ABSTRACT: Ten brands of hydrogel contact lenses were selected from five of the six British Approved Name lens classification groups to test in vitro the effects of repeated heat disinfection by means of microwave irradiation. Each lens type was tested over a number of cycles corresponding to its scheduled number of wearing days. The total diameter and back vertex power of all 80 test and 12 control lenses were measured at the end of their relevant cycling period. The back optic zone radius, center thickness, and water content were measured for 40 test and 8 control lenses. No clinically significant change was found in any of the 10 brands tested. Statistically significant changes were found in the back optic zone radius of the Frequency 55 group and water content of the Precision UV group. Some discoloration was noted in Ciba Visitint lenses. (Optom Vis Sci 2001;78:610–615)

Key Words: microwave, disinfection, lens materials

Microwave irradiation has been shown to be a highly effective means of rapidly disinfecting hydrogel contact lenses.1–3 The disinfecting method appears, essentially, to be one of moist heat,4 wherein the typical practice has been to immerse each lens in a volume of 6 to 12 ml of saline solution contained within a thermoplastic or glass vessel and then irradiate the vessel in a domestic microwave oven. Moist heat achieves three different levels of disinfection: pasteurization, complete disinfection, and sterilization. Treatment times and temperatures vary considerably according the materials being disinfected, levels of saturation, and the profile of the thermal curve used.5 Liquids held at temperatures in the range 60 to 72°C for periods ranging from 30 s to 30 min are termed pasteurized, which means a series of log reductions in the cell populations of certain species of challenge microorganisms. Complete disinfection, meaning a reduction to 0 colony forming units (cfu)/ml in all species of challenge cells, requires holding items at 100°C for periods of up to 5 min. The killing of spores, to achieve sterilization, requires holding a temperature of 121°C for 10 to 12 min.

Apart from Meridiano et al.,4 previous investigators have not discussed whether they were using pasteurizing or complete disinfecting methods. Some investigators have consciously held to a form of pasteurization by preventing the boiling of lens solution,1, 6 and others have found complete disinfection after allowing solution to visibly boil inside partially sealed cases.2, 3 Regarding the testing the effects of moist microwave disinfecting on the properties and parameters of hydrogel lenses, the two prior major studies have both allowed the visible boiling of lens solution during irradiation for periods of unspecified length.7, 8 Most of the prior research has concentrated on the potential of microwave treatment for practice use, where batches of lenses are to be treated. The present research forms part of a larger study concerning the efficacy of a daily regimen, wherein patients completely disinfect their lenses in their own microwave ovens. A patient-operated treatment will involve quite different irradiation times from those reported in prior research. Previous researchers admit to difficulties and inaccuracies in measuring the actual solution temperatures during irradiation.2 In the present study, the temperature inside irradiated storage cases was measured over successive intervals by inserting Thermax B heat strips into the solution. It was found that in an 800 W oven operating at full power, the solution temperature inside a single 10-ml thermoplastic storage case will reach \(+110°C\) in 10 to 12 s vs. the reported 120 s necessary when treating 20 cases in a 650 W oven.2 Use of heat strips also showed that in vented storage cases of the type described in earlier work, the visible boiling of saline actually occurred at temperatures in the range of 104 to 106°C. If irradiation of a single case is then stopped at 12 s, the solution temperature will still be higher than 72°C 3 min later, which means that the lenses would certainly be pasteurized, but not necessarily completely disinfected. Other important considerations are as follows. Individual ovens manifest different patterns of radiation distribution, which can vary according to the age and cleanliness of the oven.9 Because the treatment times required for microwave disinfecting are relatively short, the starting temperature of the solution may also be a significant factor.
In the present study, two groups of researchers used three methods of batch irradiating test lenses for cycle periods ranging between 120 and 150 s. The methods were devised to facilitate testing of the full spectrum of lens types most commonly used by patients, which involved irradiating large numbers of lenses up to a maximum of 730 2-min cycles. The criteria for lens selection were that lenses should be taken from each main polymer class, replacement frequency type, and with a variety of different prescription values. These criteria produced an assortment of 92 lenses for testing, comprising a test group of 80 lenses and a control group of 12 lenses.

