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Clinical comparison of optimum and large diameter soft contact lenses

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ABSTRACT

Purpose: To compare the clinical performance of large diameter lenses with optimally fit lenses in the same material and moncurve back surface design.

Method: In a four-visit, randomised, bilateral, crossover, study, 25 myopic subjects wore optimum diameter lenses (control) and large diameter lenses (test) in random succession for 1 week each. Both study lenses were made of methafilcon A and of an identical design. Trial fittings with Frequency 55 (Cooperation) lenses modified with a design algorithm were used to determine the appropriate custom-made study lenses.

Results: The least squares mean scores (\pm SE) for overall comfort and end-of-day comfort (0–10 scale) were 7.57 ± 0.33 vs. 7.42 ± 0.33 ($P = 0.59$) and 7.00 ± 0.31 vs. 7.27 ± 0.32 ($P > 0.05$) for the optimum and large diameter lenses, respectively. There were no significant differences in mean (\pm SE) gradings for limbal hyperaemia (1.23 ± 0.11 vs. 1.19 ± 0.11 , 0–4 scale, $P = 0.60$) and corneal staining (1.79 ± 0.25 vs. 2.04 ± 0.25 , $P = 0.39$). Conjunctival staining was greater for the optimum lens: 1.80 ± 0.28 vs. 0.93 ± 0.28 (0–4 scale, $P = 0.001$). With regard to lens fit, the large diameter lenses showed significantly less post-blink movement (0.22 ± 0.01 vs. 0.16 ± 0.01 mm, $P = 0.004$), and greater total decentration (0.15 ± 0.02 vs. 0.21 ± 0.02 mm, $P = 0.010$). However, there was no significant difference in the key fit variable of tightness on push-up ($46 \pm 0.69\%$ vs. $48 \pm 0.69\%$, 0–100 scale, $P = 0.12$).

Discussion: The findings suggest that larger than optimal soft lenses may be worn without detriment to either comfort or ocular physiology, provided an optimal fit is otherwise maintained.

1. Introduction

Corneal diameter (CD) varies widely in a typical population, for instance, horizontal CD has been measured by ocular coherence tomography (OCT) to range from 12.1 to 14.4 mm [1]. The importance of the relationship between lens and corneal diameter is clinically accepted and textbooks typically suggest that lenses should overlap the limbus by at least 1–2 mm [2,3].

Lenses that are too small for a given eye cause irritation due to the edge encroaching onto the cornea. However, the clinical effects of lenses which are too large are uncertain and there has been little previous work in this area [4]. Theoretical calculations suggest that relatively large lenses can cause excess peripheral pressure [5]. Since many soft lens types are only available in a single diameter, it is inevitable that a significant proportion of lenses dispensed are larger than optimum. It would therefore be useful to have a better understanding of the impact of large diameter lenses on comfort and ocular physiology. The purpose of this study was to evaluate the clinical effect of relatively large diameter soft lenses compared with the effects of optimally fit lenses.

2. Method

This was a randomised, bilateral, unmasked, crossover, study that compared the clinical performance of optimally fit methafilcon A lenses with larger diameter lenses of the same moncurve design and material for 1 week each. The study was undertaken at two investigational sites in the United Kingdom (Aston University, Birmingham; Visioncare Research, Farnham) between January and May 2015.

Twenty-five subjects, aged between 18 and 70 years, were enrolled and dispensed with lenses. Subjects were required to have a spherical contact lens requirement in the range -0.50 to -6.00 D and astigmatism less than 1.50D in both eyes. Subjects were excluded if they demonstrated any signs of ocular infection, allergy, disease or corneal irregularity that could interfere with contact lens wear. Subjects were also excluded who had undergone corneal refractive surgery or any anterior segment surgery or had recently worn rigid contact lenses. Neophyte subjects were allowed, although most were existing soft contact lens wearers.

Both lens types were lathecut methafilcon A hydrogel lenses which

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Table 1
Lens Details.

	Trial Lenses	Controls	Test
Manufacturer	CooperVision	Ultravision	
Material	methafilcon A	methafilcon A	
Water content (%)	55	55	
Design	Frequency [®] 55	Custom manufactured, moncurve back surface, tricurve front surface	
Base curve (mm)	8.60	8.20 to 9.00 in 0.2 steps	
Diameter (mm)	14.2	13.5 to 16.0 in 0.1 steps	
Fitting	–	Optimal	Optimal diameter + 1.2 mm; optimal base curve + 0.6 mm
Sphere powers (D)	–0.50 to –6.00	–0.50 to –6.00	

Frequency[®] 55 lenses were used as trial lenses to determine the optimum diameter for a given subject by using photography to determine the limbal overlap.

