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Lateral interactions: size does matter $\stackrel{\text{tr}}{\sim}$

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6 Abstract

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7 Usually a high-contrast, co-local mask increases contrast threshold (inhibition). Interestingly, a laterally displaced mask (flanker) 8 can facilitate contrast detection (Vision Research 33 (1993) 993; 34 (1994) 73). When spatial scaling of these flanker effects was 9 implied, stimulus bandwidth was confounded with spatial frequency (λ^{-1}). Under conditions where at lower spatial frequencies, the 10 size (standard deviation, σ) of the Gabor patch was smaller ($\sigma < \lambda$) than higher spatial frequencies ($\sigma = \lambda$), the effect appeared scale 11 invariant. We replicated the original results for all conditions. However, when Gabor size was fixed ($\sigma = \lambda$), facilitation changed 12 with spatial frequency (range 2–13 cycles/deg.). When Gabor size was varied ($\sigma = 0.5-2\lambda$), usually the combination of larger patch 13 sizes and lower spatial frequencies caused inhibition. We were unable to find any conditions that demonstrated spatial scaling. The 14 size, both λ and σ , of both stimulus and flankers, influenced contrast threshold. Also, facilitation reduced as contrast of the flankers 15 was reduced to detection threshold. Some facilitation was apparent with sub-threshold flankers. These results need to be reconciled 16 with current models of lateral interactions. © 2002 Published by Elsevier Science Ltd.

17 Keywords: Lateral interactions; Contrast sensitivity; Stimulus bandwidth; Contrast

18 1. Introduction

19 Object detection can be affected by spatial context, 20 other objects facilitating or inhibiting detection. Incre-21 ment thresholds can be considered as the minimum de-22 tectable change in the characteristics of one sub-23 threshold target superimposed on a second target with 24 matching spatial characteristics. Both sub-threshold and 25 supra-threshold masks can influence contrast detection 26 (Kulikowski & King-Smith, 1973; Legge, 1979; Tolhurst 27 & Barfield, 1978). The second target is called a mask 28 because of its effect at high contrast. At high mask 29 contrasts typically there is inhibition, but as mask con-30 trast decreases detection may be facilitated (threshold 31 lower than with no mask) (Legge, 1979; Tolhurst & 32 Barfield, 1978). More generally, spatial masking is the 33 impact of one target on the detection of another, where

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the two targets may or may not have matching spatial 34 characteristics. For example, a large, co-centric mask 35 (e.g. a pedestal) may alter contrast threshold, with mask 36 size one of the important parameters (Legge, 1978; 37 38 Westheimer, 1965, 1967; Yu & Levi, 1997a,b). Using such increment-threshold paradigms, the spatial fre-39 quency tuning (Legge, 1978; Tolhurst & Barfield, 1978; 40 Wilson, McFarlane, & Phillips, 1983; Yu & Levi, 1998) 41 and orientation tuning (Phillips & Wilson, 1984; Yu & 42 Levi, 1998) of the mechanisms detecting sine-wave 43 gratings have been investigated. Most masking condi-44 tions inhibit (worsen) contrast detection. Contextual 45 effects of masks on contrast detection may be mediated 46 47 by short-range cortical connections (Das & Gilbert, 1999). 48

49 Interestingly, an appropriate flanker—a mask that is laterally displaced from the target (i.e. no longer co-50 centric)-may facilitate (improve) contrast detection 51 (Morgan & Dresp, 1995; Polat & Sagi, 1993, 1994a; 52 Wehrhahn & Dresp, 1998; Yu & Levi, 1997d). Polat and 53 Sagi (1993, 1994a) reported that the detection threshold 54 of a Gabor patch was lower when the patch was flanked 55 by high contrast Gabor patches. Maximum facilitation 56 (approximately half the non-flanked threshold) was 57 noted when the flankers were laterally displaced from 58

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59 the target patch by a distance equal to two to three 60 wavelengths (λ). Larger displacements (up to 8λ or 12λ) 61 produced measurable facilitation, while short displace-62 ments (e.g. 0λ or 1λ) produced inhibition (short dis-63 placements are similar to co-centric masking, as the 64 flankers and stimulus overlap). These effects were reported to be spatial frequency independent (which im-65 66 plies spatial scaling) (Polat & Sagi, 1993). Spatial scaling 67 is important as it implies a general principle of uniform operation of the visual system across all scales. Similar 68 69 facilitation by laterally displaced objects (flankers) have 70 been noted for other spatially localised (but less well 71 spatial frequency defined) objects (Morgan & Dresp, 72 1995; Westheimer, 1965; Yu & Levi, 1997d). Flanker 73 effects have been ascribed to long-range connections in 74 the visual cortex (Das & Gilbert, 1995; Gilbert, Das, Ito, 75 Kapadia, & Westheimer, 1996; Kapadia, Ito, Gilbert, & 76 Westheimer, 1995; Polat, Mizobe, Pettet, Kasamatsu, & 77 Norcia, 1998; Polat & Norcia, 1996; Yu & Levi, 1997b). 78 Careful examination of Polat and Sagi's experimental 79 conditions shows that stimulus spatial frequency and 80 bandwidth were confounded in their demonstrations of 81 the spatial scaling of the facilitation effects (Polat & 82 Sagi, 1993). For example, proportionally the Gaussian 83 envelope used for the high spatial frequency objects and 84 flankers was larger (standard deviation, $\sigma = \lambda$) than for 85 the lower spatial frequency objects and flankers 86 $(\sigma = 0.5\lambda)$, thereby altering the bandwidth of both 87 stimulus and flanker. Previously mask size has been 88 shown to alter contrast detection (Legge, 1978; Yu & 89 Levi, 1997c). As the bandwidth of Polat and Sagi's 90 stimuli may have interacted with the change in spatial 91 frequency, we examined size effects by systematically altering spatial frequency (λ^{-1}) and bandwidth (σ) . 92 93 Spatial scaling of these effects has implications for visual 94 processing at low spatial frequencies. Low spatial fre-95 quencies are important to people with visual impairment 96 through foveal (or macular) vision reduction, as high 97 spatial frequencies are not detected and many use ec-98 centric retinal locations to view. Low to medium spatial 99 frequencies may be sufficient to mediate many important 100 visual tasks such as letter and face recognition (Parish & 101 Sperling, 1991; Peli, Goldstein, Young, Trempe, & 102 Buzney, 1991; Peli, Lee, Trempe, & Buzney, 1994; Sol-103 omon & Pelli, 1994). We report that stimulus size, in 104 terms of grating spatial frequency and Gaussian enve-105 lope, influenced measured lateral interactions. The lack 106 of scaling highlights limitations of current models for 107 lateral interactions (Polat, 1999; Solomon & Morgan, 108 2000; Solomon, Watson, & Morgan, 1999; Yu & Levi, 109 1997c; Zenger & Sagi, 1996). Also we studied the impact of flanker contrast. Some

