Left self-motion
 Head coordinate task: ■— Right self-motion, □— Left self-motion
 Object + Visual condition
 World coordinate task: ●— Right self-motion, ○— Left self-motion
 Head coordinate task: ■— Right self-motion, □— Left self-motion

Object + Combined condition



Object + Visual condition

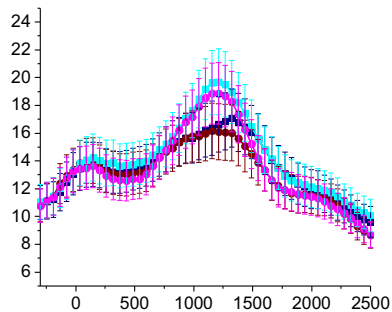
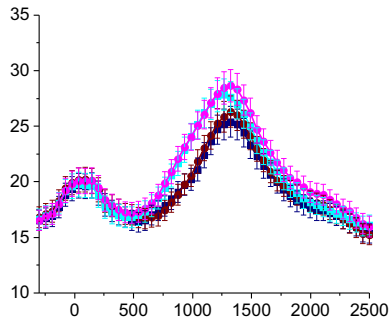


a

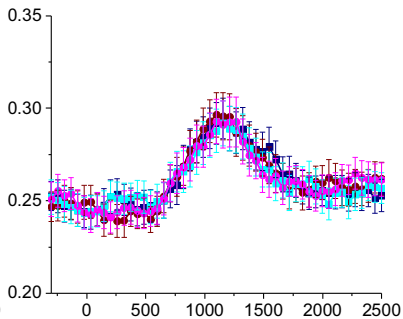
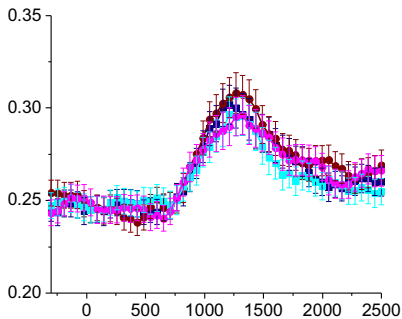
VIP

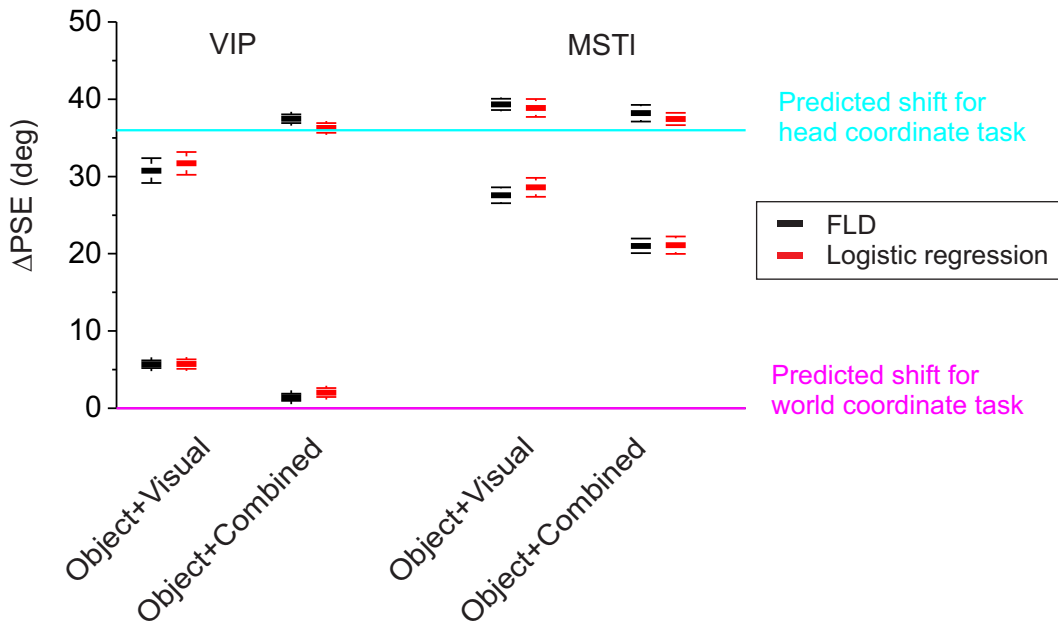
MSTl

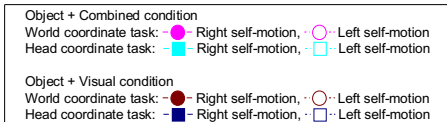
Responses (spikes/sec)

