1	Reading Modality Modifies Reading Network:
2	Insights from Neural basis of
3	Braille in Proficient Blind Readers
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11 Abstract

12 The neural basis of reading is highly consistent across a variety of languages and visual scripts. 13 An unanswered question is whether the sensory modality of symbols influences the neural basis 14 of reading. According to the modality-invariant view, reading depends on the same neural 15 mechanisms regardless of the sensory input modality. Consistent with this idea, previous studies 16 find that the visual word form area (VWFA) within the ventral occipitotemporal cortex 17 (vOTC) is active when blind individuals read Braille by touch. However, connectivity-based 18 theories of brain function suggest that the neural entry point of written symbols (touch vs. vision) 19 may influence the neural architecture of reading. We compared the neural basis of the visual 20 print (sighted n=15) and tactile Braille (congenitally blind n=19) in proficient readers using 21 analogous reading and listening tasks. Written stimuli varied in word-likeness from real words to 22 consonant strings and non-letter shape strings. Auditory stimuli consisted of words and backward 23 speech sounds. Consistent with prior work, vOTC was active during Braille and visual reading. 24 However, in sighted readers, visual print elicited a posterior/anterior vOTC word-form gradient: 25 anterior vOTC preferred larger orthographic units (words), middle vOTC preferring consonant 26 strings, and posterior vOTC responded to shapes (i.e., lower-level physical features). No such 27 gradient was observed in blind readers of Braille. Consistent with connectivity predictions, in 28 blind Braille readers, posterior parietal cortices (PPC) and parieto-occipital areas were recruited 29 to a greater degree and PPC contained word-preferring patches. Lateralization of Braille in blind 30 readers was predicted by laterality of spoken language, as well as by reading hand. These 31 results suggested that the neural basis of reading is influenced by symbol modality and support 32 connectivity-based views of cortical function.

33 Highlights

- Only sighted but not blind (Braille) readers show a posterior/anterior vOTC lexicality
 gradient
- 36 2. Posterior parietal cortex distinctively contributes to Braille reading.
- 37 3. Lateralization of spoken language and reading hand predict lateralization of Braille
- 38 4. The sensory modality of written symbols influences the neural basis of reading

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39 Keywords

40 Braille reading; blindness; reading; VWFA; posterior parietal cortex

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42

43 Introduction

44 Written language is among the most impressive human cultural achievements. The capacity to

45 record and transmit information over space and time has enabled the accumulation of scientific,

- 46 technological, and historical knowledge across generations and continents. How does the human
- 47 brain accommodate this cultural invention, which emerged only approximately 5,000 years ago?
- 48 Despite being a recent cultural invention, the neural basis of reading is highly consistent across a
- 49 variety of languages and visual scripts, including alphabetic, logographic (e.g., Chinese), and
- 50 syllabic writing systems (e.g., Japanese Kana) (Bolger, Perfetti, & Schneider, 2005; Feng et al.,
- 51 2020; Hu et al., 2010; Krafnick et al., 2016; Nakamura et al., 2012; Rueckl et al., 2015). All of
- 52 these reading systems engage regions within the left lateral ventral occipitotemporal cortex
- 53 (vOTC) (Baker et al., 2007; Cohen et al., 2000; Dehaene & Cohen, 2011; Dehaene et al., 2010).
- 54 A region in the left lateral vOTC has been termed the 'visual word form area' (VWFA) because

of its preferential response to written words and letter combinations over other visual stimuli.

- 56 The VWFA is situated within a posterior/anterior processing gradient. During reading, visual
- 57 symbols are first processed by early visual cortices and posterior portions of vOTC, which
- 58 represent simple visual features (e.g., line junctions) (Dehaene, Cohen, Sigman, & Vinckier,
- 59 2005; DiCarlo & Cox, 2007). By contrast, the middle and anterior potions of lateral vOTC are
- 60 specialized for progressively larger orthographic units, from written letters, letter
- 61 combinations/bigrams, and finally whole words (Binder, Medler, Westbury, Liebenthal, &
- 62 Buchanan, 2006; Cohen et al., 2000; Dehaene, Cohen, Sigman, & Vinckier, 2005; Dehaene et
- al., 2004; Glezer, Jiang, & Riesenhuber, 2009; Lerma-Usabiaga, Carreiras, & Paz-Alonso, 2018;
- 64 Purcell, Shea, & Rapp, 2014; Vinckier et al., 2007).
- 65 An open question is whether the vOTC posterior/anterior processing stream is the only way for
- 66 the brain to implement reading and, relatedly, why the neural basis of reading takes this
- 67 particular form. Examining the neural basis of tactile Braille offers unique insights into these
- 68 questions. Specifically, we can ask whether and how the sensory modality of written symbols
- 69 influences the neural basis of reading.

70 Tactile Braille reading achieves similar behavioral goals for people who are blind as visual print

71 reading does for the sighted: rapid access to linguistic meaning from a temporally stable

72 symbolic record. Proficient blind readers can read upwards of 200 words per minute by passing

the fingers along lines of Braille text, in which words are written as patterns of raised dots

74 (Millar, 2003). Each Braille character consists of dots positioned in a three-rows-by-two-

75 columns matrix. A single Braille character can be used to represent a letter, number, or

76 punctuation mark. In the most commonly used form of English Braille (Grade 2 Braille), Braille

characters also stand for frequent letter combinations (e.g., EA *; , OW *;) and whole words (e.g.,

78 e = every, tm = tomorrow) (<u>http://www.brl.org</u>) (Millar, 2003).

79 Consistent with a modality-invariant view of reading, several recent studies have reported that

80 the neural basis of Braille reading and that of visual print reading depend on similar vOTC

81 mechanisms (Büchel, Price, & Friston, 1998a; Debowska et al., 2016; Rączy et al., 2019; Reich,

82 Szwed, Cohen, & Amedi, 2011). Visual print and tactile Braille reading elicit activation peaks at

the anatomical location of the 'VWFA' in both sighted and blind readers (Debowska et al., 2016;

84 Dzięgiel-Fivet et al., 2021; Kim, Kanjlia, Merabet, & Bedny, 2017; Rączy et al., 2019; Reich et

al., 2011; Siuda-Krzywicka et al., 2016). In sighted adults who are trained to recognized Braille

86 words, transcranial magnetic stimulation (TMS) to the VWFA disrupts reading accuracy (Bola et

87 al., 2019; Siuda-Krzywicka et al., 2016). A recent study also found similar repetition suppression

88 effects in vOTC for tactile (blind readers) and visual (sighted readers) pseudowords (Rączy et al.,

89 2019). This evidence supports the idea that reading depends on the same neural mechanisms in

90 vOTC, regardless of symbol modality (i.e., touch vs. vision).

91 At the same time, both theoretical considerations and empirical evidence suggest that the neural 92 basis of tactile Braille and visual print reading may differ in important ways that have not been 93 fully tested. In sighted readers, posterior portions of vOTC receive visual written forms from 94 early visual cortices and pass this information along the posterior/anterior orthographic gradient 95 (e.g., Dehaene et al., 2005). By contrast, in people who are blind, Braille information enters the 96 cortex at primary somatosensory cortex (S1), making a posterior/anterior gradient unlikely. A 97 number of imaging studies also find that Braille reading activates visual areas outside of vOTC 98 in people who are blind, including V1 and dorsal occipital areas (Cohen et al., 1997, 1999;

99 Gizewski, Gasser, De Greiff, Boehm, & Forsting, 2003; Kupers et al., 2007; Melzer et al., 2001;

100 Sadato et al., 1998, 1996). TMS to the occipital pole and midoccipital cortex disrupts Braille

101 reading (Cohen et al., 1997, 1999; Kupers et al., 2007). This suggests that vOTC may not make a

102 unique contribution to Braille reading in the same way that it does to visual reading.

103 Moreover, visual cortices of people who are born blind, including vOTC and early visual areas

104 (V1-V3), are recruited for non-visual functions apart from Braille (e.g., Amedi, Raz, Pianka,

105 Malach, & Zohary, 2003; Büchel et al., 1998b; Burton, Snyder, Diamond, & Raichle, 2002;

106 Gougoux, Zatorre, Lassonde, Voss, & Lepore, 2005; Kanjlia, Loiotile, Harhen, & Bedny, 2021;

107 Kujala, Alho, Paavilainen, Summala, & Naatanen, 1992; Sathian, 2005). Particularly relevant for

108 the neural basis of Braille, large swaths of blind 'visual' cortex, including portions of V1,

109 participate in processing spoken language, including high-level semantic and grammatical

110 information (Bedny, Pascual-Leone, Dodell-Feder, Fedorenko, & Saxe, 2011; Bedny,

111 Richardson, & Saxe, 2015; Burton, Snyder, Diamond, & Raichle, 2002; Noppeney, Friston, &

112 Price, 2003; Röder, Stock, Bien, Neville, & Rösler, 2002; Watkins et al., 2012). Indeed, there is

113 evidence that the anatomical location of the 'VWFA' shows larger responses to spoken language

and responds to the grammatical structure of spoken sentences in people who are blind, more so

115 than in people who are sighted (Dzięgiel-Fivet et al., 2021; Kim et al., 2017). This pattern

116 suggests possible involvement in high-order language processing, rather than a reading-specific

117 role in blindness. Furthermore, since the anatomical distribution of written language is believed

118 to be influenced by the anatomical distribution of spoken language (Behrmann & Plaut, 2013;

119 Hannagan & Grainger, 2013; Saygin et al., 2016; Stevens, Kravitz, Peng, Tessler, & Martin,

120 2017a), recruitment of the visual cortex for language processing could itself modify the neural

121 basis of Braille reading in blind people. For example, we might expect Braille to recruit occipital

122 regions that are connected to visual networks recruited for spoken language. Together this

123 evidence suggests that the anatomical distribution and function role of visual cortices in blind

124 Braille readers and sighted visual readers may not be equivalent and merits further investigation.

125 There are also reasons to hypothesize that tactile Braille reading may differentially recruit

126 networks outside of the visual system, specifically the posterior parietal cortex (PPC). The vOTC

127 occupies a key connectivity position in sighted readers, in that it is connected to visual input on

128 the one hand and linguistic representations on the other (Barttfeld et al., 2018; Bouhali et al.,

129 2014; Hannagan, Amedi, Cohen, Dehaene-Lambertz, & Dehaene, 2015; Li, Osher, Hansen, &

130 Saygin, 2020; Saygin et al., 2016; Stevens, Kravitz, Peng, Tessler, & Martin, 2017b; Yeatman, 131 Rauschecker, & Wandell, 2013). The PPC arguably occupies an analogous connectivity-based 132 position for tactile Braille. Not only is PPC anatomically proximal and densely connected to 133 early somatosensory cortices (SMC) but like anterior/lateral vOTC, it is connected to language 134 and working memory systems (Burks et al., 2017; Duhamel, Colby, & Goldberg, 1998; Kaas, 135 2012; Lewis & Van Essen, 2000; Ruschel et al., 2014). Analogous to the functional role of 136 vOTC in visual shape recognition, the PPC furthermore plays a key role in tactile shape and 137 texture perception, pertinent to Braille recognition (Bauer et al., 2015; Hegner, Lee, Grodd, & 138 Braun, 2010). For example, stronger PPC activity is observed during tactile pattern 139 discrimination compared to vibrotactile detection (Hegner et al., 2010). We therefore 140 hypothesized that portions of PPC may specialize for tactile Braille letter and word recognition, 141 analogous to specialization for visual word form recognition within the vOTC of sighted print 142 readers. To our knowledge, the hypothesis of selective responses to Braille words in PPC has not 143 previously been tested. Although previous studies have examined activity in early SMC and 144 found expanded finger representations in proficient Braille readers, there is no evidence that this 145 plasticity reflects specialization for Braille letters and words (Burton, Snyder, Conturo, et al., 146 2002; Burton, Sinclair, & McLaren, 2004; Kupers et al., 2007; Pascual-Leone et al., 1993; 147 Pascual-Leone & Torres, 1993; Sadato et al., 1998). One goal of the current study was therefore 148 to test whether any portion of PPC shows preferential responses to Braille letters and words in 149 blind readers of Braille, akin to specialization for visual letters and words found in vOTC of 150 sighted readers.