Also we studied the impact of flanker contrast. Some
facilitation was apparent with flankers that were at or
slightly below detection threshold. This extends earlier
reports of facilitation with a sub-threshold co-located
mask (Kulikowski & King-Smith, 1973; Legge, 1979;

Tolhurst & Barfield, 1978) and may be related to the115report that sub-threshold stimulation is apparent in cells116outside the conventional receptive field and beyond the117zone that has spiking activity (Das & Gilbert, 1995).118

2. Methods

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1	2	20	20

Five subjects (four males and one female, aged 23-40 121 122 years) with normal or corrected-to-normal vision participated in the study. Two of the observers (AN and 123 RW) had extensive previous experience as psychophys-124 ical subjects and were aware of the purpose of the ex-125 periments while the other three subjects had no prior 126 experience of contrast sensitivity measures or discrimi-127 nation tasks and were naïve to the purpose of the study. 128

2.2. Apparatus 129

The stimuli were generated using a VisionWorksTM 130 computer graphics system (Vision Research Graphics 131 Inc., Durham, NH) and were presented on a NanaoTM 132 EIZO® monitor. The video format was 120 Hz non-133 interlaced. The video resolution was 1024×600 pixels 134 occupying an area of 23.4 cm (vertical) \times 40 cm 135 (horizontal) $(13.2^{\circ} \times 21.8^{\circ} \text{ at } 100 \text{ cm})$. Luminance of the 136 monitor was controlled with a 12-bit look-up table. 137 Mean display luminance was 37.5 cd/m² in an otherwise 138 dark environment. 139

2.3. Stimuli 140

Sinusoidal gratings in a Gaussian envelope (Gabor 141 function) were presented in a collinear arrangement on 142 the vertical meridian (Figs. 1A and 2A), except where 143 specified otherwise. The luminance of each Gabor patch, 144 $L(x, y, \theta)$, was defined by 145

$$L(x, y, \theta) = L_0 \left\{ 1. + C \cos\left(\frac{2\pi}{\lambda} \left[(x_0 - x_0) \cos \theta + (y_0) \sin \theta \right] \right) \exp\left(- (x_0 - x_0)^2 + (y_0 - y_0)^2 \right) / \sigma^2 \right\}$$

147 where x is the value of the horizontal axis, y of the vertical axis, (x_0, y_0) is the centre of the Gabor patch, λ 148 is the wavelength and θ is the orientation of the carrier, 149 and σ is the standard deviation of the Gaussian enve-150 lope, L_0 is the background luminance and C is the 151 contrast of the patch. A test patch was placed between 152 two flanking patches of 40% contrast (except for control 153 conditions in Experiment 3). Each patch was displayed 154 within a rectangle that was $4\sigma_x$ by $4\sigma_y$. For all experi-155 ments, except as indicated, the gratings were vertical and 156 the flanking patches were equidistant above and below 157

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158 the test patch. Test-flanker distance was defined as the 159 distance between the centre of the test patch and the 160 centre of a flanking patch. As our system did not allow 161 us to overlap the flanker patches in any one display, the 162 minimum test-flanker distance that we could display was 163 2σ . To achieve this test-flanker separation it was neces-164 sary to display the test patch and the flanking patches in 165 separate interlaced frames (stereo mode: Yu & Levi, 166 1997c; Williams & Hess, 1998). We did this for all conditions. In this configuration, only half of the max-167 168 imum contrast was available for any patch because each 169 patch was presented in every other frame only (Williams 170 & Hess, 1998; Yu & Levi, 1997c). Contrast detection 171 with the flanking patches present was compared to a 172 'standard' condition where detection threshold was de-173 termined for a test patch with no flanking patches

174 2.4. Procedure

175 A two-alternative temporal forced-choice staircase 176 procedure was used. Each trial consisted of two 100 ms 177 presentations, each a temporal square wave (immediate 178 onset and offset), accompanied by audible tones, and 179 separated by 867 ms. Only one presentation contained a 180 test patch, but both presentations contained flanking 181 patches. Audio feedback was given as an indication of 182 an incorrect response. These conditions replicated, as 183 closely as possible, the conditions of the earlier experi-184 ments (Polat & Sagi, 1993, 1994a). During the trials 185 subjects were asked to fixate at the position at which the 186 test patch was presented (centre of monitor) without the 187 aid of a fixation point, and report the interval in which 188 they saw the test patch appear. A fixation target could 189 cause masking effects. In pilot studies, we examined the 190 impact of fixation guides using two small dots posi-191 tioned on both sides of the test patch location or a single 192 moderate contrast dot presented in the location of the 193 test-patch until 500 ms before the test presentation. 194 Neither fixation guide produced a difference in contrast 195 thresholds for the 13.3 cycles/deg. condition (the small-196 est stimulus we used, and for which spatial uncertainty 197 and unsteady fixation and accommodation would be 198 expected to be greatest).