**b**

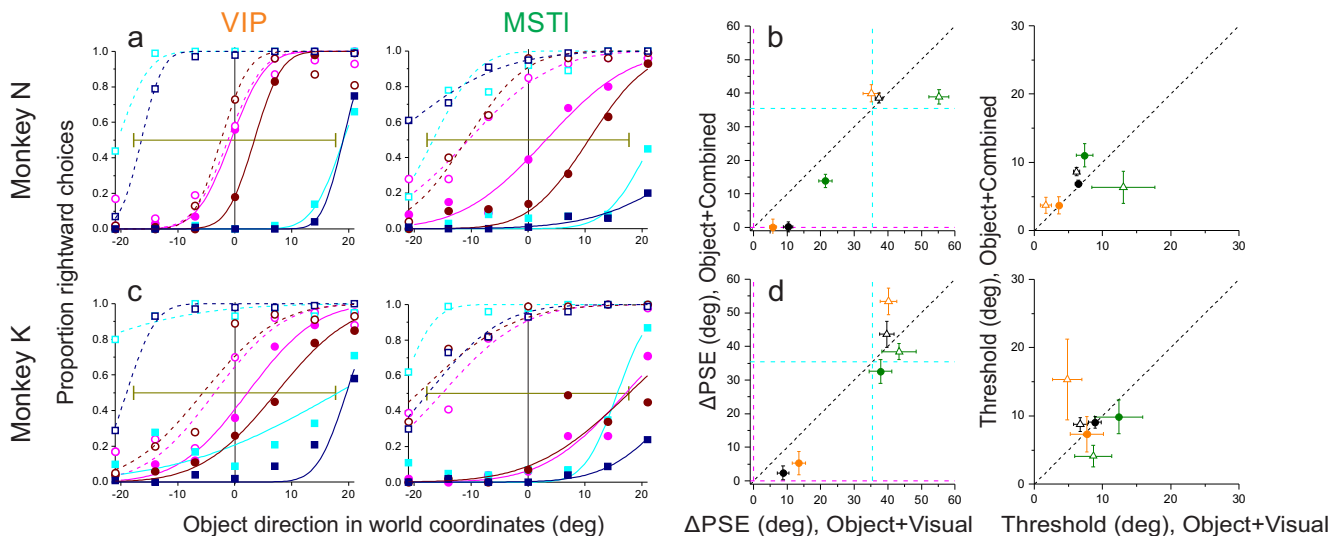
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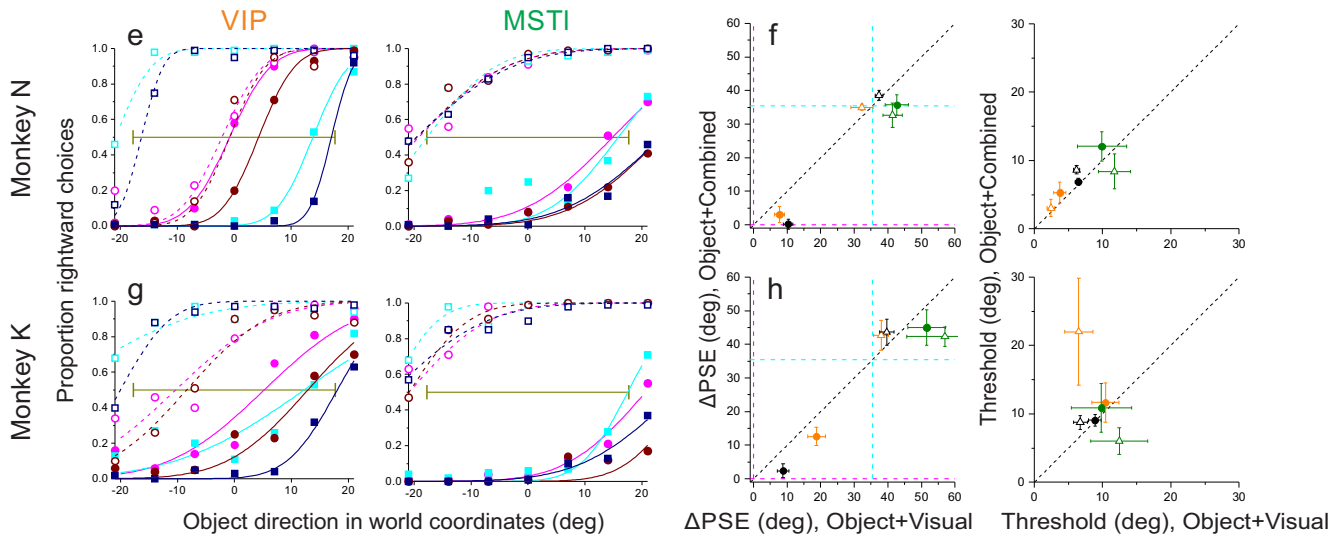


