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Evaluat on of safety and ef cacy of a new
multipurpose disinfecting solution on silicone hydrogel
contact lenses

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Abstract
Purpose: To evaluate the safety and ef cacy of a new multipurpose disinfecting solution (MPDS)
with a formulation that includes aloe vera on its composition.
Methods: This is a prospective, randomized, double-masked clinical trial with a crossover design
that included seven examinations. Two different MPDSs, Avizor Alvera® (study solution) and All
Clean Soft® (control solution), each were used for 1 month. Comilcon A silicone hydrogel
contact lenses were used during the trial. The main outcome variables were corneal staining
and deposits on the surfaces of the contact lenses. Other parameters including ocular surface
response, contact lens wettability, user satisfaction, and adverse events, were analyzed according
to the International Organization for Standardization (ISO) 11980:2010 guidance for clinical
investigation.
Results: Twenty subjects (10 women, 10 men) (mean age, 27.7 ± 5.6 years; range, 20-41) were
included. No differences between both MPDSs were found in the percentage of subjects with
corneal staining >0 at day 30 (study: 35%, control: 50%; p = 0.46); neither in the percentage of
subjects with deposits on the surface of the contact lens >0 at day 30 (study: 26.32%, control:
52.63%; p = 0.18). The study MPDS received higher rates in comfort (study: 8.14 ± 1.09, con-
trol: 7.94 ± 0.92; p = 0.56) and satisfaction at day 30 (study: 8.63 ± 0.91, control: 8.29 ± 0.80;
p = 0.19), however the scores were not signi cantly different with the control MPDS.
Conclusions: The clinical trial showed that the study MPDS is safe, ef cient, and has acceptable
physiologic tolerance, according to the ISO 11980:2010 guidance for clinical investigation.
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Evaluación de la seguridad y eficacia de una nueva solución multipropósito en lentes de contacto de hidrogel de silicona

Resumen

Objetivo: Evaluar la seguridad y la eficacia de una nueva solución única desinfectante (MPDS), con una formulación que incluye aloe vera en su composición.

Métodos: El estudio consistió en un ensayo clínico prospectivo, aleatorio y de doble ciego, con un diseño cruzado que incluyó siete exámenes. Se utilizaron durante un mes dos MPDS diferentes, Avizor Alvera® (solución en estudio) y All Clean Soft® (solución de control). Durante el ensayo se utilizaron lentes de contacto de hidrogel de silicona Comficon A. Las principales variables evaluadas fueron la tinción corneal y los depósitos sobre la superficie de las lentes de contacto. Se analizaron otros parámetros, que incluían la respuesta de la superficie ocular, la humectabilidad de las lentes de contacto, y los eventos adversos, de acuerdo con la normativa 11980:2010 de la Organización Internacional de Normalización (ISO) para investigación clínica.

Resultados: El estudio incluyó a veinte sujetos (10 mujeres, 10 hombres) (edad media, 27,7 ± 5,6 años; rango, 20-41). No se hallaron diferencias entre ambas MPDS en cuanto al porcentaje de sujetos con tinción corneal >0 al mes (estudio: 35%, control: 50%; p = 0,46), ni el porcentaje de sujetos con depósitos en la superficie de las lentes de contacto >0 al mes (estudio: 26,32%, control: 52,63%; p = 0,18). La MPDS en estudio obtuvo valoraciones superiores en cuanto a confort (estudio: 8,14 ± 1,09, control: 7,94 ± 0,92; p = 0,56) y satisfacción (estudio: 8,63 ± 0,91, control: 8,29 ± 0,80; p = 0,19); sin embargo, las puntuaciones no fueron significativamente diferentes en relación a la MPDS de control.

Conclusiones: El ensayo clínico reflejó que la MPDS en estudio es segura y eficaz, mostrando una tolerancia oculistica aceptable, de acuerdo con la normativa ISO 11980:2010 para investigación clínica.

