LIGHTSITE I: A Double-Masked, Randomized, Sham-Controlled Study with Photobiomodulation in Dry Age-Related Macular Degeneration Subjects



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0.40

0.35

0.30

0.25

0.20

0.15

0.05

0.00

-0.05

-0.10

-0 15

77 G

mprov

Baseline

ШO 0.10

Background

Age-related macular degeneration (AMD) is the leading contributor to vision loss and blindness in the developed world. There are no current treatments for the dry form except lifestyle modification and antioxidant vitamins. There is a growing body of evidence to support Photobiomodulation (PBM) in the 500-1000 nm spectrum, as a novel treatment for dry AMD. The purpose of the LIGHTSITE I study is to assess the functional and anatomical benefits of PBM using the LT-300 in a randomized, sham-controlled pilot study. We report on the interim analysis of measures of ETDRS best corrected visual acuity (BCVA) and Contrast Sensitivity (CS) as well as changes in retinal drusen volume and thickness and Quality of Life (QOL).

Methods

The LIGHTSITE I study enrolled 30 dry AMD subjects. Subjects were randomized (1:1) and received either PBM or sham treatment over 3-4 weeks with a second series 6 months from baseline (BL). Data are presented from the interim analysis out to 3 months following the initial series of PBM treatment. LT-300 uses a multi-wavelength treatment comprised of 590 nm, 670 nm and 850 nm applied to the subjects eyes for a total of 4-5 minutes per treatment per eye.

Statistical Analysis

The Sponsor and Investigators remain masked to individual treatment assignments and only group data is provided. Change from baseline is the preferred outcome metric and a linear mixed effects model by ranks was used for the statistical analysis. Some data was unavailable at the 2 or 3 month timepoints so group mean data may not reflect equal numbers at each visit. Data were compared to a previous study (TORPA II) that investigated PBM in subjects with dry AMD (Merry et al., Acta Ophthalmologica, 2016).

Table 1. Patient comparison between TORPA II and LIGHTSITE I studies.

	TORPA II	LIGHTSITE I
Patients (n)	24	30
Gender [female, male]	15, 9	18, 12
Total # eyes	42	46
Mean Age [range]	78 [66-95]	76 [52-90]
VA Letter Score	86	71.86 (Sham) 74 (PBM)

Results

Score

Letter

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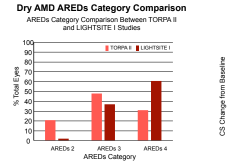


Figure 1. The AREDs classification for dry AMD subjects were compared between the TORPA II and LIGHTSITE I studies. Compared to TORPA II, LIGHTSITE I enrolled higher numbers of AREDs 4 subjects indicating more advanced stages of dry AMD.



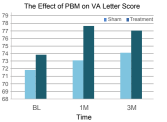


Figure 2. Subjects showed an increase in VA letter score out to 3 months following PBM from BL, (p< 0.05 paired t-test). A positive trend in VA change from BL in the PBM group versus the sham treatment group was seen, (Linear mixed effects model using ranks, p = 0.079)

The Effect of PBM on VA change from BL at 3M in the Low and High Vision Groups

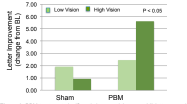


Figure 3. PBM treatment significantly improved the mean VA letter score in high vision (HV) subjects compared to sham at the 3M timepoint. Subjects ere divided into either high or low vision groups depending on whether their BL vision was above or below the mean BL VA score (~74) for each treatment group, (Linear mixed effects model using ranks, p<0.05). No significant VA letter score benefit was seen in the LV natients

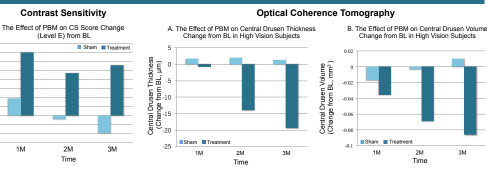
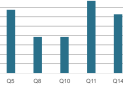


Figure 4. There was a statistically significant effect of PBM treatment on the CS score change of Level E (18 CPD) of analysis compared to BL, (Linear mixed effects model using ranks, p<0.05)

Visual Function Questionnaire-25 The Effect of PBM on VFQ Difficulty with Activities Scale



VEQ-25 Question

Figure 5. VFQ-25 overall composite score showed a statistically significant improvement with PBM (p = 0.003). Subjects showed a statistically significant improvement on Q8 and Q10 and 10-15% NS improvement in Q5, Q11 and Q14. Paired t-test, p < 0.05.

VFQ-25 questions [Difficulty with Daily Activity]

Q5: How much difficulty do you have reading ordinary print in newspapers?

Q8: How much difficulty do you have reading street signs or the names of stores?

Q10: Because of your eyesight, how much difficulty do you have noticing objects off to the side while you are walking along?

Q11: Because of your eyesight, how much difficulty do you have seeing how people react to things you say? Q14: Because of your eyesight, how much difficulty do you have going out to see movies, plays, or sports events?

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Figure 6. There was a statistically significant effect of PBM treatment on central drusen thickness (A) and central drusen volume (B) in the high vision subgroup compared to BL, (Linear mixed effects model using ranks, p<0.05). The development of drusen is a hallmark feature of dry AMD and significant reductions in drusen volume and thicknes demonstrate disease-modifying effects.



Summary & Conclusions

LIGHTSITE I Comparisons to TORPA II

· More dry AMD subjects with AREDs category 4 were enrolled in LIGHTSITE I Dry AMD subject's VA BL was more compromised in LIGHTSITE I versus TORPA II

LIGHTSITE I Conclusions:

Dry AMD patients treated with PBM demonstrated functional and anatomical improvements following PBM treatments. Over 42% of the treated dry AMD subjects had >1 line improvement in VA at 3 months. Moreover, patients classified as High Vision demonstrated enhanced PBM improvements compared to Low Vision patients suggesting PBM may be more effective in patients that are treated early. PBM improved contrast sensitivity and demonstrated reduced central drusen volume and thickness. Finally, guality of life measures were improved in subjects treated with PBM as determined by the VFQ-25 validated questionnaire. No device related adverse effects were seen. These LIGHTSITE I interim results support further clinical testing of PBM as a non-invasive treatment for dry AMD.