199 Each staircase consisted of two practice and 10 ex-200 perimental reversals or two practice and 40 experimental 201 reversals. The 10-reversal staircases were conducted in 202 sets of four. The geometric mean of 40 experimental 203 reversal contrasts was taken as the contrast threshold 204 (i.e. the average of four 10-reversal staircases or the 40-205 reversal staircase). Each subject completed 3-6 stair-206 cases for each experimental condition. Data is presented 207 as the mean of those staircases, and error bars are the 208 standard error of the mean. The initial contrast of the 209 test stimulus (25%) was easily visible at 2 cycles/deg. but 210 just above threshold at 13.3 cycles/deg. During the 211 staircase procedure, the contrast of the test stimulus was decreased 0.30 log units following three consecutive212correct responses and increased 0.405 log units for each213incorrect response. The staircases are expected to have214converged towards the 83% correct point (García-Pérez,2151998).216

While replicating some of the earlier experiments, we 217 found that non-linearities in the monitor response 218 (García-Pérez & Peli, 2001; Klein, Hu, & Carney, 1996; 219 Pelli, 1997) can alter the measured effects. In particular, 220 using gratings with few pixels per cycle (e.g. Polat & 221 222 Sagi, 1993, 1994a, b used 4 pixels per cycle for their 13.3 223 cycles/deg. targets) we found that the relative orienta-224 tion of the raster and the grating was an important 225 factor on our monitor. As discussed in Appendix A, we found it necessary to use many pixels per cycle (λ) or to 226 227 arrange stimuli so that relatively large changes in lumi-228 nance were not required along a raster line. This was achieved by fixing the size of the stimuli on the monitor, 229 so that there were about 23 pixels per cycle, and varying 230 the viewing distance to change grating spatial frequency. 231

3. Experiment 1: the effect of spatial frequency

To test the spatial scaling reported by Polat and Sagi 233 (1993), the lateral interactions at four spatial frequencies 234 235 (2, 4, 8 and 13.3 cycles/deg.) were tested with test-flanker 236 distances of 2–6 λ with all test and flanker Gabors scaled so that $\sigma = \lambda$ (Fig. 1A). Viewing distances were varied 237 from 100 to 654 cm. If there is spatial scaling then 238 similar facilitation or inhibition should be found for 239 each spatial frequency at each test-flanker distance. This 240 will not occur if the different bandwidths of the different 241 spatial frequencies used by Polat and Sagi had an impact 242 243 on their results.

244 Facilitation was found for all four spatial frequencies (2 to 13.3 cycles/deg.) for all four subjects (Fig. 1). The 245 results for subject BW differed from the other three 246 subjects at the higher spatial frequencies (8 and 13.3) 247 cycles/deg.). This is examined in detail in Experiment 3. 248 In general, for test-to-flanker distances of three, four 249 and six λ the facilitation was very similar for all four 250 251 spatial frequencies. However, substantially different effects were found at the 2λ test-flanker distance, unlike 252 Polat and Sagi (1993). Facilitation was greatest for the 253 254 higher spatial frequencies (8 and 13.3 cycles/deg.), with less facilitation at 2 and 4 cycles/deg.. The lateral in-255 teraction effects at 8 and 13.3 cycles/deg. were very 256 similar to that found by Polat and Sagi (1993) at 6.7 and 257 13.3 cycles/deg. when $\sigma = \lambda$. Any small differences were 258 probably a consequence of monitor non-linearity prob-259 260 lems (see Appendix A) and inter-subject differences (Fig. 1). Our results at 2 and 4 cycles/deg. were substantially 261 different to those of Polat and Sagi (1993) who used 262 $\sigma = 0.67\lambda$ at 4.4 cycles/deg. and $\sigma = 0.5\lambda$ at 3.3 cycles/ 263 264 deg.

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Fig. 1. Investigated spatial scaling of the lateral interaction between the central test stimulus and flankers above and below (A). Test-flanker distance was varied from 2λ to 6λ , where λ was the wavelength of the contrast grating. Facilitation (lower relative contrast threshold) was found at most test-flanker separations. Relative contrast threshold changed with test-flanker separation and grating spatial frequency in a similar manner for (B) subject AN, (C) subject AS, (D) subject BW, and (E) subject RW. As the contrast threshold differed for the four spatial frequencies, particularly at 2λ , spatial scaling does not occur. Error bars are standard error of the mean.

265 As we varied viewing distance to vary the spatial 266 content (visual angle) of our targets, the size of the illuminated field (i.e. the monitor) varied in visual angle. 267 In a control experiment we found that variation in the 268 269 field size had no effect on the results. This is not sur-270 prising given the spatial frequency, position and orientation tuning of these lateral interactions (Polat & Sagi, 271 272 1993, 1994a; Yu & Levi, 1998).

273 Examination of our data did not reveal any learning 274 effects of the sort reported by Polat and Sagi (1994b) 275 that could have confounded the differences between the four spatial frequencies that we tested. However, our 276 277 study was not designed to evaluate learning effects. 278 Further, when parts of this experiment were replicated a 279 few months later (during Experiments 2 and 3) the lateral interactions had not changed in any systematic 280 281 manner. However, it was clear that intra-subject vari-282 ability was larger than we would have liked, but not dissimilar to other groups (Solomon et al., 1999; Wil-
liams & Hess, 1998). In conclusion, the lateral interac-
tions did not spatially scale as reported by Polat and
Sagi (1993), suggesting that the bandwidth of the stimuli
had an important impact.283
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4. Experiment 2: the effect of test and flanker size 288

Since, the results at low spatial frequencies found 289 when $\sigma = \lambda$ (Experiment 1) were different from earlier 290 results when $\sigma \neq \lambda$ (Polat & Sagi, 1993), we examined 291 the lateral interactions when Gabor patch size was sys-292 293 tematically varied from $\sigma = 0.5 - 1.5\lambda$. The Gaussian envelope of both flanker and stimulus were varied si-294 multaneously (i.e. $\sigma_{\text{flanker}} = \sigma_{\text{test}}$) as shown in Fig. 2A. As 295 before, test-flanker distance was defined as the centre to 296 centre distance. Spatial frequencies of 2 and 8 cycles/deg. 297

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298 (viewing distances of 100 and 400 cm respectively) were 299 used to examine whether any effects of patch size scaled 300 with spatial frequency. If patch size is a factor in the 301 lateral interactions, then we expect changes in the lateral 302 interactions as σ is varied.

303 Contrast sensitivity of the control condition (no 304 flankers) varied with patch size as shown previously 305 (Peli, Arend, Young, & Goldstein, 1993). Facilitation 306 was found for both spatial frequencies at all test-flanker distances tested. In general, for greater test-flanker dis-307 308 tances (4 λ and 6 λ) facilitation was similar for all four 309 patch sizes (Fig. 2). However, at shorter test-flanker 310 distances (2λ and 3λ) substantial effects of patch size are 311 apparent, with facilitation for smaller patches ($\sigma = 0.5\lambda$ 312 and 0.75λ) and inhibition for larger patches. For smaller 313 patches (wider bandwidth) maximal facilitation oc-314 curred with a test-flanker distance of 2λ . For the larger 315 patches (narrower bandwidth) maximal facilitation was 316 found at larger test-flanker distances (3 λ or 4 λ), and 317 inhibition increased with patch size. While these effects 318 of patch size were similar for 2 and 8 cycles/deg., the 319 inhibitory effects for larger patches were greater for the 320 lower spatial frequency. The lateral interactions found 321 with 2 cycles/deg. and $\sigma = 0.5\lambda$ were very similar to 322 those reported by Polat and Sagi (1993) at 3.3 cycles/ 323 deg. and $\sigma = 0.5\lambda$.