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1 year of habitual CL wear (at least 5 days weekly for at least 5 h daily). Both eyes were examined, but only data from one eye was chosen randomly for analysis. Subjects were excluded if they had an active ocular surface disease or a systemic disease that could affect ocular physiology or worsen with CL wear. The exclusion criteria included tolerance or sensitivity to the study products, ocular infection or a history of herpetic keratitis, severe inflammation during the 6 months before the study, use of an ocular topical medication in the previous 3 months (except artificial tears) or a systemic medication that could possibly affect the tear film, refractive surgery, corneal irregularities, pregnancy and breastfeeding, corneal edema, corneal neovascularization or infiltrates graded 1 or higher, tarsal or conjunctival hyperemia graded 2 or higher, fluorescein corneal staining graded 2 or higher in any area, the sum of all the areas graded 4 or higher, or greater than 20% staining.

**CLs and solutions**

Four pairs of silicone Si-HCLs (Biofinity™ and Biofinity Toric™, Comfilcon A, Cooper Vision Inc., Fairport, NY, USA, base curve: 8.60 mm, diameter: 14.00 mm, power range from +8.00 to −10.00, and cylinder up to 2.25 D) were dispensed to each subject during the trial: two pairs for the two washout periods (1 week each), and two pairs for the study or control solution periods (30 days each). Subjects were instructed not to sleep or nap while wearing the CLs during the study and to wear the provided CLs a minimum of 5 h daily and 5 days weekly during the study.

The study (Avizor Alvera, Avizor S.A.) and control (All Clean Soft, Avizor S.A.) solutions were randomly assigned. The use of the MPDS was randomized and double-masked during the study. Both solutions were masked by the manufacturer. The packaging of both solutions was identical. A 5-digit code was randomly assigned to each bottle. Two codes were assigned to each subject, one corresponding to the control solution and other one to the study solution. The solutions delivery and the evaluation of subjects were performed by different evaluators. Patients received written instructions regarding the use and maintenance of their Si-HCLs. The composition of both MPDSs is shown in Table 1. A hydrogen peroxide system was provided (Ever Clean, Avizor S.A.) for washout periods.
Table 1 Composition of the multipurpose disinfecting solutions used in the study.

<table>
<thead>
<tr>
<th>Brand</th>
<th>Components</th>
<th>Other</th>
</tr>
</thead>
<tbody>
<tr>
<td>Avizar Alvera</td>
<td>Borax/boric acid</td>
<td>EDTA, polyhexanide 0.0002% Poloxamer</td>
</tr>
<tr>
<td>All Clean Soft</td>
<td>Borax/boric acid</td>
<td>EDTA, polyhexanide 0.0002% Poloxamer</td>
</tr>
<tr>
<td>Ever Clean</td>
<td>Hydrogen peroxide 3%</td>
<td>Polvinylypyrrolidone aloe vera Phosphates</td>
</tr>
</tbody>
</table>

Study schedule

All subjects completed seven examinations with two washout periods. The study design followed the requirements of the International Organization for Standardization (ISO) 11980:2010 guidance for clinical investigation (Ophthalmic optics, CL, and CL care products).

Baseline examination

The subjects attended this examination without wearing CLs for at least 24 h. The clinical trial procedures were explained to the subjects, and each subject provided informed consent and signed a data protection form.

Medical and CL histories were obtained and the subjects underwent an ocular examination. Subjective refraction and LogMAR visual acuity (LogMAR VA) with an ETDRS chart was measured; keratometry and slit-lamp evaluation of the ocular surface were performed to determine if they met the inclusion and exclusion criteria. Women of childbearing age who wished to participate in the study should perform a pregnancy test. All subjects who complied with the inclusion and exclusion criteria were scheduled for the next examination within the following 10 days after baseline examination.

Examination 1

Subjects attended the examination without wearing CLs for at least 24 h. Changes in health status or medication and possible adverse events since the previous examination were scored; if the medication change might affect the ocular surface the patient discontinued the study. A new Si-HCL with the subjects parameters was inserted, and after waiting 30 min the following procedures were performed: measurement of the LogMAR VA, assessment of the Si-HCL fit (corneal coverage, position, and mobility), and evaluation of limbal and bulbar hyperemia. The subjects wore this Si-HCL for 1 week during the first washout period and received a hydrogen peroxide solution (Ever Clean) for cleaning and maintaining the Si-HCLs.

