Spontaneous and training-induced cortical plasticity in MD patients: Hints from lateral masking

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Macular degeneration (MD) affects central vision and represents the leading cause of visual diseases in elderly population worldwide. As a consequence of central vision loss, MD patients develop a preferred retinal locus (PRL), an eccentric fixation point that replaces the fovea. Here, our aim was to determine whether and to what extent spontaneous plasticity takes place in the cortical regions formerly responding to central vision and whether a visual training based on perceptual learning (PL) can boost this plasticity within the PRL area. Spontaneous and PL-induced cortical plasticity were characterized by using lateral masking, a contrast sensitivity modulation induced by collinear flankers. This configuration is known to be sensitive to neural plasticity and underlies several rehabilitation trainings. Results in a group of 4 MD patients showed that collinear facilitation was similar to what observed in age- and eccentricity-matched controls. However, MD patients exhibited significantly reduced collinear inhibition, a sign of neural plasticity, consistent with the hypothesis of partial cortical reorganization. Three AMD patients from the same group showed a further reduction of inhibition after training, but not controls. This result suggests that PL might further boost neural plasticity, opening promising perspectives for the development of rehabilitation protocols for MD patients.
that this effect exists outside foveal vision, with the facilitatory target-to-flankers separation (and consequently, the PF size) increasing with eccentricity, most likely following the cortical magnification factor (see Fig. 1).

This distinction between foveal and peripheral collinear facilitation is of critical importance in MD patients. Indeed, evidence for facilitation in the PRL at shorter separations than in healthy peripheral vision would be consistent with the idea of a spontaneous cortical reorganization after retinal lesion, similar to what is reported in crowding4. While they share similar features, such as the increase in magnitude with eccentricity, and resemblance between target and flanking elements17,22, there is no consensus in the literature regarding the relationship between crowding and lateral masking23. Consequently, the evidence for spontaneous cortical reorganization in reshaping the crowding zone does not necessarily mean that the neural substrates underlying cortical inhibition are equally affected.

Horizontal connections between units sharing similar orientation and spatial frequency tunings in early visual cortex24–26 are known to be sensitive to PL-induced plasticity as suggested by lateral masking paradigm 27. Enhancing these first stages improves in turn higher level visual processing that can rely on more efficient inputs after training15,21.

The question is still open whether collinear facilitation exists in MD patients’ PRL and whether it can be modulated through PL, as reported for other clinical populations14. Indeed, a previous study did not find collinear facilitation in MD patients28, possibly because given the tested target-to-flankers distance (4λ), the flankers remained within the inhibitory region of the PF. In the present paper, we aimed at addressing these questions in a small sample of AMD patients (and age-matched controls) using a lateral masking configuration as a baseline measure for spontaneous cortical reorganization (Experiment 1) and as a training configuration for guided cortical reorganization (Experiment 2). Previous PL studies based on collinear facilitation showed a reduction of inhibition after training 21,27, an effect attributed to neural plasticity in early visual areas24–26. Consequently, reduced inhibition at baseline would be consistent with the hypothesis of spontaneous cortical reorganization similar, to what was observed in previous studies after training.

Method
In Experiment 1, we measured collinear facilitation (and inhibition) over a range of target-to-flankers distances in the PRL of 5 MD patients and 5 age- and eccentricity-matched controls. In Experiment 2 we trained a subgroup of participants from Experiment 1 (4 MD patients and 3 controls) to test whether facilitation and/or inhibition can be modulated through practice as observed in previous studies21,27. Because training based on PL are very long, a small sample size is not uncommon in PL studies. Nonetheless, a number of studies showed that reliable and robust results could still be obtained by comparing performances before and after training29–34.