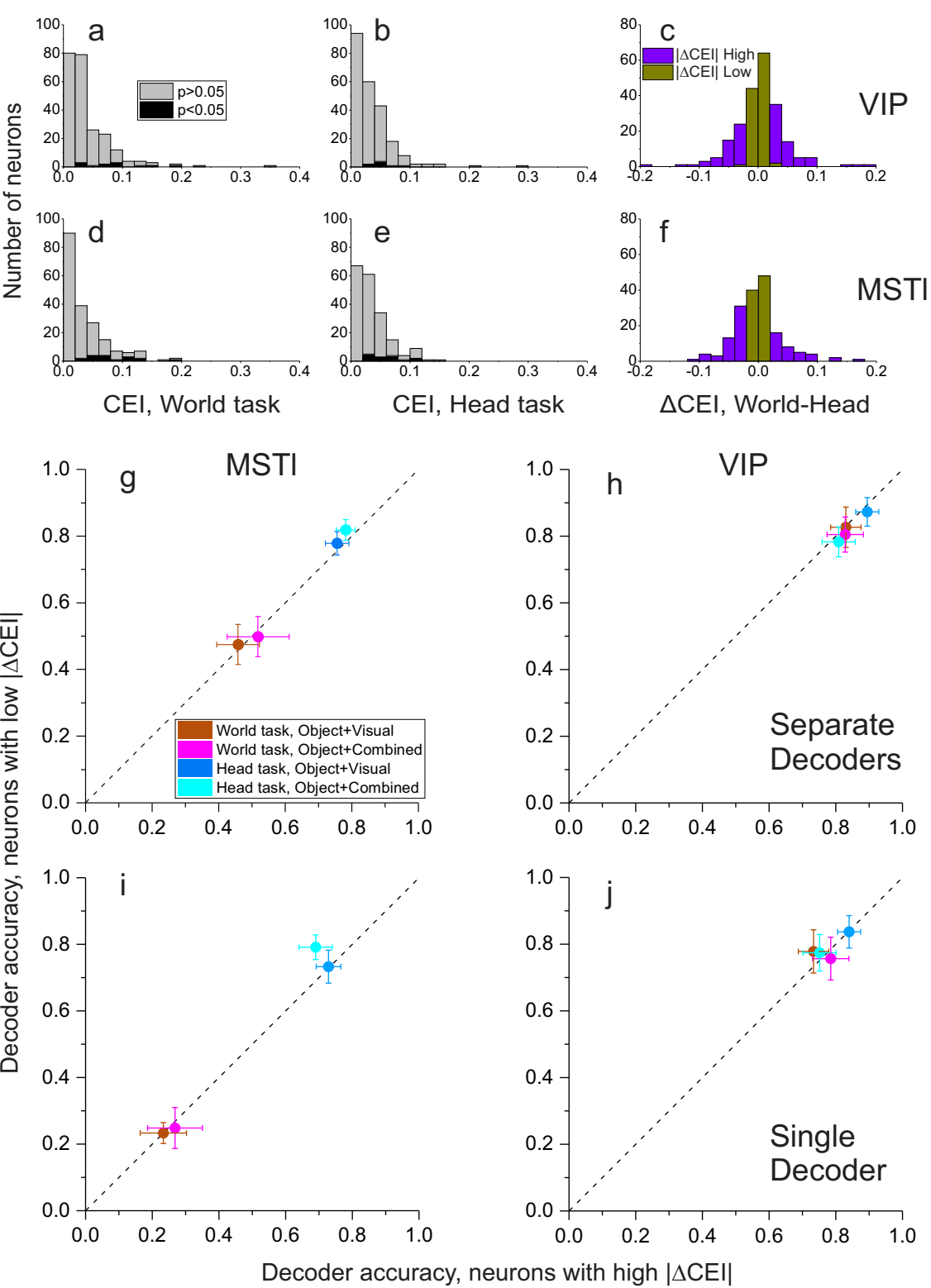


Separate decoders

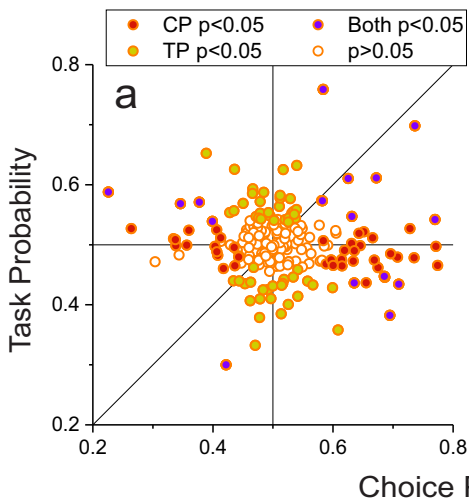


Single decoder

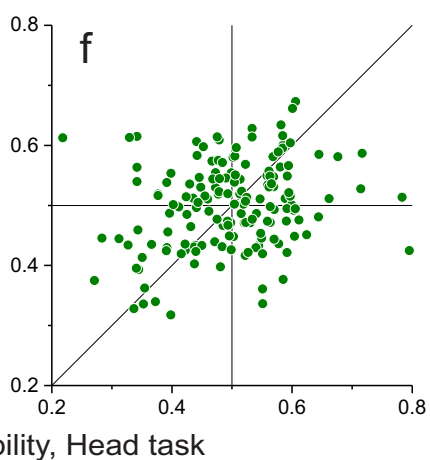
