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Visual Field Outcomes from LiGHT: Laser in Glaucoma and Ocular Hypertension, a multicentre, randomised controlled trial

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1 Visual Field Outcomes from LiGHT: Laser in Glaucoma and Ocular

- ² Hypertension, a multicentre, randomised controlled trial.
- 3
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- 24 Running head: Eye drops vs SLT: visual field progression in glaucoma.

25 Acronyms

- 26 GPA Guided progression analysis.
- 27 IOP Intra-ocular pressure.
- 28 MD Mean deviation.
- 29 OAG Open angle glaucoma.
- 30 OHT Ocular hypertension.
- 31 OR Odds Ratio.
- 32 PSD Pattern standard deviation.
- 33 PD Pattern deviation.
- 34 SLT Selective laser trabeculoplasty.
- 35 TD Total deviation.
- 36 VF Visual field.

37 Abstract

- 38 Objective
- 39 To compare visual field outcomes of ocular hypertensive and glaucoma patients treated with
- 40 Medicine-1st against those treated with selective laser trabeculoplasty (SLT, Laser-1st).
- 41 Design
- 42 Secondary analysis of patients from Laser in Glaucoma and Ocular Hypertension (LiGHT), a
- 43 multicentre randomised controlled trial.
- 44 Participants and controls
- 45 344 patients (588 eyes) treated with Medicine-1st, 344 patients (590 eyes) treated with Laser-1st.
- 46 Methods
- 47 Visual fields (VFs) were measured using standard automated perimetry and arranged in series
- 48 (median length and duration: 9 VFs over 48 months). Hierarchical linear models were used to
- 49 estimate pointwise VF progression rates, which were then averaged to produce a global progression
- 50 estimate for each eye. Proportions of points and patients in each treatment group with fast (< -1
- 51 dB/y) or moderate (< -0.5 dB/y) progression were compared using log-binomial regression.
- 52 Main outcome measures
- 53 Pointwise and global progression rates of total deviation (TD) and pattern deviation (PD).
- 54 Results

A greater proportion of eyes underwent moderate or fast TD progression in the Medicine-1st group compared with the Laser-1st group (26.2% vs. 16.9%; Risk Ratio, RR = 1.55 [1.23, 1.93], *P* < 0.001). A similar pattern was observed for pointwise rates (Medicine-1st 26.1% vs. Laser-1st 19.0%, RR = 1.37

- 58 [1.33, 1.42], *P* < 0.001). A greater proportion of pointwise PD rates were categorised as moderate or
- 59 fast in the Medicine-1st group (Medicine-1st 11.5% vs. Laser-1st 8.3%, RR = 1.39 [1.32, 1.46], P <

- 60 0.001). There was no statistical difference in the proportion of eyes that underwent moderate or
- 61 fast PD progression (Medicine- 1^{st} 9.9% vs. Laser- 1^{st} 7.1%, RR = 1.39 [0.95, 2.03], P = 0.0928).
- 62 Conclusion
- 63 A slightly larger proportion of ocular hypertensive and glaucoma patients treated with Medicine-1st
- 64 underwent rapid VF progression compared with those treated with Laser-1st.

Journal Pre-proof

65 Introduction

Glaucoma is a progressive optic neuropathy, that left untreated can lead to loss of vision. Glaucoma can have significant implications for patients and is associated with worse vision related quality of life¹⁻⁴. Assessing visual function, typically done by visual field (VF) examination, is vital for clinical management, especially for assessing the effectiveness of treatment in controlling the disease. VF progression will usually drive treatment intensity, as lowering intra-ocular pressure (IOP) is the only currently available treatment to slow the progression of glaucoma⁵.