Examination 2

This examination was 7 ± 1 days after examination 1. The subjects attended the clinic after having worn the Si-HCLs for at least 5 h. Changes in health status or medication and possible adverse events since the previous examination were scored. Biomicroscopy signs evaluated were bulbar and limbal hyperemia, presence of papillae and follicles, epithelial and stromal edema, corneal infiltrates, endothelial regularity, corneal vascularization, anterior segment inflammation and bulbar and corneal staining. After the eyes were rinsed, a new pair of Si-HCLs was inserted and after 60 min, the LogMAR VA, CL fit, bulbar and limbal hyperemia, and deposits on the anterior and posterior CL surfaces were evaluated. This pair of Si-HCL was worn for 30 days (at least 5 h daily and 5 days weekly) using the assigned MPDS solution (study or control, randomly assigned) according to the oral and written instructions provided.

Examination 3

This examination was 15 ± 2 days after examination 2. At this visit the same procedure as in the previous visit was followed. Tests performed at this visit were: subject comfort and satisfaction, LogMAR VA, non-invasive break-up time (NIBUT), CL fit and grade of deposits and biomicroscopy signs.

Examination 4

This examination was 30 ± 2 days after examination 3. The same process as examination 3 was performed. The Si-HCLs and MPDSs for this part of the trial were collected, and a new pair of Si-HCLs was inserted. After 30 min the LogMAR VA, Si-HCL fit, and bulbar and limbal hyperemia were evaluated. This pair of lenses was worn for 1 week, corresponding to the second washout period. The care solution for this phase was the same hydrogen peroxide system used in the first washout period between examinations 1 and 2.

Examinations 5, 6, and 7

The same procedures and same schedule were followed as for examinations 2, 3, and 4, respectively. The subjects received the second randomized study or control solution.

Clinical measurements

The main safety variable was defined as the percentage of subjects without corneal fluorescein staining or a maximum of grade 1 staining at the end of the month using the study MPDS versus the control MPDS.

To assess the corneal fluorescein staining, 5 µl of 2% sodium fluorescein were instilled in the lower conjunctival cul-de-sac with a micropipette (Finnpipette® 0.5–10 µl; Vantaa, Finland). Two minutes later, the anterior corneal surface was examined by slit-lamp with a 16 × magnification lens and the cobalt blue filter and through a Wratten #12
yellow filter. The cornea was evaluated using a grid with five areas (central, temporal, nasal, superior, and inferior). The intensity of the corneal fluorescein staining was evaluated in each area and scored on a scale from 0 to 4.9

The main efficacy variable was defined as the difference in the percentage of subjects without deposits on the anterior or posterior CL surfaces or maximum of grade 2 at the end of the month using the study MPDS versus the control MPDS. Deposits were evaluated by slit-lamp with 20× magnification and white light, evaluating the CL anterior and posterior surfaces while the CL was on the eye, and the deposits were graded on a scale of 0–4.9

Secondary variables evaluated were: subject comfort and satisfaction, CL anterior surface wettability (NIBUT), lens fitting characteristics, and ocular surface health status evaluated by slit-lamp.

Patient comfort and satisfaction were evaluated using a visual analog scale,10 which consisted of a 10-centimeter-long vertical line on which each centimeter was marked with a horizontal line and numbered from bottom to top with 0 to 10. With a horizontal line, the subjects indicated the degree of comfort and satisfaction achieved with their CLs at the time of the evaluation. Scores at the lower end indicated extreme discomfort or extreme dissatisfaction with the CLs; scores at the upper end indicated extreme comfort or extreme satisfaction with the CLs. The evaluation was performed by measuring in millimeters the space from the bottom (0) up to the top (10). User comfort also was analyzed by assessing the use of artificial tears and the hours of daily CL use.

To analyze the CL anterior surface wettability, the NIBUT was measured using TearScope Plus® (Keele, Windsor, Berkshire, England, UK) by projecting its light grid on the anterior CL surface.11 The subjects were instructed to blink three times and then keep their eyes open until instructed to blink again. The time in seconds that elapsed was recorded from when the subject stopped blinking until the reflected image was deformed, indicating tear film discontinuity. This measurement was performed three times, and the average of the three measurements was recorded.

