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Amblyopia and the binocular approach to its therapy

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ABSTRACT

There is growing evidence that abnormal binocular interactions play a key role in amblyopia. In particular, stronger suppression of the amblyopic eye has been associated with poorer amblyopic eye visual acuity and a new therapy has been described that directly targets binocular function and has been found to improve both monocular and binocular vision in adults and children with amblyopia. Furthermore, non-invasive brain stimulation techniques that alter excitation and inhibition within the visual cortex have been shown to improve vision in the amblyopic eye. The aim of this review is to summarize this previous work and interpret the therapeutic effects of binocular therapy and non-invasive brain stimulation in the context of three potential neural mechanisms; active inhibition of signals from the amblyopic eye, attenuation of information from the amblyopic eye and metaplasticity of synaptic long term potentiation and long term depression.

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39 1. Introduction

Amblyopia therapy is a large area as many different treatments 40 have been proposed over the last 100 years. One promising 41 42 approach for the treatment of adults with amblyopia is the combination of patching and perceptual learning in its many varied 43 forms, for which both monocular and binocular benefits have been 44 documented. More recently, the focus of research in this area has 45 46 shifted from monocular interventions that involve patching of 47 the fellow eye to approaches that directly target binocular visual function and as the primary therapeutic step. The emerging field 48 of binocular approaches to amblyopia therapy is the topic of this 49 review. 50

It is accepted that abnormal binocular visual experience in early 51 52 childhood causes amblyopia and that suppression (typically measured using the worth 4 dot test) plays an important part of the 53 clinical diagnostic picture. It has also been shown that loss of 54 binocularity is one of the defining features of amblyopia (McKee, 55 56 Levi, & Movshon, 2003) However the potential importance of 57 binocular approaches to amblyopia therapy has only recently 58 received widespread attention (Birch et al., 2014; Cleary et al., 59 2009; Hess, Mansouri, & Thompson, 2010; Hess, Thompson, & Baker, 2014; Hess et al., 2014; Li, Thompson, et al., 2013; Li 60 61 et al., 2014; Mansouri et al., 2014; Ooiemail, Su, Natale, & He,

http://dx.doi.org/10.1016/j.visres.2015.02.009 0042-6989/© 2015 Published by Elsevier Ltd. 2013; Spiegel, Li, et al., 2013; To et al., 2011). This has led to increased interest in the development of amblyopia treatments that directly address binocular dysfunction by promoting binocular vision and reducing inhibitory interactions within the visual cortex. In this review, we first summarize emerging approaches to the treatment of amblyopia that emphasize binocular visual function. We then describe the relationship between suppression of the amblyopic eye and the depth of amblyopia and explore whether suppression is due to active inhibition of information from the amblyopic eye or is simply the result of attenuated amblyopic eye signals. The concept of metaplasticity is then introduced and applied to the recovery of visual function in amblyopia. Finally, the results of studies into the application of non-invasive visual cortex stimulation to amblyopia are summarized and placed in the context of inhibition, attenuation and metaplasticity.

2. Emerging treatment options for amblyopia

Patching therapy has been used to treat amblyopia for hundreds 78 of years even though its shortcomings are many; compliance is 79 poor (Searle et al., 2002) because of the social and psychological 80 difficulty of forcing a child to wear a patch combined with the 81 impaired vision experienced by the child when the patch is in place 82 (Holmes et al., 2003; Webber et al., 2008). Although 79% of chil-83 dren show at least a 2 line improvement after 4 months of patching 84 (Repka et al., 2003), 25% of these children will regress to some 85 degree once the patch is removed (Holmes et al., 2004). More 86

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87 importantly, the binocular outcome is often poor regardless of the 88 improved amblyopic eye acuity (Birch, 2012). One reason for this is likely to be the nature of the viewing conditions during patching 89 (i.e. monocular) compared with those after patching, namely 90 binocular viewing. We do not yet know how patching works, 91 92 although possible mechanisms include a reduction of interocular suppression or a purely monocular improvement in the processing 93 94 of signals from the amblyopic eye. Since there is such a poor binocular outcome from patching, it may be safe to conclude that the 95 effects of patching primarily involve monocular mechanisms. 96

There have been a number of suggestions for improving the therapeutic approach to amblyopia. Some of these are purely monocular, some are monocular under otherwise binocular conditions and one is purely binocular, involving dichoptic stimu-100 lation and a dichoptic manipulation of contrast to enable simulta-101 neous use of both eyes. A summary of different treatment 102 suggestions is shown in Fig. 1. The first attempt to provide the 103 combination of short-term occlusion (20 min), controlled visual 104 stimulation and attentive game play (noughts and crosses) was 105 the CAM treatment (Campbell et al., 1978). Its beneficial effects 106 were later isolated to the short term nature of the occlusion and 107 the attentive game play (Mitchell, Howell, & Keith, 1983). 108 Another step in terms of the monocular approach was 109 Neurovision in which perceptual learning for threshold detection 110 was combined with short-term patching (Bonneh, Sagi, & Polat, 111 2004; Polat et al., 2004, 2005). There is no doubt that perceptual 112



Amblyopia therapies

Fig. 1. A summary of different principled approaches to the treatment of amblyopia, some purely monocular, some containing a binocular element and others purely binocular with dichoptic manipulation of parameters. Because the literature on monocular perceptual learning is large, only representative examples are shown. Also, there are a number of behavioral optometric approaches (Press, 1981) that are not included as these are beyond the scope of this review.