72 Thus far, IOP lowering eye drops have been used as a 1st-line treatment for glaucoma and ocular

73 hypertension (OHT), but a recent report from the Laser in Glaucoma and Ocular Hypertension

74 (LiGHT) trial showed that selective laser trabeculoplasty (SLT), an outpatient laser procedure for the

reduction of IOP, provides better clinical effectiveness and lower treatment intensity among newly

76 diagnosed glaucoma and OHT patients compared to IOP lowering eye drops, and comparable health

related quality of life, whilst also being cost-effective ⁶.

78 Although the IOP lowering efficacy of SLT has been extensively compared to that of eye drops^{7–11}

and despite a substantial body of research into VF progression in glaucomatous patients, little

80 evidence exists comparing SLT and IOP lowering eye drops in terms of VF outcomes. This study aims

81 to compare VF progression between patients who received SLT to those who received IOP lowering

82 eye drops, as a 1st-line treatment for glaucoma and OHT in the LiGHT trial.

83 Methods

84 Analysis cohort

Details of the LiGHT trial design and baseline characteristics are described elsewhere^{12,13}. Briefly, the LiGHT trial is a multi-centre, randomised controlled trial comparing IOP lowering eye drops to SLT. A total of 718 newly diagnosed, previously untreated OHT or open angle glaucoma (OAG) patients were randomised to one of two treatment pathways. Patients in the Medicine-1st group received

topical IOP lowering eye drops to reduce IOP, whereas patients in the Laser-1st group received SLT 89 (followed by medication if required as the trial progressed). Subsequent treatment decisions 90 surrounding treatment escalations, repeated SLT or trabeculectomy were conducted according to 91 92 the study protocol with the aid of a computerised decision algorithm to avoid bias in clinical decision 93 making. The decision support algorithm used in the LiGHT trial has been described in detail previously ^{12,14}. Patients were treated to eye-specific IOP targets that were determined according to 94 95 the computer algorithm. Recruitment lasted two years and ended in October 2014. Primary outcomes were reported at three years and additional funding allowed the trial to extend for a 96 97 further three years.

98 At each study visit, visual fields (VFs) were measured using the Humphrey Field Analyzer (HFA) with Swedish interactive threshold algorithm standard 24-2 programme (Carl Zeiss Meditec, Dublin, CA, 99 100 USA). VF measurements were used primarily as an input (along with IOP and optic disc imaging 101 measurements) into decision support software (DSS), which generated eye-specific treatment 102 recommendations at each study visit. The secondary analysis reported here used VFs extracted from the DSS database on 13th December 2018, as the trial approached the six-year mark. We constructed 103 a longitudinal series of VFs for each study eye and these formed the basis for all analyses. A total of 104 105 11,823 VFs were extracted from the database. Of these, we excluded 86 VFs with false positive rates 106 > 14% as potentially unreliable, and 56 eyes with very short series (< 5 VFs) as these contained little information from which to estimate progression. Following these exclusions there remained 11,563 107 108 VFs, approximately equally distributed between treatment groups. A total of 1178 eyes from 688 109 patients (95.8% of those randomised) were included in this analysis; treatment groups had similar patient baseline characteristics both to each other and to previously reported analyses^{6,13} (Table 1). 110 Median follow-up time (Medicine-1st 47 months, Laser-1st 49 months) and VF series length 111 (Medicine-1st 5630 VFs, 9 VFs per eye; Laser-1st 5933 VFs, 10 VFs per eye) were similar across 112 treatment groups. 113