The characteristics of lens fitting (corneal coverage, position, and mobility) were assessed according to the standardized scale.9 With a slit-lamp it was determined if the CL completely covered the cornea. The scale used to assess the position ranged from 0 to 2, with 0 indicating a suitable and centered position, 1 a slightly off-centered position, and 2 an excessively off-centered position. Regarding mobility, the scale used ranged from −2 to +2, with negative values indicating lack of mobility, 0 suitable mobility, and positive values excessive CL mobility.

Ocular surface health was evaluated at the slit-lamp. All signs were evaluated on a scale of 0 to 4.9

The same experienced practitioner examined all patients.

**Adverse ocular events and severe adverse events**

Adverse ocular events were defined as reduction of at least one line of LogMAR VA that could not be improved by the instillation of artificial tears, the presence of corneal edema, corneal infiltrates, corneal vascularization, ulcers, any corneal event resulting in permanent opacity, or any severe adverse ophthalmic event. A severe adverse event was defined as any event resulting in death or threat to the life of the subject, permanent disability, required an extended hospital stay, involved cancer and congenital abnormalities, or was the result of an overdose (administration of a dose higher than prescribed).12

**Statistical analysis**

Statistical analyses were performed using SPSS 18.0 statistical software for Windows (IBM Corporation, Armonk, NY, USA) and R version 3.1.0. The Shapiro–Wilks test was used to check the normality of distribution. $p \leq 0.05$ was considered statistically significant.

Quantitative variables were described using means and standard deviations (SD). Qualitative ones were summarized by percentages.

For the primary safety and efficacy variables, the differences in the percentage of subjects were evaluated by equality proportion test for paired groups. The same statistical method was used to compare differences in qualitative variables between control and study groups. Mean differences for quantitative variables were checked by $t$-test for paired samples or its non-parametric alternative (Wilcoxon test) when the normality assumption could not be assumed.

**Results**

Twenty subjects (10 men, 10 women) were included in the study. The average patient age was $27.7 \pm 5.6$ years (range, 20–41 years). All subjects met the inclusion criteria, and no subjects discontinued the study. All CL were exhibiting good fit throughout the study.

**Primary safety variables**

Most subjects had fluorescein staining values of grade 0 or a maximum of grade 1 staining, except for one subject with grade 2 staining after 15 days using the control MPDS and four subjects with grade 2 staining after 1 month using the study MPDS (Fig. 2). These differences were not significant. Staining higher than grade 2 did not occur during the study.

By assessing the percentage of subjects with corneal staining exceeding grade 0 with both MPDSs and for the three examinations, we observed a higher percentage after 15 and 30 days of use of the control MPDS (50% in both cases) compared with the test MPDS (40% and 35%, respectively). These differences were not significant.

No adverse events were detected that were either related directly to the study or control MPDS or the CL used, and no severe adverse events developed throughout the study.

**Primary efficacy variable**

Most subjects had anterior surface deposits that were grade 0 or grade 1 maximally, except for one subject with grade 2 deposits after 1 month using the control MPDS and two subjects with grade 2 deposits after 1 month using the study
Safety and efficacy of a new multipurpose disinfecting solution

Corneal staining (primary safety variable). The data show the percentage of subjects with corneal fluorescein staining on 0, 15, and 30 days of use of each multipurpose disinfecting solution (MPDS) (study and control). No staining higher than grade 2 was seen throughout the study.

Figure 2

Deposits on the front surface of the hydrogel contact lens (Si-HCL) (primary efficacy variable). The data show the percentage of subjects with deposits on the front surface of the Si-HCL on 0, 15, and 30 days of use of each multipurpose disinfecting solution (MPDS) (study and control). No staining higher than grade 2 occurred throughout the study.

Figure 3

Subject comfort and satisfaction with the study and control multipurpose disinfecting solutions (MPDSs). The data show the average comfort and satisfaction values of the subjects on days 15 and 30 of use of each MPDS. VAS: visual analog scale.

Figure 4

Secondary variables

Subject comfort and satisfaction were assessed after 15 and 30 days of use of both MPDSs. We observed higher average values of comfort and satisfaction for the study MPDS than for the control MPDS throughout the study; however, these differences were not significant (Fig. 4).

No subject reported the need for artificial tears while wearing the CLs throughout the study. No differences were found in the hours or days of CL use between the MPDSs (hours of use/week, study MPDS, 50.3 ± 14.93; control MPDS, 58.16 ± 12.33, p = 0.2453).