113 learning combined with short-term patching is much better than 114 longer-term patching with passive stimulation in terms of improv-115 ing monocular acuity (Li et al., 2005), however its usefulness for re-116 establishing binocular vision and stereopsis is less clear. A number 117 of hybrid-binocular approaches have been suggested, which are all directed to recovering monocular function but rather than doing 118 119 this under monocular conditions they do it under binocular viewing. The aim is to involve the fixing eye in recovery of vision 120 through intensive training/detection of targets presented exclu-121 sively to the amblyopic eye. These approaches are not designed 122 to reduce suppression, strengthen fusion and re-establish binocu-123 lar vision. The iBit system (Cleary et al., 2009), the "Push-Pull" 124 (Ooiemail et al., 2013) and the recent gaming approach by Noah 125 et al., 2014 (Fig. 1) fall into this category. An altogether different 126 127 principle was introduced by Hess, Mansouri, and Thompson 128 (2010) (Fig. 1). In this approach the primary aim is to restore binoc-129 ular fusion and stereopsis with an expected secondary consequence of improved vision of the amblyopic eye. To achieve this, 130 complementary dichoptic stimuli are used such that the visual task 131 can only be solved if both left and right information eye is com-132 133 bined (the binocular criterion). To achieve this, the contrast of 134 the signal seen by the fixing eye is reduced (to negate suppression) to a point where binocular combination is achievable. This "bal-135 ance point" is determined individually for each patient. Over time, 136 137 the treatment strengthens and extends the contrast range over 138 which binocular fusion can occur until it includes images of the 139 same contrast in each eye (comparable to natural viewing). There are no circumstances under which the treatment becomes monoc-140 ular because without binocular combination, the visual tasks used 141 142 for treatment are impossible. This approach is based on the theory that the amblyopic visual system retains the capacity for binocular 143 function and that suppression of the amblyopic eye plays an 144 important role in both the binocular and monocular functional 145 losses associated with amblyopia. It is important to note that 146 147 Evidence to support this theory is outlined below.

148 **3. Clinical suppression**

149 Clinical suppression refers to the lack of contribution of an amblyopic and/or strabismic eye under binocular viewing condi-150 tions. The most common tool for assessing this clinically is the 151 152 worth 4 dot test in which stimuli of different colors are presented anaglyphically and the degree to which each eye contributes to 153 perception is assessed subjectively. This allows for the diagnosis 154 155 of suppression and for it to be categorized as mild or severe. 156 Although there have been a variety of more quantitative proce-157 dures suggested (Zhou, Huang, & Hess, 2013) there is no gold standard for suppression measurement and in fact it is currently not an 158 important part of the standard clinical assessment. For this reason, 159 the relationship between clinical suppression and the degree of 160 161 amblyopia has, until recently, not been known. One of the first 162 attempts to address this question was a laboratory study con-163 ducted by Holopigian, Blake, and Greenwald (1988). Their sample 164 was small (n = 9) and it included patients with anisometropic 165 amblyopia, strabismic (esotropic) amblyopia and alternating strabismus with no amblyopia. They reported an inverse relationship 166 167 between acuity and depth of suppression, which they quantified 168 in terms of contrast (weaker suppression was associated with 169 poorer acuity).

More recently, new approaches have been developed to quantify the degree of suppression and these have been applied to larger samples of patients with amblyopia. They all come to a similar conclusion, namely that there is a <u>direct</u> relationship between the strength of suppression and the depth of amblyopia. Fig. 2 shows pooled data for 106 patients with amblyopia from three recent



Fig. 2. The relationship between contrast in the fellow fixing eye at the balance point (suppression) and acuity difference between the eyes (n = 106). *Dashed line:* the best linear fit to the data. The relationship shows that the lower the balance point contrast in the fellow fixing eye (i.e., the greater the difference in contrast between the eyes required for binocular function indicating stronger suppression; smaller values on the X-axis), the greater the difference in acuity between the two eyes (larger values on the Y-axis). Data from (Li, Hess, et al., 2013; Li et al., 2011).

studies (Li, Hess, et al., 2013; Li, Thompson, et al., 2013; Li et al., 2011) where the degree of suppression measured using a dichoptic motion coherence task (Mansouri, Thompson, & Hess, 2008) is plotted against the interocular LogMar acuity difference. Although there is variability between the three different clinically distinct subgroups (anisometropic, strabismic and mixed amblyopia), the overall result is clear; the greater the suppression (lower values on the *x*-axes), the greater the amblyopia (larger values on the *y*-axis) ($r^2 = 0.38$, p < 0.0001). This relationship is present for each subgroup separately (anisometropic amblyopia, n = 80, $r^2 = 0.25$, p < 0.001; mixed amblyopia, n = 9, $r^2 = 0.39$, p = 0.07; strabismic amblyopia, n = 17, $r^2 = 0.67$, p < 0.001).

In Fig. 3 we see a comparison of three different experimental approaches, each using a different visual stimulus, to further address the relationship between suppression and acuity in ambly-opia (Zhou, Huang, & Hess, 2013). Each stimulus is likely to reflect the function of a different cortical area; a local phase discrimination task reflecting mainly V1 function, a global orientation task reflecting ventral extra-striate function and a global motion task (also see Fig. 2) reflecting dorsal extra-striate function. One thing that these different measures have in common is that they all indicate that stronger suppression (though here because of the small *n*, the correlations are not statistically significant) is associated with poorer amblyopic eye acuity.

Measurements of suppression have also been collected in young children using an adaptation of the global motion task previously used in adults (Narasimhan, Harrison, & Giaschi, 2012). These results lend support to a direct relationship between suppression and amblyopia in children. Further support comes from a study of children, teens and adults using a different task where the interocular phase of a low spatial frequency sinusoid was used to measure suppression (Kwon et al., 2014).

