INTRODUCTION AND GOALS OF THE MATERIALS, DESIGN, AND CARE SUBCOMMITTEE

Examining the role of the contact lens material, design, and the care system is fundamental to understanding contact lens discomfort (CLD). However, a systematic review that tries to determine the governing factors is fraught with difficulties. A lack of a validated "instrument" (or single validated questionnaire) for measuring discomfort makes it impossible to compare between studies because reported levels of comfort (or discomfort) are inconsistent. Subject classifications can vary widely, from studies that include only neophytes or asymptomatic contact lens (CL) wearers to studies including only those contact lens–wearing subjects who experience marked dryness or symptoms of discomfort. Also, it is difficult to measure issues of importance in isolation because changing one factor in a contact lens or care solution can invariably affect another. An illustration of this relates to a change in hydrogel water content, which also affects oxygen permeability, oxygen transmissibility, modulus, and possibly lens thickness. Finally, various confounding factors between studies also make true comparisons problematic. Typical examples would include differences between brands of lenses made from the same material (which may have differing geometric designs, edge configuration, or production methods); wearing modality (lenses may be worn on a daily wear [DW] basis, overnight occasionally, or for up to 30 nights on a continuous wear [CW] basis); duration of use prior to replacement, wearing time during the day (from just a few hours to most of the day); and care product differences or exposures (which could range from no exposure in the case of daily disposable [DD] materials to a preserved system that has extensive uptake and release from the contact lens material being examined).

The purpose of this report is to summarize evidence-linking associations, mechanistic and etiological factors between contact lens materials, designs, and care solutions with CLD. The potential factors associated with this are many and varied, and graphically display the complexity of this issue.

Contact Lens Materials

Given the fact that approximately 90% of the world’s contact lens wearers are wearing soft lenses with no recent change in this figure,1 this report primarily concerns itself with the role of soft lens materials and designs and care solutions in CLD, with some discussion of rigid gas permeable lens (RGP) materials or designs where appropriate.

Conventional Hydrogel Materials

The pioneering work of Wichterle and colleagues2,3 is well known as a basis for the development of hydrogel polymers for soft contact lenses, including lightly cross-linked polymers of 2-
hydroxyethyl methacrylate (polyHEMA, 38% equilibrium water content [EWC]). Subsequent versions of polyHEMA-based materials with increased EWGs were made by copolymerizing it with both hydrophobic monomers (e.g., methyl methacrylate [MMA]) or other monomers of varying hydrophilicities (e.g., N-vinyl pyrrolidone [NVP]; methacrylic acid [MAA]).

Appendix A provides an overview of the characteristics of some commonly prescribed hydrogel materials. It was believed that a higher EWC would lead to a more wettable and comfortable lens (and increased oxygen transmissibility). However, it soon became apparent that dehydration is more pronounced with higher water hydrogels, particularly those with higher amounts of free water. This was sometimes associated with corneal desiccation staining and ultimately reduced end-of-day comfort.

In an attempt to enhance the biocompatibility of soft lenses, a novel material combining polyHEMA with a synthetic analogue of phosphorylcholine (PC) with water content of approximately 60% (omafilcon A) was developed in the early 1990s. The introduction of hizosilicon A in the late 1990s, where a nonionic copolymer of polyHEMA and glycerol methacrylate (GMA) were combined, was claimed to achieve excellent biomimicry by imitating the wetting properties of mucin. These approaches were designed in part to resist on-eye dehydration and deposition, although improvements in comfort were varied.

### Silicone Hydrogel Materials

Despite many attempts to harness the oxygen permeability of silicone rubber in contact lens materials, it was not until the late 1990s that two low-water content silicone hydrogel (SiHy) contact materials, lotrafilcon A (24% water) and balafilcon A (36% water), were released. The original intent for silicone hydrogels (due to their very high oxygen permeability) was for use as extended wear (EW) materials, but their use for daily wear has since become dominant (including their use as daily disposables). Silicone hydrogel development typically focused on compositions or macromers based on silicone-containing monomers (TRIS, siloxy macromer) that are sufficiently compatible with a range of hydrophilic monomers (including N,N-dimethyl acrylamide, NVP, polyHEMA).

Although the siloxane groups confer high oxygen permeability, they also give rise to inherent wettability issues, so several strategies have been employed to render SiHy surfaces more hydrophilic. Appendix B provides an overview of the characteristics of some SiHy materials.

### Bulk Properties of Soft Lens Materials

#### Water Content and Ionicity

Equilibrium water content and ionicity are used to classify lens materials by the Food and Drug Administration (FDA) and International Organization for Standardization, because of their impact on clinical performance. Although the relevance of such a grouping has been confirmed for such factors as dehydration and deposition, the relation with wearing comfort is less clear.

Nichols and Sinnott reported higher odds (odds ratio [OR]: 2.25) for CL-related dry eye in patients wearing high EWC lenses, but ionicity was not related with dryness symptoms. In a follow-up analysis, Nichols and colleagues showed that when compared with FDA Group I materials (the referent material), both FDA Groups II and IV were associated with a 2 to 3 times increased odds of contact lens dry eye. Further, in a small study with 10 subjects, Wilson and colleagues reported better comfort for patients fitted randomly with an FDA Group II lens (omofilcon A, non-ionic high EWC) compared with an FDA Group IV (etafilcon A, ionic high EWC) lens. Similar results were found by Guillon and colleagues when comparing the same materials in 22 patients in a crossover study wearing the lenses for 1 week in a random order. However, these studies ignore the effect of lens design, and the differences cannot be exclusively attributed to the different material properties.

Efron and colleagues compared the initial comfort of low (38%), medium (55%), and high (70%) EWC lenses and concluded that lower water content materials were more comfortable than higher EWC lenses in a nondispensing study where comfort was rated after 5 minutes of wear. This study excluded potential confounding factors such as edge design or surface finishing, as all lenses were lathe-cut in an identical design by the same manufacturer. Young also evaluated comfort in a study aiming to predict the success of fitting low (38%), medium (54%-58%), and high EWC (69%-74%) contact lenses. The results suggested improved comfort for low EWC lenses: First, the average comfort score was higher (8.4 for low, 8.2 for medium, and 8.2 for high EWC lenses). Second, low EWC flat-fitting lenses were significantly more comfortable than medium and high EWC lenses.

Silicone hydrogel materials have an additional confounding factor to the understanding of the potential role of EWC on comfort. The majority of these lenses have a low EWC, but they have substantially differing oxygen transmissibilities, modulus values, and surface wetting properties from traditional hydrogel materials. Dumbleton and colleagues conducted a study to evaluate the comfort of five different SiHy materials randomly fitted for 1-month periods, using a crossover design. All lenses generally performed similarly at the end of each period, although there was a slight difference for the ionic lens material to be associated with lower comfort at dispensing. Thus, the potential influence of material properties other than EWC or ionicity prevent any solid conclusions being drawn regarding the potential influence of these factors in SiHy material comfort.

In summary, several studies point to the increased comfort of low EWC lenses, with no direct impact of ionicity for conventional hydrogel materials. To date, no studies have been able to adequately draw any conclusions on the direct impact of these two factors for silicone hydrogels.

#### Oxygen Transmissibility

There has been a temptation to presume that oxygen transmissibility (Dk/t) is a key factor in contact lens comfort, and some of the circumstantial evidence and clinical dogma hints in this direction. Studies to determine the impact of oxygen transmissibility may be conducted either using lens materials of varying Dk/t or using sealed goggles in which the oxygen tension is varied. Millodot found reduced corneal sensitivity after exposing the cornea to hypoxic gaseous environments for up to 10 hours, following short-term wear of impermeable PMMA contact lenses and low Dk/t hydrogel lenses, and also cumulatively over years of wear of PMMA lenses. Contrary to the position that a greater supply of oxygen to the cornea might improve comfort, Millodot suggested that “a diminution of sensation with the wear of contact lenses is obviously beneficial as it helps the subject adapt more easily to the lenses.” To further this argument, the use of a topical anesthetic has been suggested as a means to assist adaptation to rigid contact lenses. Measurement of comfort while exposing the cornea to gaseous environments would be the obvious method of choice to discern the impact of hypoxia on comfort, but none of the studies in which this method has been employed has done so. It should be appreciated that many of the studies listed as evidence for or against an influence of Dk/t on comfort were not necessarily designed with that specific purpose in mind (Table 1). Highlighting the shortfalls in such study designs to meet this end does not necessarily mean that they do not
contain valuable information, but rather that using these studies to infer comfort is linked to oxygen supply is fraught with difficulties. For example, Brennan and colleagues\(^{44}\) and Malet and colleagues\(^{45}\) conducted open-label, multicenter, prospective, randomized studies with balafilcon A (\(\nu = 212\)) and lotrafilcon A (\(\nu = 134\)) SiHy lenses, respectively, worn in continuous wear (up to 29 nights without removal) with etafilcon A lenses as controls worn in extended wear (up to 6 nights). The Brennan study\(^{44}\) was contralateral eye and the Malet study\(^{45}\) was crossover. In both studies, the SiHy lenses compared with the hydrogels. However, the other common feature of these studies is the omission of a concurrent randomized, masked, control (e.g., “switching” back into a hydrogel lens) that would enable confirmation of a claimed improvement in comfort.

A multitude of studies have reported switching of existing hydrogel wearers out of their habitual lenses and into silicone hydrogels.\(^{57,48,49,51-53,55,57,61}\) Consistently, these studies have reported improvements in subjective response with the SiHy lenses compared with the hydrogels. However, the other common feature of these studies is the omission of a concurrent randomized, masked, control (e.g., “switching” back into a hydrogel lens) that would enable confirmation of a claimed improvement in comfort.

Six investigations that were randomized, partially controlled studies and were at least subject-masked have considered comfort differences between hydrogels and silicone hydrogels. Fonn and Dumbleton\(^{47}\) conducted a double-masked, contralateral, 7-hour, open-eye, nondispensing trial on 39 symptomatic and asymptomatic subjects. They found no difference between the hydrogel and SiHy lenses in comfort and dryness ratings. Cheung and colleagues\(^{43}\) conducted a prospective, double-masked, contralateral eye study in which they compared the comfort of two weekly replacement SiHy and hydrogel lenses in 33 subjects over 1 month of daily wear. They were unable to detect a significant difference in subjective comfort scores between lens material types. In the only extended wear trial of this group, Martin and colleagues\(^{66}\) measured comfort after 7 days of contralateral eye contact lens wear of a SiHy and hydrogel in 20 subjects. They found that the SiHy lens was more comfortable and led to less dryness than the hydrogel lens. In a single-center, double-masked, randomized, crossover, pilot clinical trial, Ousler and colleagues exposed 11 masked subjects to a controlled adverse environment for 75 minutes while wearing SiHy and habitual soft lenses.\(^{58}\) They found greater relief of subjective ocular discomfort associated with lens wear in adverse environmental conditions whilst wearing the SiHy. Ozkan and Papas,\(^{59}\) in a prospective, contralateral eye trial, compared comfort of a SiHy and hydrogel lens on 15 experienced lens wearers over 6 hours. Overall comfort was slightly (but significantly) higher for the low Dk hydrogel compared with the SiHy over this short time frame. Recently, Maissa and colleagues\(^{64}\) compared the comfort of four silicone hydrogels and one hydrogel in a prospective, crossover, double-masked, 10 day, daily wear trial. In rank order of comfort, the hydrogel was scored highest by...
the subjects and was statistically superior in comfort to one of the SiHy lenses at both the beginning and end of day. The controls in each of these six prospective, randomized, subject-masked studies are inadequate to test whether Dk/t alone is linked to lens comfort, as properties other than Dk/t that may affect comfort, such as surface properties and edge design, vary between the SiHy and hydrogel lenses under test. Nonetheless, the experimental designs are “more robust” than those other studies listed above, where subjects were swapped out of their habitual lenses to test lenses alone or where masking was inadequate. Interestingly, and in contrast to those studies, four of these better-executed studies did not find that SiHy lenses were superior in comfort to hydrogel lenses and indeed, in two of these studies, the hydrogel was more comfortable. Study design differences should be kept in mind when reconciling this apparent discrepancy. Overall, hydrogels seem to produce a more favorable comfort response in daily wear and shorter-term studies. During open eye wear, hydrogel lenses have sufficient Dk/t to provide near normal oxygen supply; at least to the central cornea.67 The impact of the lower Dk/t of hydrogels will be exaggerated during eye closure, or extended or continuous wear. Importantly, the study by Martin and colleagues66 was under extended wear conditions and this may partially explain the difference in results between that and the other “more robust” designs. A further confounding factor is study duration. The longest time of follow-up for the randomized, subject-masked studies was 30 days, where the “inferior” study designs saw patients followed, in some cases, for 3 years.

In recent years, four large cross-sectional studies have compared comfort between SiHy and hydrogel lenses. As noted above, Ramamooorthy and colleagues32 presented detailed statistical analysis of a cross-sectional and nested case-control study of 360 participants. The authors found FDA material classification to be a strong predictor of contact lens-related dry eye classification. Silicone hydrogel lens wear was found to be significantly protective from dryness symptoms in a univariate regression, but dropped out in the final multivariate model with FDA Group. The authors suggested that this could be because silicone hydrogels are low in EWC and correlate with FDA grouping. This finding may be highly relevant in discussions on the influence of Dk/t on comfort, as it points to the confounding effects of other material and lens properties. Chalmers and colleagues60 report on the analysis of a baseline self-administered questionnaire completed by 882 contact lens wearers comprised of 699 wearing hydrogel and 183 SiHy lenses at 84 clinical sites. Diagnosis of dry eye increased with age in the hydrogel wearers from 10.6% (at 18–24 years) to 21.1% (at 30–35 years), while it remained steady at 19% in the SiHy wearers. While not precluding selection or survival bias, this study suggests that wearers of hydrogel materials find their lenses at least the equal of silicone hydrogels (with regard to symptoms of dryness) up to 30 years of age. Young and colleagues were involved in two prospective, multicenter, nonrandomized, cross-sectional, observational studies of soft contact lens-related dryness that were partially controlled for Dk/t and where subjects responded to a questionnaire based on the Contact Lens Dry Eye Questionnaire (CLDEQ), from which they were classified as either having dry eye or not. In the first study of 932 daily wearers of soft contact lenses, the proportion of subjects scoring positive for contact lens dry eye did not differ by lens material (hydrogel versus SiHy).63 There were, however, differences in specific questions of typical and end-of-day comfort where the hydrogel performed less favorably. In their follow-up paper, they identified 226 soft contact lens wearers with self-reported contact lens-related dryness as classified by a questionnaire and 48 asymptomatic control subjects.65 This study showed that participants with and without symptoms of CL-related dryness did not differ by SiHy lens use. Interpretation of these cross-sectional studies must be made in the context of aforementioned limitations, but overall, the studies provide somewhat equivocal results. Collectively, the above studies cast doubt on the suggestion that the higher Dk/t is associated with greater comfort. Therefore, it is not surprising that studies in which different brands of SiHy lenses are compared have been unable to discern a greater comfort response for those lenses with higher Dk/t. Morgan and Efron conducted a single-center, randomized, subject-masked, crossover study on 30 subjects wearing two brands of SiHy lenses each for 8 weeks.68 The brand with the higher Dk/t did not show superior comfort to the brand with the lower Dk/t. Santodomingo and colleagues considered the same two brands used in both daily wear and continuous wear in a total of 45 subjects for 18 months.69 Again, they found no superiority of comfort in the lens with higher Dk/t. Brennan and colleagues followed 45 subjects in a prospective, partially controlled randomized, partially masked contralateral eye study of three different SiHy lens brands for a period of 12 months.70 Again, the lens with the highest Dk/t was not found to provide superior comfort.