It is clear from Fig. 2 that the lateral interactions were not scale invariant when defined in terms of the spatial frequency (λ^{-1}) of the stimulus and flanker. It is possible that the greater overlap of test and flanker for larger patch sizes (Fig. 2A) might be an important factor. As shown in Fig. 3A, the lateral interactions appear to have greater similarities when considered in terms of the size330of the patch (σ) for 2 cycles/deg.. However, this apparent331spatial scaling is probably an artefact, since it is not332apparent for 8 cycles/deg. (Fig. 3B).333

As we used a cosine-phase function, the average lu-334 minance of the Gabor patch became slightly brighter 335 than the average monitor luminance for the smaller 336 patches (i.e. when $\sigma < \lambda$) (Cannon & Fullenkamp, 1991; 337 Peli et al., 1993). Therefore, the measured detection 338 threshold may have been luminance detection rather 339 than contrast detection (i.e. detecting the patch rather 340 than the grating). To evaluate this possible artefact, the 341 experiment was repeated at 2 cycles/deg. and $\sigma = 0.5\lambda$ 342 using a sine-phase function for test stimulus and flank-343 ers. The sine-phase function had an average patch lu-344 minance equal to the average monitor luminance. No 345 difference in the lateral interactions was found between 346 the cosine- and the sine-phase function Gabor patches. 347

Stimulus bandwidth (σ) had an effect on the measured 348 lateral interactions. Facilitation similar to that found 349 with higher spatial frequencies and $\sigma = \lambda$ (i.e. contrast 350 detection with only half the contrast of the patch alone) 351 can be found for lower spatial frequencies when the 352 patch size is reduced (e.g. $\sigma = 0.5\lambda$). This effect of 353 stimulus size explains the difference between our results 354 in Experiment 1 and those of Polat and Sagi (1993) for 355 lower spatial frequencies. Also, this is further evidence 356 of the lack of spatial scaling of these lateral interaction 357 effects at close test-flanker distances ($\leq 3\lambda$). Since we 358 altered stimulus and flanker bandwidth simultaneously, 359 it is possible that the effects were not a consequence of 360 the bandwidth alone. The results may have been influ-361



Fig. 2. Investigated the effect of stimulus and flanker size (defined as the standard deviation, σ , of the Gaussian envelope) on contrast detection. As before, test-flanker distance was varied from 2λ to 6λ . Illustrations of stimuli with a test-flanker distance of 3λ and $\sigma = 0.5-1.5\lambda$ are shown in panel A. At both (B) 2 cycles/deg. and (C) 8 cycles/deg., a strong interaction between the test-flanker separation and σ (stimulus size) was found for subject RW. Similar results were found for subject AN (Fig. 3). Error bars are standard error of the mean.

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Fig. 3. Results, similar to those shown in Fig. 2 (for subject RW), were found for subject AN. Here the results are plotted against the test-flanker distance in units of the σ of the stimulus. When plotted in this manner, the results at (A) 2 cycles/deg. could be interpreted as demonstrating spatial scaling. However the results at (B) 8 cycles/deg, were less convincing of spatial scaling. Error bars are standard error of the mean.

362 enced by the greater overlap of the test stimulus and 363 flankers when the patches were larger (Fig. 2A). This 364 possibly confounding effect is greatest at the smaller 365 test-flanker distances; at a test-flanker distance of 2λ , 366 when $\sigma = 0.5\lambda$ there was no overlap of the test and 367 flanker, while for $\sigma = 1.5\lambda$ there was substantial overlap. 368 This overlap makes the task into a contrast increment detection task rather than the easier contrast detection 369 370 task (Legge, 1979). Substantial inhibition has been re-371 ported when there was no displacement of the flanker 372 (i.e. flanker and test stimulus were co-located) (Polat & 373 Sagi, 1993, 1994a; Williams & Hess, 1998).

374 **5. Experiment 3: the effect of flanker contrast**

375 As noted in Experiment 1, subject BW had little fa-376 cilitation at the two higher spatial frequencies, 8 and 377 13.3 cycles/deg., compared to the other three subjects 378 (Fig. 1). Control experiments ruled out uncorrected a-379 metropia and poor fixation as possible explanations. A 380 third possible explanation was that subject BW might 381 not have detected the higher spatial frequency flankers. 382 Subject BW had higher central and peripheral contrast 383 thresholds than subjects AN and RW, and though 40% 384 flankers would have been visible for all conditions, some 385 flankers were very close to her contrast threshold.

386 As flanker contrast appeared to be a factor in these 387 lateral interactions, we conducted a third experiment in 388 which the flanker contrast was systematically varied from above to below peripheral detection threshold. 389 390 This was of interest also because Polat and Sagi (1993, 391 1994a) increased flanker contrast with increasing flanker 392 eccentricity, after stating that the lateral interactions 393 were independent of flanker contrasts between 20% and 394 80% (Polat & Sagi, 1993). We were able to confirm that 395 independence for 20% and 40% contrast flankers as 396 described below. Finally, the effect of flanker contrast is 397 of interest as increment threshold experiments have

demonstrated inhibition at high mask contrasts that reduced to become facilitation at low contrasts (near mask threshold) (Tolhurst & Barfield, 1978; Legge, 1979). 401

Lateral interactions were measured using the same 402 403 paradigm as employed in the previous experiments. Flanker contrast was varied from 2.5% to 40% for 4 and 404 405 8 cycles/deg. (viewing distances of 200 and 400 cm respectively) and from 1.25% to 20% for 2 cycles/deg. 406 (viewing distance of 100 cm). Test and flanker Gabor 407 patches were scaled so that $\sigma = \lambda$, and test-flanker dis-408 tance was fixed at 3λ , a distance at which there were 409 substantial lateral interactions (Experiment 1). So that 410 any effects of flanker contrast could be evaluated in 411 terms of the threshold for the flanker, detection 412 thresholds were determined for Gabor patches presented 413 in the two peripheral locations of the flankers $(\pm 3\lambda)$ and 414 at fixation. A temporal two-alternative forced choice 415 procedure randomly interleaved the threshold determi-416 nations for the three tested locations. The same staircase 417 procedure as used for the lateral interaction measure-418 419 ments was employed. A small, low-contrast (31%) fixation target was presented until 300 ms before 420 421 presentation of the stimulus. This task required a fixation target, as there were no flankers in each presenta-422 423 tion to guide fixation.