Higher NIBUT values for the control MPDS were observed after 15 and 30 days, with a significant (p = 0.0481) difference at 15 days and a difference between the MPDSs of 1.62 ± 3.436 seconds. This difference decreased after 30 days and was no longer significant (0.75 ± 3.54 seconds; p = 0.3934) (Fig. 5).

Epithelial edema, infiltrates, endothelial regularity, follicles and anterior segment inflammation did not change throughout the study; all subjects had grade 0 at all examinations.

Analysis of the data obtained for bulbar hyperemia, limbal hyperemia, conjunctival staining, papillae, stromal edema, and corneal vascularization showed no significant differences in any variables at any examinations between the MPDSs (Table 2). Nevertheless, we observed trends in the percentage of subjects with values exceeding grade 0. We observed fewer subjects with conjunctival (bulbar) hyperemia and staining higher than grade 0 after 15 and 30 days with the study MPDS than with the control MPDS. Regarding limbal hyperemia, the number of subjects with a score higher than grade 0 after 15 days using the study MPDS tended to increase, a trend that reversed after 30 days. The trend toward an increase was maintained throughout the examinations for the control MPDS.

Discussion

To guarantee the safety and efficacy of new CL care solutions before they become commercially available it is necessary
to perform a clinical trial. The main purpose of this paper is to show the results of this clinical trial as is required by the clinical trials regulations.

Aloe vera is well known for its protection effect on the epithelium against potentially toxic substances.\textsuperscript{13} It also has bactericidal and bacteriostatic properties and can absorb ultraviolet light (UV) radiation.\textsuperscript{14} These properties are ideal as components of a solution for cleaning and maintaining CLs. Aloe vera is also a component that is added in different ophthalmic formulations.\textsuperscript{6,13–16}

The effect of aloe vera on ocular health has been studied because of its humectant properties. Burgalassi et al.\textsuperscript{15} concluded that ophthalmic solutions containing aloe vera were suitable to relieve dry eye symptoms. Ji and Jia\textsuperscript{16} found in an in vitro experiment that by incorporating aloe and polysaccharides (found in aloe vera extracts) into a binary solution, eyes were protected from bacterial infection and UV radiation. This implies that aloe vera may reduce the toxic effect of irritants, thereby preserving the viability of corneal epithelium cells against potential long-term toxicity caused by continuous exposure of the ocular surface to MPDSs. This effect is highest in aloe vera concentrations close to 1% (Avizor S.A.).

This is a crossover study in which each subject used two MPDSs (study and control). Each solution was used for 1 month with the same type of Si-HCL (Comfilcon A). Only one material was used to avoid interference resulting from changes in material, and this guaranteed that all differences were related to the MPDSs. Also, we compared the average values of weekly use of the Si-HCLs with each of the MPDSs evaluated. Small differences were seen that were possibly related to different habits of each subject. In any case, these differences were not significant, and this factor did not affect the results obtained for the others variables studied.

The new MPDS was not associated with adverse ocular effects for any of the safety variables included in the ISO 11980:2010 guidance for clinical investigation recommendations.\textsuperscript{9} No significant differences were detected between the study MPDS and the control MPDS, which confirmed the safety of the new MPDS.

No significant differences were found between both solutions in the evaluation of lens performance and acceptance. Results showed no interference in the cleaning ability of