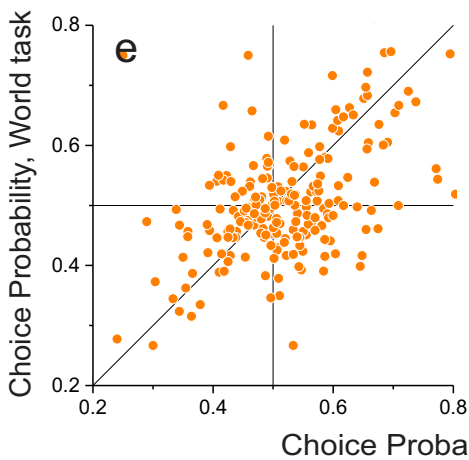
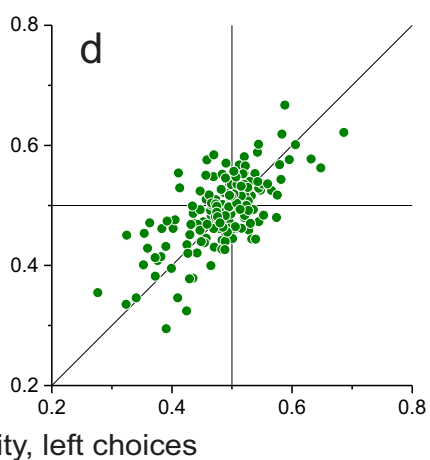
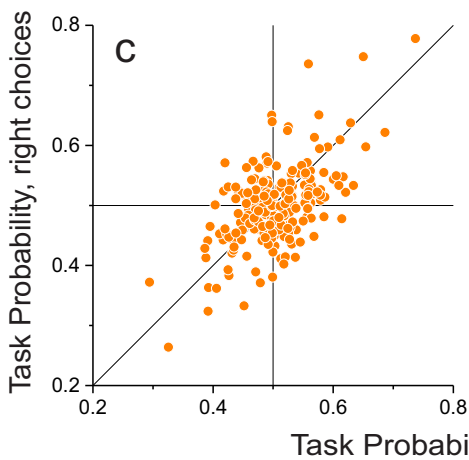
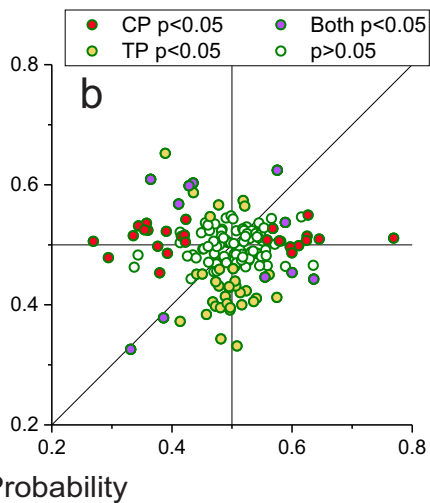