While it may be appealing to attribute measured comfort benefits to the higher oxygen transmissibility of the SiHy lenses, a host of other confounding factors will drive the comfort response. Inadequacies of the control lens in experimental studies, including lens design, modulus, surface characteristics, and the modality of wear (daily disposable, reusable frequent replacement, flexible wear, extended wear, continuous wear) rather than Dk/t per se, may be responsible for the outcomes achieved. Duration of wear, where adaptation to the higher modulus lenses takes place, might interact with the lens material effect. Other experimental biases in the form of selection, novelty, halo, Hawthorne, survival, or similar effects resulting from study designs that are inadequate to randomize or mask the control product may also influence the outcome.

In summary, there have been no Level I evidence studies that can provide an answer to the question of whether oxygen levels influence comfort. What can be said is the following:

1. Where lenses of higher Dk/t are found to be more comfortable than lenses of lower Dk/t, there are deficiencies in experimental design or inadequacies in the control lenses that prevent definite attribution of such differences to oxygen.
2. There are circumstances where lenses of lower Dk/t have been found to be more comfortable than lenses with higher Dk/t; therefore, any effect that oxygen may have on comfort is being overshadowed by other factors or there simply may be no or a converse relation.
3. Where comfort differences between higher and lower Dk/t lenses are found to be statistically insignificant, the method used to measure comfort may not be sufficiently sensitive to detect differences.

**Modulus and Mechanical Factors.** The two most important quantifiable mechanical properties are tear strength (elongation at break) and modulus, which can be measured in stretching (tensile or elastic) or compression (rigidity) mode. While modulus is a specific material parameter, the effective “stiffness” of a contact lens will also be influenced by its specific geometry (lens thickness profile) as a thick lens made from a low modulus material may still be considered relatively inflexible or stiff. A thinner lens made from a low modulus material will drape over the cornea, distributing itself evenly on the ocular surface with minimum lid interaction. In some
instances, an increase in stiffness will help mask corneal astigmatism but possibly at the expense of initial comfort. Although the rigidity modulus has historically been useful for RGP materials, it is the tensile modulus that has primarily been most often quoted for soft lens materials.

The first generation silicone hydrogels (lotrafilcon A, balafilcon A) had tensile moduli that were significantly greater than most conventional hydrogels,22,25,71 such that for some wearers a comfort or wearer adaptation period was needed and there was an increased potential for mechanically induced ocular complications.19,21,73,75 Subsequent SiHy development has progressed to lower modulus materials through chemical structure modification and/or increased EWC.4,19,21–23,25,73,74

The higher modulus of SiHy materials was initially seen as an issue when refitting hydrogel wearers into these more oxygen permeable materials. However, when Riley and colleagues49 refitted 257 patients wearing hydrogel materials with a SiHy with a relatively low modulus (senofilcon A), they reported that 50% of subjects reported no contact lens–related discomfort. Most of the studies reporting refitting hydrogel wearers into silicone hydrogels report similar or higher levels of success, even when materials with a high modulus are employed.51 However, as noted recently by Guillon,76 the study design may partly explain these findings, as most refitting studies lack a concurrent control group or adequate masking. When comparing the ability of material to predict contact lens dry eye, Ramamoorthy and colleagues32 were unable to show any difference between the 11 (or more) individual materials being compared, including at least two SiHy materials. In the few studies reported where study bias was minimized by using a control group using low modulus hydrogel lenses, no differences in comfort between hydrogel and silicone hydrogels could be identified or attributed to modulus.47,54

Dehydration. Subjective reports of “dryness” and “discomfort” are well recognized as the main factors for contact lens discontinuation77,78 and has remained unchanged over the last decade, regardless of the new lens materials introduced.79 This has led to an intuitive relationship being proposed between soft lens dehydration and discomfort, particularly at the end of the day. A connection between dehydration and discomfort seems plausible given: (1) the potential correlation between lens thickness and desiccation staining;80 (2) the potential correlation between corneal staining and discomfort;81,82 and (3) the increased friction presumably induced by dehydrated, dry lens surfaces.83 However, a proven relationship between dehydration and discomfort has been supported by relatively few studies.15,84,85 This is perhaps not surprising, given the difficulties in evaluating material dehydration and types of dehydration (e.g., initial temperature–induced dehydration followed by evaporative dehydration).86–88 It is this latter dehydration that is potentially problematic, as it produces a water gradient99,80 that draws water through the lens and, ultimately, results in corneal desiccation staining.79,91,92 Evaporative dehydration tends to be localized and therefore may result in only a small change in a given lens’ overall water content. Likewise, evaporative dehydration may be less apparent with higher power lenses and, therefore, may be even more difficult to monitor in a subject group of varying prescription.

In addition to patient and environmental factors, differences in dehydration do exist between materials. A number of in vitro studies have shown that bulk water loss is closely related to initial EWC, with low EWC lenses (including silicone hydrogels) dehydrating less than higher EWC hydrogels.6,93–96 In studies that have evaluated lens dehydration13,15,12,93,99,100 and also recorded comfort ratings, a significant relationship between the two has not been consistently shown.

Hall and colleagues11 fitted four contact lens materials to 10 subjects and recorded dehydration and comfort after 4, 8, and 12 hours. At the 12-hour time-point there was a moderate negative correlation between comfort and dehydration for etafilcon A lenses ($r = -0.64, P = 0.04$), but no correlation for the remaining three materials.13 In a study in which omafilcon A was shown to dehydrate significantly less than other lenses of similar EWC (etafilcon A), Lemp and colleagues103 concluded from their 76-subject crossover study that the increased comfort found with the omafilcon A lenses was related to decreased on-eye dehydration.

In contrast with the work by Hall and Lemp, Fonn and colleagues102 found no correlation, either in symptomatic ($r = 0.33, P > 0.05$) or asymptomatic subjects, between the change in lens water content for omafilcon A and etafilcon A and change in comfort over 7 hours of lens wear in a contralateral, double-masked, nondispensing study. Maldonado-Codina and Efron107 conducted a crossover study with 34 subjects to evaluate the impact of manufacturing technology and material composition on the clinical performance of five hydrogel lenses worn for 1 month each. Despite a significantly higher dehydration of the ionic (FDA Group IV) material after 6 hours and after 1 month of lens wear, there was no significant difference in overall comfort between lens types. Lastly, in perhaps the largest analyses of the relationship between material dehydration and comfort, Nichols and Sinnott10 and Ramamoorthy and colleagues106 showed that while indeed higher EWC hydrogel lenses tend to dehydrate to a greater degree than lower water lenses, the degree of dehydration was not associated with contact lens dry eye classification of the subjects.

In conclusion, considering the body of literature available, including several well-designed studies that attempted to address this topic, it is not likely that a causative or associative relationship exists between on-eye bulk dehydration of materials and discomfort using the current methods used to capture either dehydration or subjective comfort.

Surface Properties of Soft Lens Materials

Friction and Lubricity. Lubrication, which can be defined as any means capable of controlling friction and wear of interacting surfaces in relative motion, provides defense against wear (the loss of material from interacting surfaces in relative motion usually related to friction). Materials with low friction and low wear are thought of as being well lubricated, or having good lubricity.

Friction coefficient measurements are most often made as an indicator of the quality of lubrication or lubricity, since wear measurements of biological surfaces are challenging. A friction coefficient is the ratio of the frictional force between two contacting surfaces in relative motion to the normal force between those surfaces. A variety of in vitro test setups with different test characteristics (scale, geometry, counter surface) and parameters (protocol, environment, lubricant, lens condition) have been used to assess friction coefficients of contact lenses. While each in vitro test setup has advantages and disadvantages, it remains unclear which, if any, is representative of in vivo function and/or friction and there are no standards on the techniques as such.

Several contact lens friction studies exist in the peer-reviewed literature.53,108–115 Collectively, these studies demon-strate that friction associated with contact lenses is a challenging field of study, and support the notion that reported friction coefficients must always be considered in the context of the experimental parameters in which they were measured, which is outside the scope of this report. It is also important to note that while this is an expanding area of scientific interest,
design—is common to the various lenses of that manufacturer and that one of these is the defining determinant of comfort. 

**Wettability.** The term “wettability” is traditionally used to describe the tendency for a liquid to spread over a solid surface and consequently has been widely adopted by the contact lens industry to describe the ability of the tear film to spread and remain on the surface of a contact lens. When a lens is applied to the eye, it fundamentally disrupts the normal tear film structure and physiology in a number of ways, including increasing the evaporation rate and decreasing tear film stability. The quality of the tear film over a lens is thought to play a key role in the lubrication of the lens/ocular surface system and will ultimately influence how much friction and “wear and tear” will result.

Despite widespread use of the term, no physical measurement exists that can completely quantify wetting. Notwithstanding this limitation, a number of different laboratory (in vitro) and clinical (in vivo) techniques have been adopted to investigate the wetting properties of contact lens surfaces, details of which are outside the scope of this report. Wettability is thought to be important for all types of contact lenses, but in particular for silicone hydrogels, which tend to be more hydrophobic compared with their conventional hydrogel counterparts, at least in the laboratory.

**In Vitro Wettability.** In vitro investigations of wettability have provided us with a wealth of information about lens surfaces and what factors affect them in the laboratory. Overwhelmingly, reports in the literature document the investigation of soft lenses, particularly in recent years. Those that do investigate rigid lenses have shown that the contact angles obtained are significantly affected by the methodology and since no recent reports exist that have used more current automated techniques, it is difficult to make any kind of meaningful comparisons with soft lenses. Studies have reported contact angles for unworn lenses in water or saline or other components and have shown that angles obtained for the same lenses can vary due to the differences in methodology or experimental conditions. Despite all of these data, none has been able to show any relationship between in vitro measurements and on-eye clinical wetting and, further, whether these laboratory measurements are in any way related to comfort. For example, both Nichols and colleagues and Thai and colleagues investigated the effect of adding hydroxypropyl methylcellulose (HPMC) to a multipurpose contact lens solution and, despite differences in in vitro or in vivo wettability and tear film thickness, there was no overwhelming preference for either care solution.

**Ex Vitro Wettability.** In an attempt to make in vitro measures of contact angle more relevant, researchers have attempted to perform contact angle analysis on lenses postremoval, but there are surprisingly few publications that have measured the wettability of ex vivo contact lenses and related it to comfort. Tonge and colleagues measured dynamic contact angles of etafilcon A lenses after various periods of wear in lenses that had been presoaked in either saline or a surfactant; the surfactant-exposed lenses showed significantly lower-advancing contact angles than the saline-treated lenses (however, there was no statistically significant difference for the receding contact angle between the two treatments). Of particular note in the work was that comfort was reported as being better for the surfactant-soaked lenses compared with the saline-soaked lenses at all time points measured, although only six subjects were included. There appear to be no other studies that have related ex vivo wettability to comfort associated with contact lens wear.

**In Vivo Wettability.** In vivo wettability has been investigated using a range of relatively simple slit-lamp-based procedures...
and grading scales, in addition to more indirect techniques such as the prelens noninvasive tear breakup time (NITBUT), tear thinning time, and techniques based around specular reflection.

One investigation compared comfort and NITBUT in nelfilcon A and nelfilcon A AquaRelease lenses. The authors reported that subjective ratings of comfort over a 16-hour period were consistently higher for the eye wearing the AquaRelease compared with the eye wearing the conventional nelfilcon A lens. NITBUT was greater with the AquaRelease compared with the eye wearing the conventional nelfilcon A lenses. In multivariate modeling, Nichols and Sinnott showed that prelens tear film thinning time was highly predictive of contact lens dry eye status, even when including EWC, osmolality, and lipid layer thickness (both significant themselves) in the multivariate statistical model.

Conclusive evidence that laboratory measures of contact angle can predict the wetting performance of a contact lens on-eye is lacking. Furthermore, the link between clinical measures of wettability and contact lens comfort remains not understood, with some evidence that surrogate measures do show a relation. It is likely that the assessment of wettability provides us with an indirect method of investigating the lubrication present in the lens/eye “system” and conclusive results across numerous studies have eluded us because the techniques we have employed to probe the tear film do not accurately reflect its complex and dynamic nature.

**Wetting Agent Incorporation**

The wetting agents discussed in this section of the review are limited to agents that are releasable and incorporated into contact lenses. Wetting agents in multipurpose solutions or contact lens packaging solutions will be addressed in another section. Wetting agents may be firmly embedded and provide enhanced wettability due to the materials being exposed at the lens interface or may be progressively released from the material over the course of the day.

**Polyvinyl Alcohol.** The nelfilcon A material is a polyvinyl alcohol (PVA)-based hydrogel specifically developed for use in a daily disposable lens. Maissa and colleagues suggested that the comfort level achieved with this lens “may result from a slow release of some residual entangled PVA” from the cross-linked PVA lens matrix. Using an in vitro release model, Tighe and colleagues suggested that the mechanical effect of the eyelid greatly accelerates soluble PVA release from the lens surface, which implies that the release is mechanically triggered or “blink activated” when placed on the eye. The next iteration of this material exploited this effect by intentionally adding nonfunctionalized PVA of appropriate molecular weight to enhance the elution of PVA, thereby increasing the comfort of these lenses. It was demonstrated in a contralateral eye study that adding this nonfunctionalized PVA enhanced tear stability and subjective comfort over a 16-hour wearing period relative to the original nelfilcon A product. A further enhancement incorporated an optimized blend of nonfunctionalized PVA in the lens matrix coupled with hydroxypropyl methylcellulose (HPMC) and polyethylene glycol in the packaging saline. Tear film stability was significantly greater with DACP than with its predecessor and was comparable to tear film stability without lenses. However, comfort data were not reported in this study, which brings into question whether there was any subjective improvement in reported in-eye comfort.

**Hyaluronic Acid.** Hyaluronic acid (HA), a hydrophilic glycosaminoglycan found throughout the human body, has been used in contact lens rewetting drops and in a range of artificial tear products to treat mild, moderate, and severe dry eye and has been used as a novel internal wetting agent for contact lens materials. However, to our knowledge, HA has not been shown in any clinical studies to directly improve comfort associated with contact lens wear.

**Comparison Between Rigid and Soft Lens Materials**

At first glance, there is a considerable difference between the comfort associated with rigid and soft lenses. However, while this is true in the short term, there is little evidence that medium and long-term comfort is substantially different between them. Fonn and colleagues found no significant differences in ratings of comfort after 6 months between eyes of 27 patients fitted contralaterally with a soft and a rigid lens. However, average comfort was significantly lower for the eye wearing the rigid lens over the initial 3-month period. For the 16 patients who remained in the study for an additional period of 3 months, comfort between both eyes was reduced but remained only marginally lower for the rigid lens-wearing eye.

Morgan and colleagues were unable to find a difference in comfort between a group of adapted rigid lens wearers and soft lens wearers using lenses on a continuous wear basis. Maldonado-Codina and colleagues compared the comfort scores reported by subjects successfully wearing rigid or soft lenses on a daily wear basis with comfort reported by neophytes fitted with high oxygen transmission rigid lenses and silicone hydrogels on a daily wear basis for 2 weeks, followed by 11.5 months of continuous wear. Their results showed that, while neophytes in the SiHy group presented with a high comfort score from the very beginning, the rigid lenses group reported significantly improved comfort scores over the first 2 weeks, remaining at the same level as the silicone hydrogels over the 12 months in continuous wear. Subjects who were experienced rigid lens wearers actually reported the highest comfort levels of all wearers, suggesting that long-term rigid lens wearers may ultimately be the “most comfortable” of all lens wearers.