Stimuli were three aligned Gabor patches which were displayed either in a collinear (flankers having the same local orientation as the target) or in an orthogonal (flankers having an orthogonal orientation with respect to the target i.e., a 90 deg offset) configuration (see20). The stimuli were aligned vertically for all participants except for MD3 and her matched control, for whom they were oriented horizontally, because of the shape of MD3’s scotoma. Each Gabor patch had a spatial frequency of 1 cpd. In a previous study, we showed that this frequency maximizes peripheral facilitation magnitude35. The target-to-flankers separations were 3λ, 4λ, 6λ and 8λ. Stimuli were displayed on a 17” Dell M770 CRT monitor with a resolution of 1024 × 768 pixels, a refresh rate of 60 Hz and mean
### Table 1. Summary of characteristics of participants in the Experiments. PRL coordinates with SD for the three consecutive measures and scotoma size correspond to the tested eye. Clinical annotation: OS = left eye, OD = right eye.

| Participant | Sex | Age at the onset (years) | Time since the onset (years) | Year of birth | Diagnosis | Scotoma size | PRL coordinates (x axis) | PRL coordinates (y axis) | Tested eye |
|-------------|-----|--------------------------|-------------------------------|--------------|-----------|-------------|------------------------|------------------------|------------|
| MD1         | M   | 61                       | 7                             | 1947         | AMD       | 18.4°±1.5°   | 8.15°±0.1°               | 0.1°±0.2°              | OS         |
| MD2         | F   | 59                       | 7                             | 1949         | AMD       | 17°±2.3°     | 5.43°±1°                 | 1.14°±0.7°             | OS         |
| MD3         | F   | 82                       | 4                             | 1929         | AMD       | 25.3°±20.2°  | 2.6°±0.1°               | 9.12°±0.7°             | OS         |
| MD4         | F   | 53                       | 7                             | 1958         | Atrophic MD | 12.1°±7.4°  | 7.95°±0.1°              | 1.6°±0.3°              | OD         |
| MD5         | F   | 77                       | 3                             | 1935         | AMD       | 11.7°±13.3° | 9.25°±1.6°             | 4.8°±0.65°             | OD         |
| C1          | F   | —                        | —                             | 1949         | —         | —           | —                      | —                      | OS         |
| C2          | F   | —                        | —                             | 1951         | —         | —           | —                      | —                      | OS         |
| C3          | F   | —                        | —                             | 1930         | —         | —           | —                      | —                      | OD         |
| C4          | M   | —                        | —                             | 1955         | —         | —           | —                      | —                      | OS         |
| C5          | M   | —                        | —                             | 1945         | —         | —           | —                      | —                      | OD         |

Luminance of 47.6 cd/m². Stimuli were generated with Matlab Psychtoolbox36,37. Each pixel subtended 2.14 arcmin. A digital-to-analogue converter (Bits®, Cambridge Research Systems, Cambridge UK) was used to increase the dynamic contrast range (13-bit luminance resolution). A 12-bit gamma-corrected lookup table (LUT) was used to linearize the monitor. Experiments were carried on at the Centre de la Retine, Hôpital Pierre-Paul Riquet, Purpan Hospital, Toulouse (France).

**Participants.** Patients and controls’ information is detailed in Table 1. Importantly, in order to keep our sample of patients homogeneous, we only included in the study patients with a single and well-defined PRL. MD patients were instructed to fixate with their PRL the center of the screen where a fixation point was always present. They were asked to adapt their head position in order to fixate as naturally as possible. All participants were volunteers and gave written informed consent prior to their inclusion in the study. They received an allowance for their participation. This study was conducted in accordance with the Declaration of Helsinki (1964). The experimental protocol was approved by a national ethical committee before the beginning of the study (CPP, Comité de Protection des Personnes, protocole 13018–14/04/2014). Four patients (MD1, MD2, MD3 and MD4) and 3 controls (C1, C2 and C4) took part in Experiment 2. MD5, C3 and C5 were not able to participate in the training study for logistical reasons.

**Inclusion criteria.** Absolute central binocular scotoma (as measured by 30° and 12° central visual fields analysed with Octopus® 300 perimeter; Haag-Streit, Köniz, Switzerland), age >60, residual vision in both eyes <2/10, presence of a single PRL (as detected with OCT measurement, see the ‘PRL location’ section). Exclusion criteria: Concomitant presence of other visual diseases (cataract, retinal detachment, glaucoma), cognitive impairment (Mini-Mental State Examination (MMSE) score <25), monocular MD.