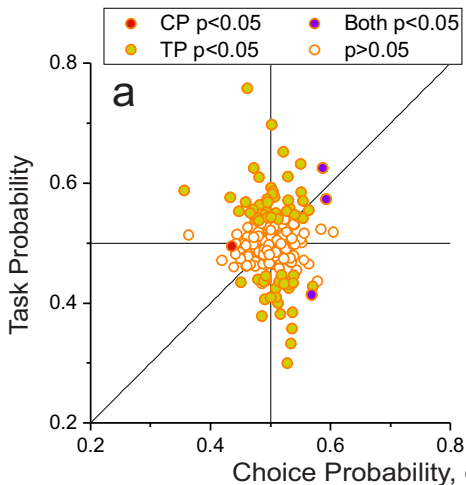
VIP



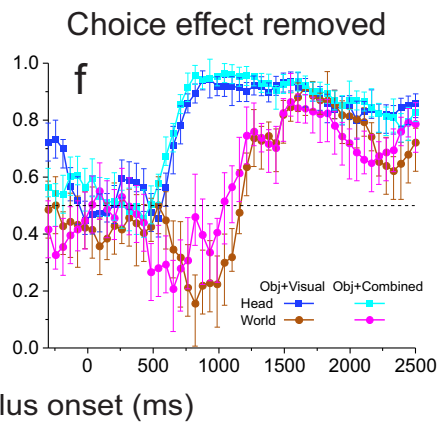
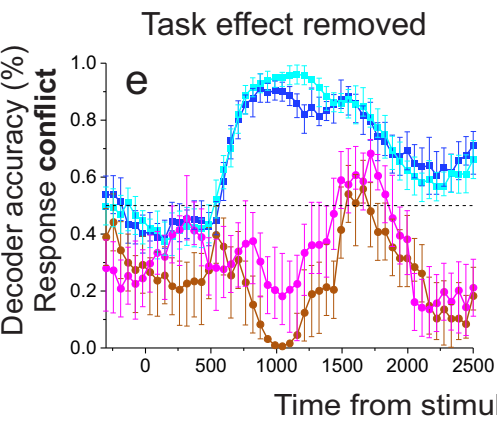
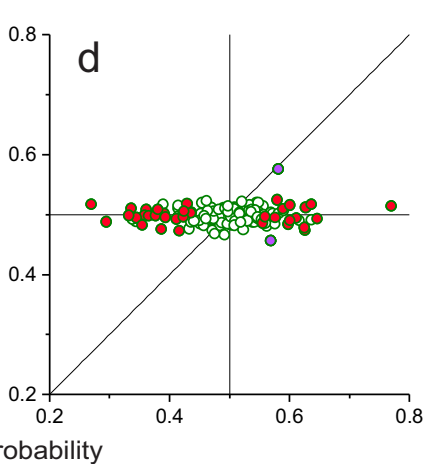
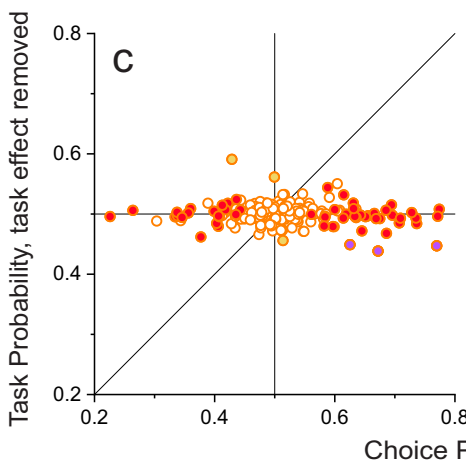
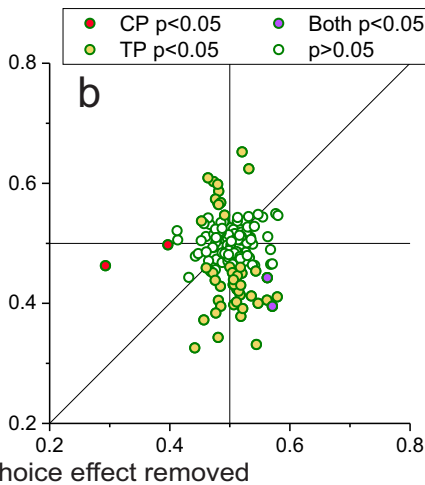
MSTl

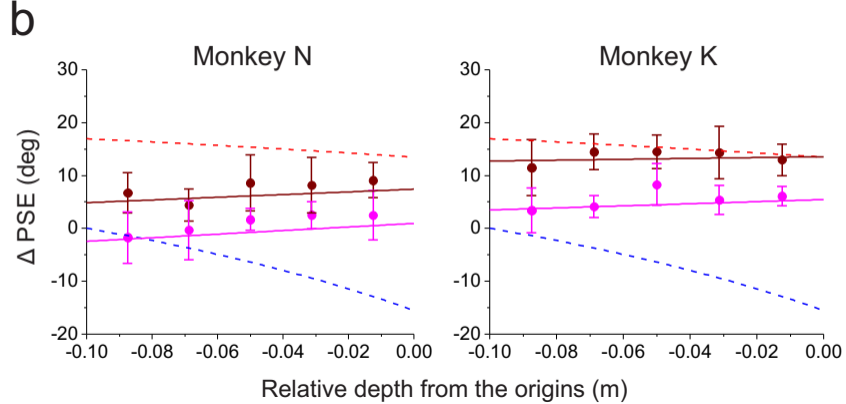
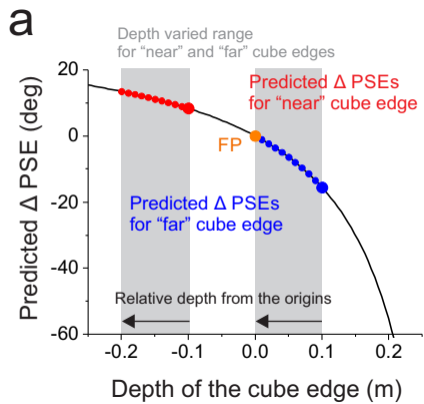


VIP



MSTl





Inventory of Supporting Information

Manuscript #: [NN-A69354B](#)

Corresponding author name(s): [Ryo Sasaki](#)

Instructions:

Please complete each of the Inventory Tables below to outline your Extended Data and Supplementary Information items.