151 Finally, we hypothesized that lateralization patterns of Braille (blind) and visual print (sighted)

reading would be analogous but distinct. The reading network is typically strongly left-

153 lateralized in sighted people, like the spoken language network (Behrmann & Plaut, 2020;

154 Ossowski & Behrmann, 2015; Schlaggar & McCandliss, 2007; Seghier & Price, 2011; Vinckier

155 et al., 2007). Studies with sighted people who have right-lateralized spoken language responses

156 find that reading 'follows' spoken language into the right hemisphere (Behrmann & Plaut, 2020;

157 Cai, Lavidor, Brysbaert, Paulignan, & Nazir, 2008; Cai, Paulignan, Brysbaert, Ibarrola, & Nazir,

158 2010; Cai & Van der Haegen, 2015; Van der Haegen, Cai, & Brysbaert, 2012). In people who

are blind, left-lateralization of spoken language is reduced and highly variable across individuals

160 (Lane et al., 2017; Röder, Rösler, & Neville, 2000; Röder et al., 2002) We therefore

hypothesized that responses to Braille would be likewise less left-lateralized in blind readers and
would show co-lateralization with spoken language across individuals.

163 A further potential determining factor of Braille lateralization that does not arise for visual print, 164 is reading hand. In visual reading, information typically enters through both eyes and is projected 165 to both hemispheres. By contrast, in the case of Braille, it is possible for the information to enter 166 the left or the right hemisphere first, depending on the reading hand. Reading hand preferences 167 and reading styles differ widely across proficient blind Braille readers (Millar, 1984, 2003). 168 Many blind readers use both hands during naturalistic reading, however, one hand is thought to 169 track position on the page, while the other is used for word recognition (Millar, 2003). We 170 hypothesized that during single hand Braille reading, lateralization in early somatosensory 171 cortices would depend on which hand was used during word recognition, but that the effect of 172 reading hand would weaken in posterior parietal reading regions and would disappear in 173 language regions (Lane et al., 2017).

174 To test these predictions, we compared the neural basis of reading in proficient congenitally blind and sighted readers using analogous reading and spoken language tasks. In the reading 175 176 tasks, participants were presented with words, consonant strings, and non-letter shapes/false 177 fonts. Reading stimuli were visual (print) for the sighted participants and tactile (Braille) for the 178 blind participants. In the spoken language task, both groups listened to audio words and 179 backward speech sounds. First, we tested the prediction that there is a posterior-to-anterior 180 gradient in preference from false-fonts to consonant strings and finally words in the vOTC of 181 sighted but not blind readers. Previous studies find that posterior vOTC responds as much or 182 more to false fonts as to letters and words, with only a small lateral/anterior portion (so-called 183 VWFA) being selective to written words and letters (Vinckier et al., 2007). By contrast, we 184 predicted that in blind readers, the entire extent of vOTC would show a preference for words, 185 consistent with its involvement in language processing (Kim et al., 2017; Lane, Kanjlia, Omaki, 186 & Bedny, 2015; Röder et al., 2002; Watkins et al., 2012). Next, we tested the hypothesis that the 187 PPC of blind Braille readers shows a functional profile analogous to the vOTC of the sighted: 188 selective responses to written words as opposed to tactile shapes in a subset of PPC, surrounded 189 by equal or greater responses to tactile shapes. We compared responses in PPC with those of 190 early SMC, where we would expect larger or equal responses to tactile shapes. Moreover, we

191 hypothesized that regions of PPC most distal from S1 and posterior to it are most likely to show 192 specialization for Braille letters and words, an anterior/posterior gradient analogous to the 193 posterior/anterior gradient observed in the vOTC of sighted readers. We also examined responses 194 across groups in other cortical areas previously implicated in reading: left inferior frontal cortex 195 (IFC) and primary visual cortex (V1) and used whole-cortex analyses to quantify the anatomical 196 distribution of visual and Braille reading (Burton, Snyder, Conturo, et al., 2002; Harold Burton, 197 Sinclair, & Agato, 2012; Rueckl et al., 2015; Sadato et al., 1998). Finally, we used laterality 198 index analyses to compare lateralization patterns across written and spoken word comprehension 199 in the two groups. We tested the prediction that lateralization of reading would be driven by the 200 lateralization of spoken language in higher-order language regions (left IFC), by reading hand in

201 early SMC, and by both factors in reading-related areas (PPC).

202 Method

203 Participants

204 Nineteen congenitally blind (12 females, mean age = 40.36 years, SD = 14.82) and 15 sighted 205 control (9 females, mean age = 23 years, SD = 6) participants took part in the task-based fMRI 206 experiment (see Table 1 for participant characteristics). The data from 10 blind and 15 sighted 207 participants have been reported previously (Kim et al., 2017). All participants were native 208 English speakers, and none had suffered from any known cognitive or neurological disabilities 209 (screened through self-report). Sighted participants had normal or corrected to normal vision. All 210 the blind participants had at most minimal light perception from birth. Blindness was caused by 211 pathology anterior to the optic chiasm (i.e., not due to brain damage). All blind participants were 212 fluent Braille readers who began learning Braille at an average age of 4.6 years (SD = 1.49) and 213 rated their reading ability as proficient to expert (mean = 4.57, SD =0.69 on a scale of 1 to 5) and 214 reported reading on average 20 hours per week (SD=19). We obtained information on Braille-215 reading hand dominance, whether they read bimanually, and reading frequency through a post-216 experimental survey conducted over the telephone with 17 of the 19 blind adult participants 217 (Table 1). All participants gave informed consent according to procedures approved by the Johns 218 Hopkins Medicine Institutional Review Board.

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221 Table 1 Participants information

Dontiningant	Age (y)	Gender	Handedness	Reading	Cause of blind	Cause of blinds	Age started reading	Reading hours	Self-reported Braille
Participant no				handedness		Cause of blindness	Braille (y)	per week	reading ability (1-5)
B1	21	М	L	Bi-R	SC	LCA	4	14	5
B2	64	F	R	Bi-R	BA	ROP	6	56	5
В3	53	М	R	Bi-R	JD	LCA	6	7	4
B4	34	М	R	L	SC	Born without optic nerve	3	21	5
В5	42	М	Am	L	BA	ROP	3	21	5
B6	29	М	R	Bi-L	SC	LCA	4	<1	4
B7	39	F	R	L	BA	ROP	4	2	5
B8	34	F	R		SC	Optic Nerve Detached	3		5
В9	49	М	R	Bi-R	BA	unknown	8	<1	3
B10	26	F	R	Bi-R	MA	ROP	3	56	3
B11	49	F	L	R	MA	LCA	7	14	5
B12	39	F	R	L	MA	ROP	5	14	5
B13	35	F	R	Bi-L	MA	LCA	4	14	5
B14	46	F	R		BA	ROP	4		5
B15	33	F	R	L	BA	ROP	4	14	4
B16	25	F	Am	Bi-R	MA	LCA	5	56	5
B17	23	М	R	Bi-R	BA	LCA	4	28	5
B18	70	F	R	R	HS	ROP	7	7	4
B19	68	F	R	Bi-R	MA	ROP	5	7	5
Average									
Blind (n=19)	41 (SD=14.82)	12F	2L/2Am		BA		4.68 (SD=1.49)	19.47 (SD=18.97)	4.57 (SD=0.69)
Sighted (n=15)	23 (SD=6)	9F	1 L		BA				

Handedness: left (L), ambidextrous (Am), or right (R), based on Edinburgh Handedness Inventory. BA = Bachelor of Arts; MA = Master of Arts; HS = High School; JD = Juris Doctor; SC = Some College; ROP = Retinopathy of prematurity; LCA = Leber's congenital amaurosis. For Braille ability, participants were asked: "On a scale of 1 to 5, how well are you able to read Braille, where 1 is 'not at all', 2 is 'very little', 3 is 'reasonably well', 4 is 'proficiently', and 5 is 'expert'?"

222 Stimuli

223 The fMRI task including reading and listening tasks (Figure 1). There were three stimulus

224 conditions for the reading task: words, non-word consonant strings, and non-letter shapes

225 (control condition). During the reading task, stimuli were visual for the sighted participants and

tactile for the blind participants. For the listening task, there were two conditions: words andbackward speech sounds (control condition).

228 The word stimuli consisted of 240 common nouns, verbs, and adjectives. For the tactile reading 229 task (blind group), the Braille words were written in Grade-II contracted English Braille, which 230 is the most common form of Braille in the United States. Braille characters contain between 1-6 231 raised pins in set positions within a 2 x 3 array (see Figure 1). In Grade-II contracted English 232 Braille, there are contractions such that single Braille characters represent frequent letter combinations (e.g., "th") or frequent whole words (e.g., the "c" can stand for "can"). With 233 234 contractions, the Braille words were on average 4 Braille characters (range = 1-8 Braille 235 characters, SD = 2.1 characters) and 11 tactile pins per word. Note that each participant was 236 presented with 120 of the 240 words during the reading task; the other 120 words were presented 237 auditorily during the listening task (see below). The word lists were counterbalanced across 238 participants. In the tactile consonant string condition, there were 24 strings repeated 5 times 239 throughout the experiment. Each string stimulus consisted of 4 Braille letters, which were 240 created using 20 English consonants. Last, the tactile control stimuli consisted of 24 unique 241 strings of 4 non-letter shapes made of Braille pins (see Figure 1). Note that any dot array within a 242 2 x 3 grid could be part of a Braille character. Therefore, to prevent participants from processing 243 the shapes as Braille letters, the shapes varied in size and pin number within arrays ranging in size from 4×5 to 7×7 . The average number of Braille pins per string in the control condition 244 245 was 58.

246 For the sighted group, the word stimuli consisted of 240 common nouns, verbs, and adjectives that were on average 4 letters long (range = 3-5 letters, SD = 0.7 letters). Visual word stimuli 247 248 consisted of a new set of words matched to the Braille words on average character length (i.e., 4 249 visual letters matched to 4 Braille characters), raw frequency per million, averaged frequency per 250 million of orthographic neighbors, and averaged bigram frequency (all comparisons p > 0.4, 251 obtained from the MCWord Orthographic Wordform Database; Medler & Binder, 2005). 252 Different groups of words were used for the visual and Braille experiment to enable character 253 length matching since Braille contractions represent two or more English letters with a single 254 Braille character. Like the blind participants, sighted participants encountered half (120) of the 255 words during reading trials and the other half during auditory trials, counterbalanced across

256 participants. The visual consonant strings were the same 24 consonant letter combinations from

- the tactile consonant strings described above. Lastly, the control stimuli in the visual reading task
- 258 were 24 unique strings, each comprised of 4 characters, which were false fonts. There were 20
- 259 false font characters in total, which matched the 20 English consonants on the number of strokes,
- 260 presence of ascenders and descenders, and the stroke thickness.
- 261 The stimuli for the listening task were taken from each group's respective word list. For the
- audio word condition, stimuli were 120 words taken from the reading task described above. For a
- 263 given word, half of the participants received it in the reading task and half received it in the
- listening task. The auditory words were recorded by a female native English speaker. The
- average word length was 5 letters long (SD = 1.4 letters). The average playtime for the auditory
- stimuli was 0.41 s long (SD = 0.3 s). The control auditory stimuli comprised backward speech
- sounds, which were created by playing each audio word in reverse.