were ordered following trial fitting with a cast moulded lens of the same material (Frequency[®] 55, CooperVision, Pleasanton, CA, USA). The lathecut lenses were custom manufactured to match the thickness and edge profile of the cast moulded lens (Ultravision CLPL, Leighton Buzzard, UK). The lens used for trial fitting was a single diameter and base curve design (Table 1) and, therefore, in order to select the optimum design for a given eye, an algorithm was used to: i) compensate for non-optimum tightness (i.e. tight or loose), ii) adjust for non-optimal lens diameters (Appendix A). For a lens fitting to be judged as optimum, it was required to cover the cornea in all directions of gaze, be central to the cornea with around 1.2 mm of conjunctival overlap, show sufficient post-blink movement with no edge stand-off, and to show optimal tightness by the push-up test [6,7]. The methods for assessing lens fit have previously been described [7].

Horizontal visible iris diameter was measured with a 0.1 mm increment graticule using a slit lamp biomicroscope and horizontal corneal diameter with an Anterior Segment Optical Coherence Tomographer (AS-OCT; Visante, Carl-Zeiss, Oberkochen Germany). Corneal topography was also conducted (E300, Medmont, Nunawading, VIC, Australia).

The large diameter lens was specified as being 1.2 mm larger in diameter than the optimal lens and 0.6 mm flatter in base curve so as to give a clinically equivalent fitting (e.g. Optimal lens = 8.6/14.2; Large diameter lens = 9.2/15.4) [5]. Since the lenses were custom made, the first pair was dispensed at a second visit at which the lens fit and visual performance were assessed and confirmed to be satisfactory.

Subjects were issued with the AOSept (Alcon, Fort Worth, TX, USA) hydrogen peroxide disinfection system. The use of saline for rinsing prior to insertion and rewetting drops was allowed, only if necessary.

A range of clinical variables was assessed at baseline and then reassessed at the 1-week follow-up visit (Table 2) with the subjects having worn the lenses for at least 2 h on those visit days. Slit lamp findings were graded with reference to the CCLRU grading scales [8]. For assessment of corneal staining, a yellow filter was used to enhance the appearance of any staining and this was graded for each of five corneal sectors. Similarly, for conjunctival staining, this was graded for each of four segments.

Lens comfort (insertion, during day and end-of-day) was graded by subjects on a 0–10 scale. Symptoms were monitored with the CLDEQ-8 questionnaire [9]. The CLDEQ-8 results were consolidated to produce a total score on a 0–33 scale. Subjects reported their typical insertion time and, if there was a reduction in comfort, the time that this typically occurred so that their comfortable wearing time could be determined.

Between follow-up visits, subjective comfort was monitored by SMS text messaging. Subjects were contacted four times a day (08:00, 12:00, 16:00, 20:00) on Days 2 and 6 of each lens wear period and asked to grade current lens comfort, also on a 0–10 scale. The SMS messages were pre-scheduled to be sent and received via an internet-based

Table 2
Summary of Clinical Assessments.

Comfort & Symptoms
Comfort (0–10, where 10 = cannot be felt)
CLDEQ-8 (0–33 scale, 0 = no problems)
Lens Fit
- Lens centration (mm, –ve value = inferior or temporal)
- Corneal coverage (Y/N)
- Post blink movement (mm)
- Primary-gaze lag (mm)
- Tightness on push-up (0–100, 50 = optimal, < 50 loose, > 50 tight)
- Overall fit acceptance (0–5, Grade 3–5 = acceptable)
Slit lamp Examination
- Limbal hyperaemia (0–4, 0.1 steps)
- Bulbar hyperaemia (0–4, 0.1 steps)
- Palpebral hyperaemia (0–4, 0.1 steps)
- Palpebral roughness (0–4, 0.1 steps)
- Corneal staining (0–4 in 5 sectors, i.e. 0–20)
- Conjunctival fluorescein staining (0–4 in 4 segments, i.e. 0–16)
- Conjunctival indentation (0–4)
- Other findings (0–4).

messaging service, FASTSMS (Worcestershire, UK, <http://www.fastsms.co.uk/>).

The study followed the tenets of the Declaration of Helsinki (2013). The protocol was reviewed by the Aston University Ethics Committee and a favourable opinion was received prior to undertaking the study. All subjects received detailed information about the study and signed an informed consent form before participation.

2.1. Statistical analysis

The statistical analysis was undertaken using SAS software Version 9.4 (SAS Institute, Cary, NC, USA). Four hypotheses were tested, specifically, that the following four variables would be significantly poorer with the large diameter lenses compared with the optimal lenses: overall comfort (at visit), end-of-day comfort, limbal hyperaemia, and conjunctival fluorescein staining. Each of these was tested using mixed linear models. The models included the following fixed effects: lens, order, visit, and site; and the random effect subject nested in site. Non-inferiority was concluded if the lower limit of the 95% confidence interval of the difference (test-control) was greater than X and superiority if the lower bound was greater than zero (X = –0.5, –1 and +0.5 for comfort, limbal hyperaemia and conjunctival staining, respectively). Due to the repeated measures study design, the recommended 15° of freedom could be achieved with at least 16 subjects completing the study [10]. Additional variables were tested for statistically significant differences using the mixed model analysis.