**PRL localization.** For each MD patient, we determined the PRL position using Spectral-Domain Optical Coherence Tomography (Spectralis OCT, Heidelberg Engineering, Heidelberg, Germany). First, high resolution scans of the retinal fundus, centered on the atrophic macular area, were acquired via OCT in order to localize the fovea (Fig. 2a). Three additional landmarks (i.e., crossing blood vessels) were identified on the infrared image of the macula and their coordinates were measured with respect to the fovea. In a successive acquisition, patients were asked to fixate with their PRL the central fixation point presented by the OCT system while high speed X-line OCT B-scans were acquired (Fig. 2b). To avoid the inclusion of patients with more than one PRL, this operation was repeated at least three times, and between each acquisition, the OCT system was defocalized while the patient was asked to fixate elsewhere. The same acquisition protocol was systematically conducted at each examination. As noted previously, patients with more than one PRL were excluded from the study. Afterwards, the distance between the PRL and the fovea was derived by comparing its position with respect to the three landmarks (Table 1 and Fig. 3). The small SD range of the three separate measures of PRL coordinates (Table 1) is a further indication that the patients were using a single PRL for fixation tasks.

**Procedure.** In Experiment 1, we tested eight conditions per participant, one for each target-to-flankers distance (3λ, 4λ, 6λ, 8λ) and orientation (collinear and orthogonal). MD patients performed the task monocularly with their dominant eye. They fixated with their PRL while the configuration was presented at the centre of the screen. A fixation point was always present to indicate the center of the configuration. Patients were instructed to keep their fixation on the central dot with their PRL and to use the three discs to adjust their gaze on the central dot (Fig. 4). A similar approach was already taken in previous studies that used visual aids to stabilize fixation in this clinical population38. Each control participant was assigned to a patient of similar age and trained at the eccentricity corresponding to the PRL position of his/her paired MD patient. Control participants had to fixate a central dot while the position of the testing configuration was randomized along the x axis. Position along the y-axis was fixed. Participants had to perform a contrast detection task with a temporal-2AFC (two alternative forced choice) procedure, with one interval containing both target and flankers and the other interval containing
only the flankers (see Fig. 4). Each interval lasted 133 ms and the ISI was 500 ms. Participants had to report in which interval they detected the target. Target contrast varied according to a 3down/1up staircase, in which three consecutive correct responses reduced the target contrast of 0.1 log units and each wrong response increased the contrast of the same amount. The staircase terminated after 120 trials or 14 reversals. Contrast thresholds (Michelson contrast), corresponding to 79% of correct detection, were estimated from the algebraic mean of the last 6 reversals. Acoustic signals (50 ms tone of 1000 Hz) were provided to signal the beginning of each interval and as a feedback for uncorrect responses (50 ms tone of 500 Hz). Participants sat in a dark room at a distance of 57 cm from the screen. Testings were performed monocularly. The testing configuration was always vertical (see Fig. 4), except for MD3 and her corresponding control, for whom the global and local orientation of the triplet was horizontal. This was due to the position of the PRL and the shape of the scotoma of MD3 that would have made it difficult to allow for the three Gabor patches to be visible all the time in a horizontal configuration. All 8 conditions (4 target-to-flankers separations x2 orientations) were performed within a single day. In Experiment 2, participants were trained for 12 sessions (4 blocks per day, 3 sessions per week) on the same configuration as in Experiment 1. Target to flankers distances were varied between blocks in each daily session, starting from the largest separation (8λ, 6λ, 4λ, 3λ). Spatial frequency of the stimuli was 1cpd. Participants were trained only with the collinear configuration and tested on the orthogonal configuration before and after the training. Before each experiment, we ran a practice block of 15 trials to ensure that the patients were able to see the two flankers while fixating in the center of the screen with their PRL. All participants (patients and controls) reported that both the target and the flankers were clearly visible under these experimental conditions.
Target-to-flankers distances (3λ, 4λ, 6λ, 8λ) showed a significant main effect of Group (F_{1,8} = 10.604, p < 0.011) and of Target-to-flankers distances (F_{1,28} = 9.77, p < 0.001). As reported earlier, we could not test for the interaction effects (see Statistical analyses section). Post-hoc analysis, in the form of differences of differences, showed that the difference between collinear and orthogonal thresholds was significantly different for 3λ with respect to 8λ (p < 0.001, iso-ortho|3λ vs iso-ortho|8λ) and for 3λ vs 6λ (p < 0.001, iso-ortho|3λ vs iso-ortho|6λ).