114 Statistical analysis

We compared VF outcomes between groups by constructing hierarchical linear models describing 115 116 change in VF measures over time using the visual field data described above. A trend based method 117 of comparison was chosen because it is potentially more sensitive than event based methods such as Guided Progression Analysis (GPA) for detecting progression ^{15,16}, especially where the number of 118 119 events is expected to be small as in these early cases. We examined change at each of the 52 measured locations (excluding the blind spot) in each VF series, specifying a random effects 120 structure nesting locations within eyes, within individuals¹⁷. This accounted for variation in response 121 among locations, due to eye level variation and correlation between eyes within individuals, 122 123 respectively, whilst pooling information across the entire cohort to produce the most accurate 124 estimates. Fixed effects terms represented baseline values (equivalent to y-axis intercept [dB]) and 125 rate of change per year (slope; dB/year) in each treatment group, enabling us to simultaneously 126 evaluate (using the slope by group interaction term) the statistical evidence for a difference in progression rates between groups and to estimate effect size (i.e. difference in slopes)^{16,18}. 127 Two outcome variables were modelled. Total deviation (TD) is the difference of the measured 128 129 sensitivity at each location from that expected for a patient of that age with no pathology. Pattern 130 deviation (PD) is the TD value at each location adjusted for generalised depression of sensitivity across the VF¹⁹. Both PD and TD values were extracted from the HFA. Generalised depression and 131 132 changes in TD may be caused by several non-glaucomatous conditions including cataract, whereas 133 PD is designed to highlight the more localised VF changes found in glaucoma. However, glaucoma almost always has a diffuse component which is ignored by PD, so it is a less sensitive measure than 134 TD and is prone to underestimation of glaucomatous damage than TD^{20} . Models were fitted in R 135 136 version 3.5 (R Development Core Team, R Foundation for Statistical Computing, Vienna, Austria). 137 Alongside pointwise estimates, global estimates of TD and PD progression for each study eye were 138 extracted from the models. For each eye, the estimated rate at each location was extracted; the

mean of these pointwise rates was calculated to give the global estimate for that eye. Pointwise
estimates enable better detection of spatially localised changes, whereas global estimates are useful
for describing diffuse changes in sensitivity.

142 To assess the clinical importance of differences between treatment groups, we categorised 143 estimated progression rates of each location and eye into one of six categories (fast progression: -1 > 144 slope dB/y, moderate progression: $-1 \le \text{slope} < -0.5 \text{ dB/y}$, slow progression: $-0.5 \le \text{slope} < 0 \text{ dB/y}$, 145 slow improvement: $0 \le \log < 0.5 \text{ dB/y}$, moderate improvement: $0.5 \le \log < 1 \text{ dB/y}$, fast improvement: slope ≥ 1 dB/y. Category boundaries in the progression end (i.e. slope < 0) of the rate 146 distribution were based on those previously reported in studies of glaucoma progression in clinical 147 populations^{21,22}. A symmetrical set of boundaries were applied to the improvement end of the 148 149 distribution as a measure of variability. A tendency towards faster progression and also faster 150 improvement in one treatment group (i.e. a fatter tailed distribution) would indicate greater 151 variability in rates rather than a shift towards faster progression. We used log-binomial (relative risk) 152 regression to compare the proportion of locations and eyes in each group undergoing fast or 153 moderate progression, representing patients at the greatest risk of vision loss. These models were 154 non-hierarchical, with treatment group as the predictor and the outcome being a binary variable 155 indicating whether the estimated rates (from the hierarchical model) were above or below -0.5 dB/y. 156 At the other end of the rate distribution, the proportions of locations and eyes undergoing fast or moderate improvement were compared in a similar manner. 157

We conducted a sensitivity analysis to further investigate the influence of cataract, refitting our
models to exclude eyes that underwent cataract removal. Similarly, eyes that underwent
trabeculectomy may have experienced a step increase in sensitivity after surgery. We censored VF
series for these eyes at time of surgery and refitted the models.

The study adhered to the tenets of the Declaration of Helsinki. Ethical approval was obtained from
 local boards at each participating centre. All patients provided written informed consent before

participation. The study is registered at controlled-trials.com (ISRCTN32038223) and the protocol is
 available online¹².