Animal studies in which strabismic amblyopia is induced prismatically also argue for a direct relationship between the degree of suppression and the degree of amblyopia in different neuronal populations in visual cortex. The results of Bi et al. (2011) show that stronger suppression is associated with deeper amblyopia in areas V1 and V2 of monkey cortex (Fig. 4).

If suppression was simply a secondary consequence of the monocular loss of function in amblyopia, one would expect weaker suppression to be associated with poorer monocular vision in the amblyopic eye (Holopigian, Blake, & Greenwald, 1988). This is because there would be less information to suppress in patients with deeper amblyopia. The results described above demonstrate the opposite relationship whereby stronger suppression is

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associated with a greater loss of monocular vision. This indicates
that binocular deficits play a key role in amblyopia and suggests
a different approach to therapy, one that tackles the primary
binocular problem as a first step.

225 3.1. A binocular therapeutic approach

226 A number of laboratory observations led to a way of treating the 227 binocular vision deficit that is associated with amblyopia. First, it 228 was demonstrated that if the interocular contrast was suitably 229 adjusted to compensate for the amblyopic contrast threshold def-230 icit, binocular summation at threshold became normal (Baker 231 et al., 2007). This indicated that strabismic and anisometropic amblyopes were capable of normal binocular function at specially 232 233 selected interocular contrasts. Second, it was found that normal binocular combination could be achieved at suprathreshold 234 contrasts if the interocular stimulation was suitably balanced 235 236 between the two eyes (Baker, Meese, & Hess, 2008; Mansouri, Thompson, & Hess, 2008). Thus, even for strabismic adults, if the 237

images of the two eyes are properly aligned and the contrast in 238 the two eyes suitably balanced, information from the two eyes 239 could be combined normally. This demonstrated that humans with 240 amblyopia had latent binocular capabilities and had not been ren-241 dered structurally monocular, as previously thought on the basis of 242 the early animal deprivation literature. It was subsequently found 243 that allowing the eyes to combine information under these 244 balanced conditions resulted in a progressive strengthening of 245 binocular fusion and a correspondingly greater tolerance in the 246 interocular contrast differences required to support fusion (i.e. 247 repeated exposure to binocularly balanced stimuli allowed fusion 248 to occur at smaller interocular contrast differences). 249

This work led to a new dichoptic approach to treatment based on providing viewing conditions that allowed the two eyes to work together and the gradual alteration of interocular contrast differences until binocular combination occurred for all viewing conditions. The treatment, which typically involves 1 h a day for at least 4 days a week over a 4–6 week period, resulted in a re-establishment of binocular vision in the vast majority of cases



Acuity difference between the eyes (log units)

Fig. 3. The relationship between the degree of suppression and acuity difference between the eyes for dichoptic tasks requiring global orientation (top panel), global motion (middle panel) and local phase (bottom panel) judgements. In all panels, different symbols represent different subjects. The solid line represents the best linear fit to the data. On the right of each figure is an illustration of the stimuli used. (Modified from Zhou, Huang, & Hess, 2013).

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Fig. 4. Relationships between the extent of facilitatory/suppressive binocular interactions (10 log Peak B/M) of V1 (top) and V2 (bottom) neurons in individual strabismic monkeys and the depth of their amblyopia (Amblyopia index values were calculated for each monkey by integrating the area between the contrast sensitivity functions for the operated and fellow eyes and dividing it by the area under the function for the operated eye. This index ranges from 0 (no deficit) to 1.0 (no measurable sensitivity in the operated eye). Relationships are shown between the proportion of binocularly suppressive V1 (i.e., Peak B/M < 0 db) (top) and V2 (bottom) neurons and the depth of amblyopia (A1) (right columns) (from Bi et al., 2011).

257 regardless of the type of amblyopia or the age of the patient. 258 Furthermore, in the majority of adults, both stereopsis and monocular acuity improved (Hess et al., 2014) though there is not a strong 259 260 correlation between these two measures. This is not unexpected 261 because the reduction in stereopsis in amblyopia is not solely 262 due to the acuity reduction. To date 192 adults and children have been treated using this approach (Birch et al., 2014; Hess, 263 Mansouri, & Thompson, 2010; Hess et al., 2014; Li, Thompson, 264 265 et al., 2013; Li et al., 2014; Mansouri et al., 2014; Spiegel, Li, 266 et al., 2013; To et al., 2011) and the results (summarized in 267 Table 1) are promising. For adults (17 years and over), the average 268 improvement in amblyopic eye visual acuity is 0.24 LogMAR 269 (n = 84, 95%) CI = 0.04 LogMAR, p < 0.001). This is shown in Fig. 5A. For compliant children, the average improvement is 0.16 270 271 LogMAR (n = 91, 95% CI = 0.02, p < 0.001). For adults (17 years and over), the average improvement in amblyopic eye stereo is 272 2.55 log units (n = 65, 95% CI – 0.16, p < 0.001). This is shown in 273 Fig. 6A. For compliant children, the average stereo improvement 274 is 0.19 log units (*n* = 84, 95% CI = 0.11, *p* = 0.001). This corresponds 275 276 to an average improvement of 1175 arc s and is shown in Fig. 6B. We have recently shown that the improvements in visual function 277 that result from binocular training cannot be accounted for only by 278 the act of playing a videogame. In particular, binocular training 279 280 using the falling blocks game results in significantly larger 281 improvements visual acuity and stereopsis than monocular train-282 ing on the same game (Li, Thompson, et al., 2013).