Since the effect of the group was significant, we conducted separated analysis on MD participants and controls. In patients, a linear model ART with Orientation (Collinear vs Orthogonal) and Target-to-flankers distances (3λ, 4λ, 6λ, 8λ) as factors, showed a significant effect of Orientation (F_{1,28} = 3.11, p = 0.042). Post-hoc analysis showed a significant difference between 3λ and 6λ (p = 0.025).

In controls, a linear model ART with Orientation (Collinear vs Orthogonal) and Target-to-flankers distances (3λ, 4λ, 6λ, 8λ) as factors showed a significant main effect of Orientation (F_{1,28} = 6.06, p = 0.021) and Target-to-flankers distances (F_{1,28} = 14.642, p < 0.001).

Post-hoc analysis showed a significant difference between thresholds at 3λ and 6λ (p < 0.001) and between 3λ and 8λ (p < 0.001).

Threshold elevation. Following previous studies^{17,21}, collinear inhibition and facilitation were computed as ratios of contrast thresholds (CT), using the following formula, separately for each target-to-flankers separation:
We used the orthogonal condition rather than the Gabor alone as a baseline measure of facilitation to distinguish actual orientation-specific increase in contrast sensitivity from reduction of spatial uncertainty. This approach is standard for peripheral collinear facilitation measurements\(^{19,21,35}\).

For Experiment 1, we conducted a mixed model Aligned Rank Test on threshold elevations with between factor Group (MDs vs Controls) and within factor Target-to-flankers distances (3\(\lambda\), 4\(\lambda\), 6\(\lambda\), 8\(\lambda\)). The interaction was calculated using the r package npInt for Nonparametric Interaction Tests for Factorial Designs with Repeated Measures (Feys, 2015). Results showed a main effect of Target-to-flankers distance (\(F_{3,24} = 17.808, p < 0.001\)) and the interaction between factors (\(F_{3,24} = 5.39, p = 0.00554\)).

Post-hoc analysis (ART pairwise comparisons, holm adjusted) showed that overall threshold elevation at 3\(\lambda\) was significantly higher than all the other separations (\(p = 0.0391, p < 0.001\) and \(p < 0.001\) for 4\(\lambda\), 6\(\lambda\) and 8\(\lambda\), respectively). Additionally, we explored post-hoc tests for the interaction. Since tests of differences of combinations of levels between factors have issues in ART (Kay, 2014), it is recommendable to test for differences of differences; (i.e., for the interaction Group x Target-to-flankers distance, we might test whether the difference AMD - control is different for 3\(\lambda\) vs 4\(\lambda\), Wobbrock et al., 2011). This analysis showed that the difference

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TE = \log_{10}\left(\frac{CT_{\text{collinear}}}{CT_{\text{orthogonal}}}\right) \tag{1}
\]

Table 2. Individual contrast thresholds for Experiment 1.