Finally, Nichols and Sinnott and Ramamoorthy and colleagues used a variety of statistical modeling approaches in a cross-sectional sample of 360 contact lens-wearing subjects to evaluate rigid lenses, compared with soft lens wearers, in predicting contact lens dry eye. Similar to other studies, rigid lens wear was not associated with a difference in predicting classification of subjects with or without contact lens dry eye.

In summary, there is little published evidence of a significant difference in the reported comfort between soft and rigid lenses in the long term, once the initial adaptation phase is complete. However, clinicians are aware of the fact that many RGP-wearing patients report increased comfort when they are switched into a soft lens, so this lack of evidence may relate more to the fact that such a study has not been conducted.

**Lens Design and Fit**

**Soft Lens Design and Fit.** The fact that soft contact lens fit can affect contact lens wearing comfort is supported by the practical experience of every contact lens practitioner. Moreover, it seems logical that a soft lens showing excessive movement or failing to cover the cornea will cause irritation through interaction between the cornea and edge of the lens. Nevertheless, few clinical studies have shed light on correlations between the subtleties of soft lens fit and comfort responses. The reasons for this are probably 2-fold: first,
although many clinical trials have identified differences in comfort between lens types, only a few have systematically varied design parameters. Many clinical studies compare lenses of differing material as well as design. However, those studies that have sought to systematically evaluate lens design have tended to involve relatively low EWC designs and thin materials. Second, since so many factors affect contact lens–wearing comfort, it is difficult to control for all potentially confounding factors (e.g., modulus, thickness, and edge profile).

**Corneal Coverage and Lens Diameter.** An essential requirement of successful rigid contact lens fit is for the edge to stay clear of the cornea in all positions. Due to their inherent flexibility, the reverse is true with soft lenses. Since the lens edge rests against the eye and, due to stretching, exerts pressure, it has to overlap the relatively sensitive cornea to avoid discomfort. In order to allow for blink-induced movement, an overlap of at least 0.5 mm is assumed to be necessary. However, there appears to have been no work to confirm the optimum amount of overlap. A potential source of confusion is the fact that the true cornea is >1 mm larger than the visible iris diameter (“white-to-white diameter”). Furthermore, the limbal transition zone varies between individuals and, thus, visible iris diameter is a poor predictor of corneal diameter.

The pioneers of soft lens design assumed that the smallest acceptable lens diameter was optimal. But this was soon reassessed and early 12.5- and 13.5-mm designs were replaced by larger lenses. Early one study found that 13.5-mm diameter lenses showed full corneal coverage in approximately 50% of eyes, while 12.5-mm lenses gave total coverage in less than one-third of eyes. One study of tear replacement found an interrelation between lens diameter, comfort, lens movement, and “tear replenishment,” with smaller diameter lenses giving reduced comfort, greater movement, and faster tear replenishment. However, these apparent interrelations are confounded by the fact that the lens diameters were relatively small (12.0–13.5 mm) and that their closeness to the typical corneal diameter is likely to have impacted comfort.

Most current soft contact lenses fall within the diameter range 13.8 to 14.2 mm. However, since all lenses shrink when raised from room temperature to eye temperature, the diameter on-eye is quite different, being 13.54 to 14.16 mm in one study. It is possible, therefore, that some lenses are uncomfortable due to them being smaller than expected when placed on the eye.

**Lens Movement and Base Curve.** It is assumed that some lens movement is necessary in order to encourage postlens lubrication and, in turn, avoid mechanical irritation and corneal desiccation. However, there has been little or no research to examine the consequences of zero soft lens movement. Of perhaps greater interest to lens wearing comfort is the opposite situation of excessive movement. The conventional belief is that excessive movement causes discomfort through encroachment of the lens edge onto the cornea. However, an alternative explanation might be irritation through excessive interaction between the lens and the lids. It is difficult to quantify the true amount of lens movement as some movement takes place when the lids are closed.

An extensive evaluation of soft lens fit reviewed more than 2000 contact lens fittings that had been classified as loose, tight, or optimal. Unacceptable fittings categorized as “loose” tended to show more movement and poorer comfort, with the mean comfort score for loose fittings being 7.4 compared with 8.5 for optimal fittings (on a 10-point scale). A relatively large proportion (63%) of loose fittings were found to be less than comfortable (<9 on 0–10 scale). Only one study has noted a significant correlation between lens movement and comfort, with less mobile lenses being rated more comfortable. Conventional contact lens practice assumes that the greatest influence on tightness of fit is back surface radius of curvature (base curve; BC). Most soft lens designs incorporate a single spherical curve on the back surface, although some soft lenses have utilized bicornic and aspheric designs. One of the most systematic clinical evaluations of BC variation was undertaken with relatively thick, low EWC lenses; therefore, the findings have to be treated with some caution. Lowther and Tomlinson attempted to determine the minimum change in BC required to effect a significant change in various clinical outcomes such as vision, corneal edema, lens movement, and comfort. A change in BC of 0.95 mm was required to have a significant effect on comfort. A later study of midwater lenses found that a 0.6 mm flattening of BC resulted in significantly poorer comfort. A study with first generation SiHy lenses found improved comfort with 24% of patients by switching from an 8.60- to an 8.40-mm BC. However, the clinical picture is clouded by a proportion of the flatter lens fittings showing edge stand-off due to the relatively stiff material characteristics. Other studies of individual lens designs available in two BCs have found no significant difference in comfort when subjects were fitted with both BCs.

**Lens Centration and the Lens-Eye Relationship.** It is assumed that soft lenses center in order to reach an equilibrium state that balances the various forces from the lids as well as the lens–ocular surface interaction. It seems unlikely that small amounts of decentration (e.g., <0.3 mm) are likely to affect comfort as this does not significantly alter the interaction of the lens with the cornea or the lids. However, in some cases there may be confounding factors, with centration being influenced by a factor that also affects comfort (e.g., looseness of fit). To our knowledge, no studies have examined the effect of minor changes in lens centration on comfort during lens wear.

**Edge Alignment and Lens Edge Profile.** The design parameters related to lens edge profile are less easy to specify as they encompass the thickness at various points near the edge and the actual shape of the edge profile. A study of lathe-cut low EWC lenses found no significant difference in comfort when the edge thickness was systematically varied between 0.08 and 0.16 mm. Similarly, no significant difference was found in a study of high-water lenses varying in edge thickness from 0.12 to 0.24 mm. However, these edge thicknesses are relatively thick compared with molded designs and it is possible that the range was not wide enough to detect differences.

Modern molded designs generally taper to a thinner edge than lathe-cut and older molded designs. Several edge shapes have been identified in the literature, including so-called “rounded,” “knife,” and “chisel” edges. In a study by Maissa and colleagues, the lens with the thickest edge shape (rounded) gave poorer comfort than one of the chisel edge designs and two of the knife edge designs. This rounded edge profile was also slightly less comfortable in the work by Hubner and colleagues. It is plausible that the thinner designs sit closer to the bulbar conjunctiva and have less interaction with the lids than the rounded design. Alternatively, since the lens types were of varying materials, it is possible that the relatively high modulus of the rounded design may also have been a factor influencing comfort. Evidence for the reduced lid interaction theory is provided by ocular coherence tomography (OCT) imaging. These show that thin, tapered edge designs show a smoother transition between the conjunctiva and the lens surface and produce less disruption (“buildup”) of conjunctival tissue at the lens
edge.\textsuperscript{187} Sharper, pointed edge designs also show less movement than thicker, rounded edges and induce more pronounced conjunctival staining.\textsuperscript{185}

Another finding that may relate to edge fit comes from one study that found better comfort with a bicurve back surface design compared with a monocusc design, even though all other parameters were held constant, including sagittal depth and edge thickness.\textsuperscript{169} It is possible that the bicurve design afforded better alignment with the eye at the periphery of the lens, reducing localized pressure at this point.

**Toric and Multifocal Designs.** More sophisticated lens designs such as those incorporating prescriptions to correct astigmatism and presbyopia are thicker than spherical designs and this may affect comfort during wear.

**Toric Contact Lenses.** A study of contact lens dropouts found a disproportionately high number of astigmatic lapsed wearers.\textsuperscript{77} When, as part of the same study, these lapsed found a disproportionately high number of astigmatic lapsed wearers, they were refitted with contact lenses, there was a higher failure rate with toric soft lenses than spherical lenses. In a recent survey of soft lens wearers, symptoms of dryness were more frequent among toric soft lens wearers (45% vs. 30%, \( P = 0.04 \)).\textsuperscript{63} This mirrored the findings of a 1989 study that also found more symptoms of dryness in toric than spherical soft lens wearers (40% vs. 13%, \( P < 0.01 \)).\textsuperscript{187} There is evidence that CLD from a variety of sources is often misinterpreted as dryness.\textsuperscript{65} The interaction of the lid margin with front surface irregularities may be difficult to distinguish from the interaction with a dry lens surface.

An early study of toric soft contact lenses systematically evaluated the clinical performance of toric designs of varying prism and truncation.\textsuperscript{189} Although comfort was assessed and contributed to the outcome variable of “overall acceptance,” the comfort results were not reported separately. Not surprisingly, there was a tendency for the designs with thicker prism and more truncation to be less acceptable. Using a more recent prism ballasted, but non-truncated, design, Cho and colleagues found no significant difference in comfort between this and its spherical equivalent.\textsuperscript{191}

**Multifocal Soft Lenses.** Clinical trials that evaluate multifocals in comparison with spherical soft lenses give some insight into possible effects on comfort from multifocal optical designs. However, there are few such studies. One study from 1990 found no significant difference in comfort, even though the lens was a diffractive multifocal incorporating optic zone with a cushioning effect that reduces the edge clearance of the rigid lens. The fact that the discomfort reduces when the eyes are held closed would also tend to refute this.\textsuperscript{193}

Three important factors relating to the edge of rigid lenses govern comfort; these are the thickness and shape of the edge and the amount of clearance from the cornea. The greater the edge clearance, the greater the interaction with the eyelids and, in turn, poorer comfort.\textsuperscript{199} Cornish and Sulaiman\textsuperscript{199} evaluated the effect of rigid lenses of varying center thickness (\([CT], 0.10–0.21 \text{ mm}\)) on comfort and found that the thinnest lenses were more comfortable, which was attributed to greater on-eye lens flexure. Mandell\textsuperscript{200} attempted to characterize edge shape by specifying the location of the edge apex and lens thickness at various distances from the edge, and found less comfortable edges tended to incorporate an apex close to the lens front surface. Shanks evaluated 13 edge shapes and noted differences in comfort between lenses, but came to no overall conclusion on the optimum shape.\textsuperscript{201} Korb and Exford\textsuperscript{202} proposed an alternative strategy for maximizing comfort. Rather than finishing the peripheral contour of the lens so as to allow the upper lid to slide easily over the lens, they modified the edge and periphery to encourage “lid attachment.” The most systematic clinical evaluation of rigid lens edges was by La Hood,\textsuperscript{203} who assessed the comfort of four representative edge designs in four subjects: round, square, rounded anterior with square posterior, and square anterior with round posterior. The two designs with rounded anterior edges were significantly more comfortable than the other two. There was no significant difference in comfort between the lenses with square and rounded posterior edge profiles. The results confirm that the interaction of the lens edge with the eyelid is the most important factor in determining comfort in rigid lens wear.

**Large Diameter Lenses.** Large diameter RGP lenses might improve comfort by reducing lens movement and reducing the interaction of the lid with the edge of the lens. One potential classification for rigid lenses according to their overall diameter is “corneal” (<12.5 mm), “corneoscleral” (12.5–15.0 mm), and “scleral” lenses (>15.0 mm).\textsuperscript{204}

While there is clinical evidence of the short-term improved comfort with corneoscleral and scleral lenses compared with corneal RGP lenses, few well-controlled studies have addressed this point. Sorbara and Mueller\textsuperscript{205} compared the comfort of RGP lenses with different overall diameters in a nondispensing study in patients with keratoconus. The authors concluded that smaller diameter lenses (8.7 and 9.0 mm) were initially more comfortable in central cones, while larger lenses (10.1 and 10.4 mm) were preferred in oval cones. According to their results, lens movement was not directly related to comfort.

To date, there is no solid evidence of the benefits of corneoscleral and scleral lenses compared with corneal RGP lenses in terms of comfort, beyond clinical intuition. The average comfort ratings reported by some authors for scleral lenses (with diameters from 18.0–25.0 mm) fitted to patients with several ocular surface diseases\textsuperscript{206} are in the same range as comfort values already reported for corneal lenses in healthy subjects.\textsuperscript{163,165} Nevertheless, a direct comparison cannot be drawn, as most of the corneoscleral lenses were prescribed for eyes with serious eye disease.

In conclusion, considering the growing interest in the use of corneoscleral and scleral rigid lenses for eyes that do not exhibit disease or abnormal surface profiles, it is clearly necessary to conduct well-controlled, randomized studies where the potential for enhanced comfort of these lenses over standard diameter rigid lenses is investigated.

**Tear Exchange.** Placement of a contact lens on the eye leads to disruption of the tear film and to stagnation of the postlens tear layer during soft contact lens wear.\textsuperscript{207,208} Liberal exchange of this layer is generally considered preferable because it more closely represents the natural free flow of tears when no contact lens is in place and because buildup of debris behind the lens has been anecdotally associated with increased likelihood of corneal inflammatory events.\textsuperscript{209–214}
Measurement of tear exchange is almost exclusively conducted by determining the expulsion of a “marker” of some sort from behind the lens, which is typically sodium fluorescein, using a technique called fluorophotometry. McNama and colleagues \cite{16} used fluorophotometry to measure tear exchange in 23 subjects while they wore lenses of four different diameters in separate 30-minute wearing trials. Increased tear exchange was accomplished by decreased comfort, although intuitively the effects are linked and both are the result of decreasing lens diameter. Paugh and colleagues \cite{215} found fluorescence decay over 30 minutes to be greater with a prototype lotrafilcon A contact lens than thinner, less mobile etafilcon lenses in 11 subjects, but found no significant correlation between lens comfort and tear exchange rate.

Lin and colleagues \cite{216} investigated the effect of scalloped microchannels on the posterior surface of contact lenses on tear exchange over a 30-minute period, measuring comfort concurrently. They theorized that the channels might lead to increased tear exchange and were able to show this in Asian, but not in non-Asian, subjects. The microchannels do not induce any discernible change in comfort during the relatively short wearing times for which this design has been studied. \cite{216,217} One further method that may increase tear exchange is to fenestrate lenses, which would increase the flow of tears from the back surface of the lens to the front surface. To date, no studies appear to have investigated tear exchange with fenestrated soft lenses, but one paper \cite{218} was able to demonstrate that such a procedure does result in markedly reduced comfort due to interactions between the palpebral conjunctiva and the fenestrations.

The above methods of achieving increased tear exchange all involved increased lens movement, which is the likely associated factor. Unfortunately, increased lens movement is also associated with decreased comfort. The conventional view and some evidence is that tightly fitting lenses are comfortable and that loose-fitting lenses are likely to be less comfortable than well-fitting lenses. \cite{36,77}

In summary, there is little evidence that increasing tear exchange will have a positive effect on lens comfort and, to the contrary, changes to lens parameters that may bring about increased tear exchange are likely to have a simultaneous negative impact on lens comfort.