No adverse effects have been reported from this approach and no patients have reported diplopia because they are always working under conditions where fusion is operating. Over a matter of a few weeks of training, binocular fusion could be extended to all contrasts even when the fixing eye was viewing stimuli at 100% (i.e. natural viewing). To date, this approach has been limited to patients with anisometropic amblyopia or strabismic amblyopia with a small angle of strabismus (<10PD). While it is known that 290 the treatment gains in acuity and stereo are sustained, less is 291 known about the effect of treatment on the motor status of 292 patients with a strabismus. For example, we do not yet know 293 whether these gains in binocular function are the consequence of 294 an ocular re-alignment or in spite of the ocular misalignment. 295

3.2. Binocular re-balancing; inhibition, attenuation or metaplasticity? 296

As described above, there is evidence that binocular re-balancing therapy works. However, its neural basis is still a matter of some debate. The most obvious explanation is that reducing the active inhibition of cortical inputs from the amblyopic eye allows for latent binocular function to be realized. Based on what we know about the excitatory and inhibitory circuits involved in binocular combination, the obvious site of this inhibition would be the point at which contralateral inhibitory signals contribute to contrast gain control prior to excitatory binocular combination (Meese, Georgeson, & Baker, 2006; Meese & Hess, 2004). This is shown in schematic form in Fig. 7, which depicts the first stage of a two-stage contrast gain control system. However other explanations include contrast attenuation of the information from the amblyopic eye and synaptic metaplasticity.

3.2.1. Signal inhibition

Support for an active inhibitory process comes mainly from the 312 physiological literature. Mower et al. (1984) showed that the 313 binocularity of over 50% of cortical neurons in strabismic cats could 314 be restored with microiontophoretic injections of bicuculline, a 315 GABA antagonist. Furthermore, primate studies have observed 316 non-specific inhibitory interactions between the eyes of strabismic 317 animals (Sengpiel & Blakemore, 1996; Smith et al., 1997) and 318 Sengpiel et al. (2006) showed that strabismic suppression was 319

Table 1

	Study	Ν	Age (yrs)	Tx hours	Amblyopia type	Design	Intervention	Display	Acuity improvement (LogMAR)	Stereopsis improvement	Side effects	Compliance	Treatment location	Follow up
Adults	Hess, Mansouri, and Thompson	9	24-49	5-52	Strab, mixed	Prospective case series	Dichoptic global motion	Stereoscope	0.26 (<i>p</i> = 0.003)	8/9 improved (<i>p</i> = 0.01)	None	Supervised	Laboratory	N/A
	(2010) To et al. (2011)	9	17–51	6-35	Aniso, strab,	Prospective case series	Falling blocks	iPod (lenticular)	0.19 (<i>p</i> = 0.02)	5/9 improved (<i>p</i> = 0.04)	None	Supervised	Laboratory	N/A
	Li et al. (2013)	18	19–26	10	Aniso, strab, mixed	Patching controlled, crossover	Falling blocks	Video goggles	0.18 (<i>p</i> < 0.001)	15/18 improved (<i>p</i> < 0.001)	None	Supervised	Laboratory	Stable at 3 months (n = 5)
	Spiegel et al. (2013)	16	17–31	11	Aniso, strab, mixed	Sham controlled crossover for tDCS. Dichoptic treatment consistent across groups	Falling blocks + tDCS	iPod (lenticular)	0.34 (p < 0.001)	14/16 improved (<i>p</i> = 0.004)	None	Supervised	Laboratory	Stable at 3 months $(n = 8)$
Children & adults	Hess et al. (2014)	14	13–50	22-108	Aniso, strab, mixed	Prospective case series	Falling blocks	iPod (lenticular or anaglyphic)	0.11 (<i>p</i> < 0.001)	11/14 improved (<i>p</i> < 0.001)	Transient asthenopia N = 1	On average patients played for 64% of the prescribed treatment time	Home	N/A
	Mansouri et al. (2014)	22	5-73	10-64	Aniso, strab	Prospective case series	Dichoptic global motion	Video goggles	0.34 (<i>p</i> < 0.001)	Not measured	None	Supervised	Laboratory	Stable at 6 months (<i>n</i> = 17)
Children	Knox et al. (2012)	14	5–14	5	Aniso, strab, mixed	Prospective case series. Participants had plateaued with patching and had stable VA	Falling blocks	Video goggles	0.09 (<i>p</i> < 0.001)	7/14 improved (<i>p</i> = 0.02)	None	Supervised	School (lunch break)	N/A
	Li et al. (2014)	45	4–12	16-32	Aniso, strab, mixed	Sham controlled	4 dichoptic games including falling blocks	iPad (anaglyphic)	0.08 (<i>p</i> < 0.001) compliant only: 0.1 (<i>p</i> < 0.001)	5/45 improved (p > 0.05) Not significant	None	34/45 played for 4 h or more	Home	Stable at 3 months (n = 21)
	Birch et al. (2014)	45	3–7	16–32	Aniso, strab, mixed	Sham controlled	4 dichoptic games including falling blocks	iPad (anaglyphic)	(p < 0.001) (p < 0.001) compliant only: 0.14 (p < 0.001)	3/45 improved ($p = 0.2$) Not significant Compliant children from Li et al. and Birch et al. $12/70$ improved. $p = 0.001$	None	28/45 played for 8 h or more	Home	N/A

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Fig. 5. (A) The combined acuity outcome data from 82 adults with amblyopia across a number of studies (see Table 1). An improvement of 1 line or more on the LogMar chart (0.1 LogMar) is considered significant. The large black triangle (±95% CI) indicates the average improvement. (B) The combined acuity outcome data from 90 children with amblyopia (see Table 1). An improvement of 1 line on the LogMar chart (0.1 LogMar) is considered significant. The large black diamond (±95% CL) indicates the averaged improvement. Only children who complied with treatment are included from the Li et al. (2014) and Birch et al. (2014) papers. Data points are jittered slightly to allow overlapping points to be seen.