|                  | Collinear 3\(\lambda\) | Collinear 4\(\lambda\) | Collinear 6\(\lambda\) | Collinear 8\(\lambda\) | Orthogonal 3\(\lambda\) | Orthogonal 4\(\lambda\) | Orthogonal 6\(\lambda\) | Orthogonal 8\(\lambda\) |
|------------------|------------------------|------------------------|------------------------|------------------------|------------------------|------------------------|------------------------|------------------------|
| AMD1             | 0.217                  | 0.170                  | 0.210                  | 0.238                  | 0.229                  | 0.270                  | 0.215                  | 0.280                  |
| AMD2             | 0.486                  | 0.316                  | 0.193                  | 0.230                  | 0.242                  | 0.257                  | 0.211                  | 0.282                  |
| AMD3             | 0.355                  | 0.204                  | 0.171                  | 0.274                  | 0.246                  | 0.216                  | 0.215                  | 0.289                  |
| AMD4             | 0.122                  | 0.177                  | 0.164                  | 0.110                  | 0.071                  | 0.121                  | 0.221                  | 0.143                  |
| AMD5             | 0.492                  | 0.229                  | 0.094                  | 0.090                  | 0.201                  | 0.241                  | 0.105                  | 0.120                  |
| C1               | 0.201                  | 0.078                  | 0.048                  | 0.056                  | 0.065                  | 0.056                  | 0.071                  | 0.101                  |
| C2               | 0.140                  | 0.028                  | 0.026                  | 0.029                  | 0.026                  | 0.037                  | 0.035                  | 0.033                  |
| C3               | 0.371                  | 0.413                  | 0.141                  | 0.104                  | 0.208                  | 0.171                  | 0.209                  | 0.129                  |
| C4               | 0.112                  | 0.070                  | 0.018                  | 0.028                  | 0.041                  | 0.029                  | 0.026                  | 0.037                  |
| C5               | 0.180                  | 0.093                  | 0.056                  | 0.060                  | 0.067                  | 0.095                  | 0.134                  | 0.124                  |

Figure 5. Box-and-whisker diagrams for collinear and orthogonal contrast thresholds (Michelson contrast) for MD (red boxes) and Control participants (green boxes) as a function of the target-to-flankers distances (\(\lambda\)). In each boxplot, the central mark is the median. The edges of the box are the 25th and 75th percentiles and the whiskers are the interquartile range (i.e., Q3–Q1). The black circles represent the mean and the vertical bars are +/- SEM.
AMC-controls was significant between 3\(\lambda\) and 8\(\lambda\) \((p = 0.014973)\) and 3\(\lambda\) and 6\(\lambda\) \((p = 0.00193)\), suggesting that patients exhibited both less inhibition \((0.213 +/− 0.156 \text{ vs } 0.464 +/− 0.171)\) and less facilitation \((-0.0836 +/− 0.0391 \text{ vs } -0.165 +/− 0.109)\) than controls at baseline.

MD patients. We estimated collinear facilitation by conducting a series of one sample t-tests (threshold elevation vs zero value) similarly to previous studies\(^{20,21}\). Results showed a significant threshold reduction for target-to-flankers separations of 6\(\lambda\) \((t_4 = 3.05, p = 0.038)\) and 8\(\lambda\) \((t_4 = 4.78, p = 0.009)\) and a significant threshold elevation for 3\(\lambda\) \((t_4 = 6.04, p = 0.004)\).

Controls. Independent one sample t-tests revealed a significant threshold reduction for target-to-flankers separations of 6\(\lambda\) \((t_4 = 4.78, p = 0.008)\) and 8\(\lambda\) \((t_4 = 3.38, p = 0.027)\) and a significant threshold elevation for 3\(\lambda\) \((t_4 = 6.04, p = 0.004)\).

Finally, comparing threshold elevation in the two groups, at 3\(\lambda\) MD patients showed less inhibition than controls (two-sample t-test, \(t_8 = 2.42, p = 0.042\)), as well as less facilitation at 6\(\lambda\) \((t_8 = 1.17, p = 0.018)\).

Training Results (Experiment 2). Contrast thresholds. For Experiment 2 (Fig. 6 and 7, Table 3), a Mixed Model ART with within factors Training (Pre vs Post), Orientation (Iso vs Ortho) and Target-to-flankers distances (3\(\lambda\) vs 4\(\lambda\) vs 6\(\lambda\) vs 8\(\lambda\)) and between factor group (MDs vs controls) showed a main effect of Group \((F_{1,5} = 16.91,\)

Figure 6. Box-and-whisker diagrams for collinear (above) and orthogonal (below) Michelson contrast thresholds for MD (red boxes) and Control participants (green boxes), before and after training as a function of the target-to-flankers distances. See Fig. 5 for the details of the legend.
Figure 7. Single participant data for the training, divided by target-to-flankers distances (3 to 8 lambda, from lighter green to darker green for controls, from lighter red to darker red for MD patients).