268 **Procedure**

269 The experiment had a total of 5 runs, each with 20 task trials. In each trial, participants were

- 270 presented with a block of 6 stimuli from a single condition (e.g., tactile reading consonant strings
- 271 condition) and then performed a memory probe task. All stimulus conditions for both reading
- and listening trials were presented in every run. Each condition was repeated 4 times per run, and
- the order of conditions was counterbalanced across runs. There were 6 rest periods (16 s)
- throughout each run. One sighted participant and two blind participants were excluded from
- behavioral analysis due to failure to record their responses.
- 276 For the blind participants, each trial began with a 0.5 s auditory cue instructing participants to 277 "Touch' (reading trial), or "Listen" (listening trial). Then participants felt or heard blocks of 6 278 target items, one at a time. For 10 of the blind participants, tactile target stimuli were presented 279 on the Braille display for 2 s, followed by a 0.75 s inter-stimulus interval (ISI) (6-item list 280 duration: 16.5 s) (Kim et al., 2017). For the newly added 9 blind participants, the ISI was 281 lengthened to 1.75 s due to a coding error which caused the 6-item list duration to be prolonged 282 to 22.5 s. Control analyses revealed no effects of ISI duration on the results and the data are 283 henceforth combined. After the 6-item list had been presented, there was a short delay (0.2 s),

284 followed by a beep (0.5 s). Then a probe stimulus (2 s) was then presented, and participants 285 indicated with a key press whether or not the probe had been present in the list. Participants had 286 5.3 s to make a response. The participants were asked to read with their dominant hand and 287 responded with the other hand. The listening task was analogous in format to the reading task. 288 The audio words and backward speech were on average 0.41 s long. The timing and sequence of 289 events were identical for the listening task (6-item list duration 16.5 s). 290 For sighted participants, the trial event sequence (cue, 6-item block, beep, probe, response) was 291 analogous to above. Each trial began with an auditory cue instructing participants to "Look" 292 (reading trial) or "Listen" (listening trial). During reading trials, 6 visual stimuli appeared 293 centrally for 1 s each, followed by an ISI of 0.75, during which participants were asked to 294 maintain gaze on a black central fixation cross (total block duration: 10.5 s). Note that visual 295 reading blocks were shorter than tactile reading blocks for the blind participants because pilot 296 testing indicated that visual reading is faster under these conditions. Listening trials also had a

total stimulus block duration of 10.5 s, to be consistent with the reading trials within the sightedgroup.

299 fMRI data acquisition

300 Functional and structural images were acquired using a 3T Phillips scanner at the F. M. Kirby

- 301 Research Center. T1-weighted images were collected using a magnetization-prepared rapid
- 302 gradient-echo (MP-RAGE) in 150 axial slices with 1 mm isotropic voxels. Functional BOLD
- 303 scans were collected T1-weighted structural images were collected in 150 axial slices with 1 mm
- 304 isotropic voxels. Functional BOLD scans were collected in 36 sequential ascending axial slices.
- 305 TR = 2 s, TE = 0.03 s, flip angle = 70°, voxel size = $2.4 \times 2.4 \times 2.5$ mm, inter-slice gap = 0.5
- 306 mm, field of view (FOV) = $192 \times 172.8 \times 107.5$. Acquisition parameters were identical for the
- 307 resting-state and task fMRI experiment.

308 fMRI data analysis

309 Preprocessing and whole-cortex analysis

310 Analyses were performed using FSL (version 5.0.9), FreeSurfer (version 5.3.0), the Human

311 Connectome Project workbench (version 1.2.0), and custom in-house software. The cortical

312 surface was created for each participant using the standard FreeSurfer pipeline (Dale, Fischl, &

313 Sereno, 1999; Glasser et al., 2013; Smith et al., 2004). For task data, preprocessing of functional

314 data included motion-correction, high-pass filtering (128 s cut-off), and resampling to the

315 cortical surface. Cerebellar and subcortical structures were excluded. On the surface, the task

316 data were smoothed with a 6 mm FWHM Gaussian kernel. Two runs for blind and three runs for

317 sighted participants were dropped due to equipment failure.

318 The three conditions in the reading task and two conditions in the listening task were entered as

319 covariates of interest into general linear models. Only the six-item period in each trial was

320 entered into the model. Covariates of interest were convolved with a standard hemodynamic

321 response function, with temporal derivatives included. Probe stimulus, response periods, and the

322 trials in which participants failed to respond were entered as covariates of no interest. The effect

323 of the mean signal of white matter and CSF, as well as the motion spike, were also included as

324 the covariates of no interest. Runs were combined within subjects using fixed-effects models.

325 Data across participants were combined within groups using random-effects analysis. Reported

326 whole-cortex contrasts were run thresholded at p < 0.01 vertexwise, and p < 0.05 cluster-

327 corrected.

328 fMRI ROI analysis

Individual-subject functional regions of interest (ROIs) were defined within the vOTC and other regions previously implicated in Braille reading (V1), language (left inferior frontal cortex, IFC), and tactile perception (left posterior parietal cortex, PPC, and hand region of the left primary somatosensory-motor cortex, SMC). To construct the left vOTC search space, we first combined the left fusiform, inferior temporal, and lateral occipital parcels from Freesurfer's automated aparc parcellation and then excluded V1, V2 regions, and the vertices with y-axis greater than -30 (Lerma-Usabiaga et al., 2018). To test the posterior-to-anterior function gradient, the left 336 vOTC search space was divided to three portions: posterior (y < -64), middle (-48 > y > = -64), 337 and anterior portion ($y \ge -48$). The search space in the right hemisphere was created by 338 flipping the left vOTC masks along the x-axis. The V1 search space was defined from a 339 previously published anatomical surface-based atlas (PALS-B12; Van Essen, 2005). The left 340 inferior frontal language (IFC) search space was defined by using a sentence vs. non-words 341 contrast (Fedorenko, Hsieh, Nieto-Castañón, Whitfield-Gabrieli, & Kanwisher, 2010). The 342 parietal search space was defined by the orthogonal contrast of all tactile conditions (words, 343 consonant strings, and control) > rest in whole-cortex analysis, excluding the occipital parcels 344 from Freesurfer's automated aparc parcellation. To look for lateralization effects in vOTC across 345 groups, we examined responses separately for the right and left hemispheres.

346 Individual-subject functional ROIs were defined in group-wise search spaces (described below). 347 Each individual subject's ROI was defined as the top 5% of vertices activated for the 348 tactile/visual consonant strings > tactile/visual controls contrast within the search spaces listed 349 above. We used this consonant string contrast for the primary analysis in order to focus on 350 orthographic as opposed to semantic responses. However, all analyses were also repeated using 351 the words > control contrast and results from these analyses are reported in the supplementary 352 material (Figure S3 and Figure S5). To avoid using the same data to define ROIs and to test 353 hypotheses, a leave-one-run-out cross-validation procedure was used. ROIs were defined based 354 on data from all but one run, then the percent signal change (PSC) was extracted from the left-355 out run. This procedure was repeated iteratively across all runs and the PSC was averaged across 356 iterations.

Repeated-measured ANOVAs were used to analyze the ROI data, and two-tailed paired *t*-tests
were used for pairwise comparisons. All *p* values were Bonferroni-corrected for multiple
comparisons.

360 Topographical preference map

To explore the posterior-to-anterior gradient in left vOTC and in a data-driven way, we mapped the topographical preference of the vOTC during reading using a winner-take-all approach. We took the bilateral vOTC as the mask, and color-coded each vertex within the mask according to which stimulus condition it responded most strongly. The topographical preference map of the 365 PPC and parieto-occipital/dorsal occipital cortex was created using the same winner-take-all

366 approach. The mask was defined by the orthogonal contrast of all tactile conditions (words,

367 consonant strings, and control) > rest in the whole-cortex analysis.

368 Laterality index analysis

369 To determine whether spoken and written language co-lateralize to the same hemisphere, we

370 performed laterality index (LI) analyses. LI was calculated separately for the reading and

371 listening tasks for each participant in the SMC, PPC, vOTC, V1, IFC, and also for the whole

372 cortex. For the reading task, LI was determined based on the tactile/visual words > rest contrast.

373 For the listening task, LI was determined using the audio words > rest contrast. The LI was

374 calculated using the standard formula: (L - R) / (L + R), where L and R refer to the sums of the z

375 statistics from the relevant contrast within the left and right hemispheres, respectively. LI ranges

376 from -1 to 1, with a score of 1 indicating strong left lateralization and -1 strong right

377 lateralization.

378 The bootstrap/histogram method was used to ensure that LIs were not overly influenced by

arbitrary activation threshold choices or outlier voxels. Bootstrapped LIs were computed using

380 20 evenly spaced thresholds ranging from z = 1.28 to z = 4.26 (corresponding to one-sided p =

0.1 to p = 0.00001, uncorrected). For every threshold, each participant's z statistic map was

masked to only include the voxels exceeding the threshold within the search space. Then we

383 sampled the suprathreshold voxels 100 times with replacement in each hemisphere at a sampling

ratio k = 1.0. The LIs were then calculated using each pair of left and right hemisphere samples,

385 yielding a histogram of 10,000 threshold-specific LIs. Next, a single LI for each threshold was

calculated by averaging the values after removing the upper and lower 25% of the 10,000

threshold-specific values. Finally, the LI reported for each participant represents the averageacross all thresholds.

389 A small number of participants were excluded from the LI analysis for a particular region if they

390 did not have suprathreshold activation in both hemispheres (listening task- SMC: 2 sighted, 2

391 blind participants excluded; PPC: 1 sighted; V1: 6 sighted; IFC: 1 sighted; reading task- SMC: 4

392 sighted; PPC: 1 sighted; IFC: 1 sighted).

393 To examine the effect of spoken language lateralization and Braille reading handedness on the

394 reading lateralization, a multiple regression was conducted for each region. The LI of spoken

395 words in IFC and dominant reading hand were entered as regressors and the LI of written words

396 was the dependent variable. Although some participants reported reading Braille bimanually, the

397 participants were asked to read tactile stimuli during the experiment only with their dominant

398 reading hand. There were 7 blind participants in the left Braille-reading handed group and 10 in

399 the right Braille-reading handed group.