**Miscellaneous Factors**

**Tinted Lenses.** The tints on soft contact lenses can be translucent or opaque and in general, three types of tint are commonly used: visibility (or “handling”) tints, enhancement tints, and those with an opaque (or semiopaque) tint. Opaque tints can be applied using dot matrix printing on the lens surface, \cite{219} which can result in a relatively rough surface, \cite{220,221} and one study comparing these lenses with their clear equivalent found increased discomfort with the colored lenses. \cite{221}

Manufacturers have tried to overcome this problem by either housing the tint within the lens itself in the form of a laminate or applying a hydrophilic “coating” layer. \cite{220} The laminate construction has the advantage of encapsulating the printed matrix, but leads to increased lens thickness, \cite{222} which in turn may have a detrimental effect on comfort. Opaque lenses also have a fixed pupillary aperture that may lead to the wearer being more aware of these lenses due to constrictions of the visual field, \cite{223,224} peripheral vision blur, \cite{225}, and so-called “annular tinted contact lens syndrome” where subtle distortions of the cornea and induced astigmatism have been observed, with a subsequent reduction in vision. \cite{222,226} These visual problems may have an impact on the perceived comfort of the contact lenses, since problems with vision appear to affect comfort (Papas E, et al. IOVS 2003;44:ARVO E-Abstract 5694). In contrast, Gauthier and colleagues \cite{225} compared opaque colored lenses with their clear equivalents and observed no difference in overall comfort. Thus, evidence remains contradictory in terms of impact of lens tinting on comfort in contact lens wear.

**Indicator Markings.** Since most soft lenses are cast-molded, indicator markings on these lenses are placed onto the metal mold inserts by techniques such as electric discharge, diamond point engraving, and laser etching. The importance of the form of these markings to comfort have been the subject of various patents, with some placing much importance on these markings being composed of individual dots no greater than 90 \(\mu m\) in diameter being recessed into the lens front surface by a depth of 2 to 10 \(\mu m\). \cite{227,228} There is no published literature to date relating lens markings to in eye comfort, but anecdotal reports have occurred of lens wearers reporting increased lens awareness when lens markings have been added to or changed for existing products. In addition, anecdotal reports exist of these markings becoming filled with tear film components, \cite{229} and these deposits could act as a source of irritation.

**Contact Lens Deposition**

Since the commercialization of soft lenses in the early 1970s, clinicians have realized that contact lens materials rapidly attract tear film contaminants and that this deposition impacts lens performance. \cite{230,231} While intuitively it would appear obvious that there would be a link between comfort and contact lens deposits, proving such a link is somewhat more challenging, as many studies rely on visible deposition rather than biochemically measuring the actual degree of deposition. Visible measures of deposition are usually done either on-eye at the slit lamp biomicroscope or off-eye using various versions of the RUDKO scale, first reported by Allergan in the mid-1970s. \cite{232} This is problematic, as it is known that visible and measured deposition show a poor correlation. \cite{233,234}

**Visible Deposits and Comfort.** Roughly half of the studies conducted to date investigating comfort and its link with deposition have used visible deposition rather than biochemical analysis of deposits (Table 2).

The earliest of these studies by Nilsson and colleagues \cite{235,236} showed that lenses with the greatest level of deposition were generally less comfortable \cite{235} and that the use of a weekly enzyme cleaner resulted in increased comfort. \cite{236} While no direct correlation between comfort and deposition was reported in either paper, each surmised that increasing deposition was an important factor in reducing lens comfort. However, one study conducted at around the same time \cite{237} and two later studies \cite{238,239} were unable to demonstrate any correlation between visible deposition and comfort.

A larger multicenter study \cite{240} investigated lenses that were replaced every day versus those worn for up to 1 year. The daily disposable lenses, not surprisingly, exhibited reduced deposits over the course of the study and exhibited higher levels of comfort. The investigators linked these two factors, but many other factors (e.g., surface wettability; care solution effects) could have been the major reason for the improved comfort. Two other studies \cite{241,242} conducted at a similar point in time looking at the impact of frequent replacement of lenses on subjective performance were also able to link reduced levels of visible deposits with improved comfort. Of these, only one study \cite{241} actually reported a correlation between subjective responses and deposition, and while the correlation was weak; \((r = -0.35)\), it was statistically significant. The remaining five studies (Truong TN, et al. IOVS 2008;49:ARVO E-Abstract 4853) \cite{14,15,243,244} all included Silly materials that were replaced
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equivocal. One study253 showed no relevant correlation, while
comfort when subjects wore SiHy materials, the results were
baseline protein on the contact lens.

correlation (\(r = \frac{-0.13}{0.16} \); \(P = 0.03\) and comfort on insertion (\(r = -0.16\); \(P = 0.008\)).

In summary, the evidence-linking lipid or mucin deposition with contact lens comfort is either nonexistent or weak and future work should, perhaps, be directed at investigating mucin or lipid breakdown products rather than total lipid or mucin, since such a link is to be established.

Wearing Modality

Lens Wearing Schedule (Daily, Flexible, Extended, or Continuous Wear). Comparison of comfort differences between daily wear; flexible wear (occasional overnight wear); extended wear (regular overnight wear for up to 6 sequential nights); and continuous wear (regular overnight wear for over 6 sequential nights) of contact lenses is difficult because of numerous confounding factors. By virtue of the known average difference in comfort between contact lens wearers and nonwearers, a greater degree of discomfort on awakening is to be expected in those who sleep in lenses. Differences between daily wear comfort and comfort after sleep may be a function of hypoxia or tear disturbances rather than the wear schedule. Further, those who sleep in their lenses may self-select or survive in that modality on the basis of comfort, and prolonged wear such as extended wear means a longer wearing period and comfort differences may be related to the exposure time rather than the modality.

A number of studies have compared comfort between daily wear and extended wear of hydrogel lenses. Poggio and Abelson259 conducted a historical cohort study of 2433 cosmetic contact lens wearers. They reported that users of disposable extended wear lenses reported symptoms less frequently at routine visits than users of nonreplaced hydrogel daily wear lenses. It is important to note that the lens materials and designs and replacement frequencies were different in the two groups, meaning that the comfort difference can not be attributed to the wearing schedule in isolation.

Nichols and colleagues260 conducted a randomized, crossover, dispensing clinical trial specifically for the purpose of comparing daily disposable and disposable extended-wear modalities, using commercially available etafilcon brands. There was no significant difference between DD and EW in terms of lens comfort and awareness. However, a significant number of patients reported increased levels of ocular discomfort and irritation in the morning while in the extended-wear modality. Despite this, the subjects preferred the extended wear option overall, on the basis of convenience. The study by Aakre and colleagues261 followed 49 successful cosmetic contact lens wearers. They reported that users of disposable extended wear lenses reported symptoms less frequently at routine visits than users of nonreplaced hydrogel daily wear lenses. It is important to note that the lens materials and designs and replacement frequencies were different in the two groups, meaning that the comfort difference can not be attributed to the wearing schedule in isolation.

In summary, studies conducted using visible methods to determine lens deposition have provided poor evidence that comfort and deposits are linked, particularly over the 1 month or less that lenses are now typically worn. Future studies would be ill advised to rely on visible assessment of deposition to relate to contact lens comfort.

Quantified Protein Deposits and Comfort. To date, a
dozens of studies have used various biochemical analyses to quantify protein deposition and attempt to link it with contact lens comfort, and these are summarized in Table 3.

Of the nine studies looking at hydrogel materials alone (investigating all FDA categories), eight were not able to show any correlation between the quantity of protein deposited and symptoms, they were unable to report any significant correlation between deposits and comfort in 76 subjects wearing ionic, high EWC lenses. They found that the subjects using the daily cleaner reported improved comfort and that the lenses collected from these subjects showed reduced protein (lysozyme) deposition. No actual correlation analysis was reported, but the authors surmised that the reduced deposition and comfort were linked. Subbaraman and colleagues251 examined the correlation between symptoms and protein deposition over an 8-hour period in 30 subjects using an FDA Group IV material; while they were unable to report any significant correlation between total protein and symptoms, they were able to show a correlation (\(r \geq 0.64\); \(P < 0.001\)) between reported comfort and the amount of denatured lysozyme. They concluded that while comfort cannot be linked with quantity of protein deposited, it might be related to the amount of denatured protein on the contact lens.

Of the three studies investigating protein deposition and comfort when subjects wore SiHy materials, the results were equivocal. One study253 showed no relevant correlation, while another254 showed that a rewetting drop resulted in increased comfort and reduced total protein, total lysozyme, and increased protein activity and thus a relationship was surmised. The most recent study on a large sample of SiHy lenses was able to show a weak correlation (\(r = -0.13\)) with comfort on insertion.253 To date, no study has investigated the link between denatured protein and comfort for SiHy materials, but one in vitro study256 has reported that lysozyme activity does reduce over time following its deposition on both hydrogels and silicone hydrogels.

In summary, the main conclusion is that the amount of deposited protein appears unrelated to contact lens comfort, but protein activity may be correlated. However, further work in this area—particularly with regard to the degree of activity of proteins other than lysozyme—is required to confirm any such relation.

Quantified Lipid and Mucin Deposits and Comfort. The final area linking contact lens deposition and comfort are those quantifying the amount of lipid or mucin deposition, and these five studies are summarized in Table 4.

The two studies reporting on mucin deposition (one looking at hydrogel materials only255 and the other including silicone hydrogels256) were both unable to link the amount of mucin deposition with comfort. This lack of association was also reported for lipids in two studies looking at hydrogel materials.11,247 A more recent study investigating lipid deposition on a variety of SiHy materials was able to show weak (but significant) correlations for both overall comfort (\(r = -0.13\); \(P = 0.03\)) and comfort on insertion (\(r = -0.16\); \(P = 0.008\)).
<table>
<thead>
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<th>Author</th>
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<td>Bruce et al.²³⁹</td>
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<tr>
<td>Author</td>
<td>Study Design</td>
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</tr>
<tr>
<td>Nichols244</td>
<td>Prospective, nonrandomized, 2-phase (cleaning without and then with a rub step), daily wear, nonmasked, multicenter (phase I) and single-site (phase II)</td>
<td>2 wk</td>
<td>Galyfilcon A</td>
<td>Phase I: 118 existing wearers; Phase II: 33 “heavy depositors” from phase I</td>
<td>Graded off-eye at the slit lamp using a modified RUDKO system</td>
<td>Digital rub reduced visible deposition: No correlation</td>
<td>II</td>
</tr>
<tr>
<td>Cheung et al.54</td>
<td>Prospective, randomized (material), contralateral eye, daily wear, double-masked, single-site</td>
<td>2 wk</td>
<td>Etafilcon A; galyfilcon A</td>
<td>30 asymptomatic existing wearers</td>
<td>Graded off-eye using a modified RUDKO system</td>
<td>Galyfilcon A exhibited greater deposition; similar comfort between lenses: No correlation</td>
<td>I</td>
</tr>
<tr>
<td>Truong TN, et al. JOVS 2008; 49:ARVO E Abstract 4833</td>
<td>Retrospective, analysis of data from database of 60 single studies, single site</td>
<td>Lens age not reported; appear to be short-term fitting studies (6 h)</td>
<td>Various SilHy materials</td>
<td>275 existing wearers</td>
<td>Not described</td>
<td>Visible deposition correlated with comfort scores</td>
<td>III</td>
</tr>
<tr>
<td>Author</td>
<td>Study Design</td>
<td>Study Period/ Age of Lens at Analysis</td>
<td>SCL Materials</td>
<td>Number of Subjects</td>
<td>Visible Deposits Recorded or Graded</td>
<td>Laboratory Analytical Methods Used</td>
<td>Correlation With Comfort Scores</td>
</tr>
<tr>
<td>-------------------</td>
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<td>---------------------------------</td>
</tr>
<tr>
<td>Hydrogel material(s) only</td>
<td>Prospective, randomized (lens material), daily wear and extended wear, nonmasked, single-site</td>
<td>Various periods from 1 min to 7 d</td>
<td>Etafilcon A; polyHEMA</td>
<td>4 neophytes and 11 existing wearers</td>
<td>Graded on-eye at the slit lamp using a modified RUDKO system</td>
<td>SDS-PAGE</td>
<td>Increased protein with etafilcon A material: No correlation</td>
</tr>
<tr>
<td>Lever et al.</td>
<td>Prospective, nonrandomized, observational, daily wear, nonmasked, multicenter</td>
<td>Single visit (lens age not controlled; lens required replacement)</td>
<td>FDA groups I, II, III, and IV</td>
<td>977 lenses from 890 existing wearers</td>
<td>Graded off-eye using a modified RUDKO system</td>
<td>Ninhydrin assay</td>
<td>No correlation</td>
</tr>
<tr>
<td>Jones et al.</td>
<td>Prospective, randomized (wearing period), daily wear, investigator masked, single site</td>
<td>1 mo and 3 mo</td>
<td>Vaserfilcon A</td>
<td>12 existing wearers</td>
<td>Graded 0–3 on-eye at the slit lamp</td>
<td>UV spectroscopy</td>
<td>Protein deposits increased with wearing period; comfort no difference: No correlation</td>
</tr>
<tr>
<td>Lebow and Christensen</td>
<td>Prospective, randomized (solution system), daily wear, investigator masked, multicenter</td>
<td>3 mo</td>
<td>3× FDA group IV materials</td>
<td>76 existing wearers</td>
<td>Graded off-eye using a modified RUDKO system</td>
<td>HPLC OPTI-FREE solution</td>
<td>Reduced visible and protein deposition and increased comfort: Correlation surmised</td>
</tr>
<tr>
<td>Young et al.</td>
<td>Prospective, randomized (material), contralateral eye (material), daily wear, double-masked, single-site</td>
<td>1 mo</td>
<td>Atafilon A; etafilcon A; omafilcon A; polyHEMA; Weicon CE</td>
<td>30 (mix of neophyte and existing wearers)</td>
<td>None</td>
<td>Spectrofluorimetry</td>
<td>No correlation</td>
</tr>
<tr>
<td>Bruinsma et al.</td>
<td>Prospective, nonrandomized (wearing period), daily wear, nonmasked, single-site</td>
<td>10 d and 50 d</td>
<td>Etafilcon A</td>
<td>10 neophytes</td>
<td>None</td>
<td>SDS-PAGE</td>
<td>Comfort worse with lenses worn for longer period of time: No correlation</td>
</tr>
<tr>
<td>Michaud and Giasson</td>
<td>Prospective, randomized (replacement period), daily wear, investigator-masked, single-site</td>
<td>1 d up to 26 d</td>
<td>Etafilcon A</td>
<td>17 existing wearers</td>
<td>None</td>
<td>Total protein by UV- colorimetric assay</td>
<td>No correlation</td>
</tr>
<tr>
<td>Caron et al.</td>
<td>Prospective, randomized (rewetting drop), daily wear, double-masked, single-site</td>
<td>2 wk</td>
<td>Etafilcon A</td>
<td>22 asymptomatic existing wearers (18 analyzed)</td>
<td>None</td>
<td>Modified Lowry</td>
<td>Reduced protein with Clens 100 drops: No correlation</td>
</tr>
<tr>
<td>Author</td>
<td>Study Design</td>
<td>Study Period/ Age of Lens at Analysis</td>
<td>SCL Materials</td>
<td>Number of Subjects</td>
<td>Visible Deposits Recorded or Graded</td>
<td>Laboratory Analytical Methods Used</td>
<td>Correlation With Comfort Scores</td>
</tr>
<tr>
<td>------------------------------</td>
<td>------------------------------------------------------------------------------</td>
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</tr>
<tr>
<td>Subbaraman et al.²⁵¹</td>
<td>Prospective, randomized (wearing time), crossover, daily wear, investigator-masked, single-site</td>
<td>2, 4, 6, or 8 h</td>
<td>Etafilcon A</td>
<td>30 existing wearers (16 symptomatic of dryness)</td>
<td>None</td>
<td>SDS-PAGE; Western blot; micro-BCA; lysozyme activity assay</td>
<td>No correlation between comfort and total protein or total lysozyme, but strong correlation with denatured lysozyme ($R^2 \geq 0.64$)</td>
</tr>
<tr>
<td>Subbaraman et al.²⁵⁴</td>
<td>Prospective, randomized (drop use–surfactant-based rewetting drop vs. saline drop), crossover, extended wear, investigator-masked, single-site</td>
<td>1 mo</td>
<td>Lotrafilcon A</td>
<td>24 existing wearers</td>
<td>None</td>
<td>SDS-PAGE; Western blot; dot blot; lysozyme activity assay</td>
<td>Use of a surfactant-containing rewetting drop resulted in increased comfort on insertion, reduced total protein, reduced lysozyme and increased protein activity: Correlation surmised</td>
</tr>
<tr>
<td>Santos et al.²⁵⁵</td>
<td>Prospective, randomized (material), contralateral eye (material), daily wear, subject-masked, single-site</td>
<td>15 d (etafilcon A); 30 d (all other materials)</td>
<td>Balafilcon A; etafilcon A; galyfilcon A; tetrafilcon A; lotrafilcon B</td>
<td>31 neophytes</td>
<td>None</td>
<td>SDS-PAGE; fluorescence microscopy</td>
<td>No correlation</td>
</tr>
<tr>
<td>Zhao et al.²⁵⁵</td>
<td>Prospective, nonrandomized, parallel group, daily wear, double-masked, single-site</td>
<td>2 or 4 wk</td>
<td>Balafilcon A; galyfilcon A; lotrafilcon B; senofilcon A</td>
<td>583 lenses</td>
<td>None</td>
<td>Bicinchoninic acid or NanoOrange</td>
<td>Weak association with comfort on insertion ($R = -0.13$)</td>
</tr>
</tbody>
</table>
Table 4. Studies Comparing Comfort With Lipid and Mucin Deposits