Table 3. Individual Michelson contrast thresholds difference after training (Experiment 2).
MD Patients. A Mixed Model ART on contrast thresholds with factors Training (Pre vs Post), Orientation (Iso vs Ortho) and Target-to-flankers distances (3λ vs 4λ vs 6λ vs 8λ) showed a main effect of Orientation (F_{1,5} = 9.07, p = 0.005) and Target-to-flankers distance (F_{1,30} = 9.43, p < 0.001).

Threshold elevation. A mixed model ART with within factors Training (Pre vs Post) and target-to-flankers distances (3λ vs 4λ vs 6λ vs 8λ) and between factor Group (MD vs Control) showed a main effect of Target-to-flankers distances (F_{3,5} = 21.65, p < 0.001), a main effect of Group (F_{1,5} = 11.82, p = 0.018). The MD group showed lower threshold elevation (0.1 +/− 0.025 vs −0.041 +/− 0.021), indicating overall greater facilitation for all the separations tested. Moreover, paired t-tests comparing pre- and post- training performance separately for each target-to-flankers separation showed that controls worsened their performance at 6λ after training (p = 0.004), while pre- post- paired t-test in MD patients approached significance for 3λ (p = 0.062). However, control group showed low contrast thresholds for the large target-flanker distances, suggesting a small room for improvement during the training. This might have reduced the differences between collinear and orthogonal thresholds and could thereby be the cause of this apparent performance deterioration.

Discussion
Cortical reorganization in patients with central vision loss, either spontaneous or training induced, is a topic of great interest in visual and clinical neuroscience. Recent results from psychophysics, neurophysiology and neuroimaging studies remain unclear. Indeed, previous studies using psychophysics approaches reported evidence for and against spontaneous cortical reorganization \({}^{4,6}\), respectively. Similarly, some neuroimaging studies supported the hypothesis of a spontaneous neural reorganization in the lesion projection zone of MD patients \({}^{49}\). This effect existed for stimuli presented outside the scotoma, around the PRL or not \({}^{44}\). However, other fMRI studies did not find evidence for spontaneous cortical reorganization in MD patients \({}^{45}\) or monkeys with retinal lesion \({}^{46}\). In Experiment 1, we showed that in MD, both collinear inhibition and facilitation emerged at target-to-flankers separations that are consistent with those measured in previous studies in normal peripheral \({}^{19,21,35}\) but not foveal \({}^{17}\) vision. With respect to control participants, MD patients had overall higher contrast thresholds, and presented both reduced collinear inhibition at short separations (3λ) and reduced facilitation at larger separations (6λ and 8λ). On the one hand, reduced inhibition is consistent with the hypothesis of a spontaneous or use-dependent cortical reorganization. Previous training studies on collinear facilitation reported reduced inhibition after training \({}^{24,27}\) suggesting neural plasticity in the early visual areas as possible neural substrate of this phenomenon \({}^{25-28}\). On the other hand, the reduced facilitation and the emergence of the effect at target-to-flankers separations consistent with normal periphery rather than fovea might indicate that spontaneous cortical plasticity did not produce major reorganization in the PRL projecting zone, in agreement with recent psychophysical studies \({}^{16}\).

Overall, despite being tested at similar eccentricities, MD patients showed lower contrast sensitivity with respect to their age-matched controls. Previous studies on this population have shown that even in the absence of a clear lesion in the peripheral retina, a central scotoma might impair peripheral vision through a reduction of fixation stability and/or damages in the lateral connections of the retina \({}^{20}\). Moreover, it has been shown that the PRL chosen by patients for fine vision and fixation might not always the most sensitive or adequate region of the spared retina \({}^{30,48}\). Consistently, Van der Stigchel and colleagues \({}^{1}\) showed that in a visual search task, AMD patients had inferior performances than controls tested with an artificial scotoma, suggesting that the switch from a fovea-based to a PRL-based reference frame might impair efficiency in a number of visual tasks.