Table 3
Summary of Demographics and Ocular Topography (Medmont and AS-OCT).

Variable			
No. of Subjects/Eyes		25/50	
Age (years)	Mean (SD)	32.9 (15.9)	
	Range	18–60	
Sex	Male: Female	10:15	
Spectacle sphere (D)	Mean (SD)	−2.97 (1.07)	
	Range	−5.50 to −1.25	
Spectacle cylinder (D)	Mean (SD)	−0.43 (0.28)	
	Range	−1.0 to 0.00	
Cylinder axis (N eyes(%))	WTR	22 (44%)	
	ATR	14 (28%)	
	Oblique	14 (28%)	
Palpebral Aperture (mm)	Mean (SD)	–	10.12 (1.31)
	Range	–	8.0–16.0
Horizontal Visible	Mean (SD)	11.40 (0.31)	–
	Range	10.8–12.0	–
Iris Diameter (mm)	Mean (SD)	7.77 (0.20)	7.76 (0.20)
	Range	7.40–8.15	7.37–8.14
Corneal Apical Radius (mm)	Mean (SD)	0.46 (0.13)	0.76 (0.12)
	Range	0.09–0.65	0.40–1.00
Corneal Shape Factor	Mean (SD)	13.23 (0.54)	12.43 (0.51)
	Range	12.35–14.59	11.20–13.45
Corneal Sagittal Height (mm)	Mean (SD)	3.06 (0.25)	2.75 (0.21)
	Range	2.61–3.59	2.16–3.16
Corneo-scleral Junction Angle (°)	Mean (SD)	172.0 (2.5)	177.7 (2.2)
	Range	166–177	172–183
Corneo-scleral Junction Angle (°)	Mean (SD)	177.4 (1.6)	177.7 (1.8)
	Range	174–180	173–184

WRT = With The Rule ($180 \pm 20^\circ$); ATR = Against The Rule ($90 \pm 20^\circ$).

3. Results

The results are summarised in Tables 3–6 and the statistical analysis of key variables in Tables 8 and 9.

A total of 25 subjects were enrolled and successfully completed the study. The subjects' average age was 32.9 years (SD: 15.9, range: 18–60) and 60% (15/25) were female (Table 3). The mean sphere refractive error was 2.97 D (SD: 1.07, range: 1.25 D–5.50 D) and mean cylindrical refractive error was 0.43 D (SD: 0.28, range: Plano to −1.00 D). The mean horizontal visible iris diameter, as measured using a slit lamp graticule, was 11.40 mm (SD: 0.31, range 10.8–12.0) and mean palpebral aperture was 10.12 (SD: 1.31, range 8.0–16.0).

The mean horizontal corneal diameter, measured by AS-OCT, was 13.23 mm (SD: 0.54, range 12.4–14.6) and mean vertical corneal diameter was 12.43 mm (SD: 0.51, range 11.2–13.5). The mean corneal sagittal heights were 3.06 mm (SD: 0.24, range 2.61–3.59) and 2.75 mm (SD: 0.21, range 2.16–3.16) for the horizontal and vertical meridians, respectively.

Table 4
Summary of Subjective Assessments and Wearing Times.

Variable	Frequency 55	Dispensing		Follow-up		P-values	
		Large Diameter	Optimum Diameter	Large Diameter	Optimum Diameter		
No. of Subjects	25	25	25	25	25		
Overall Comfort (0–10)	Mean (SD)	8.86 (1.1)	9.00 (1.0)	8.95 (0.9)	7.31 (1.6)	7.46 (1.8)	0.59
	Range	6–10	7–10	7–10	5–10	3–10	
End-of-day Comfort (0–10)	Mean (SD)	–	–	–	7.27 (3.2)	7.00 (3.1)	0.55
	Range	–	–	–	2–10	4–10	
CLDEQ-8 (0–33)	Mean (SD)	–	–	–	9.16 (6.1)	8.64 (5.5)	0.65
	Range	–	–	–	2–25	2–23	
Average WT (hrs)	Mean (SD)	–	–	–	11.9 (2.4)	11.5 (2.6)	–
	Range	–	–	–	8–18	6–18	
Comfortable WT (hrs)	Mean (SD)	–	–	–	9.5 (3.4)	9.1 (3.7)	0.67
	Range	–	–	–	3–17	3–18	

3.1. Comfort

None of the assessments of overall comfort showed a significant difference and, therefore, the hypothesis, that subjective comfort would be significantly poorer with large diameter, was not met (Table 4). The least squares (LS) mean 1-week comfort scores were 7.42 and 7.57 (0–10 scale) for the large diameter and control lenses, respectively. The LS mean scores for end-of-day comfort were 7.27 and 7.00 (0–10 scale) for the large diameter and control lenses, respectively.