<table>
<thead>
<tr>
<th>Author</th>
<th>Study Design</th>
<th>Study Period/Age of Lens at Analysis</th>
<th>SCL Materials</th>
<th>Number of Subjects</th>
<th>Visible Deposits Recorded or Graded</th>
<th>Laboratory Analytical Methods Used</th>
<th>Correlation With Comfort Scores</th>
<th>Level of Evidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrogel material(s) only</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Jones et al. [247]</td>
<td>Prospective, randomized (wearing period), crossover, daily wear, investigator-masked, single-site</td>
<td>1 and 3 mo</td>
<td>Vasurfilcon A</td>
<td>12 existing wearers</td>
<td>Graded 0–3 on-eye at the slit lamp</td>
<td>Fluorescence spectrophotofluorimetry</td>
<td>Lipid deposits increased with wearing period; comfort no difference; no correlation</td>
<td>I</td>
</tr>
<tr>
<td>Young et al. [11]</td>
<td>Prospective, randomized (material), contralateral eye (material), daily wear, double-masked; single-site</td>
<td>1 mo</td>
<td>Atafilcon A; etafilcon A; omafilcon A; polyHEMA; Weicon CE</td>
<td>30 (mix of neophyte and existing wearers)</td>
<td>None</td>
<td>Spectrophotofluorimetry</td>
<td>No correlation</td>
<td>I</td>
</tr>
<tr>
<td>Berry et al. [257]</td>
<td>Prospective, observational, daily wear, nonmasked, single-site</td>
<td>Single visit (lens age not controlled; min of 3 wk old)</td>
<td>Variety of hydrogel materials</td>
<td>50 existing wearers (19 graded as being symptomatic)</td>
<td>None</td>
<td>Dot and Western blots</td>
<td>Mucin deposition similar between symptomatic and asymptomatic subjects; no correlation</td>
<td>II</td>
</tr>
<tr>
<td>Silly material(s) included</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Zhao et al. [255]</td>
<td>Prospective, nonrandomized, parallel group, daily wear, double-masked, single-site</td>
<td>2 or 4 wk</td>
<td>Balafilcon A; galyfilcon A; lotrafilcon B; senofilcon A</td>
<td>583 lenses</td>
<td>None</td>
<td>Thin-layer chromatography</td>
<td>Weak association with comfort on insertion ($R = -0.16$) and comfort overall ($R = -0.13$)</td>
<td>II</td>
</tr>
<tr>
<td>Berry et al. [258]</td>
<td>Prospective, nonrandomized, daily wear, double-masked, single-site</td>
<td>2 wk</td>
<td>Senofilcon A; vifilcon A</td>
<td>33 neophytes</td>
<td>None</td>
<td>Dot and Western blots</td>
<td>Materials can modulate mucin adherence; no correlation</td>
<td>I</td>
</tr>
</tbody>
</table>
the two wearing schedules.50 Bergenske and colleagues55 reported on 317 subjects wearing SiHy lenses in CW and compared them with 81 neophytes introduced to hydrogel lenses DW in a prospective, 3-year, open-label, nonrandomized study. They found that wearers of the hydrogel lenses reported diminished-day and end-of-day dryness more frequently. Ramamoorthy and colleagues106 considered a range of factors that might be associated with contact lens–related dry eye in a cross-sectional study. This study considered both SiHy and hydrogel lens wearers combined. Those who wore their lenses overnight were almost one-third as likely to be classified as having "dry eye" as those who did not, in a univariate model. However, a model that controlled potential confounding factors (age, sex, recent contact lens refitting, and number of weekly applications of artificial tears/rewetting drops) did not show overnight wear to be a significant factor.

As noted earlier, the wearing modality may interact with the effect of oxygen on comfort, if such an effect exists. Perhaps surprisingly, a tentative conclusion can be drawn that, apart from relatively minor dryness upon awakening, individuals who sleep in lenses are not at a disadvantage and indeed may benefit in terms of comfort and dryness compared with those who do not sleep in lenses and wear them on a daily wear basis. There have been no studies reporting superior comfort in DW, save for comfort on awakening. To date, inadequacies of study designs prevent more definitive conclusions from being drawn.

**Duration of Wear.** The comfort and dryness response to contact lens wear may not be static. Certainly, practitioners and patients are familiar with comfort adaptation to gas-permeable lenses. Fonn and colleagues165 and Morgan and colleagues164 have both tracked the dramatic change in comfort that occurs during the initial few weeks of wear. SiHy lenses are generally stiffer than hydrogel lenses and may also have surfaces with higher coefficient of friction, as discussed earlier. It is reasonable to propose that a similar, albeit lesser, adaptation may occur during wear of these lenses as takes place during the initial wear of gas-permeable lenses. Certainly, while the first week of wear may show the largest adaptation to comfort, the study of Chalmers and colleagues48 suggested that there is a gradual decrease in the percentage of subjects with more severe dryness frequency and also with moderate or severe dryness over 6 to 12 months while wearing SiHy lenses in CW mode.

The impact of duration of wear may also explain in part the differences observed between studies when comparing hydrogels and silicone hydrogels. In short-term studies (up to 30 days) particularly for DW, hydrogels would generally seem to provide equal or superior comfort.54,59,64,116 In longer-term studies, silicone hydrogels would appear to provide better comfort.44,45,48,55

A confounding and possible competing factor in considering the impact of duration of wear on comfort is the effect of lens replacement frequency. There does not appear to be any Level I evidence to support the contention that duration of wear is important in determining contact lens comfort.

**Lens Age and Replacement Frequency.** Lens age is an obvious candidate for influencing CLD, as lenses begin to attract tear film components immediately upon application to the eye.245,262–264 Complexing and denaturation of this material can then lead to potentially problematic deposits on and within the lens.265,266

In an early cross-sectional study, Brennan and Efron109 found that increasing lens age led to increased frequency of dryness, but they were only able to separate differences between lenses younger and older than 6 months that had been worn on a conventional (nonplanned replacement) basis. The move toward disposable lenses in the late 1980s was an initiative to limit the amount of deposition, in the hope of diminishing complications.267,268 Frequency of replacement of contact lenses thus becomes highly relevant as a factor affecting contact lens wearing comfort and also offers a framework to examine the effect of lens age on comfort.

A review of studies that have identified improved comfort with disposable or frequent replacement reusable lenses and, furthermore, with increasing frequency of lens replacement. However, it is important to remember that the lenses used for this purpose are not necessarily made from the same lens material or with the same design as the comparator lenses or cared for with the same system. The imperfect control in such instances leaves open the possibility that it is not the lens replacement frequency as much as these other factors that are responsible for the observed improvements in comfort. A further challenge faced by researchers is achieving appropriate masking with respect to the replacement frequency in dispensing studies. Table 5 lists studies that allow the effect of lens age or different replacement frequencies on comfort to be compared, along with an indication of the quality of the studies in conforming with the evidence-based principles adopted within this report.

Boswall and colleagues conducted a retrospective chart review at a single contact lens practice from extended wear patients, of whom 65 wore disposable (7- to 14-day replace ment) contact lenses and 61 wore nonplanned replacement lenses.270 They found severe symptoms (itching, burning/drying, and foreign body sensation) to be reduced in the disposable group, implying that increasing lens age is a factor in producing such symptoms. However, the average wearing time each day for the nonreplaced lenses was longer, presenting a possible confounding factor. Poggio and Abelson271 historical cohort study compared disposable extended wear with nonreplaced daily and extended wear lenses. They found that disposable extended wear contact lens users reported symptoms less frequently at scheduled visits than both conventional daily wear and conventional extended wear users and that they had a lower rate of unscheduled visits for symptoms.

Poggio and Abelson's271 historical cohort study of 1954 daily wearers of soft contact lenses found that those using reusable, frequently replaced lenses had a significantly lower reported frequency for symptoms (particularly grittiness, scratchiness, irritation, and pain) compared with nonplanned replacement conventional wearers. In 1996, Pritchard and colleagues241 randomly assigned 119 neophytes to either a 1- or 3-month replacement schedule or nonreplacement group while wearing thin 38% EWC polyHEMA contact lenses. While there were reduced complications with the more frequently replaced lenses, ratings of comfort and overall satisfaction were not found to be different between the groups. Potential reasons for there being no difference include the possibilities that replacement frequency does not influence comfort, replacement frequency is less important in thin low EWC lenses, their technique lacked sensitivity to measure such a difference in this population, or subject bias induced by their knowledge of how often they replaced their lenses.

Relatively few studies have compared the comfort advantage of 2-week versus 1-month replacement and there are certainly no Level I evidence that make this comparison. Malet and Schneider273 reported a prospective study of 3066 daily wearers of monthly replacement lenses who were refitted at over 300 individual practices into a 2-week replacement regime. This observational study concluded that subjective comfort was improved by reducing replacement intervals to 2 weeks. The study made a particular point in identifying that the improved subjective comfort was also dependent on a compatible lens cleaning regimen.275
Studies listed enable comparison of comfort versus age of contact lenses. The major experimental paradigm that allows such comparisons are investigations in which different lens replacement frequencies are used (Brennan and Efron 189 and Frangie et al. 269 are the exceptions). Note that prior lens usage varies between experiments. Both, both hydrogel and silicone hydrogels were worn; Conv, nonplanned replacement; q, quarter. See Table 1 for other abbreviations.

* Indicates that increasing lens age is associated with decreasing comfort.
† Details not explicitly provided in the paper.

### Table 5. Studies Comparing Comfort of Lenses of Different Age

<table>
<thead>
<tr>
<th>First Author</th>
<th>Year</th>
<th>Test</th>
<th>Control</th>
<th>Material</th>
<th>Modality</th>
<th>Study Type</th>
<th>n</th>
<th>Sites</th>
<th>Duration</th>
<th>Rand</th>
<th>Masked</th>
<th>Effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brennan</td>
<td>1989</td>
<td>&lt;6 mo</td>
<td>&gt;6 mo</td>
<td>Hyd</td>
<td>DW</td>
<td>Retro</td>
<td>71/33</td>
<td>Single</td>
<td>[µ = 22 mo]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Boswall</td>
<td>1993</td>
<td>1-2 wk</td>
<td>Conv</td>
<td>Hyd</td>
<td>EW</td>
<td>Retro</td>
<td>65/61</td>
<td>Single</td>
<td>[µ = 23/26 mo]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Poggio</td>
<td>1993a</td>
<td>2 wk</td>
<td>Conv</td>
<td>Hyd</td>
<td>DW</td>
<td>Retro</td>
<td>1258/696</td>
<td>Multi</td>
<td>8-27 mo</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Poggio</td>
<td>1993b</td>
<td>2 wk</td>
<td>Conv</td>
<td>Hyd</td>
<td>EW</td>
<td>Retro</td>
<td>905/473</td>
<td>Multi</td>
<td>8-34 mo</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nason</td>
<td>1994</td>
<td>1 d</td>
<td>Conv</td>
<td>Hyd</td>
<td>DW</td>
<td>Pros</td>
<td>70/125</td>
<td>Multi</td>
<td>1 y</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Snyder</td>
<td>1995</td>
<td>1 d</td>
<td>Conv</td>
<td>Hyd</td>
<td>DW</td>
<td>Pros</td>
<td>18</td>
<td>Single</td>
<td>2 wk</td>
<td>✓</td>
<td>✓</td>
<td>I, S</td>
</tr>
<tr>
<td>Nilsson</td>
<td>1995</td>
<td>1 d</td>
<td>Conv</td>
<td>Hyd</td>
<td>DW</td>
<td>Pros</td>
<td>20</td>
<td>Single</td>
<td>3 mo</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Solomon</td>
<td>1996</td>
<td>1 d</td>
<td>Conv</td>
<td>Hyd</td>
<td>DW</td>
<td>Pros</td>
<td>73/136</td>
<td>Multi</td>
<td>3 y</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pritchard</td>
<td>1996</td>
<td>1 mo</td>
<td>Conv</td>
<td>Hyd</td>
<td>DW</td>
<td>Pros</td>
<td>37/41</td>
<td>Single</td>
<td>2 y</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Nichols</td>
<td>2000</td>
<td>1 d</td>
<td>Conv</td>
<td>Hyd</td>
<td>DW</td>
<td>Pros</td>
<td>37/41</td>
<td>Single</td>
<td>2 y</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sulley</td>
<td>2001</td>
<td>2 wk</td>
<td>1 mo</td>
<td>Hyd</td>
<td>DW</td>
<td>Pros</td>
<td>525</td>
<td>Multi</td>
<td>1 mo</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Maler</td>
<td>2002</td>
<td>2 wk</td>
<td>1 mo</td>
<td>Hyd</td>
<td>DW</td>
<td>Pros</td>
<td>5066</td>
<td>Multi</td>
<td>1 mo</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aakre</td>
<td>2004</td>
<td>1 d</td>
<td>1 mo</td>
<td>Hyd/SiHy</td>
<td>DW/CW</td>
<td>Pros</td>
<td>19/30</td>
<td>Single</td>
<td>6 mo</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Frangie</td>
<td>2008</td>
<td>2 wk</td>
<td>1 mo</td>
<td>Both</td>
<td>DW</td>
<td>Retro</td>
<td>434</td>
<td>Survey</td>
<td>†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ramamoorthy</td>
<td>2008</td>
<td>2 wk</td>
<td>1 d</td>
<td>†</td>
<td>†</td>
<td>Retro</td>
<td>176/22</td>
<td>Single</td>
<td>†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>2 wk</td>
<td>1 mo</td>
<td>†</td>
<td>†</td>
<td>Retro</td>
<td>176/109</td>
<td>Single</td>
<td>†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>2 wk</td>
<td>1 q</td>
<td>†</td>
<td>†</td>
<td>Retro</td>
<td>176/19</td>
<td>Single</td>
<td>†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dumbleton</td>
<td>2010</td>
<td>2 wk</td>
<td>1 mo</td>
<td>SiHy</td>
<td>†</td>
<td>Retro</td>
<td>717/617</td>
<td>Survey</td>
<td>†</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lazon de la Jara</td>
<td>2013</td>
<td>1 d</td>
<td>2-4 wk</td>
<td>SiHy</td>
<td>DW</td>
<td>Pros</td>
<td>39/213</td>
<td>Single</td>
<td>3 mo</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

- Conv, nonplanned replacement
Sulley and Meyer’s study—yielded similar results. In both studies, the lens materials and designs were different between the replacement frequency groups, subjects were not masked, and there was not a concurrent control lens run alongside the 2-week replacement lenses.

