Assessment of Contact Lens with the High Definition (HD) Property

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Abstract

Background: Contact lenses have been in use since the 1930s. During these years, many materials have been considered with the aim of producing more advanced and biocompatible devices. The 1940s was the era of the first modern contact lenses, with the debut of polymethyl methacrylate (PMMA). The lack of oxygen supply to the cornea was the main disadvantage of PMMA. This oxygen impermeability was solved with the development of different contact lens materials, new generation of contact lens use hydrogel and silicone to enhance oxygen permeability. Recently, the new contact lens material known as silicone hydrogel lenses has been used to produce a new lens with the HD property.

Aim of Study: The aim of this study was to assess a new high definition silicone hydrogel contact lens.

Patients and Methods: 22 subjects have been recruited from King Saud University students (male, age 21 ± 1.5 years). The new Balafilcon A lenses (PUREVISION 2 HD); (Bausch + Lomb, Tokyo, Japan) has been assessed and compared with another (non HD) silicone hydrogel lens Balafilcon A (PureVision, Bausch & Lomb Inc., Rochester, NY).

The participants asked to wear each of the two lenses for one week (the lens was given randomly) and 48 hours wash out period was allowed between the lenses. Clinical tests including visual acuity (VA) and contrast sensitivity (CS) were applied to evaluate the lens performance at the end of each week. Additionally, contact lens dry eye questionnaire (CLDEQ-8) has been used to obtain the subjective feedback.

Result: The mean of CLDEQ-8 with PureVision was 8.1 and with Pure Vision 2 HD was 6. For contrast sensitivity the mean with PureVision was 1.74 while with Pure Vision 2 HD was 1.77. The mean of Visual Acuity was equal with the two lenses 1.00 log. Statistical tests using Wilcoxon-signed rank test showed no significant difference exist between the HD PureVision 2 and PureVision lens. For Visual Acuity (p=0.317), Contrast sensitivity (p=0. 150) (Fig. 2) and CLDEQ-8 (p=0.339).

Conclusion: The silicone hydrogel HD lens evaluated in this study showed good performance. However, no significant difference in clinical performance was observed between the HD lens and the non HD silicone hydrogel lens.

Key Words: Contact lens – Silicone hydrogel – High definition (HD) – Contrast sensitivity.

Introduction

CONTACT lenses have been in use since the 1930s. Since that time many materials have been considered with the aim of producing more advanced devices. The first soft contact lenses were made from 80% 2-hydroxyethyl methacrylate (HEMA) and 20% diethylene glycol methacrylate (DEGMA); later, pure HEMA was used, on its own, resulting in improved mechanical properties of lenses [1].

In 1979 Tanaka et al. [2] attempt to combine hydrogel with silicone. They successfully combined HEMA with a monomer (trimethylsiloxy) – methacryloxy-propylsilane (TRIS) producing a rigid lens material [2]. Tanaka also suggested inserting a polar group (hydroxyl) to modify TRIS. Co-polymerising TRIS with hydroxyl (hydrophilic monomer) resulted in a material suitable for use as a soft contact lens.

The co-polymerising of TRIS with a hydroxyl group successfully enhanced the oxygen permeability but did not address the issue of limited lens movement. This issue can be explained by referring to two approaches to improving ocular compatibility which was used in the development of the first silicone hydrogel lenses.

The first approach was the incorporation of fluorine into the hydrogel structure. This improved three characteristics of contact lens: wettability, mechanical properties and oxygen permeability [3].

The second approach was the development of macromer technology. “Macromers are large monomers formed by pre-assembly of structural units

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that are designed to bestow particular properties on the final polymer” [4]. The use of macromers and their role in enhancing oxygen permeability was explained in 1991 by Robertson et al. [5]. Their construction involved the combination of hydrophilic polyethylene oxide segments with oxygen gas permeable polysiloxane units. These two approaches, TRIS modification and macromer technology, are the basis of the principle used to develop the first commercial silicone hydrogel lenses: PureVision (Bausch & Lomb, Rochester, NY) and Focus Night & Day (now Air Optix Night & Day, Ciba Vision) [6].

Up until 2005, all silicone hydrogel contact lens materials either depended on modified TRIS (PureVision), alone, or together with silicone based macromers (AirOptix Night & Day, Acuvue Advance and Acuvue Oasys).

In 2006 a new range of silicone hydrogel contact lenses was released. This family of silicone hydrogel lenses included Biofinity (Comfilcon A, CooperVision, Inc) and PremiO (Asmofilcon A, Menicon). More recently Avaíra (Enfilcon A, CooperVision, Inc) has been released [7].

In 2007, Menicon Co, Ltd. (Japan) introduced PremiO lens for daily wear on a two-week replacement basis. The PremiO lens incorporates a plasma surface treatment (Nanoglass™ technology) to improve comfort and reduce dryness [8].

Up until now the development in contact lenses is continuous, the contact lenses manufacturers are working hard to develop lenses more compatible and less effective on cornea and tear film stability.