424 Fig. 4A and B show that, within the variability of these measurements, facilitation was not different for 425 flankers of 20% and 40% contrast, as stated by Polat and 426 Sagi (1993). As flanker contrast was reduced further, 427 428 facilitation decreased eventually to zero. Conversely, previous increment-threshold experiments have reported 429 that facilitation increased as the mask contrast was re-430 duced to threshold (Legge, 1979; Tolhurst & Barfield, 431 432 1978). The lower panel (Fig. 4C and D) shows the frequency-of-seeing data and fitted cumulative Gaussian 433 functions for the flanker locations. The vertical dashed 434 lines in the upper panel (Fig. 4A and B) represent the 435 thresholds as determined from the staircase (i.e. 83% 436 R.L. Woods et al. | Vision Research xxx (2002) xxx-xxx



Fig. 4. Investigated the effect of flanker contrast. Test-flanker distance was fixed at 3λ and flanker contrast varied between 1.25% and 40%. Results are shown for (A) subject AN and (B) subject RW. Facilitation was greatest at the highest flanker contrasts (with no difference between 20% and 40% contrast), gradually decreasing to zero as flanker contrast reduced. Contrast thresholds (defined as 83% correct) of the flankers are shown as the vertical lines (same style as for the data). The frequency-of-seeing curves for the flankers are shown for (C) subject AN and (D) subject RW. In many cases sub-threshold flankers produced some facilitation. Error bars are standard error of the mean.

437 correct: (García-Pérez, 1998)). While there are small 438 differences between the two subjects and between spatial frequencies, it appears that there was some facilitation 439 440 even with sub-threshold flankers. In most cases, as 441 flanker contrast reduced the facilitation had begun to 442 decrease before flanker threshold was reached. This 443 seems to offer an explanation for the very limited facil-444 itation found for subject BW at higher spatial frequen-445 cies (Fig. 1D). Overall, subject BW had higher contrast 446 thresholds than the other three subjects in Experiment 1, 447 and we suspect that subject BW had similarly higher 448 peripheral detection thresholds for all conditions in 449 Experiment 1. In which case, as the flanker contrast was 450 the same for all spatial frequencies in Experiment 1, the 451 higher spatial frequency flankers were close to her pe-452 ripheral contrast thresholds, thereby reducing the facil-453 itation.

The reduction in lateral interactions with decreasing 454 455 flanker contrast is not a complete explanation of the 456 failure to find lateral interactions for subject BW similar 457 to those found for the other three subjects in Experi-458 ment1 (Fig. 1). While we know that the flanker contrast 459 approached BW's peripheral detection threshold for the 460 condition at which the highest peripheral threshold 461 might be expected (13.3 cycles/deg. and test-flanker 462 distance 6λ), this explanation is less convincing for 8

cycles/deg. and for the shorter test-flanker distances (2 463 and 3λ). In Fig. 4 there is a trend for the facilitation for 464 the highest spatial frequency (8 cycles/deg.) to remain 465 reasonably stable until the flanker contrast was reduced 466 to the peripheral threshold, and then to quickly de-467 crease. Conversely there is a trend for the facilitation for 468 the lowest spatial frequency (2 cycles/deg.) to begin to 469 decrease gradually with reducing flanker contrast, even 470 when the flanker was well above the peripheral thresh-471 old. On this basis, we might expect more facilitation for 472 subject BW at smaller test-flanker distances at 8 and 473 13.3 cycles/deg. than was found. 474

6. Discussion

476 Spatial scaling of these lateral interactions as implied by Polat and Sagi (1993) and equivalent performance in 477 peripheral vision (Polat & Sagi, 1994b) would imply a 478 general principle of operation within the visual system 479 480 that could be applied across all scales and across the visual field. Such generality is very appealing. We dis-481 cuss our results in terms of this possible general principle 482 and associated visual models and in terms of the vision 483 of people with central vision impairment. 484

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485 Clearly the lateral interactions did not spatially scale 486 at all test-flanker distances (Figs. 1 and 2). The size of 487 both the carrier (λ) and the envelope (σ) of the Gabor 488 patch stimuli had an impact on the measured lateral 489 interactions. However, neither could these lateral inter-490 actions be characterised only by the separation in terms 491 of visual angle. At larger test-flanker distances (>3 λ) the 492 lateral interactions appear to be independent of spatial 493 frequency (λ^{-1}) . In other words, there may be spatial scaling at these longer test-flanker distances. The effects 494 495 of size become apparent when the flankers are close to 496 the test patch. The impact of proximity was greatest for 497 the lowest spatial frequency (2 cycles/deg.). This suggests 498 that this experimental paradigm may involve (at least) 499 two visual mechanisms: one a central (largely) inhibitory 500 zone and the second a more extensive zone that, given 501 the right relationships between the central target and the 502 flanker (often) produces facilitation. Our results suggest 503 that the spatial profiles of the visual mechanisms re-504 sponsible for these lateral interactions vary with the 505 spatial frequency tuning of the mechanism. In general, it 506 appears the central inhibitory zone is larger, relative to 507 the wavelength to which it is tuned, for the lower spatial 508 frequency mechanisms than the higher spatial frequency 509 mechanisms. This relationship is opposite to that re-510 ported by Wilson et al. (1983), who measured increment 511 thresholds, and found that lower spatial frequency sen-512 sitive mechanisms had larger bandwidths than higher 513 spatial frequency sensitive mechanisms, since this im-514 plies that the spatial extent of spatial frequency filters 515 decreases (relative to λ) with spatial frequency. This 516 difference supports a conclusion that the facilitation is 517 not simply summation within a large filter.