Overall, the comfort assessments by SMS also showed similar LS mean comfort scores: 7.60 vs. 7.73 (0–10 scale) for the large and optimal diameter lenses, respectively. When analysed by time point, two statistically significant differences were noted. Comfort was significantly better for the optimum diameter lens at the midday assessment on Day 2 (8.25 vs. 7.58, $P < 0.05$), however, the larger diameter lens was rated significantly higher at the evening assessment on Day 6 (7.52 vs. 6.76, $P < 0.05$) (Fig. 1). However, these findings must be treated with caution as they are based on only a proportion of the subject group; the overall response rate for the SMS assessments was 78.8%, and of those subjects 10.8% could not make an assessment because they were not wearing lenses at the time.

The mean comfortable wearing times reported at the follow-up visit were 9.7 and 9.4 h, for the large and optimal diameter lenses, respectively.

3.2. Symptoms: CLDEQ-8

The most frequently reported symptoms from the CLDEQ-8 questionnaire were ocular discomfort and dryness. A greater proportion of subjects reported experiencing frequent or constant discomfort while wearing the large diameter lens than for the optimum diameter lens (12 vs. 5, Fig. 2).

A similar proportion of subjects reported frequent or constant dryness with the large diameter lens compared to the optimum diameter lens (6 vs. 7, respectively).

3.3. Lens fit

The mean base curve and lens diameter dispensed were 8.57/14.15 mm for the optimal lenses and 9.17/15.35 mm, for the large diameter lenses (Table 5). The diameter of lens judged as optimum ranged from 13.6 to 14.8 mm. All of the lens fittings at dispensing were judged as acceptable by the investigators.

At the follow-up assessments, there were significant differences in lens fit with respect to centration, post-blink movement and overall lens fit acceptance.

Total decentration was calculated as the vector summation of horizontal and vertical centration. There was significantly greater total

Table 5
Summary of Lens Fit at Dispensing and Follow-up.

Variable	Frequency 55	Dispensing		Follow-up		P-value
		Large Diameter	Optimum Diameter	Large Diameter	Optimum Diameter	
No. of Eyes	50	50	50	50	50	
Tightness on Push-up (%)	Mean (SD) 44.8 (6.0)	48.6 (5.6)	48.7 (5.0)	47.3 (4.6)	45.9 (4.2)	0.12
Post-blink Movement (mm)	Range 38-	42-	43-	40-	35-	
Primary-gaze Lag (mm)	Mean (SD) 0.22 (0.1)	0.17 (0.1)	0.18 (0.1)	0.17 (0.1)	0.23 (0.1)	0.0038
Total Decentration (mm)	Range 0.1–0.6	0.1–0.3	0.1–0.4	0.0–0.4	0.0–0.5	
Horizontal Decentration (mm)	Mean (SD) 0.26 (0.1)	0.29 (0.2)	0.27 (0.1)	0.32 (0.3)	0.32 (0.1)	–
Vertical Decentration (mm)	Range 0.1–0.6	0.1–0.6	0.0–0.6	0.0–1.5	0.0–0.7	
Corneal Coverage (n eyes(%))	Mean (SD) 0.21 (0.1)	0.31 (0.2)	0.24 (0.1)	0.20 (0.2)	0.14 (0.1)	0.010
Diameter Acceptance (mm)	Range 0.0–0.6	0.1–0.8	0.1–0.5	0.0–0.5	0.0–0.3	
Overall Fit	Mean (SD) –0.08 (0.1)	–0.11 (0.2)	–0.06 (0.2)	–0.06 (0.2)	–0.04 (0.1)	0.45
Acceptance (0–5)	Range –0.3 to 0.3	–0.6 to 0.3	–0.3 to 0.3	–0.5 to 0.3	–0.3 to 0.3	
Fitting Success (n eyes(%))	Mean (SD) 0.04 (0.2)	–0.05 (0.3)	–0.05 (0.2)	–0.03 (0.2)	0.03 (0.1)	0.0044
Yes	Range –0.3 to 0.6	–0.6 to 0.4	–0.5 to 0.3	–0.5 to 0.4	–0.3 to 0.3	
No	50 (100%)	20 (100%)	20 (100%)	50 (100%)	50 (100%)	
0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)	
1.18 (0.11)	–0.01 (0.27)	0.01 (0.07)	0.01 (0.12)	0.01 (0.10)	0.01 (0.10)	< 0.0001
0.9 to 1.4	Range –0.6 to 0.7	0.9 to 1.4	–0.1 to 0.2	1.0 to 1.5	–0.2 to 0.3	
3.85 (0.56)	Mean (SD) 3.45 (0.42)	3.55 (0.43)	3.85 (0.56)	3.53 (0.61)	3.93 (0.56)	0.0005
3–4	Range 3.0–4.5	3–4	3–5	2–4.5	2–5	
20 (100%)	Yes	20 (100%)	20 (100%)	46 (92%)	48 (96%)	
0 (0%)	No	0 (0%)	0 (0%)	4 (8%) [†]	2 (4%) [†]	

* Insufficient movement on blink.