In a conference abstract, Jones and colleagues reported that subjects replacing their lenses on a daily disposable or 2-week schedule were less likely than monthly wearers to report dryness. Frangie and colleagues surveyed 271 and 165 patients wearing a variety of monthly replacement hydrogel and SiHy lens brands, respectively. A total of 68% of the hydrogel wearers and 71% of the SiHy wearers noticed a decrease in comfort during the month. Less than 10% of both groups noticed that this occurred within the first 2 weeks of the wearing period, with the remaining 90% finding that the discomfort developed in the third and fourth weeks of the month. While they did not provide statistical analysis, Long and colleagues showed data that supports the concept that lenses after a month of wear are less comfortable than after 2 weeks of wear.

One study stands in contradiction to the general trend of reports that have found shorter replacement schedules lead to better comfort. Dumbleton and colleagues conducted a survey of 1344 wearers of SiHy contact lenses through practitioners in the United States, with approximately half wearing lenses with a manufacturer’s recommended 2-week replacement schedule and the other half on a monthly schedule. Noncompliance was found to lead to lower comfort, perhaps unexpectedly. After adjusting for compliance, the authors reported modest but statistically significantly better comfort with the monthly replacement lenses compared with the lenses recommended for a 2-week modality. The study surveyed existing wearers and so selection and survival bias cannot be ruled out in addition to differences in lens materials and design.

Most of the remaining studies considering the impact of replacement frequency have looked at the effect of daily disposables on lens comfort. Early hydrogel lens studies considered the 1-day replacement modality against conventional lens replacement. Daily replacement led to a number of benefits, including improvements on a range of comfort measures. Daily disposability also led to a range of comfort benefits against lenses planned to be replaced on a 1- to 3-month basis. Solomon and colleagues also found benefits of a daily versus a 2-week replacement schedule. It is interesting to note that the single study investigating the impact of replacement frequency on comfort, which conforms to the gold standard principles of a controlled, randomized, masked study did not find a difference between daily and two weekly replacements. However, it is not clear in this study whether the method for measuring comfort was adequate to detect differences, whether the sample size (n = 18) was sufficiently powered to detect a difference, whether lens care influenced comfort, and whether the lens type used was not resistant to the effects of aging.

The potential comfort advantage of daily disposal of SiHy lenses versus other replacement schedules has not been widely studied. In the one available study to-date, Lazon de la Jara and colleagues reported that end-of-day comfort and dryness ratings were significantly better for daily disposable wear than when the same material and design was used in a reusable manner with either hydrogen peroxide or multi-purpose care systems.

Two studies have compared daily disposable lenses with extended or continuous wear. In the first, daily disposable hydrogel lenses were compared with a 2-week replacement hydrogel. There was no difference between the daily disposable and extended wear modalities with respect to subjective responses, except that those sleeping in their lenses found comfort on awakening to be inferior. In the second study, daily disposable hydrogel lenses were compared with monthly replacement silicone hydrogels in continuous wear. There were no differences of note between the two groups. A final study looking at the effect of a variety of replacement intervals on lens comfort was conducted by Ramamoorthy and colleagues. Individuals (n = 360) were surveyed with the CLDEQ and categorized as either having or not having dry eye. Daily, 2-week, monthly, and quarterly replacement schedules were all represented in the sample, as were both hydrogel and SiHy lenses. Replacement schedule was not found to be predictive of how the subjects were classified. However, it is not clear that the study was powered to make determinations with respect to the different schedules and to what extent selection and survival bias were influential.

In summary, there is almost a complete absence of masked, randomized, controlled studies that consider the impact of replacement schedule on comfort and dryness, preventing a definitive statement being made on the topic. Nonetheless, there is a tendency for the studies that provide circumstantial evidence regarding replacement schedules to suggest that replacement that is more frequent is conducive to greater comfort.

Time of Day. End-of-day dryness and discomfort arguably represent the most challenging issue for the contact lens industry today. While discomfort is the major reason cited for contact lens discontinuation, a breakdown by the exact nature of discomfort is illuminating. Chalmers and Begley studied responses to a questionnaire of 1054 patients who presented for eye care in a multicenter cross-sectional study. The leading causes of discontinuation among the 167 former contact lens wearers in their sample were dryness and end-of-day discomfort, which were cited by 41.9% and 38.3%, respectively.

Begley and colleagues surveyed 367 unselected contact lens wearers in their 2001 cross-sectional study of North American optometric clinical practices, finding that the percentage reporting moderate to intense ocular discomfort increased from 19% in the morning to 56% in the evening. The frequency of dryness among the 367 contact lens wearers in the group was significantly higher late in the wearing day compared with earlier in the day, with an increase from 12.7% who reported moderate to intense dryness in the first 2 hours of wear to 28.5% late in the day. An analysis of questionnaire responses from 84 clinical sites in North America, Chalmers and colleagues found that between 3% and 15% of subjects, depending on age and type of lens material worn, reported end-of-day discomfort (“extreme” or “very”) and between 7% and 24% report end-of-day dryness. Young and colleagues reported severity of end-of-day dryness to be of sufficient significance to categorize subjects as having “contact lens-associated dry eye” in 31% of wearers.

While noticed even in nonwearers and spectacles wearers, all symptoms—but particularly end-of-day dryness—are more pronounced in contact lens wearers. Further, contact lens wearers report markedly fewer symptoms without the lenses in place: for example, only 1.5% report moderate to intense late-day dryness according to Chalmers and Begley.

Evidence in support of decreased comfort toward the end of the day necessarily cannot arise from Level I studies, as it is virtually impossible to conduct a controlled, masked, randomized study where time of day is the key independent variable. Nonetheless, a large volume and variety of experiments, often investigating other phenomena or as part of multivariable analyses, provide undeviating data to support the hypothesis.
that comfort decreases during the day and is exacerbated by contact lens wear.

The original empirical demonstration of decreasing comfort toward the end of the day appears to have been by Pritchard and Fonn. In their 1995 study, which sought to link lens dehydration with symptoms, 19 subjects rated dryness on visual analog scales during 7 hours' wear of three different hydrogel lenses. Dryness ratings rose consistently for all three lens types at 1, 3, and 7 hours after lens insertion. In a follow-up study, Fonn and colleagues sought to determine whether lens dehydration correlated with discomfort and dryness in two different lens types in 40 subjects, but this time they separated subjects into symptomatic and asymptomatic groups. They replicated the findings of increased dryness and decreased comfort over a 7-hour period of hydrogel lens wear in the asymptomatic group, but interestingly there was no significant variation over time in the symptomatic group. Others have confirmed the finding among hydrogel lens wearers.

Fonn and Dumbleton published the first paper reporting reduced end-of-day comfort with SiHy lenses in a 7-hour nondispensing study of 20 asymptomatic and 19 asymptomatic wearers. Again, like the 1999 study from the same group, the reduced end-of-day comfort was isolated to the symptomatic group. The paper suggested that the degree to which comfort diminishes and dryness increases is very similar to that observed with hydrogel lenses. Many others have reported decreased comfort over the course of the day among SiHy wearers.

Wear of daily disposable contact lenses is not protective from the effect of time of day on comfort and dryness. Walker and colleagues measured a drop in mean comfort across the day when comparing two different hydrogel daily disposable lenses in a 20-site study of 282 subjects. Diec and colleagues tracked decreasing comfort across the day for a series of hydrogel and SiHy lenses worn on a daily disposable basis. They found no difference between lenses but noticed decreased comfort across the course of the day.

While there may be differing degrees of discomfort between daily wearers and those who sleep in their contact lenses, extended or continuous wear also does not seem to prevent the phenomenon of decreased comfort and increased dryness toward the end of the day. In their 2005 study looking at wearers switched out of hydrogels and into silicone hydrogels, Chalmers and colleagues observed among the baseline data for those using hydrogel lenses on an extended basis, more than 50% greater frequency and severity of end-of-day dryness. Subbaraman and colleagues and Schafer and colleagues have made supporting observations with extended wear of silicone hydrogels. These consistent findings also seem to hold up across different ethnicities. Long and McNally studied symptoms in 88 Asian subjects that were switched from hydrogel lens wear to SiHy lens wear. Decreased comfort and increased dryness toward the end of the day was evident before and after switching lens type. Indeed, 84% of subjects reported end-of-day dryness either occasionally, frequently, or all the time with their habitual lenses.

Investigators have used variables based around wearing time both to demonstrate the consequence of end-of-day dryness and also as a measure of comfort for testing experimental hypotheses. As an example, Young and colleagues found mean comfortable wearing time in a group of UK subjects classified as having contact lens associated dry eye to be 9.1 hours compared with 12.4 hours in those without dry eye and, in a sample of North American wearers, 9.4 hours in the dry eye group compared with 12.1 hours in those without dry eye.

The interaction between the lens and the lid wiper appears to play a significant role in end-of-day dryness and discomfort. Lens surface coefficient of friction appears to be correlated to overall lens comfort and in particular to end-of-day comfort (see previous section on friction and lubricity). Certainly the simple addition of lubricants yields immediate, if short-term, benefits in comfort. However, the mechanism by which coefficient of friction might be linked to end-of-day comfort is uncertain. Daily accumulation of lens surface buildup, or diminution of tear quality during the course of the day, may lead to higher lens surface friction later in the day. Alternatively, lens surface friction may remain relatively stable, although at a raised level during wear compared with the bare cornea, but the lid-wiper region becomes irritated or damaged with erosion of cells during the course of a day's wear as a result of the rubbing between the lid and the front surface of the lens and so becomes uncomfortable. Overnight, the affected epithelium repairs and, on awakening and reaplication of the contact lens, the cycle begins again.

So far, there are no records of measurement of lens surface friction changes over the course of a day. The more modest changes of comfort across days and weeks of wear as evidenced by the data on replacement schedules above compared with the dramatic change over the course of a day would argue more strongly for the latter hypothesis of “wear and tear and then repair” to explain end-of-day discomfort. One further piece of evidence relates to the fact that replacement of a lens during the middle of the day appears to have minimal impact on end-of-day comfort, suggesting that a fatigue-like response in one or more ocular tissues or stimulation of ocular surface nociceptors induced by the presence of the contact lens occurs. Without doubt, further research is needed to ascertain the true origin of end-of-day dryness with contact lens wear.

From a scientific standpoint, the hypothesis that contact lens comfort decreases toward the end of the day can never be tested in a controlled, randomized, masked study because normal subjects will always have awareness of the duration for which they have worn the lenses and naturally the sequence of such measures. Thus, while the various studies reported here may be controlled, randomized, and masked with respect to lenses, care systems, or some other independent variable, they are not with respect to time of day and daily duration of wear. Nonetheless, the wealth of reporting on the matter and the overwhelming consistency of the data lead us to identify this problem as one of the major, if not the leading, issue with contact lens wear today.

**Care Products and Packaging Solutions**

**Compositions of Care Solutions.** A contact lens care solution is composed of several important components, including preservatives (or biocides), surfactants, chelating agents, and buffering agents. All these components have different functions and are incorporated into a lens care system to provide adequate disinfection efficacy and enhanced comfort. The difference in clinical performance observed between various lens care solutions may be due to the differing components and concentrations in the care products and the manner in which these components interact with the lens material. An essential point to consider is that the care system can result in reduced comfort or enhanced comfort—two very different outcomes; one is likely caused by uptake and subsequent release of the components of the care system and the other by the adsorption of a comfort “additive” to the lens material from the care system.

**Biocides.** Various biocides are incorporated into lens care regimens at different concentrations and the effect of these
biocides on subjective comfort and how these biocides interact with different lens materials has attracted significant interest.

**Peroxide-Based Systems.** Hydrogen peroxide-based solutions are used at a concentration of 3% (30,000 ppm). The subjective sensitivity threshold for peroxide ranges between 50 and 300 ppm, and it is recommended that the solutions be neutralized to a concentration of less than 100 ppm. When present in high concentrations, residual peroxide can be toxic to the cornea and can cause discomfort/pain.

Few studies have compared the effect of peroxide-based systems on subjective symptoms in a comparison with other care systems that are preserved with a different biocide. A randomized, single-masked, crossover design study evaluated the degree and frequency of corneal fluorescein staining and subjective responses in 85 hydrogel lens wearers following the use of PHMB-based (ReNu) and peroxide-based (AO Sept) systems for 1 month. It was found that the overall comfort and comfort in the evening were significantly better when the subjects used the peroxide-based system (P = 0.02 for both occasions). Another study investigated the clinical and subjective performance of a peroxide-based lens care system (ClearCare) in comparison to a Polyquad/Aldox-based multipurpose solution (MPS; OPTI-FREE RepleniSH) when used with lotrafilcon B and senofilcon A SiHy lenses. This randomized, contralateral (lens type) and crossover (care system) study involved 24 participants and they found that the peroxide-based system resulted in longer reported comfortable wearing times than the MPS (10.93 vs. 9.84 hours; P < 0.01). However, no significant difference was found between solutions in overall ratings of subjective comfort, or dryness. While these two studies taken in isolation would appear to suggest that peroxide-based systems are superior to preserved systems, there are multiple other components that differ between the products and this makes it impossible to support the fact that it is merely the biocide alone that resulted in the reported comfort differences.

**PHMB-Based Versus Polyquad-Based Systems.** Several studies have compared the effect of using a PHMB-based system when compared with a perquaternium-based MPS. A multisite, 231-subject, double-masked, crossover study was performed to evaluate the subjective comfort and satisfaction and clinical signs with two MPSs used with alphafilcon A and etafilcon A lenses. Subjects used each of the two MPSs, Polyquad/Aldox-based (OPTI-FREE Express) and PHMB-based (ReNu MultiPlus), for 28 days and found that subjective ratings of comfort and satisfaction were in favor of the polyquad-based system (ClearCare). Epstein found that the users of the PHMB-preserved product reported decreased comfort over the course of the day. Interestingly, it was also reported that the PHMB-based system was also associated with a reduction in relative corneal sensitivity (P = 0.004). However, a subsequent letter to the editor questioned this latter finding. A randomized, controlled, and investigator-masked clinical study compared the clinical performance of a PHMB-based (MeniCare Soft) and Polyquad/Aldox-based (OPTI-FREE Express) MPS with two SiHy lenses (lotrafilcon A and galyfilcon A), and found no significant difference between the two solutions.