400

401 **Results**

402 Behavioral Results

403 Higher accuracy and shorter reaction times for word-like stimuli

404 Because the two groups differed in age, we regressed out the effect of age on accuracy and 405 reaction times and performed analyses on the residuals (see Figure S1 in Supplementary 406 materials, results from raw data are also included in Figure S1). In the reading task, there was a 407 significant effect of age on accuracy (main effect of age, $F_{(1, 85)} = 5.681$, p < 0.05). A two-way 408 lexicality (words, consonant strings, control) by group (sighted, blind) ANOVA performed on 409 the residuals revealed higher accuracy on more word-like stimuli (words and consonant strings > 410 control) in both blind and sighted groups (main effect of lexicality: $F_{(2, 54)} = 13.963$, p < 0.001). 411 There was no lexicality by group interaction ($F_{(2,54)} = 0.872$, p = 0.737). The group effect was 412 marginal (sighted > blind, $F_{(1,27)} = 3.603$, p = 0.068). For the listening task, there was a trending effect of age on accuracy ($F_{(1,56)} = 2.907$, p = 0.094). A two-way lexicality (words, control) by 413 414 group (sighted, blind) ANOVA on the residuals revealed a lexicality effect (words > control; $F_{(1)}$ 415 $_{29} = 50.944, p < 0.001$), no group effect ($F_{(1, 27)} = 0.843, p = 0.367$) or group by lexicality interaction $(F_{(1, 27)} = 0.549, p = 0.465)$. 416

417 Likewise, for reaction times during the reading task, there was a significant effect of age ($F_{(1, 85)}$

418 = 39.089, p < 0.001). A two-way lexicality (words, consonant strings, control) by group (sighted,

419 blind) ANOVA on the residuals revealed a lexicality effect (words and consonant strings <

- 420 control; $F_{(2, 54)} = 8.09, p < 0.001$). There was no group effect ($F_{(1, 27)} = 8.09, p = 0.297$). The
- 421 group by lexicality interaction effect was marginal ($F_{(2, 54)} = 2.763, p = 0.072$). Pairwise
- 422 comparisons showed the shorter reaction times on more word-like stimuli in blind group, but
- 423 there was no difference across stimuli in the sighted group (blind: words vs. control, $t_{(16)} = -2.91$,
- 424 p < 0.01; consonant strings vs. control, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, p < 0.01; words vs. consonant strings, $t_{(16)} = -2.604$, $t_{(16)}$
- 425 -0.686, p > 0.99; sighted: all pairwise comparisons p > 0.05; the *p*-values were Bonferroni-
- 426 corrected).
- 427 During the listening task, the main effect of age on reaction time was significant $(F_{(1, 85)} =$
- 428 15.892, p < 0.001). A two-way lexicality (words, control) by group (sighted, blind) ANOVA on
- 429 the residuals revealed a lexicality effect (words < control; $F_{(1,29)} = 50.944$, p < 0.001). There was
- 430 no group effect ($F_{(1, 27)} = 0.071$, p = 0.792) or group by lexicality interaction ($F_{(1, 29)} < 0.001$, p > 0.001
- 431 0.99).

432 fMRI Results

433 Visual (sighted) but not tactile Braille reading (blind) elicits a posterior-to-anterior functional 434 gradient in left vOTC and shows left-lateralization

Two signatures of visual reading responses in vOTC are 1) a posterior-to-anterior word form
gradient and 2) left-hemisphere lateralization. We asked whether Braille reading in blind
individuals shows similar posterior-to-anterior and laterality effects as visual reading in sighted

- 438 people. We divided the left and right vOTC each into the posterior, middle, and anterior
- 439 subregions (ROIs) and compared responses in these subregions across hemispheres and groups
- 440 (see Methods, Figure 1). We first conducted a four-way hemisphere (left, right) by
- 441 posterior/anterior subregion (posterior, middle, anterior) by lexicality (words, consonant strings,
- 442 control) by group (sighted, blind) ANOVA to examine reading responses across groups. This
- 443 ANOVA revealed a four-way interaction ($F_{(4, 128)} = 3.028, p < 0.05$), indicating that lexicality,
- 444 hemisphere, and posterior/anterior subregion interact with group. Next, we used separate
- 445 ANOVAs for each group to unpack the 4-way interaction. Because of the large number of factors
- and to preserve readability, we report only hypothesis-relevant effects in this section. A complete
- 447 summary of all effects can be found in the Supplemental Materials.

448 For the sighted group, we found the expected three-way interaction between hemisphere (left,

449 right), posterior/anterior subregion (posterior, middle, anterior) and lexicality (words, consonant

450 strings, control; $F_{(4, 56)} = 4.287$, p < 0.01). Next, we looked at each hemisphere separately in the 451 sighted group.

452 In the left vOTC, there was a two-way interaction between lexicality (words, consonant strings,

- 453 control) and posterior/anterior subregion (posterior, middle, anterior; $F_{(4, 56)} = 9.69, p < 0.001$),
- 454 reflecting the expected posterior-to-anterior functional gradient. Pairwise comparisons revealed
- 455 that the posterior vOTC responded similarly to all visual stimuli (all pairwise comparisons p > 1
- 456 0.05). By contrast, in middle vOTC, consonant strings elicited higher responses than both words
- 457 and control stimuli (Bonferroni-corrected paired *t*-test for words vs. consonant strings: $t_{(14)} = -$
- 458 3.918, p < 0.05; consonant strings vs. control: $t_{(14)} = 4.106$, p < 0.01). In anterior vOTC,
- 459 responses to words and consonant strings were both higher than control and not different from
- 460 each other (Bonferroni-corrected paired *t*-test for words vs. control: $t_{(14)} = 3.461$, p < 0.05;

461 consonant strings vs. control: $t_{(14)} = 3.327$, p < 0.05, all other pairwise comparisons p > 0.05).

- 462 In the right vOTC of the sighted group, a two-way lexicality (words, consonant strings, control)
- 463 by posterior/anterior subregion (posterior, middle, anterior) ANOVA revealed no main effect of

464 lexicality ($F_{(2,28)} = 0.448, p > 0.05$) and no interaction ($F_{(4,56)} = 0.987, p > 0.05$). To

summarize, these results demonstrate that in the sighted group, there was a posterior-to-anterior

466 functional gradient for processing word form during reading in the left but not right vOTC.

467 Next, we examined these effects in the blind group. We conducted a three-way hemisphere (left,

468 right) by posterior/anterior subregion (posterior, middle, anterior) by lexicality (words,

469 consonant strings, control) ANOVA. Unlike in the sighted, there was no significant three-way

470 interaction ($F_{(4, 56)} = 0.877$, p = 0.482). Although there was no interaction, we conducted a

- 471 separate ANOVA testing for a lexicality effect across the posterior/anterior subregions for each
- 472 hemisphere separately in order to match the analysis of the sighted group.

473 In the left vOTC of the blind group, all three (posterior, middle, anterior) subregions responded

474 most to words, followed by consonant strings followed by tactile shapes (Figure 1). There was a

- 475 two-way interaction between lexicality (words, consonant strings, control) and posterior/anterior
- 476 subregion (posterior, middle, anterior; $F_{(4, 72)} = 3.198$, p < 0.05). However, the nature of this

477 interaction was different from that observed in the sighted group. All pairwise-comparisons

- 478 between conditions were significant in all three subregions (words > consonant strings >
- 479 control), except the difference between words and consonant strings did not reach significance in
- 480 the anterior vOTC (Bonferroni-corrected paired *t*-test for words vs. consonant strings: posterior

481 vOTC $t_{(18)} = 2.678, p < 0.05$; middle vOTC: $t_{(18)} = 3.166, p < 0.05$; anterior vOTC: $t_{(18)} = 2.016, p$

- 482 = 0.177; words vs. control: posterior vOTC: $t_{(18)} = 5.463$, p < 0.001; middle vOTC $t_{(18)} = 8.547$, p
- 483 < 0.001; anterior vOTC: $t_{(18)} = 5.874$, p < 0.001; consonant strings vs. control: posterior vOTC:
- 484 $t_{(18)} = 3.413, p < 0.01$; middle vOTC $t_{(18)} = 4.696, p < 0.01$; anterior vOTC: $t_{(18)} = 5.034, p < 0.01$
- 485 0.001).

486 Unlike in the sighted group, in the right hemisphere of the blind group, lexicality effects were 487 similar to the left hemisphere. All three (posterior, middle, anterior) subregions responded most 488 to words, followed by consonant strings followed by tactile shapes. There was also a two-way 489 interaction between lexicality (words, consonant strings, control) and subregion (posterior, 490 middle, anterior; $F_{(4,72)} = 7.064$, p < 0.001). Pairwise comparisons showed that the posterior 491 right vOTC responded more to words than control ($t_{(18)} = 4.112, p < 0.01$); the middle vOTC 492 responded more to words than both consonant strings ($t_{(18)} = 4.011$, p < 0.01) and control ($t_{(18)} =$ 493 4.819, p < 0.001); and the anterior vOTC responded most strongly to words and consonant 494 strings than control stimuli (words vs. consonant strings: $t_{(18)} = 2.429$, p = 0.07; words vs. 495 control: $t_{(18)} = 5.561$, p < 0.001; consonant strings vs. control, $t_{(18)} = 4.522$, p < 0.01). Other 496 pairwise comparisons did not reach significance (posterior vOTC: words vs. consonant strings, 497 $t_{(18)} = 2.349$, p = 0.091; consonant strings vs. control, $t_{(18)} = 2.16$, p = 0.134; middle vOTC: 498 consonant strings vs control, $t_{(18)} = 2.073$; p = 0.159).

In summary, in the blind group, the entire posterior/anterior extent of the vOTC responded more to words than either consonant strings or tactile shapes. Unlike in the sighted, we did not observe the posterior-to-anterior functional gradient or a left hemisphere dominance for written words.

- 502 For the listening task, similar to the reading task, we conducted a four-way hemisphere (left,
- 503 right) by subregion (posterior, middle, anterior) by lexicality (words, control) by group (sighted,
- 504 blind) ANOVA. The four-way interaction effect with group was marginal and we, therefore, did
- not proceed to further analyses ($F_{(2, 64)} = 2.717$, p = 0.074). It is worth noting that in sighted

506 group, responses to auditory stimuli were below rest in posterior vOTC and above rest in the 507 more anterior regions. This pattern was not observed in the blind group (see Figure S2).

508 Topographical preference map of vOTC: gradient only in sighted readers

509 In order to explore the posterior-to-anterior gradient in a data-driven way, we mapped the

510 topographical preferences of the blind and sighted vOTC during reading using a winner-take-all

511 approach (Figure 1B). We coded the vertex-wise preferences in different colors for words,

512 consonant strings, and control stimuli (see Methods). In the sighted group, a clear posterior-to-

513 anterior gradient in the left vOTC was observed. The posterior section shows a preference for the

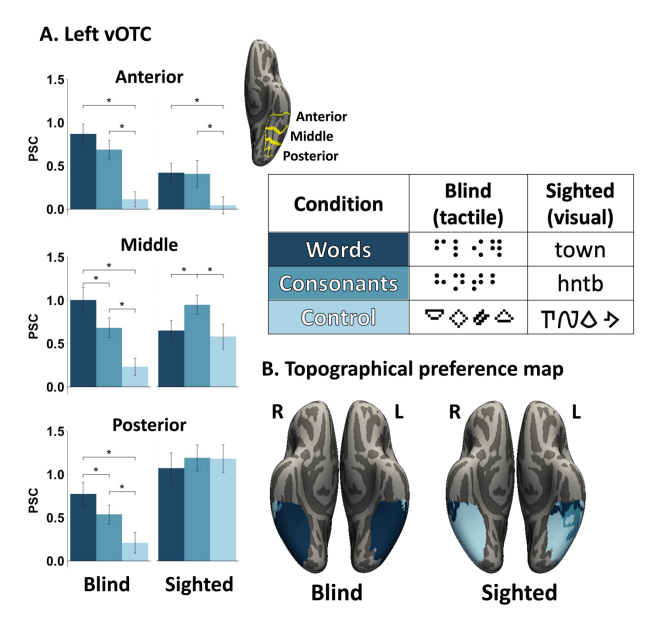
514 visual control false font stimuli whereas anteriorly, most vertices preferred consonant strings or

515 words. In the sighted group's right vOTC, almost all vertices responded most strongly to the

516 control stimuli. These patterns contrast starkly with the blind vOTC maps, which show a clear,

517 bilateral preference for tactile words throughout the entire extent of both left and right vOTC.