Table 6
Summary of Slit Lamp Findings.

Variable	Baseline	Large Diameter	Optimum Diameter	P-values
No. of Eyes	50	50	50	
Limbal Hyperaemia (0–4)	Mean (SD) 1.04 (0.51)	1.18 (0.53)	1.23 (0.52)	0.60
Min	0–2.4	0.1–2.2	0.2–2.3	
Bulbar Hyperaemia (0–4)	Mean (SD) 1.22 (0.43)	1.30 (0.57)	1.37 (0.60)	0.39
Range	0.4–2.5	0.3–2.8	0.2–2.7	
Upper Palpebral Hyperaemia (0–4)	Mean (SD) 1.16 (0.37)	1.29 (0.53)	1.30 (0.40)	–
Range	0.5–2.4	0–2.4	0.6–2.5	
Upper Palpebral Roughness (0–4)	Mean (SD) 1.00 (0.34)	0.83 (0.38)	1.02 (0.40)	–
Range	0.4–2.5	0.3–1.8	0.3–2.5	
Lower Palpebral Hyperaemia (0–4)	Mean (SD) 1.17 (0.43)	1.34 (0.59)	1.34 (0.57)	–
Range	0.4–2.4	0.4–2.6	0.2–2.5	
Lower Palpebral Roughness (0–4)	Mean (SD) 1.28 (0.56)	1.20 (0.49)	1.21 (0.51)	–
Range	0.2–2.8	0.3–2.3	0.5–2.6	
Corneal Staining Type–Total (0–20)	Mean (SD) 0.62 (1.10)	1.94 (1.75)	1.66 (1.66)	–
Range	0–4	0–7	0–7	
Conjunctival Staining–Total (0–16)	Mean (SD) 0.84 (1.81)	0.76 (1.41)	1.62 (2.11)	0.0006
Range	0–7	0–6	0–8	

Table 7
Summary of Study Contact Lens Parameters.

	Large Diameter	Optimum Diameter
No. of Eyes	50	50
Base Curve (mm)	Mean (SD) 9.17 (0.19)	8.57 (0.19)
Range	8.70–9.70	8.10–9.10
Diameter (mm)	Mean (SD) 15.35 (0.29)	14.15 (0.29)
Range	14.8–16.0	13.6–14.8
Back Vertex Power (D)	Mean (SD) –3.04 (1.02)	–3.04 (1.02)
Range	–1.25 to –5.25	–1.25 to –5.25
Centre Thickness* (µm)	Mean (SD) 93 (± 17)	92 (± 14)
Peripheral junction thickness* (µm)	Mean (SD) 164 (SD: ± 14)	157 (SD: ± 10)
Edge thickness* (µm)	Mean (SD) 150 (SD: ± 17)	142 (SD: ± 15)

* Lens thickness measurements taken from nine pairs optimum and large diameter lenses using Rehder gauge (n = 36).

decentration with the large diameter lens compared to the optimum lens, 0.21 vs. 0.15 mm ($P = 0.01$). There were also significant differences in vertical decentration and absolute horizontal decentration, -0.03 vs. $+0.03$ mm, ($P = 0.004$) and 0.14 vs. 0.07 mm, ($P = 0.002$) respectively, for the large and optimum diameter lenses. As expected, diameter acceptance was assessed as significantly greater for the large diameter lenses: 1.20 vs. 0.01 mm ($P < 0.0001$).

The large diameter lens showed significantly less post-blink movement than the optimum lens, 0.16 vs. 0.22 mm, ($P = 0.004$). There was no significant difference in lens tightness between the two lenses, 48% vs. 46% for the large and optimum diameter lenses, respectively ($P = 0.12$).

Despite the optimisation of fit, investigators rated overall fit acceptance significantly poorer for the large diameter lens compared with the optimum diameter lens, 3.48 vs. 3.88 , (0–5 scale, $P = 0.0005$). Six of the 100 lens fittings were judged as unacceptable due to insufficient movement on blink; four were large diameter lenses and two were optimum diameter lenses.

Table 8
Tests of Fixed Effects from the Analysis of Primary Variables.