166 Results

167 Total deviation

Estimated mean pointwise total deviation decreased in both the Medicine-1st and Laser-1st groups 168 over time (mean and 95%CI: Medicine-1st = -0.25 dB/y [-0.31, -0.19]; SLT = -0.19 dB/y [-0.25, -0.13]). 169 170 There was little evidence for a difference in mean rates of progression between groups (slope by group interaction term, t = 1.41, P = 0.157) but the distribution of estimated progression rates did 171 vary by group. Distributions of both pointwise and global estimates were more strongly left skewed 172 in the Medicine-1st group than in the Laser-1st group (Figure 1, global estimates), indicating that 173 greater proportions of locations and eyes in the Medicine-1st group showed evidence of more rapid 174 175 progression (Table 2).

One in four eyes underwent moderate or fast progression in the Medicine-1st group compared with 176 177 approximately one in six eyes in the Laser-1st group (Risk Ratio, RR = 1.55 [1.23, 1.93], P < 0.001). Similarly, a greater proportion of locations was categorised as having moderate or fast progression 178 in the Medicine-1st group (RR = 1.37 [1.33, 1.42], *P* < 0.001). There was no evidence for a difference 179 between treatment groups in the proportion of eyes that underwent moderate or fast improvement 180 (RR 1.29 [0.83, 2.04], P = 0.266). A greater proportion of locations was categorised as having 181 moderate or fast improvement in the Medicine- 1^{st} group (RR = 1.31 [1.24, 1.39], P < 0.001). 182 183 Following exclusion of eyes that underwent cataract removal, the differences between treatment groups were attenuated: eyes that underwent moderate or fast progression (RR = 1.43 [1.11, 1.83], 184 P = 0.005; locations (RR = 1.25 [1.21, 1.29], P < 0.001). Censoring VF series at trabeculectomy had 185 186 almost no influence on estimated differences between treatment groups (RRs not shown).

187 Pattern deviation

188	The distribution of progression estimates was similar for pattern deviation but estimated rates were
189	lower and differences between treatment groups were less pronounced than for total deviation.
190	Estimated mean pointwise pattern deviation decreased in both the Medicine-1 st and Laser-1 st groups
191	over time (mean and 95%CI: Medicine-1 st = -0.12 dB/y [-0.16, -0.09]; Laser-1 st = -0.09 dB/y [-0.13, -
192	0.06]). There was no evidence for a difference in mean rates of progression between groups (t =
193	1.19, $P = 0.236$) but both pointwise and global estimates were more strongly left skewed in the
194	Medicine-1 st group than in the Laser-1 st group (Figure 2).
195	There was no evidence for a statistical difference between treatment groups in the proportion of

eyes that underwent moderate or fast progression (Table 3, RR = 1.39 [0.95, 2.03], P = 0.0928). A

197 greater proportion of locations was categorised as having moderate or fast progression in the

198 Medicine-1st group (Table 3, RR = 1.39 [1.32, 1.46], *P* < 0.001). There was no evidence for a

difference between treatment groups in the proportion of eyes that underwent moderate or fast

improvement (RR 1.86 [0.75, 4.64], P = 0.181). A greater proportion of locations were categorised as

having moderate or fast improvement in the Medicine- 1^{st} group (RR = 1.37 [1.24, 1.51], P < 0.001).

202 Following exclusion of eyes that underwent cataract removal, the differences between treatment

groups were attenuated: eyes that underwent moderate or fast progression (RR = 1.18 [0.78, 1.77],

204 *P* = 0.436); locations (RR = 1.29 [1.22, 1.35], *P* < 0.001). Censoring VF series at trabeculectomy had

almost no influence on estimated differences between treatment groups (RRs not shown).

206 Baseline sensitivity, IOP and progression rates

Eyes that underwent fast progression or improvement had lower average sensitivity at baseline than
 those with intermediate progression or improvement rates (Figure 3). Similarly, eyes that underwent
 fast progression or improvement had slightly lower IOP targets set at baseline than those with

intermediate rates (Figure 4). There was no evidence that the distributions of baseline sensitivity orIOP targets differed between treatment groups (Table 1).