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519 Figure 1. (A) Responses in left vOTC across the posterior, middle, and anterior subregions for 520 blind and sighted groups during the reading tasks (left column). Bars show results from 521 consonant string > false fonts leave-one-run out individual subject ROI analysis. Error bars 522 denote standard errors +/- the mean. Asterisks (*) denote significant Bonferroni-corrected pairwise comparisons within the task (p < 0.05). The right upper row showed example stimuli for 523 524 the reading trials for the blind and sighted groups. (B) Maps of the blind and sighted vOTC 525 showing topographical preference to words, consonant strings, or control stimuli during the 526 reading task.

527 The posterior parietal cortex (PPC) but not S1 of blind readers shows a preference for written 528 Braille words and consonant strings

529 We tested the hypothesis that the PPC shows preferential involvement in Braille reading, analogous to vOTC preference for visual print in the sighted group (Figure 2A). A two-way 530 531 lexicality (words, consonant strings, control) by group (sighted, blind) ANOVA in the reading task showed a main effect of lexicality ($F_{(2, 64)} = 13.206$, p < 0.001) and a group by lexicality 532 533 interaction ($F_{(2.64)} = 5.123$, p < 0.01; functional ROIs were defined using the words > controls 534 contrast). There was no main effect of group ($F_{(1,32)} = 1.452$, p = 0.237). In the sighted group, 535 consonant strings elicited higher responses than both words and control stimuli (Bonferroni-536 corrected paired *t*-test, words vs. consonant strings: $t_{(14)} = -3.805$, p < 0.01; consonant strings vs. 537 control: $t_{(14)} = 6.922$, p < 0.001; words vs. control: $t_{(14)} = 1.406$, p > 0.99). By contrast, in the 538 blind group the PPC responded more to both tactile words and consonant strings relative to 539 control stimuli (Bonferroni-corrected paired *t*-test, words vs. consonant strings: $t_{(18)} = 1.571$, p =540 0.298; consonant strings vs. control: $t_{(18)} = 3.028$, p < 0.01; words vs. control: $t_{(18)} = 3.165$, p < 0.01; words vs. control: $t_{(18)} = 3.165$, p < 0.01; words vs. control: $t_{(18)} = 3.165$, p < 0.01; words vs. control: $t_{(18)} = 3.165$, p < 0.01; words vs. control: $t_{(18)} = 3.165$, p < 0.01; words vs. control: $t_{(18)} = 3.165$, p < 0.01; words vs. control: $t_{(18)} = 3.165$, p < 0.01; words vs. control: $t_{(18)} = 3.165$, p < 0.01; words vs. control: $t_{(18)} = 3.165$, p < 0.01; words vs. control: $t_{(18)} = 0.01$; words vs. control: $t_{(18)} = 0.0$ 541 0.01). Note that when the posterior parietal ROI was defined instead using the words > controls 542 contrast, the blind group continued to show a larger lexicality preference than the sighted (see 543 Supplemental Materials for details; Figure S5). These results suggest a specific involvement of 544 the PPC in tactile Braille reading.

- 545 For the listening task, the two-way lexicality (words, control) by group (sighted, blind) ANOVA
- 546 revealed a significant main effect of lexicality in the PPC (words > control, $F_{(1,32)} = 11.112$, p < 100
- 547 0.01; see Figure S4). There was no main effect of group ($F_{(1,32)} = 3.275$, p = 0.08) and no
- 548 interaction between group and lexicality ($F_{(1,32)} = 2.372, p = 0.133$).

We examined responses of the left SMC hand region to test whether it showed a similar preference for Braille words and consonant strings as the PPC (Figure 2A). For the reading task, the two-way lexicality (words, consonant strings, control) by group (sighted, blind) ANOVA showed a main effect of lexicality ($F_{(2, 64)} = 7.265$, p < 0.001; functional ROIs were defined using the words > controls contrast), with higher responses to the consonant strings than control stimuli. Noting that the responses to all stimuli were below rest in SMC in blind group. There was no main effect of group ($F_{(1, 32)} = 0.604$, p = 0.443) and no group by condition interaction (F 556 $_{(2, 64)} = 1.501$, p = 0.231). For the listening task, the two-way lexicality (words, control) by group

- (sighted, blind) ANOVA revealed a main effect of group ($F_{(1,32)} = 15.622, p < 0.001$), with
- 558 overall greater responses in sighted group than blind group. There was no main effect of
- 559 lexicality ($F_{(1, 32)} = 1.933$, p = 0.174) and no interaction ($F_{(1, 32)} = 0.658$, p = 0.423). Results
- 560 were similar when the SMC ROIs were instead defined using the words > controls contrast. In
- sum, unlike in the PPC, we found no evidence for specialization of SMC for Braille reading as
- 562 compared to perception of control tactile shapes.

563 **Topographical preference map of parieto-occipital stream: shift in preference from shapes to** 564 word-like Braille stimuli along anterior-to-posterior axis

565 Finally, we constructed a data-driven preference map in PPC and parieto-occipital/dorsal

566 occipital cortex analogous to the one created for vOTC (see Figure 2B). In the blind group, this

567 map shows preferential responses to tactile shapes in anterior portions of PPC, immediately

adjacent to S1. A small middle region in left and right PPC showed a preference for consonant

569 strings, whereas the most posterior portion of PPC, as well as parieto-occipital and dorsal

570 occipital regions responded preferentially to words. To summarize, the overall pattern suggests

an anterior-to-posterior decoding pattern in the parieto-occipital stream in the blind group,

572 analogous to the posterior-to-anterior vOTC gradient observed in sighted readers.

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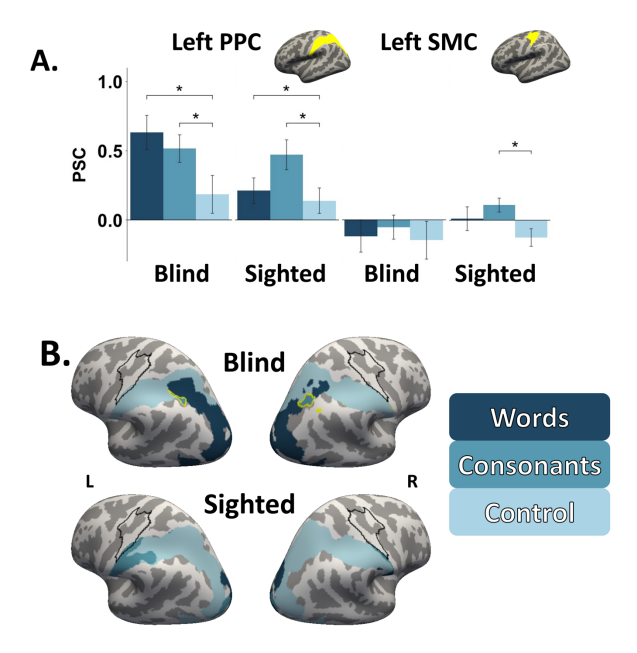


Figure 2. (A) Responses in left PPC, and SMC ROIs for blind and sighted groups during the reading (blue colors) tasks. Error bars denote standard errors +/- the mean. Asterisks (*) denote significant Bonferroni-corrected pairwise comparisons within the task (p < 0.05). T = tactile, V = visual. (B) Maps of the blind and sighted vOTC showing topographical preference to words, consonant strings, or control stimuli during the reading task in the posterior parietal and parietooccipital cortex. Black line: hand region of the primary sensory-motor cortex; yellow line: vertices showed the preference for consonant strings.

Left vOTC responds to linguistic stimuli in blind and sighted readers, but differently to words and consonant strings across groups

583 We examined the effects of lexicality across groups on left vOTC responses during the reading 584 tasks using a two-way lexicality (words, consonant strings, control) by group (sighted, blind) 585 ANOVA (functional ROIs were defined using the words > controls contrast, Figure 3). We 586 observed a main effect of lexicality ($F_{(2, 64)} = 42.293$, p < 0.001) and no main effect of group (F587 (1, 32) = 0.004, p = 0.948). A lexicality by group interaction revealed different response patterns across sighted and blind individuals ($F_{(2, 64)} = 10.272, p < 0.001$). While in the blind group 588 589 words elicited larger responses than consonant strings, responses to consonant strings were 590 numerically but not significantly larger than to words in the sighted group (Bonferroni-corrected 591 paired t-test words vs. consonant strings blind: $t_{(18)} = 3.027$, p < 0.05; sighted: $t_{(18)} = -1.317$, p =592 0.614). In both groups, words and consonant strings showed larger responses than control stimuli

593 (all pairwise comparisons p < 0.05) (Figure 3).

594 For the listening task, a two-way lexicality (words, control) by group (sighted, blind) ANOVA 595 revealed greater overall responses to words than control stimuli (main effect of lexicality, $F_{(1,32)}$ 596 = 35.919, p < 0.001; functional ROIs were defined using the words > controls contrast). There was no main effect of group ($F_{(1,32)} = 1.362$, p = 0.252). The lexicality by group interaction was 597 598 marginal ($F_{(1,32)} = 3.785$, p = 0.061), indicating a larger difference between audio words and 599 audio control stimuli in the blind group than in the sighted group. A similar pattern was observed 600 when the vOTC functional ROI was instead defined using the words > control contrast (see 601 Supplemental Materials; Figure S3).

The left inferior frontal cortex (IFC) prefers word-like written and spoken stimuli across blind and sighted readers

604 We analyzed responses in the left IFC across groups with the prediction that this high-level

605 language region would show similar response patterns across blind and sighted readers.

606 Consistent with this prediction, responses were similar across groups for both tasks in the left

607 IFC. For the reading task, a two-way lexicality (words, consonant strings, control) by group

608 (sighted, blind) ANOVA revealed a significant main effect of lexicality, with larger responses

for words and consonant strings over the control condition ($F_{(2, 64)} = 46.313, p < 0.001$; Figure 3;

610 functional ROIs were defined using the words > controls contrast). Neither the main effect of 611 group $(F_{(1,32)} = 0.004, p = 0.947)$ nor the interaction $(F_{(2,64)} = 1.017, p = 0.367)$ were 612 significant. Likewise, for the listening task, the two-way lexicality (words, control) by group 613 (sighted, blind) ANOVA revealed the expected main effect of lexicality (words > control; $F_{(1,32)}$ 614 = 23.778, p < 0.001). There was no main effect of group ($F_{(1,32)} = 0.753$, p = 0.392) and no 615 lexicality by group interaction ($F_{(1,32)} = 0.357$, p = 0.554). There was also no group-by-616 condition interaction when functional ROIs were defined using the words > controls contrast. 617 Both groups still showed a preference for words over control stimuli and in this case, there was 618 also a larger response to words over consonant strings in both groups (see Supplemental 619 Materials for details; Figure S5). These results are consistent with prior studies showing similar 620 responses to spoken and written language in the left inferior frontal cortex of blind and sighted 621 adults.