Variable	Model Term	Numerator Degrees of Freedom	Denominator Degrees of Freedom	F-Value	P-Value
Comfort	Lens Type	1	23.0	0.29	0.5931
	Lens Order	1	22.0	0.75	0.3962
	Pair	1	23.0	0.11	0.7379
	Site	1	22.0	2.82	0.1073
Comfort (SMS)	Lens Type	1	242.8	1.07	0.3030
	Day	1	242.5	0.24	0.6223
	Time	3	242.0	10.08	< 0.0001
	Lens Order	1	21.1	0.02	0.8783
	Pair	1	242.7	0.92	0.3384
	Day × Type	1	241.8	1.72	0.1913
	Type × Time	3	242.0	2.26	0.0823
	Day × Time	3	241.7	0.52	0.6716
	Day × Type × Time	3	242.1	1.09	0.3550
	Site	1	21.0	2.99	0.0985
Limbal Hyperaemia	Lens Type	1	23.0	0.29	0.5971
	Lens Order	1	22.0	1.27	0.2724
	Pair	1	23.0	0.52	0.4762
	Site	1	22.0	0.00	0.9444
Conjunctival Staining	Lens Type	1	23.0	15.98	0.0006
	Lens Order	1	22.0	0.34	0.5651
	Pair	1	23.0	0.98	0.3333
	Site	1	22.0	10.66	0.0035

Table 9
Least Square Mean Differences Estimates and 95% Confidence Intervals for Primary Variables at the 1-Week Follow-Up Visit.

Variable	Difference	LS Mean Difference	Std. Err	95% CL	Non-Inferiority Met?	Superiority Met?
Comfort	Test-Control	-0.2	0.28	-0.7 to 0.4	No	No
SMS comfort- overall	Test-Control	-0.13	0.12	-0.4 to 0.1	Yes	No
Limbal Hyperaemia	Test-Control	-0.0	0.09	-0.2 to 0.1	Yes	No
Conjunctival Staining	Test-Control	-0.9	0.22	-1.3 to -0.4	Yes	Yes

LS-Means: least-square means, Std. Err: standard error, CL: confidence limits. Non-inferiority is established if the upper confidence limit is less than +0.5. Superiority is established if the upper confidence limit is less than 0.

3.4. Slit lamp findings

The slit lamp findings are summarised in Table 6. There was a significant difference between the optimum and large diameter lenses for conjunctival fluorescein staining; however, there were no significant differences for any of the other slit lamp variables.

The primary hypothesis, that limbal hyperaemia will be significantly greater with large diameter soft lenses compared with optimally fit lenses, was not met: 1.19 vs. 1.23 (0-4 scale) [Least Square Mean Difference (LSMD): 0.0, 95% CL: (-0.2, 0.1)]. The lower

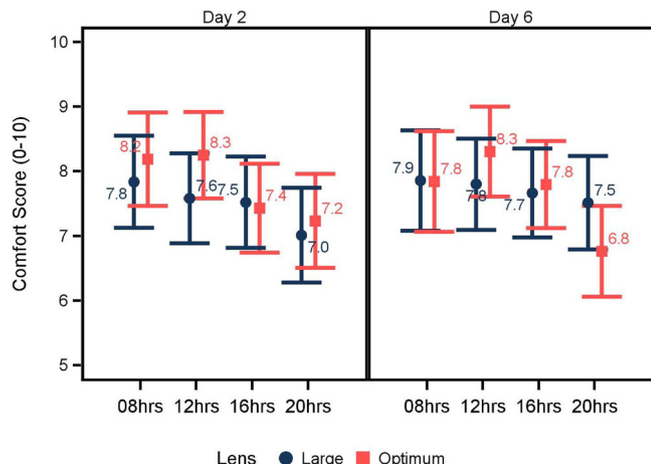


Fig. 1. Least square mean estimates for SMS comfort by day and time (and 95% confidence intervals).

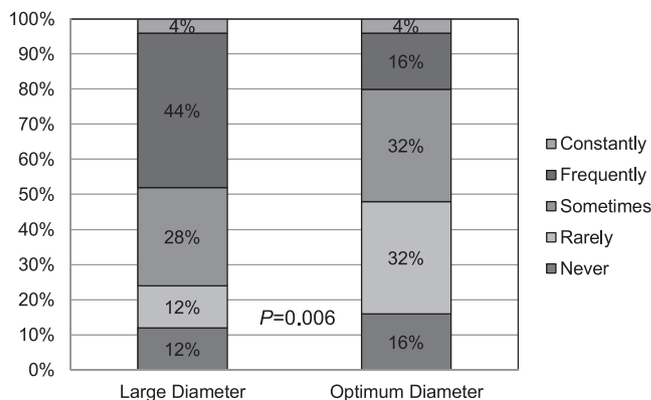


Fig. 2. Frequency of Eye Discomfort from CLDEQ-8 responses.

confidence interval is greater than the lower confidence bound. Hence it can be concluded that the large diameter and optimum lenses were equivalent with respect to limbal hyperaemia (Fig. 3).