212 Discussion

213 This study reports on the VF progression differences between glaucoma/OHT patients treated with Medicine-1st and patients treated with Laser-1st in the LiGHT trial. Using TD values, we estimated 214 that one in four eyes had moderate or fast VF progression in the Medicine-1st group whereas in the 215 Laser-1st group this value was about one in six. The difference between groups was less pronounced, 216 with no statistical evidence for a difference, when using PD values. The proportion of pointwise rates 217 that were moderate or fast was slightly greater in the Medicine-1st group using both PD and TD. 218 219 These differences were not reflected at the upper ends of the rate distributions for either eyes or locations, indicating that our findings were not the result of greater variability in one or other 220 221 treatment group.

The results of this study suggest that treating patients with Laser-1st may delay VF progression in 222 comparison to Medicine-1st. IOP control with eye drops may rely upon patient concordance with 223 224 treatment; indeed IOP lowering drops have been reportedly available to patients only 69% of the 225 time, whilst concordance may range between 76-86% with even lower figures reported for more complex instillation regimes^{23–25}. Although self-reported concordance in the LiGHT trial has been 226 high¹⁴, the possibility of poor concordance having a significant adverse effect on disease control 227 228 cannot be ruled out as actual dose monitoring was not carried out. However, patients in clinical trials are reported to have higher rates of concordance than those in routine care²⁶. Thus the true 229 230 magnitude and clinical importance of the slowing of VF progression in the Laser-1st group may be much greater. SLT has also been proposed to provide better diurnal IOP stability, as a result of a 231 continuous effect on the trabecular meshwork^{27–30}. This is in contrast to the episodic (and sometimes 232 erratic) administration of medication that may allow greater diurnal fluctuation in IOP, and in turn 233

faster disease progression. Even with exact concordance with instillation regimes, there are likely tobe long gaps between doses overnight, during which IOP may rise.

236 We observed differences in VF progression between treatment groups despite the fact that both 237 groups were treated to similar IOP targets. This indicates that monitoring of IOP reduction alone (usually measured during office hours and so potentially unrepresentative of diurnal pressure 238 239 variation) may be insufficient to predict functional changes indicative of progression. This suggests 240 that clinical trials of new glaucoma treatments should include both IOP and VF related outcomes. 241 Greater differences were observed for TD, hinting that non-glaucomatous changes may have also 242 contributed towards differences between groups. Changes in TD may be caused by a number of nonglaucomatous conditions, such as cataract. Were there higher rates of cataract in the Medicine-1st 243 group it could partially explain the tendency towards faster TD progression. During the period 244 245 covered by this analysis, cataracts were removed from 10.9% of eyes in the Medicine-1st group and 246 7.1% of eyes in the Laser-1st group. Assuming that cataracts not yet requiring surgery follow this 247 distribution, generalised depression of sensitivity due to lens opacity have contributed towards the 248 differences in TD rate between the two treatment groups. This is consistent with the higher rates of cataract after topical medical treatment of glaucoma previously reported by landmark glaucoma 249 studies^{31–34} and itself may contribute to a significant clinical advantage of a Laser-1st compared to a 250 251 Medicine-1st protocol. Our sensitivity analysis showed that differences between treatment groups were narrowed when eyes that underwent cataract removals were excluded. PD models were as 252 253 strongly influenced by the exclusions as TD models. For example, following the exclusions there was 254 no statistical evidence for a difference in the proportion of eyes undergoing fast or moderate PD 255 progression (there remained strong evidence for a difference in the proportion of locations with moderate or fast progression). This may indicate that as well as having lower sensitivity than TD²⁰, 256 257 PD may not be immune to the influence of cataract. Alternatively, the similar responses of TD and PD following exclusions may indicate that cataract was not driving the between group differences. 258 Instead, cataract formation may be associated with faster glaucoma progression (with oxidative 259

stress a potential biological basis for the association) and by excluding cataract removal eyes much
of the glaucoma signal may have been excluded also. Considering that we still found clinically
relevant differences between treatment groups following exclusion of eyes from which cataracts
were removed, and recognising the limitations of both TD and PD, we conclude that greater
incidence of both cataract-related and glaucomatous progression in the Medicine-1st group is likely
to have contributed towards the observed differences between treatment groups.