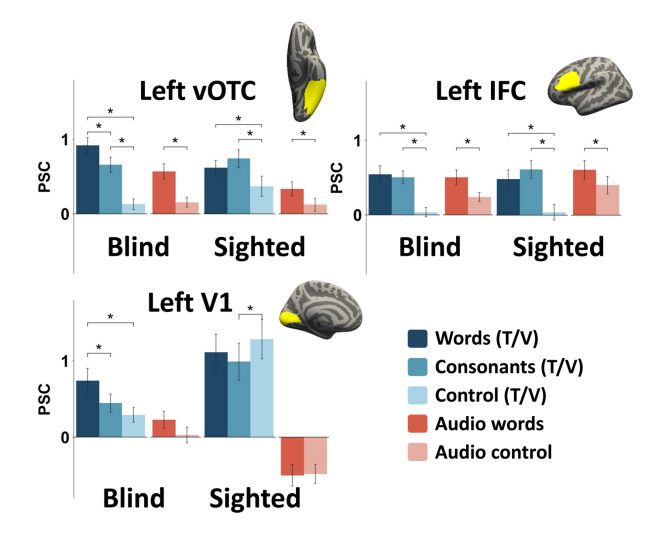
622 V1 shows a preference for words in blind readers

623 We investigated the effects of lexicality across groups in V1 (Figure 3), because it was 624 previously identified as relevant to Braille reading (Sadato et al., 1996; Cohen et al. 1997). As 625 with vOTC, we first examined responses in left V1 during the reading task using the consonant 626 strings > control functional ROIs. A two-way lexicality (words, consonant strings, control) by 627 group (sighted, blind) ANOVA revealed main effects of lexicality ($F_{(2, 64)} = 4.247, p < 0.05$) and 628 group (sighted > blind, $F_{(1,32)} = 6.964$, p < 0.05). There was also a significant lexicality by group 629 interaction ($F_{(2, 64)} = 9.487$, p < 0.001). In the blind group, V1 responded most to words and 630 there was no difference between consonant strings and control (Bonferroni-corrected paired t-631 test, words vs. consonant strings: $t_{(18)} = 2.641$, p < 0.05; words vs. control: $t_{(18)} = 3.691$, p < 0.01; 632 consonant strings vs. control: $t_{(18)} = 2.367$, p = 0.214). In the sighted group, V1 responded more 633 to control stimuli than consonant strings (Bonferroni-corrected paired *t*-test, $t_{(14)} = 2.652$, $p < 10^{-10}$ 634 0.01). There was no difference between other conditions (pairwise comparisons p > 0.05.) V1 635 responses in the blind group were similar when functional ROIs were defined using words > 636 control (see Supplemental Materials for details; Figure S5). In the sighted group, however, a 637 marginal preference for words over false fonts emerged in this alternative analysis (Bonferroni-638 corrected paired *t*-test, $t_{(14)} = 2.573$, p = 0.067; Figure S5). This latter result is consistent with 639 some previous studies showing that V1/V2 responded more to words than non-letter control

640 stimuli like scrambled words (Szwed et al., 2011; Szwed, Qiao, Jobert, Dehaene, & Cohen,

641 2014).

- 642 For the listening task, the two-way lexicality (words, control) by group (sighted, blind) ANOVA
- 643 showed a main effect of group ($F_{(1, 32)} = 16.067, p < 0.001$), with overall greater activation seen
- 644 in blind than sighted V1. There was no main effect of lexicality ($F_{(1,32)} = 2.344, p = 0.316$) and
- no interaction between the factors ($F_{(1, 32)} = 1.589$, p = 0.217). Notably in the sighted but not
- 646 blind group, responses to both words and audio control were below rest. This pattern of results
- 647 was the same in words > control ROI (See Supplemental Materials for details; Figure S5).



649 **Figure 3.** Response of left vOTC (upper left), left IFC (upper right), left V1 (lower left) during

- reading (blue colors) and listening (orange colors) tasks for blind (left) and sighted groups
- 651 (right). Bars show PSC for tactile stimuli (blind group), visual stimuli (sighted group), and audio
- 652 stimuli (both groups). Error bars denote standard errors +/- the mean. Asterisks (*) denote

653 significant Bonferroni-corrected pairwise comparisons within the task (p < 0.05). T = tactile, V = 654 visual.

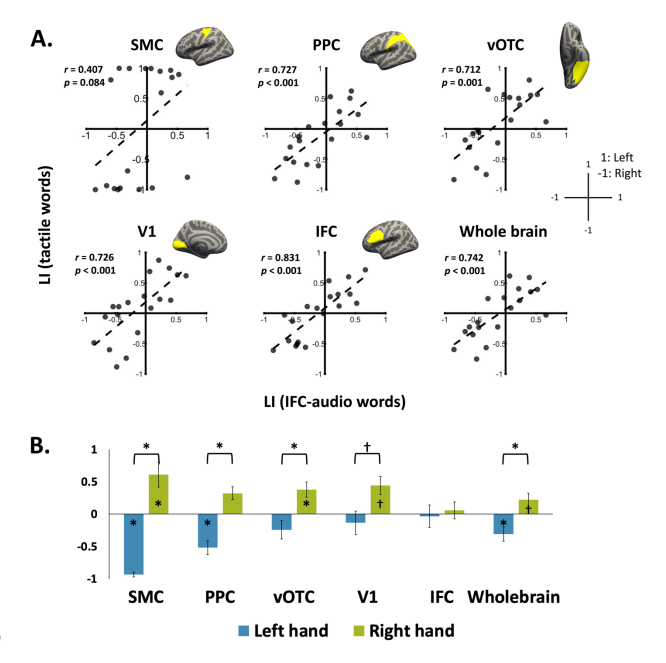
Lateralization of Braille correlates with spoken language lateralization and Braille-reading hand

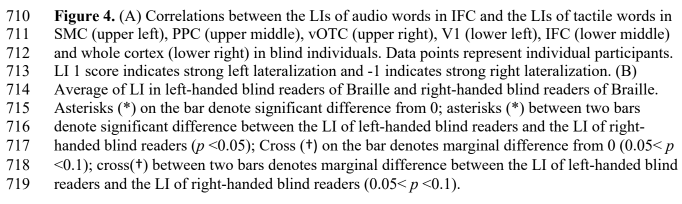
657 We used a lateralization index (LI) analysis to investigate the lateralization of spoken and written 658 language across blind and sighted readers. First, we computed LIs separately for written 659 (tactile/visual words > rest) and spoken (audio words > rest) language in the SMC, PPC, vOTC, 660 V1, IFC and whole cortex in sighted and blind groups. On average, the blind group showed no 661 systematic lateralization for written or spoken words in any region (one-sample t tests of LI = 0, 662 reading: SMC: $t_{(18)} = 0.167$, p = 0.869; PPC: $t_{(18)} = -1.257$, p = 0.225; vOTC: $t_{(18)} = 0.799$, p = 0.799, p = 0.79663 0.435; V1: $t_{(18)} = 0.735$, p = 0.472; IFC: $t_{(18)} = -0.054$, p = 0.958; whole cortex: $t_{(18)} = -0.166$, p = -0.1660.87; listening: SMC: $t_{(13)} = -1.332$, p = 0.206; PPC: $t_{(18)} = 0.051$, p = 0.96; vOTC: $t_{(18)} = 0.322$, p 664 = 0.751; V1: $t_{(18)}$ = -0.506, p = 0.619; IFC: $t_{(18)}$ = -1.135, p = 0.271; whole cortex: $t_{(18)}$ = 0.395, p665 666 = 0.697). For the sighted group, we found left-lateralized activation in vOTC, IFC and whole 667 cortex for written words (one-sample t tests of LI = 0, vOTC: $t_{(14)} = 5.31$, p < 0.001; IFC: $t_{(13)} =$ 668 5.776, p < 0.001; whole cortex: $t_{(14)} = 5.748$, p < 0.001). The sighted group's SMC, PPC and V1 669 activity was not systematically lateralized for written words (one-sample t tests of LI = 0, SMC: 670 $t_{(10)} = 1.172$, p = 0.268; PPC: $t_{(13)} = 0.404$, p = 0.692; V1: $t_{(14)} = 1.614$, p = 0.129). For spoken 671 words, the sighted group's vOTC and IFC activity was left lateralized (one-sample t tests of LI = 672 0, vOTC: $t_{(14)} = 3.42$, p < 0.01; IFC: $t_{(13)} = 3.767$, p < 0.01). We found right-lateralized activation 673 in PPC and V1 for spoken words in the sighted group (one-sample t tests of LI = 0, PPC: $t_{(13)} = -$ 674 3.161, p < 0.01; V1: $t_{(8)} = -3.872$, p < 0.01). There were no systematic lateralization in SMC and 675 whole cortex for the listening task (one-sample t tests of LI = 0, SMC: $t_{(13)} = -0.848$, p = 0.412; 676 whole cortex: $t_{(14)} = 1.449$, p = 0.169). To summarize, we found left-lateralized activity in vOTC 677 and IFC for written and spoken words in the sighted group. By contrast, the blind group did not 678 show systematic lateralization in any of the regions or the whole cortex for written or spoken 679 words. Among blind participants there was substantial variability in lateralization, with some 680 participants showing strong left and others strong right lateralization, consistent with previous 681 studies of lateralization of spoken language in this population (Figure 4, see also Lane et al., 682 2017 and Roder et al., 2002).

683 Next, we determined if lateralization of the Braille reading network could be predicted by the 684 laterality of spoken language and Braille reading hand across blind individuals. A multiple 685 regression analysis was conducted in each region, with the LI of spoken words in IFC and 686 dominant reading hand entered as the regressors and the LI of written words as the dependent 687 variable. First, both the dominant reading hand and the LI of spoken words in IFC predicted the LI 688 of written words in PPC, vOTC and whole cortex (PPC: dominant reading hand: $\beta = 0.55$, p < 0.55689 0.001; LI of spoken words in IFC: $\beta = 0.55$, p < 0.001; adjust $r^2 = 0.843$; vOTC: dominant reading 690 hand: $\beta = 0.468$, p < 0.01; LI of spoken words in IFC: $\beta = 0.611$, p = 0.001; adjust $r^2 = 0.727$; whole 691 cortex: dominant reading hand: $\beta = 0.399$, p = 0.001; LI of spoken words in IFC: $\beta = 0.534$, p < 0.001692 0.001; adjust $r^2 = 0.761$). Second, in V1 and the IFC, only the LI of spoken words predicted the LI 693 of written words (V1: dominant reading hand: $\beta = 0.258$, p = 0.144; LI of spoken words in IFC: $\beta =$ 694 0.734, p = 0.001; adjust $r^2 = 0.575$; IFC: dominant reading hand: $\beta = -0.112$, p = 0.359; LI of spoken 695 words in IFC: $\beta = 0.814$, p < 0.001; adjust $r^2 = 0.702$). Last, we found in SMC, only the dominant 696 reading hand predicted the LI of written words (dominant reading hand: $\beta = 1.624$, p < 0.001; LI 697 of spoken words in IFC: $\beta = 0.311$, p = 0.261; adjust $r^2 = 0.771$). To summarize, in blind 698 individuals, responses to Braille written words and spoken words were co-lateralized to the same 699 hemisphere across most of the Braille reading network, including the vOTC, V1, PPC, and the 700 IFC. Braille reading hand also had an effect on the lateralization of Braille written words in 701 vOTC, PPC, and SMC.

- In the sighted group, we did not find the co-lateralization of spoken and written language to the
- same hemisphere. The correlation between the LI of spoken words in IFC and the LI of written
- words in vOTC was not significant (r = -0.233, p = 0.423). In addition, there were no
- correlations between the LI of spoken words in IFC and the LI of written words in V1, SMC or
- 706 whole cortex (V1: r = -0.301, p = 0.296; SMC: r = 0.169, p = 0.62; whole cortex: r = 0.12, p =
- 0.683). However, the LI of spoken words in IFC was positively correlated with the LI of written
- 708 words in PPC and IFC (PPC: r = 0.55, p < 0.05; IFC: r = 0.732, p < 0.01).