The secondary hypothesis, that corneal staining will be significantly greater with large diameter soft lenses was not met: 2.04 vs. 1.79 (0-4 scale) [LSMD +0.3, 95% CL: (-1.3, -0.4)] (Fig. 3). A similar proportion of eyes showed corneal staining with the large and optimum diameter lenses (74% vs. 68%, respectively). Corneal staining type did not exceed > Grade 2 for either lens.

The primary hypothesis, that conjunctival fluorescein staining will be significantly greater with large diameter soft lenses was not met: 0.93 vs. 1.80. (0-4 scale) [LSMD: -0.9, 95% CL: (-1.3, -0.4)], P = 0.0006. Since the upper bound was less than zero but still greater

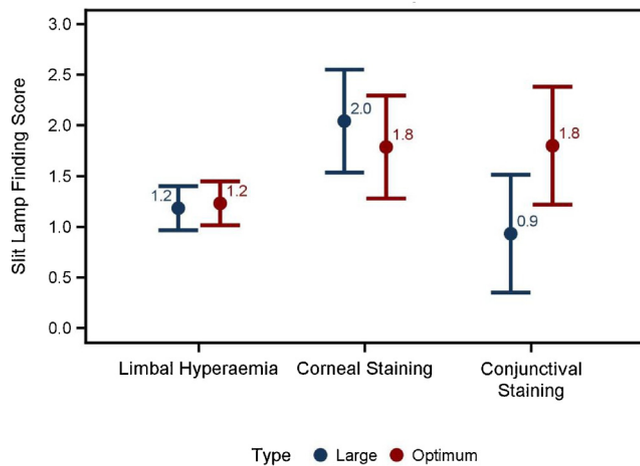


Fig. 3. Least square mean estimates for slit lamp findings at 1-week follow-up visit (and 95% confidence intervals).

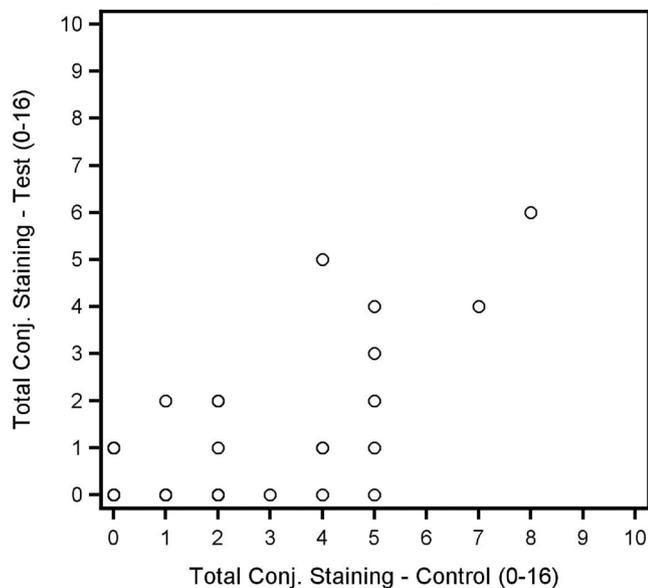


Fig. 4. Scatter plot of conjunctival staining at 1-week follow-up visit.

than the lower equivalence margin, conjunctival staining was therefore statistically significantly greater with the optimum lens than the larger diameter lens, although this was not clinically significant (Figs. 3 and 4).

In addition, a greater proportion of eyes showed conjunctival staining with the optimum lens than with the large diameter lenses (52% vs. 34%).

3.5. Lens metrology

Centre thickness, peripheral junction thickness and edge thicknesses were measured for a sample of lenses from nine subjects (Table 7). Thicknesses were measured using a Rehder thickness gauge (West Lafayette, IN, USA).

4. Discussion

4.1. Comfort

There were few differences in comfort ratings between the large and optimum diameter lenses. Interestingly, there was no significant

difference in either overall comfort or end-of-day comfort. These are unexpected findings given the increased interaction between the lids and lens edge as a result of the greater surface area of the larger lenses. The fact that there was no difference might, in part, be explained by the similar centre thickness and peripheral thickness for the two lens types. The fact that the lenses were identical material and all fitted to give optimum fit also reduces the risk of one lens type being less comfortable than the other [4].

Subjects did, however, report more frequent discomfort with the larger diameter lens, although there was no significant difference in the intensity of discomfort. Given that there was no difference in overall comfort, these findings would suggest that subjects experienced more frequent but transitory episodes of lens awareness.