There are four sections; *1. Extended Data*, *2A. Supplementary Information: Flat Files*, *2B. Supplementary Information: Additional Files*, and *3. Source Data*. Each section includes specific instructions. Please complete these tables as fully as possible. We ask that you avoid using spaces in your file names, and instead use underscores, i.e.: *Smith_ED_Fig1.jpg* **not** *Smith ED Fig1.jpg*

Please note that titles and descriptive captions will only be lightly edited, so please ensure that you are satisfied with these prior to submission.

If you have any questions about any of the information contained in this inventory, please contact the Editorial Assistant: neurosci@us.nature.com

1. Extended Data

Complete the Inventory below for all Extended Data figures.

- Keep Figure Titles to one sentence only
- File names should include the Figure Number. i.e.: *Smith_ED Fig1.jpg*
- Please be sure to include the file extension in the Filename. Note that Extended Data files must be submitted as .jpg, .tif or .eps files *only*
- All Extended Data figure legends must be provided in the Inventory below and should not exceed 300 words each (*if possible*)
- Please include Extended Data *ONLY* in this table

Figure #	Figure title One sentence only	Filename This should be the name the file is saved as when it is uploaded to our system. Please include the file extension. i.e.: <i>Smith_ED_Fig1.jpg</i>	Figure Legend If you are citing a reference for the first time in these legends, please include all new references in the Online Methods References section, and carry on the numbering from the main References section of the paper.
Extended Data Fig. 1	Summary of psychophysical thresholds (inverse of sensitivity) across task conditions.	Sasaki_ED_Fig1.eps	Extended Data Figure 1. Summary of psychophysical thresholds (inverse of sensitivity) across task conditions. (a) Average threshold for the Object Only condition (no self-motion) is plotted against average thresholds for the Object+Visual and Object+Combined conditions for the world (brown/magenta) and head (blue/cyan) coordinate tasks. Error bars represent 95% confident intervals. Averages taken over n=185 sessions across the two animals. (b) For each session, threshold in the Object+Combined condition is plotted against the corresponding threshold in the Object+Visual condition. Black symbols show mean thresholds

			and error bars represent 95% confidence intervals. Data from 128 sessions for Monkey N and 57 sessions for monkey K.
Extended Data Fig. 2	Summary of receptive field locations for populations of VIP (orange, N=66) and MSTI (green, N=44) neurons.	Sasaki_ED_Fig2.eps	Extended Data Figure 2. Summary of receptive field locations for populations of VIP (orange, N=66) and MSTI (green, N=44) neurons. Cells are included here if they had significant structure in receptive field maps obtained by reverse correlation (17% of VIP and 13% of MSTI neurons) or if they had clear hand-mapped receptive fields for which good estimates of RF center and size were obtained (13% of VIP neurons and 12% of MSTI neurons). Significant structure in reverse correlation maps was assessed by a two-sided permutation test ($p < 0.05$), in which we scrambled the relationship between response amplitude and stimulus location within the RF, as described previously ⁵⁶ . Ellipses approximate the RF dimensions and were derived either from a two-dimensional Gaussian fit (contour at half-maximal response) to receptive field maps obtained by reverse correlation (VIP: N=38; MSTI: N=23), or from hand mapping (VIP: N=28; MSTI: N=21). Coordinate (0, 0) represents the center of the visual display, where the fixation target was located. Yellow dashed lines represent the starting location of the moving object and the range of directions in head coordinates.
Extended Data Fig. 3	Data from four additional VIP neurons, illustrating diversity of effects of self-motion on tuning curves.	Sasaki_ED_Fig3.eps	Extended Data Figure 3. Data from four additional VIP neurons, illustrating diversity of effects of self-motion on tuning curves. Top: Object+Combined condition. Bottom: Object+Visual condition. Format as in Fig. 4. Error bars denote SEM ($n=10$ stimulus repetitions per datum).
Extended Data Fig. 4	Summary of time courses of average firing rates and directional selectivity.	Sasaki_ED_Fig4.eps	Extended Data Figure 4. Summary of time courses of average firing rates and directional selectivity. (a) Average response across all 223 VIP and 177 MSTI neurons is shown for each stimulus condition for both the head and world coordinate task conditions. For each neuron, responses were taken from the object motion direction that elicited the maximum firing