Fig. 6. (A) The combined stereopsis outcome data from 65 adults with amblyopia across a number of studies (see Table 1). Stereopsis was not measured in Mansouri et al. (2014) (n = 17 adults). An improvement of 0.5 log units is considered clinically significant. The large black triangle (±95% CL) indicates the average improvement. (B) The combined stereopsis outcome data from 85 children with amblyopia across a number of studies (see Table 1). Stereopsis was not measured in Mansouri et al. (2014), (n = 5 children). An improvement of 0.5 log units is considered clinically significant. Unmeasurable stereo is assigned a value of 4 log units (10,000 arc s), corresponding to *D*max (Hess, Lui and Wang, 2002). The large black triangle (±95% CL) indicates the mean improvement, which is statistically significant. Only children who complied with treatment are included from the Li et al. (2014) and Birch et al. (2014) papers. Data points are jittered slightly to allow overlapping points to be seen.

mediated by inhibitory interactions involving GABA in the cat (see 320 also Sale & et al., 2007). Recently, Scholl, Tan, and Priebe (2013) 321 322 showed that in esotropic cats, estimates of the excitatory and inhibitory input to single neurons indicated the presence of binocular 323 suppression occurring as the result of inhibition at the thalamocor-324 tical synapse. Modeling suggested that this inhibition was medi-325 by inhibitory interneurons receiving input from 326 ated thalamocortical inputs and simple cells, and results in suppression 327 of binocular responses of both simple and complex cells (inherited 328 329 from their simple cell input). This is illustrated in Fig. 8.

Sengpiel et al. (2006) suggest that the suppression is of a more global nature and possibly involves horizontal connections between same and opposite eye domains in the more superficial layers of the primary visual cortex.

3.2.2. Signal attenuation

Results from human psychophysics relating to the loss of binocular combination in amblyopia have not been as clear cut as the animal neurophysiological data described above (Hess et al., 2014). The studies of Harrad and Hess (1992) provide evidence

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Fig. 7. Excitatory (green) and inhibitory (red) circuits involved in combining information between the two eyes. The inhibitory interocular connections that cross in the center of the schematic model may underpin active suppression. The full circuit involves two stages of contrast gain control each with separate sources of additive noise (S), one before and one after excitatory summation. L = left eye, R = right eye. From Meese and Hess (2004).

for multiple types of "suppression", some involving active inhibi-339 340 tion and others not. Fig. 9 illustrates the different forms that suppression can take psychophysically. Here, thresholds are plotted 341 for a dichoptic masking task where the increment to be detected 342 (y-axis) is presented to either the amblyopic (filled symbols) or fel-343 low fixing eye (open symbols) and the pedestal that is plotted on 344 345 the x-axis is presented to the other eye. The axes have been nor-346 malized to the contrast threshold of each eye, so the monocular 347 contrast deficit for the amblyopic eye has been accounted for. 348 The solid line is the dichoptic masking expected for a normal visual 349 system from the results of Legge and Foley (1980). Results falling 350 on this line indicate normal dichoptic masking. In the results 351 shown in the top left of Fig. 9. a passive monocular attenuation 352 explanation is sufficient and this is true in some observers with 353 anisometropic amblyopia as well as some with strabismic ambly-354 opia (Harrad & Hess, 1992). However, Harrad and Hess' results sug-355 gest that there are other forms of interaction that are not amenable 356 to a simple attenuation explanation. In some cases, the strength of 357 the dichoptic influence from the amblyopic to the fixing eye is 358 weaker (top middle panel of Fig. 9) than predicted from the

monocular contrast threshold attenuation, in some cases the strength of the dichoptic influence from the fixing to the amblyopic eye is stronger (top right panel of Fig. 9) or weaker (bottom middle panel) than that predicted from the monocular contrast threshold loss. In cases of alternating strabismus, there was simply no interaction between the eyes in either direction (bottom right panel of Fig. 9). Harrad and Hess showed that these suppressive interactions depended on spatial frequency, being much more marked at high spatial frequencies.

There have been a number of subsequent studies of suppression that have provided support for a passive attenuation (or imbalance) rather than for an active inhibition (Baker, Meese, & Hess, 2008; Huang, Baker, & Hess, 2012; Zhou et al., 2014). These results argue that although the dichoptic interactions themselves are normal in amblyopes, the fact that the amblyopic eye needs more contrast to detect stimuli means that stimuli of a fixed suprathreshold contrast will produce less masking from the amblyopic to fellow fixing eye. The resultant interocular imbalance in dichoptic masking will allow the fellow fixing eye to always dominate in binocular viewing. This effect is illustrated in Fig. 10 from the results of Huang et al. (2014) in which one eye views a noise field that is sinusoidally modulated in time and the other eye is briefly presented with letter stimuli of different contrasts at varying time points. Masking is demonstrated by the sinusoidal nature (rectified) of the threshold elevation for detecting the letter stimuli. The results from observers with amblyopia (middle panel) show approximately normal (compared with left panel) masking from fixing to amblyopic eye (dashed curves) but less masking from the amblyopic to fixing eye (solid curves). This is amenable to an explanation based on the reduced contrast sensitivity of the amblyopic eye as demonstrated by the model results (right panel). However, to date this explanation has not been tested directly, a process that would entail using masks that are equi-detectable (at a constant suprathreshold contrast) for each eye. Only then would we know if a simple attenuation explanation could be applied to suppression for this particular paradigm.