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720 Whole cortex analyses

721 Tactile Braille (blind) and visual print (sighted) reading activated both common and distinctive 722 cortical areas across groups. For reading as compared to rest, both sighted (visual words) and 723 blind (Braille words) readers activated the bilateral vOTC (blind peak: -41, -57, -13; sighted 724 peak: -41, -58, -12), including the location of the classic VWFA (peak: -46, -53, -20), as well as 725 early visual cortices, specifically the foveal confluence (V1/V2/V3) (Figure 5). Within vOTC, 726 responses in the blind group extended further medially and anteriorly and were more extensive in the right hemisphere, relative to the sighted group. The vOTC activation in the blind group also 727 728 extended further laterally and superiorly, into lateral occipital, occipitotemporal, and inferior temporal cortex. Both groups also activated posterior prefrontal cortices (inferior frontal gyrus 729 730 and middle frontal gyrus). Notably, visual cortex responses (e.g., V1) are likely driven at least in 731 part, by different processes across groups, since the sighed group is performing a visual task, 732 whereas the blind group is performing a tactile task.

733 In the blind but not sighted group, reading relative to rest produced extensive activation in 734 bilateral posterior parietal cortices, including superior parietal lobule and supramarginal gyrus 735 (SMG). This parietal activation was posterior to early sensory-motor hand representations. The 736 sighted group activated only a small cluster in parietal cortex, in the left superior parietal lobule. 737 The blind, but not sighted group, also activated parieto-occipital and dorsal occipital regions 738 (middle occipital gyrus). The sighted group additionally activated a lateral temporal region that 739 was not observed in the blind group. Finally, whereas responses to written words were left-740 lateralized in the sighted group, they were bilateral in the blind group.

Listening to words (audio words > rest) likewise revealed partially overlapping responses across groups. In the blind group only, listening to words activated the bilateral vOTC (peak: -42, -44, -16), including the location of the classic VWFA, and early 'visual' cortices. Both groups activated classic fronto-temporal language regions in inferior and lateral prefrontal as well as lateral temporal cortices (Figure 5). Responses in frontal regions were left-lateralized in the sighted group and bilateral in the blind group. The sighted but not blind group activated the left sensorimotor cortex/postcentral gyrus. 748 Reading as compared to hearing words (tactile/visual words > audio words) also revealed 749 similarities and differences across groups (Figure 5). For the sighted group (visual words > audio 750 words), reading words induced greater activation in bilateral vOTC, including the typical 751 location of the VWFA and regions posterior to it, as well as bilateral early visual cortices. Like 752 the sighted, the blind group also activated a region in the left vOTC (fusiform; peak: -27, -61, -753 14), but this activation was medial to the typical VWFA location. A cluster of activity was also 754 observed lateral to the typical VWFA location in the blind group, in the inferior temporal/lateral 755 occipital cortex (peak: -45 -67 -6). Outside vOTC, a cluster of activity was also observed in the 756 blind group in left foveal early 'visual' cortices. The blind but not sighted group also showed 757 extensive activation in posterior parietal cortices, including the SMG and superior parietal 758 lobule. Blind readers also activated dorsal occipital/parieto-occipital cortices during reading. The 759 blind group additionally activated the bilateral superior frontal gyrus and right precentral gyrus. 760 In the sighted group, a small cluster was observed in the left superior parietal lobule. 761 In sum, we observed the following pattern. First, although both groups activated vOTC during

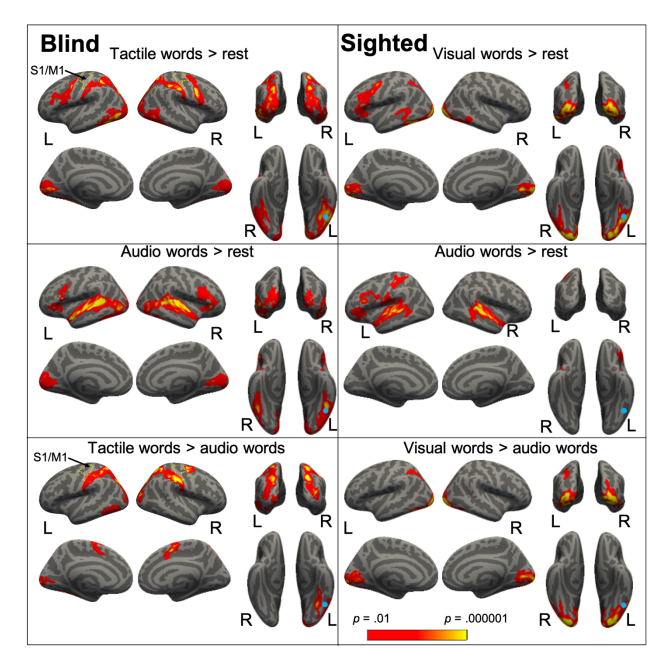
reading, the peak location, distribution and functional profile of responses in vOTC were distinct

763 across groups. Only the blind group showed robust vOTC responses during spoken word

764 comprehension. Second, in contrast to the sighted group, the blind group activated extensive

765 posterior parietal, parieto-occipital, and dorsal occipital areas during (Braille) reading. This

766 parieto-occipital stream was not engaged by spoken word comprehension in blind readers.



768Figure 5. Whole-cortex results for blind (left column) and sighted (right column) groups for the769following contrasts: tactile/visual words > rest (top row), audio words > rest (middle row),770tactile/visual words > audio words (bottom row). The blue circles marked the approximate771location of the classic VWFA (MNI coordinate: -46, -53, -20). The yellow line marked the hand772S1/M1 region. Whole cortex contrasts were thresholded at p < 0.01 vertex-wise, and at p < 0.05773cluster-corrected.

774 **Discussion**

775 Consistent with previous studies, we find that reading activates partially overlapping networks 776 across blind readers of tactile Braille and sighted readers of visual print. In particular, we 777 observed similar responses to written and spoken words and letters in the left IFC of sighted and 778 blind people. We also observed partially overlapping responses in the vOTC across groups. In 779 agreement with past findings, the highest peak of activation for Braille reading relative to rest 780 and visual reading relative to rest was near the canonical 'VWFA' location (Braille words > rest: 781 -41, -57, -13; sighted visual words > rest: -41, -58, -12) (Cohen et al., 2000; Dzięgiel-Fivet et al., 782 2021; Kim et al., 2017; Rączy et al., 2019; Reich et al., 2011). However, we also observed key 783 differences in the neural bases of Braille and visual print reading, in vOTC, V1, and posterior 784 parietal cortices, as well as in lateralization patterns.

785 vOTC of sighted but not blind readers contains a hierarchical word form gradient

786 Consistent with past research, in sighted readers, we observed a posterior-to-anterior functional 787 gradient only in the left vOTC. The posterior portion of the left vOTC responded equally to all 788 visual stimuli in the ROI analysis, the middle portion showed a preference for consonant strings, 789 while the most anterior portion responded more to words and consonant strings than to false 790 fonts. Preferential responses to consonant strings in the middle vOTC of sighted readers are 791 consistent with prior literature showing stronger activation to non-word stimuli in the VWFA 792 when longer presentation times are used or more attention is required (Bruno, Zumberge, Manis, 793 Lu, & Goldman, 2008; Cohen, Dehaene, Vinckier, Jobert, & Montavont, 2008; Dehaene & 794 Cohen, 2011; Ludersdorfer, Schurz, Richlan, Kronbichler, & Wimmer, 2013). A winner-take-all 795 map revealed a similar pattern as the ROI analysis and further showed larger responses to false 796 fonts than consonant strings or words in posterior portions of left vOTC. The larger responses to 797 false fonts in posterior vOTC likely reflect greater attention to less familiar visual stimuli, as 798 indicated by slower reaction times and poorer accuracy. This pattern is consistent with prior 799 studies with sighted readers (Ludersdorfer et al., 2013; Wang, Yang, Shu, & Zevin, 2011). In 800 addition, in sighted readers, the posterior vOTC showed a modality-specific response: above rest 801 activity during visual reading and deactivation during listening, while the most anterior aspect of 802 left vOTC responded equally to visual and auditory stimuli. These results are consistent with the

803 view that in sighted readers the middle and anterior portions of left lateral vOTC become 804 specialized for recognition of letters and words, constituting the so-called 'VWFA.'

805 By contrast, in the blind group, we found no evidence for left-lateralization, no evidence for a 806 posterior-to-anterior functional gradient, or posterior/anterior change in modality preference. In 807 blind readers, the entire posterior/anterior extent of bilateral vOTC showed a preference for 808 Braille words over consonant strings and tactile shapes during reading and a larger response to 809 spoken words than backward speech. Unlike in the sighted group, in the blind group, no portion 810 of the vOTC showed a consonant string preference over words and shapes, whereas in the 811 sighted group, the middle vOTC responded more to consonant strings than words. In addition, 812 there was no change in preference for written as opposed to spoken words along the

813 posterior/anterior extent of vOTC.

814 The Whole-cortex analysis also revealed differences in lateral/medial organization of vOTC 815 across groups. As noted above and previously documented, when Braille and spoken words were 816 compared to rest, a peak of activation was observed in the classic VWFA region along the 817 medial/lateral axis, although in blind readers, additional activity was also observed throughout 818 much of vOTC. By contrast, when Braille words were compared to spoken words in the blind 819 group, peak activity in the vOTC was medial to the classic VWFA location (peak: -27, -61, -14). 820 We did not observe such a medial peak for the same contrast in sighted readers. This medial 821 vOTC region has previously been shown to be functionally connected to dorsal parietal cortices, 822 which are involved in spatial attention and effortful letter-by-letter reading in sighted people 823 (Bouhali, Bézagu, Dehaene, & Cohen, 2019; Cohen et al., 2008; Corbetta & Shulman, 2002; 824 Henry et al., 2005; Saalmann, Pigarev, & Vidyasagar, 2007). As discussed in detail below, the 825 PPC appears to play an important role in Braille reading and may send information to medial 826 vOTC in blind readers.

Although the precise role of vOTC in Braille reading remains to be determined, the present evidence suggests that although the vOTC is involved in both tactile reading and visual reading, the anatomical distribution of responses within vOTC, the functional profile, and therefore likely the cognitive contribution differs. In sighted readers, information reaches lateral vOTC from early visual areas and is sent onward to fronto-temporal language regions, as well as receiving 832 top-down input from the language regions (Bouhali et al., 2014; Hannagan et al., 2015; Saygin et 833 al., 2016; Stevens et al., 2017b; Yeatman et al., 2013). Lateral vOTC thus contributes to 834 decoding linguistic information (phonological, semantic, and grammatical) from visual word 835 forms (Dehaene & Cohen, 2011; Price & Devlin, 2011). By contrast, we hypothesize that in 836 blind readers of Braille, the classic VWFA location in lateral vOTC receives linguistic (i.e., 837 semantic, grammatical) information from fronto-temporal language circuits and serves as one of 838 the entry points for language into posterior 'visual' circuits. This hypothesis is supported by prior 839 studies showing that in blind but not sighted people, the classic VWFA location is sensitive to 840 syntactic complexity of spoken sentences and shows enhanced responses to spoken language 841 (Burton, Snyder, Diamond, et al., 2002; Dzięgiel-Fivet et al., 2021; Kim et al., 2017; Lane et al., 842 2015). At the same time, the current data and prior evidence suggest that other parts of the 843 'visual' cortex, including a medial portion of vOTC, may play a role in Braille reading. An 844 intriguing albeit speculative possibility is that medial vOTC receives Braille-relevant input from 845 PPC. Lacking connectivity data, the present study cannot test this hypothesis directly. One way 846 to test this possibility in future work would be to use online TMS in combination with fMRI to 847 disrupt information flow to the vOTC in blind readers of Braille by stimulating parietal cortices.