518 Unlike the stimuli of Polat and Sagi (1993, 1994a) our 519 flankers were not additive. Rather our stimuli were re-520 stricted to $4\sigma \times 4\sigma$ squares. Consequently when the test-521 flanker distances were short, the contrast of the flankers 522 that was co-local with the test patch, was less than that 523 when presented by Polat and Sagi (by as much as half). 524 Given that contrast thresholds increase with spatial 525 frequency over the range tested, and the flanker contrast 526 was fixed, the contrast of the co-local elements of the 527 flankers was closer to contrast threshold of the lower 528 spatial frequency test patches. Over much of the range 529 of mask contrasts, inhibition increases (facilitation de-530 creases) as the contrast of the mask approaches the 531 contrast threshold of the test patch (Legge, 1979). This 532 may be, at least in part, an explanation of the spatial 533 frequency dependent differences in contrast threshold at 534 short test-flanker distances. Given that the contrast of 535 Polat and Sagi's flankers that were co-local with the test 536 patch at short test-flanker distances were higher than 537 ours were, one might expect a greater spatial-frequency 538 dependent effect than in our data, but this is not ap-539 parent in their data.

One possible interpretation of our data is that short-540 541 range lateral interactions are a function of spatial frequency while long-range lateral interactions are inde-542 543 pendent of spatial frequency. This differs from the 544 results of Yu and Levi (1997c) who measured increment thresholds. Yu and Levi's stimulus configuration in-545 cluded a mask co-centric with (superimposed on) the test 546 547 stimulus. Test stimulus height was fixed while mask height was varied. They found for shorter masks that the 548 549 effect of mask height was independent of spatial frequency and inhibition was greatest at a mask height that 550 551 was a multiple of the height of the test patch (i.e. a fixed visual angle). The effect of longer masks varied with 552 553 spatial frequency, with the suggestion that higher spatial frequencies had smaller (in terms of visual angle) end 554 zones. Unfortunately their masks were not long enough 555 556 at the lower spatial frequencies to make comparisons in terms of λ . Their stimuli were wide-band (clipped) in the 557 vertical dimension. It is possible that the differences in 558 the results of these two studies are a consequence of the 559 560 complex changes in spatial content with spatial frequency, test-stimulus height and mask height in their 561 stimuli, or a consequence of the task differences (con-562 trast detection versus increment threshold). 563

As noted by Polat and Sagi (1993, 1994a), these lat-564 eral interactions extend well beyond the range of the 565 classical receptive field. Recent studies (Das & Gilbert, 566 1995; Gilbert et al., 1996) have found that there are 567 substantial supra-threshold and sub-threshold neural 568 569 interactions in the primary visual cortex over ranges that 570 are consistent with the psychophysical evidence. Cortical pyramidal cells that extend up to 6-8 mm may be 571 involved (Gilbert et al., 1996). Using a 0.5° stimulus, 572 Das and Gilbert (1995) reported a 0.75 mm ($\sim 0.5^{\circ}$) 573 central zone in which spiking activity was found and a 4 574 mm ($\sim 4^{\circ}$) surrounding inhibitory zone in which there 575 576 was sub-threshold activity. Facilitation due to flanking line stimuli has been reported in parallel psychophysical 577 and electrophysiological studies (Kapadia et al., 1995). 578 Potential models for these lateral interactions include a 579 simple transducer (Solomon et al., 1999), end-stopping 580 (Yu & Levi, 1997c), a two-stage, contrast-gain model 581 582 (Zenger & Sagi, 1996) and two-stage rectified filters (Polat, 1999; Solomon & Morgan, 2000). Of these 583 models, only Yu and Levi (1997c) measured or tested 584 their model with a range of spatial frequencies and only 585 that model incorporates any specific elements that could 586 587 account for a lack of spatial scaling. It is possible that these models could be adapted by the addition of a small 588 number of additional parameters that incorporate 589 changes in spatial frequency and patch bandwidth. 590 591 More problematic is the apparent discrepancy noted above between our results and those of Yu and Levi 592 593 (1997c) who examined a wide range of spatial frequencies. They interpreted their results as spatial filters with 594 end-stopping that varied with spatial frequency, and 595

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596 length tuning determined by target length. The central 597 inhibitory zone apparent in our data (Figs. 1 and 2) 598 varied as a function of both the spatial frequency (λ^{-1}) 599 and the envelope (σ) of the Gabor test patch. We did not examine a sufficiently wide range of visual angles to 600 601 evaluate our data in terms of length tuning. Yu and 602 Levi's results can be interpreted as limited confirmatory 603 evidence for a lack of spatial scaling of these lateral 604 interactions, given the relationship between spatial frequency and strength and extent of end-stopping. As 605 606 noted above, differences between these two studies may 607 be related to stimulus characteristics (e.g. our test pat-608 ches were not clipped in the vertical dimension) and task 609 (i.e. our flankers did not overlap the target in the same 610 way). End-stopping may be an adequate explanation for 611 short-range lateral interactions (e.g. test-mask distance 612 $\leq 3\lambda$) but seems unable to explain the long-range lateral 613 interactions. It seems that new or revised models are 614 required to account for our results. Such models may 615 have to incorporate two different and competing mech-616 anisms and should consider the dynamic nature of re-617 ceptive fields found electrophysiologically (Kapadia, 618 Westheimer, & Gilbert, 1999).

619 The different lateral interactions found with changes 620 in patch size (σ) (Figs. 2 and 3) suggest that bandwidth 621 of the stimuli is an important factor for the visual mechanisms responsible for these lateral interactions. 622 623 However, the simple effect of overlap of the stimuli 624 cannot be disregarded. At the shorter test-flanker dis-625 tances the narrow-band, larger stimuli (i.e. $\sigma > \lambda$) have 626 more overlap of test stimulus and flankers. As test and 627 flanker begin to overlap, the task changes from contrast 628 detection to increment threshold. When, as in the Polat 629 and Sagi paradigm, the flanker is gradually moved rel-630 ative to the test stimulus, it becomes difficult to make the 631 classical distinction between contrast detection and in-632 crement threshold. For $\sigma = \lambda$ patches, as shown in Fig. 633 1A, overlap begins at a test-flanker distance of about 3λ . 634 The difference in overlap that was confounded with the 635 change in size of the Gabor patches in Experiment 2 636 could have had a significant contribution to this re-637 ported effect of stimulus size. To examine this possibil-638 ity, we shall need to systematically and independently 639 vary overlap and bandwidth of the flankers (Woods et 640 al., in preparation).