METHODS

Lens Materials

Previous researchers such as Harris et al. 7 and Quesnel et al. 8 have used lenses with a single common prescription and with brands selected according to the U.S. FDA system of lens classification. The present study investigated an entirely random group of different prescriptions and lens thickness, ranging in back vertex power from −13.75 to +12.00 D, to reflect the kind of diversity encountered in practice. The British Approved Name (BAN) system of lens classification was preferred to the FDA system, because BAN classifies lenses by polymer families, rather than water content and ionic nature. BAN, therefore, allows for better generalization about the mechanical effects of heating on lenses belonging to specific polymer families. The one limitation of choosing the BAN system was that there are no commonly used lens brands in

<table>
<thead>
<tr>
<th>Lens Parameter</th>
<th>Significant Change</th>
<th>SEM Test A</th>
<th>SEM Tests B and C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total diameter (mm)</td>
<td>0.25</td>
<td>0.12</td>
<td>0.15</td>
</tr>
<tr>
<td>Back vertex power (D)</td>
<td>0.25</td>
<td>0.09</td>
<td>0.12</td>
</tr>
<tr>
<td>Back optic zone radius (mm)</td>
<td>0.30</td>
<td>0.06</td>
<td>—</td>
</tr>
<tr>
<td>Center thickness (μm)</td>
<td>10.0</td>
<td>8.9</td>
<td>—</td>
</tr>
<tr>
<td>Water content (%)</td>
<td>5.0</td>
<td>0.5</td>
<td>—</td>
</tr>
</tbody>
</table>

TABLE 2.

Clinically significant changes selected for lens parameters and SEM for equipment used
After the batch had cooled, the solution was discarded and replaced with 7 ml of fresh saline before the lenses were re-cycled. The control group lenses were placed into eight more storage cases and were not irradiated.

The second method B was devised to test 42 annual replacement lenses for 365 cycles. The test group was composed of 20 74% lenses and 20 38% lenses; the control group was composed of four lenses, two 74% and two 38%. Compartmentalized polypropylene trays were used to hold the lenses; each tray was divided into 12 compartments of 20-ml capacity. One test lens was placed in each compartment, and each lens was immersed in 2 ml of isotonic saline. The trays were stacked on top of the other, and the whole assembly was capped by a tray filled only with 80 ml of tap water to increase the thermal load to 140 ml, which prevented significant evaporation of saline during irradiation. The assembly was then loaded into a Toshiba 650 W oven with a turntable and irradiated at full power for 2 min. At the end of each cycle, approximately 0.3 ml of solution had evaporated from each compartment, and the saline concentration of the remaining 1.7 ml of solution had risen accordingly. The mechanics of handling 40 test lenses over 365 cycles made it impracticable to replace each lens in fresh saline at the start of each irradiation cycle, as was done in test method A. Accordingly, in method B, the solution volume and concentration were restored to their isotonic start values by addition of 0.3 ml of purified water British Pharmacopoeia (B.P.) through a pipette. Four control lenses were put into 2 ml of saline in the compartments of another tray, which was not irradiated.

The third method C was devised to overcome difficulties encountered in method B above, which led to occasional loss or damage to lenses because of lenses sticking to surfaces within the compartments. Method C was used to test 22 of the annual replacement lenses used in method B for a further 360 cycles. The test group was composed of 10 74% lenses and eight 38% lenses; the control group comprised four lenses, two 74% and two 38%. In method C, a thermoplastic microwave container with a capacity of 1.5 liters, was filled with 1100 ml of isotonic saline solution. Twenty 25-mm holes were drilled into the lid of the container. Through these holes were inserted the lens holder assemblies of the same storage cases described in method A. Because the caps of the lens holders were larger than the holes, the lenses could be suspended under the container lid, so that when the lid was placed on top of the base, all the lens holders were fully immersed within the 1100 ml of saline. A second, identical, container base was filled with 1 liter of saline at room temperature to use as a plunge bath for cooling the lenses after each heating cycle.