			rate. Error bars represent SEM. Color coding as in Fig. 7. Results were nearly identical if the responses of neurons were normalized before averaging. (b) Average direction discrimination index (DDI) for populations of VIP (n=223) and MSTl (n=177) neurons (see Methods, Eqn. 2). DDI values were computed separately for leftward and rightward self-motion and then averaged for each neuron. Error bars represent 95% confidence intervals. For this figure, both average responses and DDI values were computed within a 300 ms sliding time window that was advanced across the stimulus epoch in steps of 50 ms.
Extended Data Fig. 5	Decoder results are robust to the type of classifier used.	Sasaki_ED_Fig5.eps	Extended Data Figure 5. Decoder results are robust to the type of classifier used. Black data points represent results from the FLD classifier used in all main figures. Red data points show results from a logistic regression decoder. For this comparison, the same population responses were used for training and testing each decoder. The results are very robust to the type of decoder used. Error bars represent 95% confidence intervals (across n=1000 bootstraps).
Extended Data Fig. 6	Comparison of decoder results across animals.	Sasaki_ED_Fig6.eps	Extended Data Figure 6. Comparison of decoder results across animals. (a-d) Results for separate decoders trained to perform the world and head coordinate tasks. Format as in Figure 6. Each row shows results separately for each animal. Pink and cyan dashed lines in panels b and d: expected Δ PSE for perfect performance in the world and head coordinate tasks, respectively. Error bars in panels b and d represent 95% confidence intervals (across n=1000 bootstraps). (e-h) Results for the single decoder, shown separately for each animal. Decoders were trained separately using responses from each animal, yet main results are conserved across subjects. Error bars represent 95% confidence intervals (across n=1000 bootstraps). Format as in panels a-d.
Extended Data Fig. 7	Effect of partial cube frame on single-unit	Sasaki_ED_Fig7.eps	Extended Data Figure 7. Effect of partial cube frame on single-unit responses and population decoding. (a, d)

	<p>responses and population decoding.</p>		<p>Distributions of the cube effect index (CEI, see Methods) for areas VIP and MSTl, respectively, in the world coordinate task. Black and gray shading denotes neurons with CEI values that are significantly different from zero and non-significant, respectively (two-sided permutation test, $p < 0.05$). (b, e) Distributions of CEI for VIP and MSTl, respectively, in the head coordinate task condition. (c, f) Distributions of the difference in CEI (ΔCEI) between world and head task conditions for VIP and MSTl, respectively. Green and purple shading indicates a median split of the data based on the absolute value, ΔCEI . (g, h) Comparison of decoder accuracy (proportion correct) for populations of neurons with above-median ΔCEI (abscissa) and below-median ΔCEI (ordinate) values, for areas VIP and MSTl, respectively. Error bars represent 95% confidence intervals (across $n=1000$ bootstraps). Data in these panels come from decoders that were trained separately for the world and head coordinate task conditions. (i, j) Same as panels g and h, except for a single decoder trained to perform the task across both reference frame conditions. Format as in g, h.</p>
<p>Extended Data Fig. 8</p>	<p>Summary of choice-related and task-related response modulations.</p>	<p>Sasaki_ED_Fig8.eps</p>	<p>Extended Data Figure 8. Summary of choice-related and task-related response modulations. (a) Scatter plot of task probability (TP) and choice probability (CP) values for VIP neurons ($N=223$). Color of the symbol centers corresponds to significance of TP and CP values as follows: blue center, both TP and CP are significantly different from 0.5 (two-sided permutation test, $p < 0.05$); red center, only CP is significantly different from 0.5; gold center, only TP is significantly different from 0.5; white center, neither TP nor CP is significant. The observation that TP and CP values are largely uncorrelated here is an empirical observation that is not enforced by the analysis. (b) Scatter plot of TP and CP values for MSTl neurons ($N=177$). Symbol center color conventions as in panel a. (c) Scatter plot of TP values for VIP neurons</p>