As we find the sign HD (High Definition) on most of the TV screens, computers, mobiles, camera etc. The Bausch & Lomb also produce soft contact lens known as PureVision 2 HD lenses.

For the screens and cameras, the HD means high definition or more pixels leading to a brighter, clear picture. For the HD contact lenses according to the manufacturer, the HD lens improves the quality of vision by using what they call “high definition Optics”.

The Pure Vision 2 HD lens is supposed to reduce spherical aberration [9], which usually affects the vision in low light. Patient with spherical aberration often suffer from impaired vision, halos or blurred vision and glare in low light [10].

The Pure Vision 2 HD was described as one of the most breathable lens available, allowing more oxygen to the cornea and they are very thin which is supposed to enhance ocular comfort [11].

The aim of this study is to investigate the performance of the new silicone hydrogel HD lens. Due to the special design of the HD lens to reduce optical aberration, clinical differences between this new HD lens and the other silicone hydrogel lens are expected to exist.

The two important goals of any contact lenses are improving vision and maintaining ocular comfort. So three common clinical tests have been applied here including visual acuity (VA) test [12], contrast sensitivity (CS) test [13] and subjective assessment with contact lens questionnaire [14-17].

**Patients and Methods**

22 subjects have been recruited from King Saud University students (male, age 21 ± 1.5 years) and asked to take part in this study. The inclusion criteria were >18 years old, no history of ocular or systemic diseases reported. Participants with systemic or ocular disease have been excluded from the study.

This study was conducted at King Saud University, Riyadh, Saudi Arabia during the period from February to April 2018.

This study followed the tenets of the Declaration of Helsinki [18]. Informed consent was obtained from all individual participants included in the study.

The new Balafilcon A lenses (PUREVISION 2 HD); (Bausch + Lomb, Tokyo, Japan) performance has been assessed and compared with another (non HD) silicone hydrogel lens Balafilcon A (PureVision, Bausch & Lomb Inc., Rochester, NY) (Table 1).

The participants are asked to wear each of the two lenses for one week (the lens was given randomly) and 48 hours wash out period was allowed between the lenses. Clinical tests including visual acuity (VA) and contrast sensitivity (CS) were applied to evaluate the lens performance at the end of each week. Additionally contact lens dry eye questionnaire (CLDEQ-8) [16] has been used to obtain the subjective feedback.

**Pure Vision HD Lens:**

PureVision contact lenses with high definition optics are designed to reduce spherical aberration
Contrast sensitivity (log unit)

1.8
1.7
1.6
1.5
1.4

across the entire power range. Aberration is a distortion acquired by a wave-front of light when it passes through an eye with irregularities.

**Visual acuity (VA test):**

The visual acuity test is used to determine the smallest letters you can read on a standardized chart (Snellen chart) or a card held 20 feet away.

**Contrast sensitivity:**

The Pelli-Robson test measures contrast sensitivity using large letters as targets (equivalent to 20/60 acuity). For each group of 3 letters, contrast is decreased from left to right and from the top to the bottom of the chart. The lowest contrast at which 2 or 3 of the letters in a group can be read determines a log contrast sensitivity score. The Pelli-Robson contrast sensitivity chart is an 86x63cm chart that consists of 16 triplets of 4.9cm (2.8° at 1m) letters, and it assesses contrast sensitivity (CS) at a spatial frequency of about 0.50 to 1 cycle/degree. Within each triplet, the letters have the same contrast, and the contrast in each successive triplet decreases by a factor of 0.15 log units (Fig. 1).

**CLDEQ-8:**

The CLDEQ-8 has 5 parts, it evaluates dryness, discomfort, and “blurry vision”; frequency of “closing eyes” due to discomfort and “removing Contact Lens to relieve discomfort. It measures the impression lenses for patients by assessing each part from 0 to 4 where 0 gives the best impression and full satisfaction.

The recent International Workshop on Contact Lens Discomfort reported that the contact lens dry eye questionnaire (CLDEQ) is the only tool validated for the evaluation of contact lens induced discomfort [14].

**Results**

The test of normality using (Kolmogorov-Smirnov) showed that the data was normally distributed ($p>0.055$). The statistical tests using Wilcoxon-signed rank test showed that no significant difference between the PureVision 2 HD and PureVision lens. For Visual Acuity ($p=0.317$), Contrast sensitivity ($p=0.150$) (Fig. 2) and CLDEQ-8 ($p=0.339$) (Fig. 3).

The mean of CLDEQ-8 with PureVision was 8.1 and with Pure Vision 2 HD was 6. For contrast sensitivity the mean with PureVision was 1.74 while with Pure Vision 2 HD was 1.77. The Visual Acuity was equal with the two lenses 1.00 log.
Discussion

This study investigates the clinical performance of new HD silicone hydrogel lens and compares it with one of the commonly use non HD silicone hydrogel.