To our knowledge this is the first study to robustly compare VF outcomes between IOP lowering 266 drops and SLT, as previous research has focused on IOP lowering alone as a surrogate for disease 267 control. In the absence of a universally accepted, standardised classification of rates of visual field 268 progression we have adopted that used by Chauhan et al.²¹: fast progressors as <-1dB/year (-269 1dB/year is approximately ten times faster than age related decay). Although statistical methods 270 271 differ among studies, our estimates of global TD progression are broadly comparable with MD rates 272 in clinical glaucoma populations, which report median progression rates ranging from -0.62dB/year to -0.05dB/year)^{21,35,36}. For the formal comparisons of Medicine-1st vs. Laser-1st we reported the 273 proportion of eyes with moderate or fast progression, combining these categories to ensure 274 reasonable data support for each outcome. These figures are not directly comparable with the 275 number of VF progressions reported in the recent paper on the primary outcomes of LiGHT⁶, where 276 277 progression was detected using GPA. The proportions reported here are larger, possibly because 278 trend based methods are more sensitive for detecting progression than event based methods such as GPA¹⁵, especially given the relatively high upper threshold of the moderate/fast classification (-279 280 0.5dB/year). Also, this analysis covers a longer follow-up period, extending beyond the 36-month 281 point reported previously and so a larger proportion of eyes would be expected to show evidence of VF progression in our study. Despite these methodological differences, both analyses report higher 282 283 risks of VF progression in the Medicine-1st group, that may be related to the higher rates of disease 284 deterioration previously reported⁶.

This VF analysis is more detailed than those previously reported for LiGHT ^{6,14,37} in that pointwise 285 rates were modelled and then averaged to produce global rate estimates, retaining more 286 information than if global VF measures such as MD or Pattern Standard Deviation (PSD) had been 287 288 used. Furthermore, we considered the overall shapes of the progression rate distributions rather 289 than using the mean of each distribution as the single point of comparison. We show that 290 differences between treatment groups were manifest only towards the more rapidly progressing 291 end of the rate distribution. If we had concentrated solely on mean TD and PD we would have found no differences between treatment groups, consistent with the MD and PSD results reported at 36-292 months¹⁴. 293

The data derived for this study were drawn from a carefully conducted, randomised controlled trial. 294 Patients were monitored according to routine clinical care; the trial used eye specific IOP targets 295 296 which were objectively defined and adjusted by a computerised decision algorithm to avoid bias¹². 297 Similarly, to avoid bias in clinical decision making, treatment escalation decisions were initiated by 298 the computerised decision algorithm, which followed a robust protocol developed according to international guidelines by the EGS, American Academy of Ophthalmology Preferred Practice Pattern 299 and the and the South-East Asia Glaucoma Interest Group^{38–40}. The decision support algorithm used 300 in the LiGHT trial has been described in detail before ^{12,14}. The success of this strategy is highlighted 301 302 by the well matched distributions of baseline damage and IOP targets between treatment groups (Table 1, Figures 3 and 4). As a result, any differences in VF progression between treatment groups 303 reflect genuine change, in the presence of identical IOP control practices between the two groups. 304 Patients treated with Laser-1st exhibited slower VF progression, as shown in this study, in addition to 305 better IOP control, less intense medical and surgical treatment and lower rates of disease 306 deterioration⁶. 307

The data presented here support the use of SLT as a first line treatment for glaucoma and OHT as
 suggested by the previously reported improved clinical outcomes, lower treatment intensity and

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- 310 cost-savings for the NHS. With slower VF deterioration SLT may delay or completely avert the need
- 311 for more intense medical and surgical intervention in a significant proportion of patients.