			computed separately for right and left choices (N=223). (d) Same as panel c but for MSTl neurons (N=177). (e) Scatter plot comparing CP values for the world and head coordinate task conditions (N=223). (f) Same as panel e but for MSTl neurons (N=177).
Extended Data Fig. 9	Effects of selectively removing choice- or task-related response modulations.	Sasaki_ED_Fig9.eps	Extended Data Figure 9. Effects of selectively removing choice- or task-related response modulations. (a) Scatter plot of TP and CP values for VIP (N=223) after selective removal of choice-related response modulations (see Methods for details). Format as in Extended Data Fig. 8a. (b) Same as panel a except for MSTl (N=177). Format as in Extended Data Fig. 8b. (c) Scatter plot of TP and CP values for VIP after selective removal of task-related response modulations. (d) Same as panel c, except for MSTl. (e) Time course of decoder performance based on activity of 223 VIP neurons on response conflict trials, after removal of task-related response modulations. Data are shown for the case of separate decoders for world and head coordinate task conditions. Format as in Fig. 7b. Error bars represent 95% confidence intervals (across n=100 bootstraps). (f) Time course of VIP decoder performance, as in panel e, but after removal of choice-related response modulations.
Extended Data Fig. 10	Results from behavioral control sessions in which the depth of the partial cube was varied across trials.	Sasaki_ED_Fig10.eps	Extended Data Figure 10. Results from behavioral control sessions in which the depth of the partial cube was varied across trials. (a) Predicted Δ PSE values are shown as a function of the depth of the partial cube. Red and blue data points show predicted Δ PSE values and depths for the near and far edges of the cube. (b) Dashed curves replot the predictions from panel a, where the horizontal axis is now depth relative to the origins for the near (red) and far (blue) cube edges (where the origins are the farthest depths for each edge). Data points represent behavioral Δ PSE values for the two monkeys (n=7 sessions for each animal); magenta and brown data points show results for the Object+Combined and

			Object+Visual conditions. Error bars show 95% confidence intervals, and lines show regression fits. The slopes of the linear fits were not significantly different from zero for either animal or either self-motion condition (two-tailed t-test, $p > 0.15$ for all four cases).
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Delete rows as needed to accommodate the number of figures (10 is the maximum allowed).

2. Supplementary Information:

A. Flat Files

Complete the Inventory below for all additional textual information and any additional Supplementary Figures, which should be supplied in one combined PDF file.

- **Row 1:** A combined, flat PDF containing any Supplementary Methods, Discussion, Equations, Notes, Additional Supplementary Figures, simple tables, and all associated legends. Only one such file is permitted.
- **Row 2:** Nature Research's Reporting Summary; please provide an updated Summary, fully completed, without any mark-ups or comments. **Please note that this is a required document.**

Item	Present?	Filename This should be the name the file is saved as when it is uploaded to our system, and should include the file extension. The extension must be .pdf	A brief, numerical description of file contents. i.e.: <i>Supplementary Figures 1-4, Supplementary Discussion, and Supplementary Tables 1-4.</i>
Supplementary Information	No		

Reporting Summary	Yes	Sasaki_nr-reporting-summary_RENEWED
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B. Additional Supplementary Files

Complete the Inventory below for all additional Supplementary Files that cannot be submitted as part of the Combined PDF.

- Do not list Supplementary Figures in this table (see section 2A)
- Where possible, include the title and description within the file itself
- Spreadsheet-based tables and data should be combined into a workbook with multiple tabs, not submitted as individual files.
- Please note that the *ONLY* allowable types of additional Supplementary Files are:
 - Supplementary Tables
 - Supplementary Audio
 - NMR Data
 - Computational Data
 - Supplementary Videos
 - Supplementary Data
 - Cryo-EM Data
 - Suppl. Software

Type	Number If there are multiple files of the same type this should be the numerical indicator. i.e. "1" for Video 1, "2" for Video 2, etc.	Filename This should be the name the file is saved as when it is uploaded to our system, and should include the file extension. i.e.: <i>Smith_Supplementary_Video_1.mov</i>	Legend or Descriptive Caption Describe the contents of the file
Choose an item.			
Choose an item.			
Choose an item.			
Choose an item.			

Choose an item.			
Choose an item.			

Add rows as needed to accommodate the number of files.

3. Source Data

Complete the Inventory below for all Source Data files.

- Acceptable types of Source Data are:
 - Statistical Source Data
 - Plain Text (ASCII, TXT) or Excel formats only
 - One file for each relevant Figure, containing all source data
 - Full-length, unprocessed Gels or Blots
 - JPG, TIF, or PDF formats only
 - One file for each relevant Figure, containing all supporting blots and/or gels

Figure	Filename This should be the name the file is saved as when it is uploaded to our system, and should include the file extension. i.e.: <i>Smith_SourceData_Fig1.xls</i> , or <i>Smith_Unmodified_Gels_Fig1.pdf</i>	Data description i.e.: Unprocessed Western Blots and/or gels, Statistical Source Data, etc.
Source Data Fig. 1		
Source Data Fig. 2		
Source Data Fig. 3		
Source Data Fig. 4		
Source Data Fig. 5		
Source Data Fig. 6		
Source Data Fig. 7		

Source Data Fig. 8		
Source Data Extended Data Fig. 1		
Source Data Extended Data Fig. 2		
Source Data Extended Data Fig. 3		
Source Data Extended Data Fig. 4		
Source Data Extended Data Fig. 5		
Source Data Extended Data Fig. 6		
Source Data Extended Data Fig. 7		
Source Data Extended Data Fig. 8		
Source Data Extended Data Fig. 9		
Source Data Extended Data Fig. 10		