4.2. Slit lamp findings

The only difference in ocular physiology was related to conjunctival staining, which was significantly greater for the optimum diameter lens than the large diameter lens. This is a surprising finding, as the pressure of the eyelids acting over a larger surface area might have been expected to produce greater mechanical interaction between the lens and conjunctiva. Two possible explanations for the greater conjunctival staining with the optimal design are: i) increased conjunctival exposure, and ii) greater lens movement.

Some mid-peripheral corneal staining (especially superior epithelial arcuate lesions [SEAL] or pre-SEAL staining), might have been expected with the larger diameter lens, but was not the case. It is likely that, in both instances, such staining may have been avoided by the lower modulus material employed in the manufacture of the study lenses.

A greater degree of limbal hyperaemia might have been expected with the larger lens as a result of greater mechanical interaction coupled with reduced oxygen supply. The fact that this was not the case may have been due to the fact that both lenses were fitted so as to give optimum tightness of fit. In relation to oxygen, although the larger lens covered a larger area of conjunctiva (15% difference between the diameters), the lens thicknesses were similar over the cornea and therefore supplied similar levels of oxygen to the cornea.

4.3. Lens fit

Post-blink movement was significantly less with the large diameter lenses, even though the larger surface area of the large lens might have been expected to encourage greater movement. However, increased surface area is also likely to increase friction between the lens and ocular surface which would discourage movement. On balance, this finding suggests that the latter effect predominates.

Despite the fact that the larger diameter lenses were optimised for fit, overall lens fit acceptability was still rated significantly poorer than for the optimum diameter lenses. This was partly due to the greater decentration seen with the larger lenses, most likely a result of the greater mass of the lens acting with gravity. Also, the reduced movement with the larger lenses resulted in four fittings being downgraded to unacceptable.

It is possible that differences in lens fit may have been evident if the large diameter lenses had not been optimised with respect to base curve; in other words, if the diameter had been increased without a compensating change to base curve. In particular, greater lens tightness might have been apparent due to the increased sagittal depth of the lens. This might also have resulted in greater peripheral pressure [5], leading to conjunctival indentation and increased conjunctival staining.

4.4. Optimum design

To the best of the authors' knowledge, this study was unique in selecting the optimum soft lens parameters to the nearest 0.2 mm and placing no limits on BC or diameter. It is notable that this led to the use

of a wide range of parameters. The range of optimum diameters was 1.2 mm, however, this was small in comparison with the range of horizontal corneal diameters (> 2.2 mm). Table 7 shows that a large proportion of the optimal lens designs selected were outside of the range of lenses typically offered. Although the present study suggests that the larger than optimal lens diameters should not be a concern, other compromises of lens fit may be problematic. Small lens diameters are known to cause discomfort [11]. In addition, a previous study has shown that relatively loose or tight fittings can lead to increased corneal staining and conjunctival hyperaemia [12].

4.5. Limitations of the study

Although objective ways of assessing soft lens fit have been developed, subjective evaluation is almost as repeatable, though the range of values is generally reduced [13]. Since there are currently no reliable objective methods for selecting an optimal soft lens design for a given

eye, a possible source of error is that this relied on the judgement of the investigator. However, the lens fit assessments with the final lenses suggest that this was relatively successful. All of the final optimal lenses were judged on-eye to be within 0.3 mm of optimal.

The larger lenses were optimised for tightness of fit whereas theoretical data suggest that, in a typical population, large diameter lenses tend to be tighter than optimum [5]. It is possible, therefore, that the results would be different when looking at large lenses coupled with a relatively tight fit.

In conclusion, this study has shown that larger than optimal soft lenses may be worn without detriment to comfort or ocular physiology provided an optimal fit is otherwise maintained.

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Appendix A

Lens selection guide.

Guide for selection of optimum lens parameters (BC/Diameter) based on Frequency[®] 55 trial lens.

Diameter Acceptance (mm)*		Loose	Optimum Fit	Tight
Large	+1.0	–	8.1/13.2	8.5/13.4
	+0.8	–	8.2/13.4	8.6/13.6
	+0.6	–	8.3/13.6	8.7/13.8
	+0.4	8.0/13.6	8.4/13.8	8.8/14.0
	+0.2	8.1/13.8	8.5/14.0	8.9/14.2
Optimum	0.0	8.2/14.0	8.6/14.2	9.0/14.4
Small	–0.2	8.3/14.2	8.7/14.4	9.1/14.6
	–0.4	8.4/14.4	8.8/14.6	9.2/14.8
	–0.6	8.5/14.6	8.9/14.8	–
	–0.8	8.6/14.8	9.0/15.0	–
	–1.0	8.7/15.0	9.1/15.2	–

For the Large diameter lens, add 1.2 mm to the diameter and flatten the base curve by

0.6 mm to give clinical equivalent; e.g. Optimal = 8.6/14.2; Large diameter = 9.2/15.4.

* +ve indicates larger than optimum for given cornea.

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