\begin{table}[h]
\centering
\caption{Subjects with biomicroscopy signs exceeding grade 0 during the study.}
\begin{tabular}{|c|c|c|c|c|c|c|}
\hline
 & \multicolumn{2}{c|}{Exam} & \multicolumn{2}{c|}{Control MPDS} & \multicolumn{2}{c|}{CI 95\% difference in proportions} \\
 & No. & \% & No. & \% & Lower & Upper & \textit{p} Value \\
\hline
Bulbar hyperemia & 0 d & 11 & 55 & 10 & 50 & 0 & 49.86 & 0.8271 \\
 & 15 d & 11 & 55 & 12 & 60 & 0 & 51.95 & 0.8347 \\
 & 30 d & 12 & 60 & 16 & 80 & 0 & 71.11 & 0.4431 \\
Limbal hyperemia & 0 d & 6 & 30 & 4 & 20 & 0 & 40.68 & 0.5229 \\
 & 15 d & 10 & 50 & 7 & 35 & 0 & 54.87 & 0.4609 \\
 & 30 d & 5 & 25 & 8 & 40 & 0 & 49.72 & 0.3971 \\
Conjunctival staining & 0 d & 4 & 20 & 5 & 25 & 0 & 34.32 & 0.7382 \\
 & 15 d & 7 & 35 & 11 & 55 & 0 & 60.64 & 0.3348 \\
 & 30 d & 10 & 50 & 12 & 60 & 0 & 55.76 & 0.6684 \\
Papillae & 0 d & 4 & 20 & 7 & 35 & 0 & 46.83 & 0.3557 \\
 & 15 d & 5 & 25 & 8 & 40 & 0 & 49.72 & 0.3971 \\
 & 30 d & 6 & 30 & 8 & 40 & 0 & 46.41 & 0.5903 \\
Stromal edema & 0 d & 0 & 0 & 1 & 5 & 0 & 14.55 & 0.3049 \\
 & 15 d & 0 & 0 & 0 & 0 & 0 & 0.00 & 0.0000 \\
 & 30 d & 0 & 0 & 0 & 0 & 0 & 0.00 & 0.0000 \\
Corneal vascularization & 0 d & 16 & 80 & 14 & 70 & 0 & 63.50 & 0.7141 \\
 & 15 d & 15 & 75 & 14 & 70 & 0 & 57.73 & 0.8526 \\
 & 30 d & 16 & 80 & 16 & 80 & 0 & 55.44 & 1.0000 \\
\hline
\end{tabular}
\textbf{MPDS:} multipurpose disinfecting solution; \textbf{CI:} confidence interval; \textbf{Exam:} examination; \textbf{No.:} number of subjects who met the criteria; \%: percentage of subjects who met the criteria. 0 d: day 0, 1 hour of Si-HCL use (examination 2 or 5); 15 d: 15 days of Si-HCL use (examination 3 or 6); 30 d: 30 days of Si-HCL use (examination 4 or 7).
\end{table}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{image.png}
\caption{Anterior surface wettability (NIBUT) of the hydrogel contact lenses with both multipurpose disinfecting solutions (MPDSs) (study and control). The data show the average NIBUT values on days 15 and 30 for each MPDS. The error bars represent the standard deviations. * indicates a significant (\textit{p} < 0.05) difference between the control and study MPDSs on day 15.}
\end{figure}
the MPDS when aloe vera was included in the formulation. The higher percentage of subjects with deposits exceeding grade 0 on the front and back surfaces of the CLs after 15 and 30 days of use of the control MPDS could have been due to a more beneficial effect of the study MPDS. The increased deposits found between days 15 and 30 for both solutions could have been due to deposits over the CL surface that increased with time of wear and CL age. These results agreed with previous reports.17–19

In the current study, no differences in symptoms were found between the two MPDSs. Higher NIBUT values for the control MPDS was observed at 15 days, but these differences may not be considered clinically relevant, and disappeared at 30 days of use. No differences were found in the ocular surface slit-lamp evaluation, indicating that both MPDSs had the same effect on the ocular surface. No variations were found throughout the study in the fitting characteristics indicating that the study MPDS did not alter CLs parameters. No significant differences in comfort or satisfaction were found between the MPDSs; however, the study MPDS obtained higher averages than the control MPDS for both parameters on days 15 and 30 of the study.

This study has some limitations like the small sample size, although according to the statistical calculation. Furthermore, some items, like the mechanism of action of aloe vera or the advantages of adding aloe vera in the MPDSs, cannot be discussed using the results of this clinical trial cause it has been designed exclusively to study the security and efficiency of the new MPDS. For that reason more studies to prove the advantages of adding aloe vera in the MPDS are warranted.

In conclusion, the study MPDS is safe, has an acceptable physiologic tolerance, and is efficient according to the ISO 11980:2010 guidance for clinical investigation.9

**Conflicts of interest**

Rubén Urbano-Rodríguez works for the company that has supported the study: Avizor. The rest of the authors don’t have any conflicts of interest to declare.

**References**