As a whole, the psychophysical and physiological explanations for suppression are not in agreement; physiologically there is evidence for active suppression between the two eyes of strabismic animals, psychophysically the picture of suppression is less clearcut. Simple attenuation of the amblyopic eye together with normal dichoptic inhibitory interactions may both play a part. However, attenuation alone is unlikely to provide a sufficient explanation for the population suppression measures discussed previously.



Fig. 8. Loss of thalamic input in a circuit model of strabismus. (A) Left (L) and right (R) eye inputs converge on layer 4 simple cells, generating disparity selectivity. Simple cell inputs converge onto complex cells in layer 2/3, which are also disparity selective. (B) In strabismic animals, simple cells receive monocular input. A loss of binocularity causes a loss of disparity selectivity, which also occurs in complex cells through feedforward inputs. Complex cells receive inputs from simple cells and thus can be binocular. Suppression of binocular responses is mediated by inhibitory interneurons receiving input from thalamocortical inputs and simple cells. In this simple model, the strabismus-induced changes are qualitatively similar for all neurons regardless of the initial difference in synaptic strength, spatial selectivity, and spatial phase between the inputs from each eye to the neuron. (From Scholl, Tan, & Priebe, 2013 – Fig. 9).

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Fig. 9. Dichoptic masking functions for amblyopic observers. The incremental contrast seen by one eye (filled symbols amblyopic eye; open symbols fixing eye) is plotted against the pedestal contrast seen by the other eye. Different categories of response are shown to demonstrate the heterogeneity of suppression in amblyopia, see main text for further information. From Harrad and Hess (1992).



Fig. 10. Dichoptic masking of a briefly presented letter stimulus (open symbols amblyopic eye; filled symbols fellow fixing eye) by the sinusoidal modulation of the contrast of a noise field in the other eye. Results are compared for normals, amblyopes and for a model simulation, see main text for further information. (From Huang et al., 2014).

403 Monocular contrast sensitivity loss of the amblyopic eye is greatest 404 at high spatial frequencies and minimal or non-existent at very low 405 spatial frequencies (Hess & Howell, 1977; Levi & Harwerth, 1977) 406 and the spatial properties of the global motion and dichoptic phase 407 measures that have been used to date are in the low spatial frequency range. This makes it less likely that monocular attenuation 408 of contrast in the amblyopic eye can account for the results shown 409 in Figs. 2 and 3. 410

411 3.2.3. *Metaplasticity*

Instead of thinking of rebalancing as a means of reducing the
interocular inhibition or compensating for signal attenuation, it
might be more useful to think about it in terms of synaptic plasticity. Our understanding of plasticity at the level of the synapse has

changed considerably over the last decade. An understanding of 416 synaptic plasticity goes well beyond the rules suggested by Hebb 417 whereby synapses "that fire together wire together". Synaptic plas-418 ticity is governed by NMDA receptors (Sawtell et al., 2003) which 419 support long-term potentiation (LTP) and long-term depression 420 (LTD) (Cho & Bear, 2010). The way in which this bidirectional 421 synaptic modification operates is itself modifiable. This is termed 422 metaplasticity. Specifically, the threshold change in synaptic input 423 that results in LTP rather than LTD depends on the history of cor-424 tical activity as described by the Bienenstock-Cooper-Munro 425 (BCM) theory (Bienenstock, Cooper, & Munro, 1982). Potentiation 426 occurs when activation exceeds this threshold, which itself is a 427 function of the history of neuronal firing. This bidirectional synap-428 tic modification is illustrated in Fig. 11 where the change in 429

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synaptic strength is plotted against the postsynaptic activity; low
levels of post-synaptic activity result in LTD, high levels in LTP.
The level of post-synaptic activity corresponding to the transition
from LTD to LTP is termed the modification threshold.

434 Instead of thinking about suppression in terms of an active inhi-435 bition or signal attenuation, it could simply be the outcome of 436 synapses with strong fixing eye activation and weak amblyopic 437 eye activation. No matter how strongly the amblyopic eye is acti-438 vated under these conditions, the synapse will be unable to take advantage of the increased neural activity because its modification 439 threshold is governed by the activity from the fixing eye. However 440 441 with, for example, dichoptic therapy, when the fixing eye activation is driven down, the modification thresholds may shift in favor 442 of LTP and the weak inputs from the amblyopic eye, that are now 443 444 more correlated with postsynaptic activity than before, may be 445 able to initiate potentiation via synaptic metaplasticity (see for 446 review, (Cooper & Bear, 2012). The longer the visual system can 447 be kept in a state where the presynaptic activity of both eyes cor-448 relates with post synaptic activity, the stronger, more permanent 449 and more balanced will be the ocular dominance. A similar argu-450 ment has been made concerning the beneficial effects of dark 451 adaptation on ocular dominance plasticity (He et al., 2007). Thought of in these terms, active inhibitory mechanisms or simple 452 signal compensation may not be the right way to conceptualize 453 454 clinical suppression or the basis of binocular therapy.