641 The decrease in facilitation with reduction in flanker 642 contrast (Fig. 4) is similar to that reported by Solomon 643 and Morgan (2000) for 13 cycles/deg. (same-phase) 644 Gabor patches measured using the Polat and Sagi par-645 adigm. They did not report the peripheral detection 646 thresholds of the flankers. While, for a 3λ test-flanker 647 distance, Polat (1999) reported no difference in facilita-648 tion between 10% and 75% flanker contrasts, his data is 649 unconvincing due to the lack of error bars or statistical 650 analysis, and the apparently lower facilitation for 40%651 than 20% or 75% flanker contrasts. Wehrhahn and

Dresp (1998) found that small line stimuli induced by a 652 larger co-linear flanking line with the same polarity 653 showed increasing facilitation as flanker contrast re-654 655 duced. If we consider our (same phase) stimuli to be equivalent to those same polarity stimuli, this is the 656 opposite of our result. The difference in results may be 657 due to the nature of their stimuli-test stimulus and 658 659 flanker were dissimilar in size and were wide-band. Interestingly, they reported that when stimulus and flanker 660 had opposite polarity, facilitation was greatest for a high 661 contrast flanker and decreased as flanker contrast re-662 duced (Wehrhahn & Dresp, 1998). This stimulus con-663 figuration seems analogous to opposite-phase Gabor 664 patch stimuli. For these opposite phase Gabor patches, 665 Solomon et al. (1999) reported inhibition that decreased 666 as flanker contrast reduced when test-flanker distance 667 was short (2λ) but no lateral interactions with larger 668 test-flanker distances (3 and 4λ ; 13.3 cycles/deg.). Simi-669 larly Williams and Hess (1998) found no effect of op-670 posite phase, 3λ flankers (4.2 cycles/deg.). Again, with 671 672 Gabor patches, the result does not appear consistent with the result found using line stimuli-well spatially 673 localised, but less well frequency-defined (wide-band) 674 (Wehrhahn & Dresp, 1998). It is possible that these 675 differences occur due to the different lateral interaction 676 fields of detectors with different spatial frequency tuning 677 as found in Experiments 1 and 2. Line stimuli would be 678 expected to stimulate a wider range of spatial frequency 679 tuned detectors than Gabor patches. Stimulus contrasts 680 are a factor also. Kapadia et al. (1999) reported that the 681 682 neuronal receptive field varied with the contrast of line stimuli, being about four times larger for low contrast 683 (10–20%) than high contrast (50–70%) stimuli. Another 684 explanation may be found in the results of Yu and Levi 685 (1997c), who used a paradigm designed to test their end-686 stopping model of lateral interactions. Yu and Levi re-687 688 ported, for a single spatial frequency (8 cycles/deg.), that the effect of phase depended on the distance of the 689 flanker from the text patch. When the flankers were 690 close, same-phase flankers produced inhibition while 691 opposite-phase flankers caused facilitation. However, 692 when the flankers were more distant, both same- and 693 694 opposite-phase flankers produced facilitation that decreased with reducing flanker contrast. This was ex-695 plained by their end-stopping model, wherein closer 696 flankers are within the "outer summation zone", while 697 more distant flankers are within the "end zone". These 698 discrepancies between results for stimuli with different 699 spatial characteristics need to be resolved for a better 700 understanding of lateral interactions in human vision. 701

It is not clear yet whether these lateral interactions 702 occur in the retinal periphery. Williams and Hess (1998) 703 found no facilitation for a limited range of conditions, 704 while in a footnote in one paper, Polat and Sagi (1994b) 705 reported facilitation similar to that found at the fovea 706 (but provided no data). Xing and Heeger (2000), using 707

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708 an annular flanker, reported differences in contrast 709 matching between fovea and periphery. In particular, in 710 peripheral vision the flanker reduced perceived contrast. 711 Since the majority of people with low vision have ma-712 cular degeneration, and it is these people who would 713 most benefit from image enhancement, further investi-714 gation of lateral interactions in peripheral vision is re-715 quired. Of particular interest in the quest to improve the 716 visual experience of people with macular degeneration 717 are the facilitatory interactions, as these effects may be 718 used to enhance images. In peripheral vision only lower 719 spatial frequencies are visible. Given that in foveal vi-720 sion these lateral interaction effects do not spatially scale 721 and are influenced by bandwidth, consideration of these 722 aspects of stimulus configuration is required when in-723 vestigating peripheral vision. In summary, while there 724 may be some general principles that describe these lat-725 eral interactions, it appears that any general principle 726 has a layer of complexity not incorporated previously 727 (e.g. it needs to include changes with spatial frequency 728 and bandwidth).

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731 Appendix A. monitor non-linearity

732 Many of the lateral interaction effects investigated 733 with this paradigm by Polat and Sagi (1993, 1994a,b) 734 and others (Solomon et al., 1999; Williams & Hess, 735 1998) were conducted with 13.3 cycles/deg. targets. 736 During our early investigations we found that the 737 number of pixels used to represent the grating appeared 738 to influence the results. Polat and Sagi tested spatial 739 frequencies ranging from 13.3 to 3.33 cycles/deg. using a 740 56 Hz non-interlaced display system with 512×512 741 pixel, viewed from 180 cm ($9.6^{\circ} \times 9.6^{\circ}$). Hence, at 13.3 742 cycles/deg., there were only 4 pixels per cycle. There was 743 a statistically significant difference between the 4-pixels 744 per cycle condition (viewing distance of 118 cm) and the 745 23 pixels per cycle condition (654 cm), and both facili-746 tation functions appeared slightly different to those re-747 ported by Polat and Sagi (1993, 1994a).

748 We suspected monitor non-linearity, in particular the 749 ability to represent accurately gratings of high contrast 750 and high frequency. Relatively high contrasts were 751 necessary in our experiment (e.g. at 13.3 cycles/deg. 752 contrast thresholds were about 10% and the flankers 753 were 40%). To obtain gratings of these nominal con-754 trasts, the required contrast of our display had to be 755 twice these amounts, since each grating was only visible 756 in every second frame (i.e. the perceived contrast was 757 half the contrast of the grating displayed in each frame).