At the start of each irradiation cycle, the saline-filled container, without lenses inserted, was brought to boiling in the microwave oven. Thereupon, the loaded lens holders, suspended under the lid, were placed immediately into the hot solution, and the whole assembly was brought back to a boil. As soon as it was boiling, the first cycle was set to 2 min. After irradiation, the lid was immediately removed and replaced with a normal lid that helped maintain the temperature of the hot saline. The lenses were then plunged into the container containing saline at room temperature and soaked for 10 s.

Having been plunge-cooled this way, the lenses were immediately returned to the hot solution, and the next irradiation cycle started. The total cycle time turned out to be approximately 2 min and 20 s—although frequent adjustments were made to guarantee...
that the total boiling time was 2 min per cycle. After each cycle, approximately 10 ml of solution had evaporated, and as in method B, the practical method of restoring the solution to isotonic values at the beginning of each cycle was to add 10 ml of fresh purified water B.P. To speed the heating and hold a much larger thermal load at 100°C, a new Panasonic 800 W oven with a turntable was used for this method C. The four control lenses were left standing in two storage cases, each containing 7 ml of saline, and were not irradiated.

**Lens Measurement**

After consultation with both optometrists and contact lens manufacturers, it was decided in method A, which was used on monthly replacement lenses, to test the parameters selected by Harris et al. and Quesnel et al. These were total diameter, back vertex power (BVP), back optic zone radius (BOZR), water content, and center thickness. For methods B and C, measurement of the lathe-cut annual lenses was carried out by the manufacturers after their standard Communité Européen quality assessment procedures. In these procedures, total diameter and BVP are measured to ±0.25 mm and ±0.25 D before labeling, rather than to ±0.30 mm and ±0.36 D, which is considered clinically significant in “The CCLRU Good Lens Guide” cited by both Harris et al. and Quesnel et al.

All lenses were randomized and then measured masked by examiners not involved in the irradiation treatment. Lenses were first measured in fresh saline before irradiation and subsequently measured in fresh saline by the same examiners within 4 h of the final treatment cycle. For method A, the following measuring equipment and techniques were used (Table 2). Total diameter and BOZR were measured at 20°C in 0.9% physiological saline using the Optimec JCF with the TC20 saline circulating system. BVP was measured using a Topcon Lm-P6 Lensmeter in air at room temperature. Center thickness was measured using a Rehder ET-1 electronic thickness gauge, and water content was measured using an Atago CL-1 hand held refractometer at room temperature.

**RESULTS**

The solution temperature in vessels used for the tests was measured using calibrated temperature strips (Thermax Range B, accuracy ±1°C) inserted into the solution during irradiation. The average temperature was 105°C for the storage cases used in method A, 104°C for the trays in method B, and 102°C for the vessel in method C. In method A, some lenses were initially found to be sticking to the lens holders, and one Frequency 55 lens was lost due to tearing during removal. In method B, two Vistagel 38% lenses were lost due to the tendency of the unrestrained lenses to move about in the turbulent boiling of the solution. At the end of testing, all five Focus Visitint lenses were found to be slightly discoloured, with a brownish hue. The results for each parameter tested are presented in Figs. 1 to 5. Clinically significant changes in individual lenses are listed in Table 3.

**Data Analysis**

Data from all three tests were recorded and analyzed using Microsoft Excel 97. Initial and final measurements were analyzed statistically by means of a two-tailed paired t-test, with a hypothesized difference = 0. The baseline provided by the control lenses then gave a point of reference about the range of error arising in the measuring procedures. The level of statistical significance was set at α = 0.05.