455 3.3. Non-invasive brain stimulation and amblyopia

456 Non-invasive brain stimulation is another way of modulating 457 excitability and inhibition/suppression within the visual cortex of 458 patients with amblyopia. A number of well established techniques 459 for safely stimulating the human brain are available. These include transcranial magnetic stimulation (TMS), which utilizes magnetic 460 461 induction to generate weak electrical currents in targeted cortical 462 areas (Barker, Jalinous, & Freeston, 1985; Hallett, 2007) and tran-463 scranial direct current stimulation (tDCS) that involves a small 464 (1–2 mA) current passed between two head mounted electrodes 465 (Nitsche & Paulus, 2000). The delivery of repeated pulses of TMS 466 (repetitive TMS; rTMS) can induce lasting increases or decreases 467 in neural excitability depending on the pattern and frequency of 468 stimulation (Fitzgerald, Fountain, & Daskalakis, 2006). tDCS can



Postsynaptic response

Fig. 11. The BCM theory of synaptic plasticity includes a sliding modification threshold that depends on the history of postsynaptic activity. The value of the modification threshold is shown for two conditions; normal viewing where the activity of the fixing eye dominates and a balanced viewing condition where the activity of the fixing eye has been reduced so that the amblyopic eye activity which was previously depressed (LTD) is now potentiated (LTP). Adapted from Cooper and Bear (2012).

also induce increases and decreases in excitability depending on 469 the direction of current flow (Nitsche & Paulus, 2000). Anodal 470 tDCS tends to increase excitability where as cathodal tDCS 471 decreases excitability. While the effects of rTMS and tDCS on neu-472 ral excitability are well documented (Dayan et al., 2013), the 473 underlying mechanisms are yet to be identified. However, a grow-474 ing number of pharmacological and neurophysiological studies are 475 shedding light on the neural mechanisms involved (Allen et al., 476 2007; Funke & Benali, 2011; Kozyrev, Eysel, & Jancke, 2014; 477 Stagg & Nitsche, 2011). For example, NMDA receptors appear to 478 be involved in the after-effects of both tDCS and rTMS (Huang 479 et al., 2007; Nitsche et al., 2003), providing a theoretical link to 480 long-term potentiation and long-term depression. 481

rTMS and tDCS have advanced our understanding of the human 482 brain and have significant potential as tools for rehabilitation. For 483 example, rTMS has been FDA approved for the treatment of depres-484 sion. Furthermore, the use of rTMS and tDCS to alter pathological 485 patterns of neural excitation and inhibition has shown promise 486 in the treatment of stroke (Hummel & Cohen, 2006; Talelli, 487 Greenwood, & Rothwell, 2007), tinnitus (Vanneste, Langguth, & 488 De Ridder, 2011), chronic pain (Fregni, Freedman, & Pascual-489 Leone, 2007) and hemispatial neglect (Muri et al., 2013). The use 490 of rTMS to alter abnormal inhibitory interactions between the 491 two cerebral hemispheres in stroke (Hummel & Cohen, 2006) 492 was the inspiration for applying non-invasive brain simulation to 493 amblyopia. As described above, signals from the amblyopic eye 494 evoke low levels of neural activity (Barnes et al., 2001) and may 495 be subject to active inhibition (suppression) within the primary 496 or extrastriate visual cortex (Bi et al., 2011; Sengpiel & 497 Blakemore, 1996). We hypothesized that rTMS would strengthen 498 the response of the visual cortex to inputs from the amblyopic 499 eye (Thompson et al., 2012). This idea was based on reports that 500 rTMS could reduce intracortical inhibition (Fitzgerald, Fountain, & 501 Daskalakis, 2006), at least within the motor cortex, and therefore 502 may reduce inhibition of information from the amblyopic eye. 503 Furthermore, rTMS had been shown to have a homeostatic effect, 504 with inhibited neural populations being more susceptible to exci-505 tatory stimulation and populations with high levels of excitation 506 being more susceptible to inhibitory stimulation (Silvanto, 507 Muggleton, & Walsh, 2008). Therefore, excitatory rTMS protocols 508 may preferentially affect inputs from the amblyopic eye whereas 509 inhibitory protocols may target fellow eye inputs. In this scenario, 510 the net effect of either an excitatory or inhibitory rTMS protocol 511 would be a reduction in the activation difference between cortical 512 inputs from the two eyes. 513

Our first study in a small group of adults with amblyopia supported this hypothesis; both excitatory and inhibitory rTMS protocols increased amblyopic eye contrast sensitivity by an average of 40%, with excitatory rTMS having a more consistent effect across participants (Thompson et al., 2008). Stimulation of the motor cortex had no effect. As part of the procedure for the calibration of stimulus intensity, we measured phosphene thresholds in both patients and controls. Phosphene thresholds are the lowest intensity of single pulse of visual cortex TMS that can elicit the percept of a phosphene and are often used as a measure of visual cortex excitability (Antal et al., 2003; Aurora, Welch, & Al-Sayed, 2003). Unexpectedly, we found that patients with amblyopia had significantly higher phosphene thresholds than controls (Fig. 12A). This preliminary finding suggests that the visual cortex of patients with amblyopia has lower overall levels of excitability that controls, possibly due to suppressive interactions.