The two lenses studied here showed good performance in term of subjective assessment with CLDEQ-8. According to Chalmers et al. [19] the CLDEQ-8 score ≥ 12 is diagnosed as SCL related symptoms. The CLDEQ-8 with PureVision 2 HD was slightly better (score 6) than that with PureVision (score 8.1). This might be due to the thin thickness [9] of the Pure Vision 2 HD (0.07mm) compared to PureVision (0.08mm) [20]. However, none of the two lenses assessed showed score more than the CLDEQ-8 cut-off value ≥ 12 and no significant difference between the lenses was observed.

The contrast sensitivity and visual acuity evaluated of the subject when they are wearing Pure Vision 2 HD and PureVision was not significantly different. The contrast sensitivity mean with PureVision 2 HD was 1.77 log and with PureVision was 1.74 log.

For patients between 20 and 50 years old, CS should be 1.80 log units and above; for patients less than 20 years old and older than 50 years, monocular CS should be 1.65 log units and above [21]. In our study, we observed that CS with PureVision 2 HD was slightly better and very close to the normal (1.80 log units).

No difference was observed regarding the visual acuity with the two lenses. Also, both lenses showed a good subjective feedback and good clinical performance.

The Pure Vision 2 HD lens (Balafilcon A) material is a first generation silicone hydrogel, plasma oxidation surface treated lens. PureVision 2 lenses with High Definition Optics are designed to reduce halos and glare and deliver clear vision. It offers design advances for outstanding comfort and breathability.

The similarity in subjective response observed between the two soft lenses evaluated in this study might be because both lenses have their surface treated with a layer of plasma which makes the surface hydrophilic, better tear film stability therefore more comfort to the patient [22].

These two lenses have good properties in term of water content, oxygen permeability, modulus, thickness and wettablity [23-25].

These good properties might reflect on the clinical performance of the two lenses. However, a longer wearing time might be required to reach the full judgment on such lens. Additionally, objective assessment of the corneal abrasion with this new lens should be targeted in future study.

Conclusion:
The silicone hydrogel HD lens evaluated in this study showed good performance. However, no significant difference was observed when the HD lens compared with a non HD silicone hydrogel lens.

Conflict of interest:
No conflict of interest to declare.

References


تقييم العدسات اللاصقة ذات خصائص الواضح العالي

الغرض: تتم الاستخدام منذ الثلاثينيات القرن العشرين, تم النظر في العديد من المواد بهدف إنتاج عدسات أكثر تقدمًا وموثوقًا جيّداً. كأن النوعية من القرن العشرين رجاء العدسات الاصطناعية الحديثة الأولى, مع ظهور مادة البوليمر. مبتكرًا, كان منع وصول الأوكسيجين إلى القرنية هو المهم الرئيسي لمادة البوليمر. مبتكرًا بسبب ضعف نقاوة المادة للأكسجين, لا حقاً تم حل تكون الأكسجين هذه مع تطور مواد عدسات لاصقة متحركة, وذلك بإضافة مادة السيليكون إلى الهيدروجين لتعزيز نقاوة العدسة للأكسجين. في الأونة الأخيرة, تم استخدام مواد العدسات الاصطناعية الجديدة المعروفة باسم عدسات السيليكون الهيدروجين لإنتاج عدسات جديدة تدعم خاصية الواضح العالي.

الطريقة: تم دعوة 22 شخصًا من طلاب جامع الملك سعود (ننزر, أعمارهم 21±15 عام للمشاركة في هذه الدراسة. تم استخدام العدسات الاصطناعية للفيلكون أي (ببور فيجن علية الواضح) (بون أد لويم, طوكو, اليابان) وممارسة أداء العدسة مع عدسات السيليكون الهيدروجين أخرى (ببور فيجن) بون أد لويم, روتشستر, نيويورك).

طلب من المشاركين ارتداء كل من العدسات لمدة أسبوع واحد (أظهرت العدسات بشكل عشوائي) وتم منح فترة 48 ساعة بين العدسات لتقدير داخل التأثير بين العدسات. تم تطبيق الاختبارات السريرية بما في ذلك حدة النظر وحساسية التباهي لتقدير أداء العدسة في نهاية كل أسبوع بالإضافة إلى ذلك, تم استخدام استبان الخاص بالأعصاب للدراسة التقييم الشخصي لمستخدم العدسات.

النتائج: تم تقييم النتائج باستخدام عدسات ببور فيجن كان (8.1) مع ببور فيجن علية الواضح كان 6. وبالنسبة لحساسية التباهي كان المتوسط (1.74) لعدسة ببور فيجن, بينما عدسة ببور فيجن علية الواضح (7.7) وكانت حدة البصر متساوية في العدسات 1.00.

أظهرت الاختبارات الإحصائية عدم وجود مخالفة إحصائية بين العدسات ببور فيجن وبور فيجن علية الواضح. حدة البصر (0.317), حساسية التباهي (0.150), والاستبان (0.399).

الخلاصة: أظهرت العدسات ذات خصائص الواضح العالي التي تم تقييمها في هذه الدراسة أداء جيداً. ولكن، لم يلاحظ أي أداء إكلينيكي مختلف بين العدسات علية الواضح والعدسة غير علية الواضح على مستوى الرؤية.