In another study, subjective symptoms and clinical signs of tolerability and comfort were compared in silicone and hydrogel lens wearers using a Polyquad/Aldox-based system (OPTI-FREE Express) and a PHMB-based system (ReNu Multi-Plus). The participants who used the Polyquad/Aldox-based system reported greater comfort than the PHMB-based system. These results should be interpreted with caution because 65% of the subjects in this study used the Polyquad/Aldox-based system while only 28% used PHMB-based MPS prior to enrollment, and this could potentially create a bias toward their habitual care system.

Another study was conducted as a prospective, bilateral, clinical trial with a single-masked investigator, and randomized crossover design with four phases to assess the compatibility of a SiHy lens material with four different MPSs (one based on Polyquad/Aldox: OPTI-FREE RepleniSH, and three based on PHMB: ReNu MultiPlus, Solo-Care Aqua, and MeniCare Soft). No difference was found in comfort between the four care systems. A recent study investigated the performance of two new MPSs (polyquaternium/alexidine-based Complete ReviTalens and polyquaternium/PHMB-based Biotrue) during a month of SiHy lens wear in neophyte volunteers. The investigators did not find statistically significant differences between the two systems.

Finally, a randomized, investigator-masked, crossover clinical trial including 51 subjects compared a Polyquad/Aldox (OPTI-FREE Express) solution to a PHMB-based solution (Complete Moisture Plus) in subjects wearing etafilcon A lenses. Each participant used the assigned care solution for 7 days, with a 1-day washout period, followed by subsequent use of the alternative solution. While interferometric differences in the prelens tear film thickness were observed (likely based on viscosity differences between the solutions), there was no overall difference in subject preference for a care solution, but “comfort” was the primary reason for a preference selection when asked their reason for preference.

**Long-Term Use of PHMB-Based Versus Polyquad-Based Systems.** Long-term users of two different preservative systems were studied to investigate whether prolonged use of these systems was associated with an increase in the frequency of dry eye. Subjects were required to have consistently used a PHMB-based or polyquad-based solution for 2 years. This investigator-masked study, involving 89 FDA Group IV hydrogel or SiHy lens wearers, found that PHMB users reported significantly more grittiness or scratchiness (67% vs. 44%; P = 0.02). However, no significant differences between the two preservative system groups were noted for the range of other dry eye evaluations or the remaining clinical assessments.

**Studies Investigating Consumer Acceptance of MPS.** One study evaluated comfort when switching to Polyquad/Aldox-based MPS (OPTI-FREE RepleniSH) when compared with two different PHMB-preserved MPSs. This multicenter, open-label study enrolled 109 contact lens wearers who were dispensed with the test solution in place of their habitual solution. Subjects assessed their experience with their habitual solution (baseline) and the test solution (day 30) using Likert-style questions. They reported that the Polyquad/Aldox MPS was associated with a statistically significant improvement in instillation comfort, end-of-day comfort, clear vision, and overall satisfaction. It is difficult to determine if a natural bias is introduced in studies such as these, as subjects given new solutions or lenses will often rate “new” products as being superior due to the mere fact that they are new, therefore, “they must be better.”

In summary, although a few studies have shown that lens wearers using a care solution that is preserved with a specific biocide show better comfort than another product, it is important to note that a lens care solution is composed of many ingredients that may also impact subjective symptoms. Therefore, it cannot be concluded that a specific biocide alone will provide improved comfort.

**Surfactants and Wetting Agents.** In contact lens solutions, surfactants are used as detergents or cleaners, removing loose debris, microorganisms, and deposits by combining with these substances to form micelles, which are then removed during the rinsing procedure. Surfactants also play a role in enhancing the wettability of contact lenses, especially when asked their reason for preference.
Silicon hydrogel lenses, which are generally more hydrophobic than conventional hydrogel lens materials.131–135

The most common surfactants found in MPS are poloxamers (Pluronic F87, Pluronic F127, Pluronic 17R4) and poloxamines (Tetronic 1304, Tetronic 1107).314,315 HPMC, which has been used for many years in rigid gas-permeable care products for its lubricating, conditioning, and cushioning functions, has also been used in soft lens solutions as a wetting agent.316 It has been shown to be effective in controlling both symptoms and signs in patients with dry eye317 and to enhance tear film stability in lens wearers.134 A block copolymer (EO-BO) containing poly(ethylene oxide) and poly(butylene oxide) has also been recently introduced and its ability to adsorb to SiHy materials has been confirmed using x-ray photoelectron spectroscopy and ultra-performance liquid chromatography.310

A randomized, controlled, double-masked, multicenter study involving 362 subjects at 19 investigational sites in the United States investigated the performance of two MPSs (tetronic-containing MPS [OPTI-FREE RepleniSH] and poloxamine-containing MPS [ReNu MultiPlus]) with habitual lenses.293 They found that the comfort and dryness mean scores were significantly better for the tetronic-containing MPS compared with the poloxamine-containing MPS at day 28 and the mean scores for scratchiness and burning were significantly lower at day 14. The improved performance of the MPS was attributed to the presence of wetting agents, including C9-E3A and propylene glycol in the tetronic-containing MPS. It was speculated that the presence of these novel wetting agents aids in cleaning, chelation, wetting, and lowering biocide lens uptake, especially in combination with Tetronic 1304.293

The wetting effect of three different lens care solutions (two care solutions with wetting agents [OPTI-FREE RepleniSH and ReNu MultiPlus] and one solution without any wetting agent [ClearCare]) on blink rate, dryness symptoms, and vision performance on 65 habitual lens wearers was studied.318 They found that solutions with wetting agents led to significantly fewer eye blinks and better ocular comfort for contact lens wearers. Moreover, the presence of wetting agents in lens care solutions also resulted in better visual performance when compared with wearing daily disposable contact lenses.

The effect of an EO-BO containing MPS (OPTI-FREE PureMoist) was compared with a MPS containing a conventional surfactant-containing MPS (ReNu Fresh) with SiHy and HEMA-based lenses.319 It was a multicenter (30-site) study involving 573 participants over several visits. The patients found that their “lenses felt moist” at day 90 when using the MPS containing the novel surfactant (P ≤ 0.02) and the “lens acceptability at day 90” was better when the MPS with the novel surfactant was used (P ≤ 0.05).319 Another study that recruited over 3000 patients from 313 ophthalmologic practices in France to participate in a 1-month prospective observational clinical study found that replacing etafilcon A lenses once every 2 weeks combined with an MPS incorporating ingredients designed for lens conditioning contributed to significant improvements in lens wearing comfort.275 Another study showed that HPMC incorporated in an MPS could form a thicker, longer-lasting layer of fluid on the hydrogel lens, leading to improvements in tear function in contact lens wearers.134,311

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Chelating Agents. Chelating agents are added to lens care regimens to act synergistically with other agents to improve disinfection efficacy and to aid in removal of tear film components.311 The common chelating agents found in lens care regimens include EDTA, citrate, and hydroxyalkylphosphonate.315 Studies published to date have compared the clinical performance of a specific care regimen in comparison with only citrate-containing regimens.252,320,321 A multicenter, investigator-masked, randomized study investigated the effect of two citrate-containing regimens on subjective comfort and deposition on a FDA Group IV lens material and compared that with a noncitrate containing MPS.320 Significant differences favoring the citrate-based regimen were observed in ocular awareness, lens awareness, visual clarity, end-of-day comfort, and end-of-day dryness.320 These findings were consistent with that of another study, where it was found that the use of a dedicated daily cleaner in conjunction with a citrate-containing system can provide patients with more comfortable and cleaner lenses.252 Another study that compared comfort when using a citrate-containing MPS versus an MPS containing HPMC found no significant difference in comfort between the two solutions.321

In summary, all the above-mentioned studies compared a few lens care solutions and attributed any increased performance to the presence of a specific component. However, this is not always possible to provide as other factors between the products may also have contributed to the perceived differences in comfort. In order to specifically determine the association between lens care solution and discomfort and dryness, two studies have conducted extensive regression analysis. In the first study,32 the relationship between contact lens characteristics, hydrogel lens materials, care solutions, and patient-related factors and dry eye status in contact lens wearers was assessed retrospectively in 360 contact lens wearers. Interestingly, there was no significant association between contact lens-related dry eye and contact lens care solutions, when grouped either by preservative type or by product brand (both P = 0.99).52 Another more recent study examined the factors associated with contact lens-related dryness symptoms in soft contact lens patients.65 Soft contact lens wearers (n = 952) from 12 clinical sites were examined and they found that neither the lens material nor the lens care systems were specifically related to contact lens-related dry eye status.

These two studies suggest that contact lens-related dryness is associated with a diverse range of underlying causes and that lens care product is not a significant factor. It should be noted that both these studies derived data from retrospective studies by pooling data from multiple studies and/or sites and conducted advanced statistical analysis to determine the association between dry eye symptoms and the lens care regimen. Nevertheless, it is important to identify how different components in a lens care solution interact with contact lens materials and if this could have an impact on the physiological and subjective performance of contact lenses.

Interaction of Contact Lenses With MPS. Silicone hydrogel materials are hydrophobic and these materials may exhibit higher attraction for certain hydrophobic/lipophilic entities, such as tear lipids322 and nonpolar active agents found in certain MPS products.323,324 When a MPS interacts with a contact lens, any of the components found in the solution can be adsorbed onto the surface or absorbed into the bulk of the lens material.310,325,326,327 Preservative uptake from lens care solutions to soft lens materials is influenced by several properties of the lens, including EW, ionicity, and hydrophobicity.325,324,327 These adsorbed components may potentially cause discomfort to contact lens wearers.
The biocide uptake into and onto various contact lenses and its subsequent influence on clinical signs and symptoms were investigated in several studies.320,328–331 The physiological and subjective responses of subjects wearing balafilcon A silicone hydrogels and the ocular response to use of a lens care product containing Polyquad/Aldox and another containing PHMB was reported by Jones and colleagues.328 The PHMB-based lens care product was associated with increased corneal and conjunctival staining and more stinging or burning on lens insertion compared with the product containing Polyquad/Aldox.328 However, the investigators were not able to relate the degree of staining with the reported symptoms. This is in contrast to two more recent publications, in which increased amounts of corneal staining led to reduced subjective comfort.329,352

A series of pilot studies was conducted over 11 months to assess combinations of three different hydrogel lenses (FDA Group II [alphafilcon A], Group IV [etafilcon A] and one SiHy [lotrafilcon A]) and four MPSs.331 New lenses were soaked overnight in one of four MPSs and were fitted on subjects who rated comfort and ocular symptoms. Corneal staining was evaluated at baseline and after lens removal. The investigators found corneal staining to be most frequent when PHMB-preserved solutions were used with Group II lenses. Instead, with the polyquad-based system, the extent of staining was low with all the lenses tested. They also found that when PHMB-based products were used with the FDA Group II material, corneal staining was evident after 1 to 4 hours of wear. However, they did not see any association between significant symptoms and the extent of staining.351

Another study investigated the physiological and subjective responses of the short-term use of various lens care products with two SiHy contact lenses (lotrafilcon B or glyfalcon A) and examined whether changes to the surface of lenses was correlated with the responses.326 Both these lens types were presoaked for 1 week in Polyquad/Aldox-based (OPTI-FREE Express) or PHMB-based (Aquify) solution and participants wore them for 6 hours. It was found that lotrafilcon B lenses soaked in PHMB caused a decrease in comfort, an increase in burning/stinging after 1 hour of wear, and an increase in lens awareness on lens insertion. When lotrafilcon B lenses were soaked in Polyquad/Aldox, they found an increase in burning/stinging after 1 and 6 hours.326 The investigators concluded that release of various components of MPS from contact lenses can have a significant influence on corneal staining and comfort responses during wear.

In summary, the results from these studies show that contact lenses interact differently with MPS depending upon their polymeric makeup. These results also show that the uptake and subsequent release of components by soft contact lenses can affect corneal staining and subjective comfort.

In conclusion, while retrospective studies suggest that the lens care product is not associated with the contact lens-related dryness and discomfort, the importance of contact lens care solutions in overall lens wear cannot be discounted and a recent publication has shown that subjective satisfaction, particularly in symptomatic wearers, can be influenced by the combination of lens and solution prescribed.353 It is critical to note that a lens care solution is composed of several components. Therefore, it would be erroneous to conclude that any individual component in a care solution will have a direct impact on subjective symptoms. Based on the evidence to date, it appears that incorporation of surfactants or wetting agents into lens care products may improve subjective comfort, possibly by improving the hydrophilicity of the lens material. However, it is difficult to isolate a specific component in a lens care product and correlate that factor with improved subjective symptom.

There is adequate evidence that suggests different lens care solutions interact differently with various contact lens materials and this depends on the properties of both the contact lens material and lens care solution. Thus, the mechanisms contributing to symptomatology during lens wear may vary based on how the components in a lens care solution interact with the lens material.

### Physical Properties of Care Solutions

Soft contact lens care solutions are made of a wide range of components, as described in the previous section. The combination and concentration of these agents will have a significant impact on the physical properties of the solution and this could potentially influence patient comfort. The following section provides a brief overview of various physical properties of contact lens care solutions and whether these properties may have an impact on contact lens comfort.

**pH.** The pH of human tears ranges between 6.6 and 7.8,354 and the human eye is capable of tolerating pH values in the range of 6.2 to 9.0 at 0.2 M strength.355 When the pH of the contact lens solution falls outside this range, patients complain of ocular discomfort and stinging.356–358 Buffering agents used in soft contact lens solutions directly affect their pH, and it is possible that the type of buffer used in a particular solution could also affect subsequent patient comfort.

A study that investigated the pH of 10 different contact lens care solutions showed that most solutions (except nonneutralized peroxide systems) had pH values that were close to neutral and fell within the reported tolerable pH range for the ocular surface.359 The large difference in the pH of peroxide-based solutions before neutralization is the principal reason for burning, stinging, and epithelial cell damage seen in patients who mistakenly insert the nonneutralized solutions directly onto the ocular surface.359

**Viscosity.** Viscosity of a solution has the potential to influence patient comfort upon lens insertion or at the end of the day, through interactions between the solution, the lens, and the patient’s tear film. Viscosity of water is 1.0 cP and that of the tear film ranges between approximately 5.0 and 1.5 cP at 25°C for normal patients.340,341

A study that investigated the viscosity of various lens care solutions found that all the solutions had viscosity values that ranged between 0.96 and 1.26 cP, but some go as high as 3 cP.359 Several studies have determined the impact of HPMC-containing solutions on patient comfort.134,136,452 One study investigated the physical properties of multipurpose contact lens solutions with and without the addition of HPMC, and also determined if there are significant differences in the tear physiology of two groups of patients wearing soft contact lenses soaked in HPMC and non-HPMC solutions.134 This study showed that the prelens tear thinning time was longer and the prelens tear film structure was improved with use of the HPMC-containing solution.134 Another clinical trial that compared the prelens tear film thickness of etafilcon lens wearers showed that the patients who used the HPMC-containing solution showed a greater prelens tear film thickness (3.02 ± 1.07 μm) when compared with those that used a non-HPMC containing solution (2.72 ± 0.86 μm).124 This study also showed no statistical difference in study subjects’ preference for either solution, but nearly every subject (90.5%) suggested “comfort” as their reason for preference.124 In summary, it appears that lens care solutions that incorporate viscosity-enhancing agents can create a thicker and longer-lasting layer of fluid on hydrogel lenses and this can potentially lead to improvements in tear function in contact lens wearers. However, if the viscosity is too high, then potential blurring effects may mitigate these comfort advantages.
Osmolarity. Osmolarity of contact lens solutions could play a role in patient comfort, as studies have demonstrated that tear film osmolarity plays a significant role in the discomfort reported by dry eye patients. A study that investigated the osmolarity of 10 different contact lens care solutions showed that the osmolarity values fell between 275 and 310 mOsm/kg, indicating that the majority of soft contact lens solutions are hypo-osmotic compared with human tears. To our knowledge, no published studies have been conducted to investigate the relationship between solution osmolarity and contact lens comfort.

