Change in objective ocular redness and symptoms after refit of weekly and monthly lenses into a water-gradient daily disposable lens material

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Introduction:
Healthy, white eyes are naturally desirable for most contact lens wearers. Self-perceived redness is often the only criterion by which wearers perceive their lenses as being suitable and their eyes “healthy” or not. Strategies and products aimed specifically at reducing ocular redness may therefore benefit ECPs. Contact lens-related hyperemia can be driven by many factors, including limited oxygen flux to the ocular surface, interpalpebral friction increase, tear film disruption, pH level changes and others [1]. Diurnal lens dehydration can steepen the lens curvature, reducing lens movement and tear film exchange, inducing conjunctival pressure marks, redness and ocular surface staining [2]. Some lens care solution ingredients can also contribute to hyperemia, through adhesion to the lens material and release on eye. Certain lens types or solutions (sometimes in combination) may be more prone to increasing hyperemia than others.

Purpose:
In this study, monthly and bi-weekly habitual CL wearers were refitted in a water gradient DD SiHy lens (deleflexon A) for four weeks of wear. Primary and secondary study objectives were the change in bulbar redness and limbal redness after refit, respectively. The exploratory endpoint related to subjective evaluation was wearing comfort.

Methods:
50 of 52 habitual, asymptomatic, full-time hydrogel lens and silicone hydrogel lens wearers (ratio 1:1) with spherical correction completed a prospective study conducted at two sites in Germany. All lens wearers used all-in-one lens care solutions (not Polyoquad-based or peroxide solutions).
Baseline measures included objective bulbar and limbal redness using the Oculus KSM Redness-Scan, and subjective comfort ratings and ocular surface staining grades. Comfort was assessed post-insertion and diurnally using a 101-point visual analog scale. Staining was measured using the 0-4-point JENVIS Grading Scale.
Participants underwent a wash-out phase of 6 ± 1 days of no lens wear before deleflexon A lenses were dispensed (at V2) for a wearing period of 26 ± 5 days. A follow-up visit (V3) was scheduled 6 ± 1 days after V2 and a final visit (V4) after an additional 20 ± 4 days. All measures were repeated at visits 2, 3 and 4. Redness and staining values for the right eye only were considered for analysis. Wilcoxon signed-rank test and t-test for paired samples were used for the statistical analyses.

Results:
Bulbar redness decreased in both subgroups over time. The mean score decreased from 0.88 to 0.77 (paired sample test p = 0.012) for previous habitual hydrogel wearers and from 0.83 to 0.71 (paired sample test p = 0.058) for previous silicone hydrogel 2/4w wearers (Figure 4).

Limbal redness decreased in both subgroups over time as well. The mean grade decreased from 0.41 to 0.23 (paired sample test p < 0.001) for previous habitual hydrogel wearers and from 0.43 to 0.30 (paired sample test p = 0.052) for previous silicone hydrogel 2/4w wearers (Figure 5).

Conclusion:
Although differences were small in the context of classic subjective grading scales, which use full grades in 1.0 steps, the Oculus KSM was able to detect fine graduated differences in a cohort of non-symptomatic wearers presenting with mild redness. The refit to a daily disposable deleflexon A lens after a wash-out phase was beneficial, with reduced bulbar and limbal redness, reduced corneal and conjunctival staining and increased wearer comfort. These benefits were slightly higher for the habitual hydrogel lens wearer subgroup.

References:
3. Sickenberger W, Dehnhard D: Validation of a novel morphing software to classify different slit lamp findings; Contact Lens & Anterior Eye 35(1), DOI: 10.1016/j.jcle.2012.08.005

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