As noted by Klein et al. (1996) and Pelli (1997) when a 758 759 raster scan requires large changes in luminance over a small distance, as occurs when writing across a high-760 contrast, high-frequency grating, the monitor may fail 761 to represent the change correctly. Presumably the elec-762 tron gun is unable to change its signal sufficiently 763 quickly (causing low pass filtering: Pelli, 1997). Conse-764 quently, the luminance of each pixel is influenced by the 765 luminance of nearby pixels. An additional calibration 766 procedure and look-up table can be used to reduce this 767 problem (Klein et al., 1996). A simpler alternative is to 768 769 have smaller changes in luminance for subsequent pixels



Fig. 5. (A) An illustration of the luminance required at each position (pixel) for configurations of 3.7, 4, and 22.6 pixels per cycle (λ) with a Gabor size $\sigma = \lambda$. The thick gray lines represent the required luminances and the thin black lines represent the nominal luminances at each location. (B) An illustration of the luminance required when the raster line is along or across a 4 pixels per cycle grating (thin black lines) or a 22.6 pixels per cycle grating (thick grey lines) with Gabor size $\sigma = \lambda$. These figures are schematic, as the actual luminance produced by each configuration is influenced by the luminance profile of each pixel and any monitor non-linearity. Error bars are standard error of the mean.

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to minimise this non-linearity. This could be achieved by(1) using more pixels to represent the grating (Fig. 5A);

(1) using more pixels to represent the grating (Fig. 5A),
or (2) having the raster write along, rather than across
the grating (Fig. 5B).

774 The raster on our monitor, as is common, wrote 775 horizontal lines starting from the upper left corner of the 776 screen. Therefore to have the raster write along the grating the gratings need to be horizontal. Alternatively, 777 778 by placing the monitor on its side vertical gratings with 779 the raster writing along the grating could be produced. 780 To investigate whether our measurements, and possibly those of Polat and Sagi (1993, 1994a), were affected by 781 782 monitor non-linearity, we measured the lateral interac-783 tions with both vertical and horizontal gratings (13.3 784 cycles/deg. Gabors with 4 pixels per cycle), both with the 785 monitor in its normal orientation and when it was on its 786 side. The facilitation produced by vertical and hori-787 zontal gratings was not different (repeated measures 788 ANOVA, p = 0.49). As shown in Fig. 6A, on average 789 the facilitation was greater when the raster wrote along 790 the grating than when the raster wrote across the grating (p = 0.006) and the shape of the facilitation functions 791 was different (interaction, p = 0.0009). In addition, the 792 measured contrast thresholds for the standard condition 793 794 (no flanking patches) were significantly lower for the raster-along conditions than the raster-across conditions 795 796 $(-0.74 \text{ versus } -0.58 \log \text{ units respectively}; p < 0.0001).$ Assuming that vision did not change, this means that 797 higher luminance differences between pixels were re-798 quired for the raster-across conditions to display the 799 same contrast as the raster-along conditions. 800

Having completed this investigation, we discovered 801 802 that when the monitor was placed on its side, there were misalignments of the three colour guns (due to the 803 earth's magnetic field) had not been removed by de-804 gaussing. The misalignments caused incorrect irradia-805 tion of the three phosphors sufficient to cause changes in 806 colour naming (e.g. dark blue became light green). These 807 effects were not obvious when viewing a screen com-808 prising only shades of grey. As we were concerned that 809 these misalignments of the pixels could make the look-810 up table inaccurate, we repeated this examination of 811



Fig. 6. (A) An evaluation of monitor non-linearity using 4 pixel per cycle, 13 cycles/deg. Gabor patches with $\sigma = \lambda$. Two of the conditions were created by placing the monitor on its side. All conditions produced facilitation (negative relative contrast threshold). There was no difference between the facilitation produced by the vertical and the horizontal gratings. However, the facilitation differed between the raster-across and raster-along conditions. (B) As for A, except the monitor was not rotated, instead the subject lay on his side to create two of the conditions. Again, the non-linearity of the monitor was apparent in the difference between the raster-across conditions. (C) An evaluation of monitor non-linearity using vertical, 13 cycles/deg. gratings. There was no difference between the facilitation produced by the raster-across and raster-along conditions for the 23 pixel per cycle gratings. (D) As for B, except with 23 pixel per cycle, 2 cycles/deg. patches. With this better spatial resolution there was no apparent monitor non-linearity there being no significant differences between the four conditions. Thus, with sufficient spatial resolution the monitor non-linearity could be avoided. Error bars are standard error of the mean.

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812 monitor non-linearity by having the subjects lie on their 813 side while viewing in lieu of rotating the monitor. We 814 found almost exactly the same effects (Fig. 6B). There 815 was a significant effect of raster orientation (repeated 816 measures ANOVA, p = 0.04) but no effect of grating 817 orientation (p = 0.51) or interaction between raster and 818 grating orientations (p = 0.27). García-Pérez and Peli 819 (2001) used a similar approach in a more systematic 820 examination of this and related artefacts associated with 821 monitors.

822 To investigate the impact of having a larger number 823 of pixels, we compared the 13.3 cycles/deg. gratings of 4 824 pixel per cycle and 23 pixel per cycle vertical gratings, 825 both with the raster-along and raster-across the grating. 826 All four conditions produced facilitation (Fig. 6C). For 827 the 23 pixel per cycle gratings, raster alignment had no 828 effect (repeated measures ANOVA, p = 0.61). This 829 suggests that when there was a sufficiently large number 830 of pixels, the monitor could produce equivalent gratings 831 when the raster was writing along or across the grating. 832 For the raster-along condition, the shape of the facili-833 tation function differed slightly between the two reso-834 lution conditions (interaction, p = 0.07). To confirm 835 this, we repeated the subject-rotating experiment when 836 using 23 pixel per cycle gratings that were 2 cycles/deg. when viewed at 100 cm. For this arrangement we found 837 838 the predicted lack of a significant difference between the 839 raster-along and raster-across conditions (Fig. 6D). 840 Hence, to avoid the non-linearity found when only small 841 numbers of pixels were used to display the grating, we 842 used a fixed target size and varied the viewing distance.

843 While the number of pixels per cycle was a problem 844 with our system, we do not know whether such prob-845 lems are found with other systems. However, the mon-846 itor that we used was reasonably new and considered of 847 good quality at the time of its purchase. Our experience 848 suggests that care should be taken when evaluating the 849 effects of stimuli created with monitors.

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