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In our original study, the effects of rTMS on contrast sensitivity were transient, returning to baseline within 24 h in most cases. In a follow up study, we found that repeated administration of visual cortex continuous theta burst stimulation (cTBS, a form of rTMS that requires only a short stimulation period) over 5 days led to



Fig. 12. Transcranial magnetic stimulation and amblyopia. Panel A shows phosphene thresholds for patients with amblyopia (n = 9) and controls (n = 5). Larger values on the *y*-axis indicate that greater intensities of single pulse TMS were required to elicit the perception of a phosphene. Patients with amblyopia had significantly higher phosphene thresholds than controls ($t_{12} = 2.8$, p = 0.02) suggesting lower levels of visual cortex excitability. Data from Thompson et al. (2008). Panel B shows contrast sensitivity data from three patients treated with 5 daily sessions of visual cortex continuous theta burst stimulation (cTBS). cTBS induced improvements in amblyopic eye contrast sensitivity that lasted up to 78 days (D0 = baseline). Figure from Clavagnier, Thompson, and Hess (2013).

long lasting improvements in contrast sensitivity that were stable
over a period of up to 78 days (Clavagnier, Thompson, & Hess,
2013) (Fig. 12B). This indicates that multiple doses of cTBS may
lead to lasting and perhaps permanent improvements in visual
function in adults with amblyopia. Only three to four repeated
(one per day) applications of cTBS were required to produce
long-term, stable improvements.

542 In a parallel series of studies, we have investigated the effect of 543 tDCS on amblyopic eye contrast sensitivity (Spiegel, byblow, et al., 2013). This work was motivated by a magnetic resonance spec-544 troscopy study, which revealed that anodal tDCS acted to reduce 545 the concentration of GABA when applied to the motor cortex 546 547 (Stagg et al., 2009). We hypothesized that anodal tDCS would have 548 a similar effect on the visual cortex and may, therefore, reduce sup-549 pression and improve vision in patients with amblyopia. Before 550 applying tDCS to patients with amblyopia, we first investigated the effects of anodal tDCS on psychophysically measured surround 551 552 suppression in observers with normal vision (Spiegel et al., 2012). Surround suppression is thought to involve GABA-mediated inhibi-553 tory interactions within the primary visual cortex (Yoon et al., 554 2010). Anodal tDCS significantly attenuated surround suppression, 555 556 but had no effect on overlay suppression, a control condition that 557 does not involve inhibition in V1. Cathodal tDCS had no effect on 558 either condition. Based on these results, anodal tDCS was applied 559 to the visual cortex of thirteen patients with amblyopia. Eight 560 out of thirteen patients experienced transient improvements in contrast sensitivity in response to anodal but not cathodal tDCS 561 562 (Spiegel, Byblow, et al., 2013). There were no obvious clinical or demographic differences between the group of patients who 563 showed improvements and those that did not, however individual 564 565 differences in the response to tDCS are well documented and have 566 been linked to a range of variables including patterns of functional connectivity within neural networks (Vanneste et al., 2011). To 567 568 ensure that the effects we observed were due to tDCS-induced 569 changes within the visual cortex, we used fMRI to measure the rel-570 ative response of V1, V2 and V3 to contrast reversing checker-571 boards shown to the amblyopic vs. the fellow eyes. After sham

tDCS, large areas of the primary and extrastriate visual cortex showed a significantly larger response to the fellow eye than the amblyopic eye in agreement with previous studies demonstrating that the amblyopic eye is less able to activate the visual cortex (Barnes et al., 2001). This bias towards stronger activation in the fellow eye was reduced by anodal tDCS, with significant effects observed in V2 and V3. Anodal tDCS may have normalized the cortical response to information from each eye, possibly by reducing suppression within the visual cortex.

The finding the anodal tDCS may act to reduce suppression in the visual cortex raised the possibility that anodal tDCS could also enhance the effects of dichoptic treatment. In a recent study we demonstrated that this was indeed the case, anodal tDCS combined with dichoptic treatment led to significantly greater improvements in stereopsis than sham tDCS combined with dichoptic treatment (Spiegel, Li, et al., 2013). This effect was not present for monocular measures of effects of anodal tDCS were limited to binocular visual function.

Non-invasive brain stimulation is now an established technique in many fields, however research into the use of brain stimulation to promote recovery of vision is sill in its infancy. Furthermore, as described above, mechanistic studies of noninvasive brain stimulation have mostly focused to the motor cortex and it is not clear how these findings translate to the visual cortex. The initial results summarized here indicate that non-invasive brain stimulation is a useful tool for investigating and potentially treating the neural basis of amblyopia. Future work will establish whether non-invasive brain stimulation has a role in amblyopia treatment, either as a stand-alone therapy or in combination with other interventions such as binocular therapy.

When considered in the context of inhibition, attenuation and metaplasticity, the effects of rTMS and tDCS on amblyopic eye function are consistent with reductions in inhibition or attenuation of information from the amblyopic eye, which may be permissive for synaptic plasticity. On the basis of current data it is not possible to definitively distinguish between changes in inhibition and attenuation. However, the preliminary data indicating abnormally

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high levels of inhibition within the amblyopic visual cortex
(Fig. 10A), combined with the ability of anodal tDCS to reduce surround suppression and GABA concentration favor a reduction in
inhibition/suppression.

613 4. Conclusions

Suppression is an important part of the amblyopia syndrome 614 and the positive correlation between suppression and the depth 615 of amblyopia indicates that binocular dysfunction is the primary 616 617 problem. Numerous studies demonstrating that balancing the 618 information seen by the two eyes can promote binocular function 619 and lead to a re-establishment of binocular vision further support 620 this idea. These advances have raised a number of questions that 621 are yet to be answered: Is the basis for the original imbalance 622 between the amblyopic and fellow eyes signal attenuation, signal 623 inhibition, metaplasticity or a combination of these? Do binocular 624 therapy and non-invasive brain stimulation lead to reduced active 625 cortical inhibition, a change in synaptic metaplasticity or the two 626 in concert? Answers to these questions will provide new insights 627 into amblyopia and the mechanisms controlling plasticity within 628 the adult human visual cortex.

629 **5. Uncited reference**

Goodman et al. (2011).

631 Acknowledgments

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