Surface Tension. The surface tension of pure water is approximately 72 mN/m and human tears have a surface tension value in the range of 40 to 46 mN/m. In a contact lens care solution, the presence/absence and type/number of surfactants will have a substantial impact on the surface tension of the solution. A study that investigated the surface tension of various care solutions showed that most multipurpose solutions have surface tension values that ranged between 29 and 40 mN/m. Among all the care solutions, the ones that do not incorporate surfactants (for example, peroxide-based systems and saline) had surface tension values that were close to that of water, whereas the solutions that had one or more surfactants had surface tension values that were closer to that of human tears. To our knowledge, no published studies have been specifically conducted to investigate the relationship between surface tension and comfort.

In summary, contact lens care solutions differ in certain physical properties and, by design, most care solutions fall within acceptable limits of ocular physiological tolerance. When properties of these solutions do not fall within the acceptable limits, clinically, this could result in burning, stinging, and epithelial cell damage. Minor shifts in the values may have the potential to influence patient comfort initially and/or at the end of the day. To date, very little has been published directly investigating the relationship between physical properties of lens care solutions with contact lens symptoms and this warrants further investigation.

Rewetting Drops

Contact lens wearers use rewetting drops for many reasons, including managing contact lens dry eye, lens dehydration and its associated dryness, general ocular lubrication, acting as a mechanical buffer between the lens and cornea, and lens surface rewetting and cleaning. Numerous formulations of lubricating eye drops exist and contain a wide variety of ingredients including cellulose derivatives, oil-based emulsions, paraffin, polyvinyl alcohol, polyacrylic acid, Polyvinylpyrrolidone, glyc erin, HA, hydroxypropyl guar, polyethylene glycol, and propylene glycol.

It has been reported that 47% of contact lens wearers use rewetting drops, but that they only provide moderate and nonsustained relief from symptoms of discomfort, in addition to inconvenience with the need for repeat instillation. There is the suggestion that a regimen of more than one type of lubricating eye drop may be needed for symptomatic contact lens wearers due to the multifactorial nature of CLD. If used in a proactive manner, the same eye drop has been shown to produce greater symptom relief than its use in a reactive manner. While there are a number of studies that demonstrate some level of symptom relief for contact lens wearers, there appears to be relatively little advantage with the use of rewetting drops or oral lubricants compared to the use of saline.

One study has shown that the use of a lubricant eye drop containing hydroxypropyl guar, propylene glycol, and polyethylene glycol (Systane) twice daily (pre- and post-contact lens wear) resulted in an increase in comfortable wear time and improved other subjective assessments of lens wear acceptability in symptomatic hydrogel lens wearers, compared with the use of a saline control drop.

Composition of Packaging Solutions

In an attempt to alleviate symptoms of dryness and discomfort, a number of manufacturers incorporate a variety of agents into both the lens material and also, more recently, the packaging solutions that are used to ship contact lenses. These so-called “blister-pack solutions (BPS)” now commonly include incorporation of water-soluble polymers, surfactants, and often unnamed “wetting agents” that have been previously described in this report. The alterations made to the BPS are to aid in preventing the lenses from sticking to the blister pack, enhance lens wettability, and improve initial comfort of the lenses after application to the eye.

One of the earliest published studies to suggest that adhering surfactants to lens materials may enhance in-eye comfort is that by Tonge and colleagues, who showed that soaking etafilcon A lenses in a solution containing poloxamine 1107 improved in-eye comfort and that the surfactant was retained for several hours after wear. To date, while several studies have examined comfort between lens materials with and without wetting agents (as previously described in this review) and studies have shown that contact angle wettability is raised in some materials once the BPS is removed, no published study has systematically examined whether the modified BPS has a direct impact on contact lens comfort. However, given the growth in the number of lens materials that are shipped in BPS that contain surface-active agents, it would seem likely that manufacturers have determined their positive impact in in-house unpublished studies.

Future Directions

Careful review of this report points toward several areas for future research that would enhance our understanding of CLD and the key factors associated with materials, design, and care systems.

One area that continues to be a frustration is the lack of association between in vitro data and their ability to predict in vivo performance. The industry would be well served by the development of in vitro methodologies that help to predict on-eye performance. An area that appears to show the most promise relates to the results from coefficient of friction evaluations. However, this field is in its relative infancy for contact lens materials and the development of methods that are more meaningful to on-eye comfort require substantial work. Current assessment methods vary widely, resulting in wide variation in the data obtained. International standardization of laboratory measurements such as frictional testing of hydrogels, as well as those for others such as contact angle and dehydration, would be a welcome addition to the industry. For frictional measurements, the relative importance of methodology, type of friction, and how these values relate to clinical observations such as lid wiper epitheliopathy, lid-parallel conjunctival folds, corneal and conjunctival staining, and in-eye wettability, are all areas worthy of future investigation. Other related areas that need substantial development relate to improved methods for determining in-eye wettability and investigation of the tear film in the vicinity of the lid wiper area and how these factors relate to comfort.

There is no doubt that contact lens materials change their hydration after being placed onto the ocular surface. It would...
be expected that a change in bulk surface hydration will impact wettability and friction. However, bulk dehydration shows only a tenuous relationship with comfort for most materials, but the development of improved methods to investigate surface dehydration (and impact on comfort) are more important than ever due to new materials that exhibit differing bulk and surface characteristics. When undertaking dehydration studies, factors such as the time intervals for assessment, standard procedures for sample handling, and eventual reinsertion in the ocular surface for subsequent measures, and control of environmental conditions and time of day all need standardization and agreement.

Much of the published clinical work in relation to soft lens material, design or solution properties has been poorly conducted, with inappropriate or missing controls, making conclusions regarding their impact on discomfort difficult. Future work investigating the impact of various characteristics must be conducted using well-controlled, randomized, cross-over studies in which all variables (replacement period, solution system, wearing time, etc.) are considered. This area requires some fundamental studies in which the isolation of a single change in a material, design, or solution characteristic is investigated. This work can only be conducted with the close cooperation of industry since this cannot be undertaken using only commercially available products. Areas of investigation of particular note relate to comparisons of some of the newer hydrogel-based materials against modern silicone hydrogels, comparisons between materials that “release” components into the tear film versus the base material without the release agent, and silicone hydrogels with standard or base surface wettability versus those with enhanced hydrogel-type coatings.

When it comes to bulk material properties, the trend thus far has been toward a modulus low enough to maximize on-eye comfort while balancing handling, durability, and tightness of fit. However, the conventional tensile modulus test involves unidirectional static loading; therefore, dynamic mechanical testing may be more appropriate, given the cyclic dynamic motion of eyelid movement coupled with the elastic and viscous flow characteristics of hydrogel materials (the cornea is also viscoelastic). For example, dynamic mechanical testing of silicone hydrogels demonstrates a characteristic rise in elastic modulus or shear-dependent elastic response that is typically not present in conventional hydrogels with similar EWC. There is the suggestion that, in order to improve comfort of current SiHy lenses, this elastic component should be similar to that for conventional hydrogels. However, this work has not been conducted in a systematic manner.

With regard to rigid lenses, increased interest in the use of RGP scleral lenses points toward a substantial number of potential studies in this area, with specific data being needed on comfort changes over the course of the day and comparative studies against corneal rigid lenses in nonpathological corneas. Large diameter RGP lenses offer an opportunity to provide a test platform with no bulk hydration changes, slower deposition, limited lens movement and minimal lid-lens edge interaction. Thus, studies comparing RGP scleral designs against both corneal rigid lenses and soft lenses may be of value. Of specific interest in the area of both rigid and soft lens design, methods to assess the impact of lens “edge” design on-eye must be developed, such that differing designs in identical materials can be compared. Methods to investigate tear replenishment and expulsion from beneath lenses must also be developed to aid us in understanding their impact on factors such as end-of-day comfort and inflammatory responses.

Further work to better understand the reasons behind the success of frequent replacement lenses is also needed, particularly with new materials. What is the optimal period of replacement for certain materials, and what are the methods used to determine this? How widely does this differ for different patients? What are the factors associated with the optimal replacement period? Is it due to changes in the material itself, the accumulation of certain tear film components, or those from the care system? What changes occur over time (over the day and over the lifetime of the lens) to both the ocular tissues in contact with the lens and the material itself? On a related note, studies to better understand the accumulation of tear film components remain to be undertaken. In particular, a better understanding of the impact of denatured proteins (other than lysozyme), lipid breakdown products and the deposition of many other tear film components are required.

There also remain large gaps in our knowledge of the role of care systems, packaging solutions and “comfort” drops on CLD. As with material-based studies, potentially valuable investigations in which systematic changes in various components are evaluated have yet to be undertaken. The short- and long-term impact of the uptake and release of lens care components and how they affect comfort are areas of future interest. Should care systems remove all tear film constituents that are deposited onto materials, or should they be designed to leave in place certain components that may help “biocompatibility”? If some components should be left in place, which ones and how much is “enough”? Finally, how effective is the delivery of wetting agents from the materials in reducing CLD and which agents are the most efficacious—and for how long?

Summary

In summary, a thorough review of the literature shows that there are surprisingly few proven links between CLD and factors related to the contact lens material, design, and care system. However, clinical acumen (in addition to recent studies) demonstrates that, in contact lens wearers who exhibit unacceptable comfort, making changes to the lens material, design, care system, and replacement schedule can improve comfort. It is also pertinent to consider, as pointed out in this review, the limitations of laboratory and academic studies, which might miss relevant variables present in the “real world.” Conclusions derived from well-conducted, well-controlled groups of subjects in a formal clinical trial might not be transferable to the thousands of patients that ultimately use the products, subjected to issues such as noncompliance, that may directly impact evaluation of comfort.

Much work remains to unravel the complexities of CLD. It is clear that a number of fundamental studies must be undertaken if an increased understanding of the role of materials, design and care regime in contact lens dryness is to occur. This will require substantial intellectual input and funding from both industry and academia alike.

Acknowledgments

Disclosure: Each workshop participant’s disclosure data can be found in the Appendix of the Introduction.

References


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APPENDIX A. Examples of Some Commonly Prescribed Hydrogel Contact Lens Materials

<table>
<thead>
<tr>
<th>Commercial Name</th>
<th>Manufacturer</th>
<th>Water Content</th>
<th>CT</th>
<th>Dk/t</th>
</tr>
</thead>
<tbody>
<tr>
<td>SofLens 38 (polymacon)</td>
<td>Bausch + Lomb</td>
<td>38</td>
<td>0.035</td>
<td>22</td>
</tr>
<tr>
<td>Biomedics 55 (ocufilcon D)</td>
<td>CooperVision</td>
<td>55</td>
<td>0.07</td>
<td>21</td>
</tr>
<tr>
<td>Acuvue 2 (etafilcon A)</td>
<td>Johnson &amp; Johnson</td>
<td>58</td>
<td>0.084</td>
<td>20</td>
</tr>
<tr>
<td>SofLens daily disposable (hilafilcon B)</td>
<td>Bausch + Lomb</td>
<td>59</td>
<td>0.09</td>
<td>19</td>
</tr>
<tr>
<td>PROCLEAR (omafilcon A)</td>
<td>CooperVision</td>
<td>62</td>
<td>0.065</td>
<td>30</td>
</tr>
<tr>
<td>Focus Dailies (nelfilcon A)</td>
<td>CIBA Vision</td>
<td>69</td>
<td>0.10</td>
<td>26</td>
</tr>
</tbody>
</table>

APPENDIX B. Examples of Some Commonly Prescribed Silicone Hydrogel Contact Lens Materials

<table>
<thead>
<tr>
<th>Commercial Name</th>
<th>Manufacturer</th>
<th>Water Content</th>
<th>CT</th>
<th>Dk/t</th>
<th>Modulus, MPa</th>
<th>Surface Treatment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air Optix Night &amp; Day Aqua (lotrafilcon A)</td>
<td>Alcon</td>
<td>24</td>
<td>0.08</td>
<td>175</td>
<td>1.4</td>
<td>25-nm plasma coating</td>
</tr>
<tr>
<td>Air Optix Aqua (lotrafilcon B)</td>
<td>Alcon</td>
<td>33</td>
<td>0.08</td>
<td>138</td>
<td>1.0</td>
<td>25-nm plasma coating</td>
</tr>
<tr>
<td>DAILIES TOTAL1 (delefilcon A)</td>
<td>Alcon</td>
<td>33</td>
<td>0.09</td>
<td>156</td>
<td>0.7</td>
<td>Soft surface gel with &gt;80% water content</td>
</tr>
<tr>
<td>PureVision (balafilcon A)</td>
<td>Bausch + Lomb</td>
<td>36</td>
<td>0.09</td>
<td>101</td>
<td>1.1</td>
<td>Plasma oxidation process</td>
</tr>
<tr>
<td>Acuvue OASYS (senofilcon A)</td>
<td>Johnson &amp; Johnson</td>
<td>38</td>
<td>0.07</td>
<td>147</td>
<td>0.72</td>
<td>None; internal wetting agent (PVP)</td>
</tr>
<tr>
<td>Acuvue Advance (galyficon A)</td>
<td>Johnson &amp; Johnson</td>
<td>47</td>
<td>0.07</td>
<td>86</td>
<td>0.40</td>
<td>None; internal wetting agent (PVP)</td>
</tr>
<tr>
<td>1 Day Acuvue TrueEye (narafilcon A)</td>
<td>Johnson &amp; Johnson</td>
<td>46</td>
<td>0.085</td>
<td>118</td>
<td>0.66</td>
<td>None; internal wetting agent (PVP)</td>
</tr>
<tr>
<td>BIOFINITY (comfilcon A)</td>
<td>CooperVision</td>
<td>48</td>
<td>0.08</td>
<td>160</td>
<td>0.75</td>
<td>None</td>
</tr>
<tr>
<td>AVaira (enfilcon A)</td>
<td>CooperVision</td>
<td>46</td>
<td>0.08</td>
<td>125</td>
<td>0.50</td>
<td>None</td>
</tr>
<tr>
<td>Menicon PremiO (asmofilcon A)</td>
<td>Menicon</td>
<td>40</td>
<td>0.08</td>
<td>161</td>
<td>0.90</td>
<td>Plasma oxidation</td>
</tr>
<tr>
<td>Clariti (Filcon II 3)</td>
<td>SaulIon</td>
<td>58</td>
<td>0.07</td>
<td>86</td>
<td>0.50</td>
<td>Nondisclosed</td>
</tr>
<tr>
<td>Definitive (erofilcon A)</td>
<td>Contamac</td>
<td>75</td>
<td>0.08*</td>
<td>76</td>
<td>0.35</td>
<td>None</td>
</tr>
</tbody>
</table>

PVP, polyvinyl pyrrolidone; USAN, United States Adopted Name.

* Estimated as lathe-cut lens designed to practitioner specification.