848 Parieto-occipital decoding stream in blind readers of Braille

849 We observed more extensive and different involvement of posterior parietal/parieto-occipital 850 cortices in Braille as opposed to visual print reading. Large segments of PPC were activated 851 during Braille reading relative to rest and spoken word comprehension. PPC activity in the blind 852 group extended inferiorly and anteriorly, into regions adjacent to and immediately posterior to 853 S1, including the supramarginal gyrus (SMG) and much of the superior parietal lobule. Notably, 854 the hand regions of S1 itself did not show robust responses during Braille reading or preferential 855 responses to Braille letters or words, consistent with prior studies (Burton, Snyder, Conturo, et 856 al., 2002; Kupers et al., 2007). Additionally, in the blind group only, parietal activation extended 857 posteriorly into parieto-occipital and dorsal occipital regions adjacent to parietal cortices and 858 ultimately into the foveal confluence. By contrast, visual print reading (relative to false fonts) by 859 sighted readers activated only a small region within the superior parietal lobule, consistent with 860 prior studies (Cohen et al., 2008; Martin, Schurz, Kronbichler, & Richlan, 2015; Reilhac, Peyrin, 861 Démonet, & Valdois, 2013).

862 The cognitive role of the wider parietal network in Braille reading is not known. The PPC has 863 strong connectivity with S1 and contains high-level tactile areas, as well as multimodal 864 representations of texture and shape (Bauer et al., 2015; Hegner et al., 2010; Kaas, 2012). Some 865 of the activation we observed likely reflects processes related to recognition of tactile patterns 866 that constitute Braille but are not specific to Braille letters or words (Boven, Hamilton, 867 Kauffman, Keenan, & Pascual-Leone, 2000; Wong, Gnanakumaran, & Goldreich, 2011), akin to 868 general responses to shapes, including false fonts, observed in vOTC of sighted readers (Grant, 869 Thiagarajah, & Sathian, 2000; Sathian & Stilla, 2010; Stilla et al., 2008). Consistent with this 870 possibility, much of the PPC, particularly its anterior portion, was more responsive to the more 871 tactilely complex and unfamiliar dot shapes than to Braille letters or words. Again, this paralleled 872 preferential responses to false fonts in posterior vOTC of sighted readers. Importantly, however, 873 within the larger swath of PPC activation, ROI analyses revealed word and letter preferring 874 subregions in the blind group, suggesting a specific involvement in Braille processing.

Word-specific activation in parieto-occipital areas extended posteriorly into dorsal occipital
cortices, only in the blind group. Unlike anterior portions of the PPC, parieto-occipital and dorsal
occipital areas showed larger responses to Braille words than Braille consonants or control
shapes. However, like anterior PPC, parieto-occipital and dorsal occipital regions responded
more to Braille words than to spoken words. This pattern suggests that parieto-occipital and
dorsal occipital areas are involved in reading-specific processing, rather than language
comprehension or tactile pattern recognition.

882 The winner-take-all map of PPC showed that the preference for words is located in the posterior 883 aspect of the PPC, adjacent to parieto-occipital and dorsal occipital areas. Interestingly, in the 884 blind group only, this map also revealed consonant preferring regions in an anatomically 885 intermediate position between shape preferring areas in anterior portions of PPC and word 886 preferring areas in parieto-occipital and dorsal occipital cortices. These regions did not emerge in 887 corrected whole-brain analyses and therefore should be interpreted with caution, requiring 888 investigation in future studies. However, the overall pattern suggests an anterior-to-posterior 889 parieto-occipital reading stream, analogous to the posterior-to-anterior vOTC gradient observed 890 in sighted readers. Within this gradient, parietal regions closer to S1, in anterior PPC represent 891 shape/texture information relevant to Braille, with posterior PPC and parieto-occipital regions

representing Braille orthography and still more posterior occipital areas representing linguisticinformation.

As noted in the Introduction, involvement of the PPC in Braille reading is predicted by

connectivity-based theories of brain function (Bedny, 2017; Hannagan et al., 2015; Mahon &

896 Caramazza, 2011; Saygin et al., 2016). One hypothesis, therefore, is that the PPC, along with

897 adjacent parieto-occipital areas, plays an analogous role in Braille orthographic processing to the

role of the vOTC in orthographic processing of visual print: conversion of tactile patterns to

899 orthographic representations (Dehaene & Cohen, 2011; Dehaene et al., 2005).

900 Further work is needed to uncover the precise cognitive contribution of PPC and parieto-901 occipital cortices to Braille reading. In sighted readers, the PPC also contributes to reading but 902 under different circumstances. The PPC is thought to participate in grapheme to phoneme 903 conversion, letter position decoding, as well as working memory processes and shows more 904 robust activity when effortful letter-by-letter reading is required (e.g., when words are degraded) 905 (Carreiras, Quiñones, Hernández-Cabrera, & Duñabeitia, 2015; Cohen et al., 2004; Costanzo, 906 Menghini, Caltagirone, Oliveri, & Vicari, 2012; Dehaene-Lambertz, Monzalvo, & Dehaene, 907 2018; Henry et al., 2005; Jonides et al., 1998; Koenigs, Barbey, Postle, & Grafman, 2009; 908 Ossmy, Ben-Shachar, & Mukamel, 2014; Taylor, Rastle, & Davis, 2013). Parietal cortex also 909 shows sensitivity to phonological rather than orthographic information during visual reading, in 910 contrast to the VWFA (Booth et al., 2003; Bouhali et al., 2019). In future studies, it would be 911 interesting to separate parietal responses to phonological as opposed to word-form information in 912 blind readers of Braille. In addition, further research is needed to explore the anatomical layout 913 of Braille-responsive parietal areas. For example, whether the parieto-occipital stream contains 914 punctate regions analogous to the VWFA, or more distributed responses to Braille letters and 915 words remains an open question. Likewise, in future studies, it will be important to test the 916 precise role of the PPC in Braille reading and to dissociate the functions of PPC, parieto-917 occipital, and dorsal occipital regions.

918 Differential role of early visual cortex in Braille and visual print reading

We observed responses to reading in V1 in the blind but not sighted group. Like dorsal occipitalareas, V1 showed a preference for words over consonant strings and control shapes. The

921 involvement of V1 in Braille reading is consistent with previous studies (Cohen et al., 1997;

- 922 Kupers et al., 2007; Sadato et al., 1996). We further found that, in whole-cortex results, a portion
- 923 of V1 (foveal aspect of left V1) responded more to Braille reading than auditory word
- 924 comprehension, whereas other portions of V1 (right hemisphere, and peripheral) did not show
- 925 such a preference. This evidence is consistent with prior work suggesting that V1 does not have a
- 926 single, homogeneous function in people who are blind but rather contains multiple anatomically
- 927 separable functional subdivisions (Amedi et al., 2003; Bedny et al., 2011; Burton, Diamond, &
- 928 McDermott, 2003; Burton, Snyder, Diamond, et al., 2002; Kanjlia et al., 2021; Kanjlia, Pant, &
- Bedny, 2019; Lane et al., 2015; Noppeney et al., 2003). Likewise, V1 may contain anatomically
- 930 separable Braille-specific and high-level language responses in blind readers of Braille.

931 Lateralization of Braille reading: effects of spoken language lateralization and reading 932 hand

933 With the exception of the primary somatosensory cortex, laterality of responses to written words 934 in the entire reading network (vOTC, PPC, V1, and IFC) is predicted by the laterality of spoken 935 word comprehension across blind individuals. On average congenitally blind individuals showed 936 reduced left-lateralization of responses to spoken and written words (see also Lane et al., 2017). Those blind individuals who show right-lateralized responses to spoken words also show right-937 938 lateralized responses to written words. Previous studies with sighted readers with right 939 hemisphere spoken language responses have likewise observed co-lateralization of spoken and 940 written language (Cai et al., 2010; Van der Haegen et al., 2012). We did not observe this pattern 941 in the current sighted sample, possibly because all sighted participants in the current study had 942 strongly left-lateralized responses to spoken language and thus there was little interindividual 943 variability. Together, these data suggest that written and spoken language tend to co-lateralize in 944 blind and sighted readers alike. This observation is consistent with the hypothesis that strong 945 connectivity to spoken language networks is one of the determining factors of which regions 946 become 'recycled' for reading.

We also found a significant effect of reading hand on the lateralization of Braille reading that
was independent of the effect of spoken language lateralization. That is, right-hand Braille
readers showed more left-lateralized activation whereas left-hand Braille readers showed a

bilateral response to Braille. In contrast to the effect of spoken language on laterality, the effect
of reading hand was strongest in the primary somatosensory cortex, persisted in PPC and vOTC,

and was absent in IFC and V1. This observation is consistent with the idea that V1 occupies the

top of a processing hierarchy for people who are blind (Buechel, 2003). Effects of reading hand

thus persist past S1 but wane at higher stages of processing, whereas effects of language

- 955 lateralization are most prominent at higher processing stages and disappear in early sensory areas
- 956 (i.e., S1).

957 In sum, the lateralization of Braille reading is jointly determined by the lateralization of spoken

958 language and the input hand that receives the initial Braille stimulus. Although specific

959 lateralization patterns differ across sighted and blind groups, an analogous connectivity principle

960 appears to govern lateralization of reading in sighted and blind readers: lateralization depends

961 jointly on connectivity to sensory input regions (unilateral S1/ bilateral V1) and language

962 networks.

963 General conclusions

964 We find that the neural basis of Braille reading differs from that of visual print reading in several 965 ways. While visual print reading recruits a posterior/anterior vOTC gradient, no such gradient is 966 observed in the vOTC of blind readers of Braille. Blind readers of Braille recruit posterior 967 parietal cortices to a greater degree and in a different way compared to visual print reading in 968 sighted people. Only blind readers show preferential responses to written words in PPC and 969 parieto-occipital cortex. We observed suggestive evidence for an anterior-to-posterior stream of 970 processing in the parietal cortex of blind Braille readers, with anterior parietal areas involved in 971 tactile pattern perception and more posterior parietal, parieto-occipital and dorsal occipital 972 regions involved in word recognition. In blind and sighted readers alike, lateralization of spoken 973 language predicts lateralization of written language. However, on average, spoken word and 974 visual word recognition is highly left-lateralized in sighted people. By contrast, neither Braille 975 reading nor spoken word recognition is strongly left-lateralized in people who are born blind. In 976 blind readers of Braille, reading hand also affects lateralization of responses to Braille.

977 Comparing the neural basis of reading across blind and sighted people suggests that there is no978 'standard reading brain.' The input modality of symbols influences the neural basis of their

- 979 recognition. At the same time, similar anatomical principles govern the localization of visual
- 980 print and tactile Braille. Connectivity patterns constrain the localization of visual print and tactile
- 981 Braille reading alike.

983

985 Credit Author Statement

- J.K., S.K., and M.B. designed research; J.K., S.K., and M.B. performed research; M.T. analyzed
 data; M.T., E.S. and M.B. wrote the paper.
- 988

989 Declaration of Competing Interest

990 The authors declare no competing interests.

991 Acknowledgments

- 992 We would like to thank all our blind and sighted participants, the blind community and the
- 993 National Federation of the Blind. Without their support, this study would not be possible. This
- work was supported by grants from the Johns Hopkins Science of Learning Institute (80034917)
- and the NIH/NEI (R01 EY027352-01). We would also like to thank the F. M. Kirby Research
- Center for Functional Brain Imaging at the Kennedy Krieger Institute for their assistance in datacollection.

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