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426 Figure legends

- 427 Figure 1. Distribution of estimated global total deviation progression rates by treatment group.
- 428 Histogram with median and 10th percentiles indicated. Curved line represents a smoothed density
- 429 estimate to the histogram.
- 430 Figure 2. Distribution of estimated global pattern deviation progression rates by treatment group.
- 431 Histogram with median and 10th percentiles indicated. Curved line represents a smoothed density
- 432 estimate to the histogram.
- 433 Figure 3. Distribution of mean deviation (MD) at baseline by estimated total deviation progression
- 434 rates.
- 435 Figure 4. Distribution of target IOP at baseline by estimated total deviation progression rates.







0.5 dB/y)

TD progression rate

dB/y)

dB/y)



Table 1. Distribution of cohort characteristics by treatment group. Values given are frequencies

unless otherwise marked.

	Medicine-1st	Laser-1st
Patients	344	344
Male	180 (52.3%)	193 (56.1%)
Female	164 (47.7%)	151 (43.9%)
Age in years, mean (SD)	62.9 (11.6)	63.4 (12.0)
OAG	271 (78.8%)	266 (77.3%)
OHT	73 (21.2%)	78 (22.7%)
Eyes	588	590
Bilateral cases	245 (71.2%)	249 (72.4%)
Follow up duration in months, median (IQR)	47 (39, 54)	49 (42, 56)
Visual fields	5630	5933
Visual fields per eye, median (IQR)	9 (8, 11)	10 (8, 12)
Interval between fields in days, median (IQR)	135 (83, 189)	140 (94, 189)
Visual field mean deviation at baseline in dB, median (IQR)	-2.0 (-4.5, -0.5)	-2.2 (-4.4, -0.6)
IOP target at baseline in mmHg, median (IQR)	18 (16, 21)	18 (16, 21)
Number of cataract removals performed	64	42

Table 2. Distribution of estimated total deviation progression rates by treatment group.

	Locations		Eyes	
Progression rate	Medicine-1st	Laser-1st	Medicine-1st	Laser-1st
Fast (-1 > slope dB/y)	10.2% (3115)	6.0% (1848)	9.5% (56)	5.4% (32)
Moderate (-1 <= slope < -0.5 dB/y)	15.9% (4864)	13.0% (3980)	16.7% (98)	11.5% (68)
Slow (-0.5 <= slope < 0 dB/y)	40.3% (12336)	43.4% (13311)	41.5% (244)	48.1% (284)
Slow improvement (0 <= slope < 0.5 dB/y)	25.7% (7863)	31.6% (9705)	25.5% (150)	29.7% (175)
Moderate improvement (0.5 <= slope < 1 dB/y)	5.9% (1798)	4.7% (1442)	5.1% (30)	4.1% (24)
Fast improvement (slope >= 1 dB/y)	2.0% (600)	1.3% (394)	1.7% (10)	1.2% (7)

Table 3. Distribution of estimated pattern deviation progression rates by treatment group.

	Locations		Eyes	
Progression rate	Medicine-1st	Laser-1st	Medicine-1st	Laser-1st
Fast (-1 > slope dB/y)	4.6% (1403)	3.2% (967)	3.4% (20)	1.7% (10)
Moderate (-1 <= slope < -0.5 dB/y)	6.9% (2103)	5.1% (1565)	6.5% (38)	5.4% (32)
Slow (-0.5 <= slope < 0 dB/y)	46.6% (14234)	48.9% (14990)	51.7% (304)	55.6% (328)
Slow improvement (0 <= slope < 0.5 dB/y)	38.9% (11900)	40.6% (12471)	36.2% (213)	36.1% (213)
Moderate improvement (0.5 <= slope < 1 dB/y)	2.6% (805)	1.8% (557)	2.2% (13)	1.0% (6)
Fast improvement (slope >= 1 dB/y)	0.4% (131)	0.4% (130)	- (0)	0.2% (1)