Spontaneous cortical reorganization and the size of the perceptual field in MD patients. As reported in the introduction, the border between collinear inhibition and facilitation can be used as an estimate of the PF size. Previous studies showed that the inhibitory region, and consequently the size of PF, increases with eccentricity \({}^{19,21}\), suggesting a relationship with the cortical magnification factor. Measuring the size of PF might thus offer hints about the cortical reorganization processes in the MD brain, i.e., whether units formerly responding to foveal stimulation have been recruited to respond to stimuli around the PRL. Consequently, a reduction in size of the PF might be consistent with a large scale, spontaneous cortical reorganization. However, results from the present study does not seem to support this hypothesis, given that facilitation emerged at the same target-to-flankers distance for both patients and controls. Nonetheless, MD patients showed a significantly reduced inhibition at short separation, an effect associated with neural plasticity in peripheral collinear facilitation \({}^{21}\). Future studies might train patients in their PRLs on shorter target-to-flankers separations with respect to those used here (<3λ), to test whether collinear inhibition (and consequently PF size) can be reduced to the spatial extent observed in the fovea. Succeeding in reducing facilitation at shorter separations than those used in the present study would actually reshape lateral interactions in the PRL in the same way as those of the healthy fovea.

Training-induced cortical reorganization. Regarding training-induced cortical reorganization, the few existing studies report small or inconsistent improvements and limited transfer \({}^{8,10-12}\). However, a recent training paradigm based on lateral interaction showed that patients improved in the trained task and learning transferred...
to visual acuity and, partially, to contrast sensitivity function (but not to crowding)\(^2\). In general, trainings based on lateral masking might be more effective because of the plasticity hypothesis and hierarchical mechanisms they rely upon\(^5\). In Experiment 2, we trained MD patients and controls for 12 sessions on the lateral interaction configuration. The rationale was that several rehabilitative approaches are based on similar training-induced modulation of collinear facilitation\(^{14,18}\). Results showed that MD patients (but not age matched controls) reduced their collinear inhibition at short flankers separations, to the point that on average, inhibition was no longer observed for all the separations tested. If the reduced inhibition observed in Experiment 1 is consistent with the hypothesis of spontaneous cortical reorganization, PL further amplified this reduction. This improvement is even larger than the one observed in younger, healthy participants\(^21\). Because the neural substrates of collinear facilitation lay in early visual cortex, a portion of which does not receive foveal input anymore, it is possible that a training specifically designed to stimulate these regions might further promote plasticity, e.g., by recruiting new neural units or by reshaping the PF size in a more fovea-like fashion. This result is particularly promising because the majority of rehabilitation approaches in MD patients currently focuses on improving reading speed and training-induced reduction of cortical inhibition is correlated with reduction of visual crowding\(^{21}\), one of factors contributing to slower letter recognition (and consequently reading speed) in peripheral vision\(^8\). However, one cannot rule out the possibility that low level neural plasticity is not the sole or main responsible of the observed performance changes. Other mechanisms, such as attentional modulation or read-out models, could also account for the training results\(^{69,38}\).

**Limitations of the study.** Given the clinical population involved, the study presents a number of limitations. The main limitation of this study is the small number of patients. This small number is largely explained by the fact that PL trainings are often long. In our case, sessions extended over several weeks. It made it more difficult to recruit a larger number of MD patients, and speculative studies showed that reliable and robust results could still be obtained in this case by comparing performances before and after training\(^{27,28–34}\). In our study, patients consistently showed reduced inhibition with respect to controls at baseline as well as larger learning effect after training with respect to controls.

Additionally, we did not directly measure eye position (e.g., by imaging the retina or tracking the eye) during the training. Nonetheless, we used visual aids to facilitate fixation. This approach was shown to improve eye stability in previous studies on MD patients\(^38\). In our case, all the patients had a unique and well-defined PRL. Thus, they could not rely on a secondary PRL to perform the task. In addition, they all reported that both the flankers and the targets were visible throughout the whole experiment. We are therefore confident that our results are very unlikely affected by fixation instability. A direct measure of eye position would however have permitted to better characterize this aspect.

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**Author Contributions**

M.M. designed, implemented the experiments, collected and analysed the data. All authors interpreted the results and wrote the main manuscript. All authors reviewed the manuscript.

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