**TABLE 3.**

<table>
<thead>
<tr>
<th>Lens Type</th>
<th>Number</th>
<th>TD (mm)</th>
<th>BVP (D)</th>
<th>BOZR (mm)</th>
<th>CT (µm)</th>
<th>WC (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequence 1</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>0.31</td>
<td>—</td>
<td>6</td>
</tr>
<tr>
<td>Sequence 1</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Surevue 74 (365 cycles)</td>
<td>1</td>
<td>—0.30</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Actifresh 1</td>
<td>—</td>
<td>0.50</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Sequence (control)</td>
<td>1</td>
<td>—</td>
<td>—</td>
<td>0.35</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

*TD, total diameter; BVP, back vertex power; BOZR, back optic zone radius; CT, center thickness; WC, water content.*
DISCUSSION

No clinically significant mean changes were found in the parameters of any lens type tested. Statistically significant changes were found in BOZR in Frequency 55 and water content in Precision UV. In common with previous research, the present work does not reveal predictable trends or patterns in the parameter shifts of lenses belonging to different FDA or BAN categories. Although there have been reservations dating back to the 1980s concerning the suitability of heat disinfection for higher water content lenses, the changes in parameters of high water content lenses appears little different to mid and low water content lenses. Looking at the 4a and 4b lenses, the changes in total diameter and water content were slightly less than for groups 1a to 3a and slightly greater in BOZR and center thickness. Prior research has found small but statistically significant increases in water content for low and mid water content lenses. In the present testing, a small but statistically significant increase was also found in water content for one lens brand, Precision UV, a high water content lens. However, greater changes were found in water content for Medalist 66 and Review 55, and these were reductions in water content. The very small increase in water content for all test lenses does not provide much support for the hypothesis that microwave treatment necessarily increases the water content of hydrogel lenses.

Figs. 1 to 5 show that the mean change in each parameter of the lens brands tested was considerably less than the corresponding limit of clinical significance. As found in prior testing, parameter changes were very close to the SEM limits for the measuring equipment used. In the testing by Harris et al., which involved treating the lenses for cycle times of 5 min, some of these small changes were statistically significant in as many as four of the five parameters tested for certain lens brands. In the work of Quesnel et al., which involved 2 min cycles, only the changes in water content were found to be statistically significant. In the present testing, the incidence of statistically significant parameter changes was lower again, with the only such changes found being in water content for Precision UV and BOZR for Frequency 55. The clinically significant changes found in four individual lenses listed in Table 3 do not appear to be statistically significant. The clinically significant change found in BOZR for one Sequence control lens further suggests that these findings could have resulted from handling and measuring errors rather than from heating effects. The present findings appear to support the views of prior researchers that microwave heating of hydrogel lenses may lead to small, statistically significant changes in some parameters of some lenses, but the changes are well within the limits of clinical acceptability.

In view of the prevalent manufacturing practice of heat sterilizing hydrogel lenses at 120°C before sale, it was surprising to discover some discoloration in the five Visitint lenses. Inquiries to the manufacturer revealed that they specifically contraindicate the use of heat disinfection for this particular lens brand, which uses a proprietary tinting agent. Because the measurements of Visitint lenses do not reveal unusual alterations in any parameter of the lenses, it is likely that heating by irradiation effects changes in the chemistry of this tinting agent rather than in the lens polymer.

CONCLUSION

In common with previous research, the present testing shows that heating of unworn hydrogel lenses by microwave irradiation does not cause clinically significant mean changes in the lens parameters of 10 previously untested lens brands, selected from 5 of 6 BAN lens polymer groups. Clinically significant changes in a single parameter were found in four of the 80 test lenses and in one of the 12 control lenses. Of the small changes found in all lens parameters, only mean changes in one parameter each of two lens brands were found to be statistically significant. Some discoloration was noted in one of the six tinted lens brands used. Although this promises well for the continuing investigation of microwave treatment of hydrogel lenses for patient use, the scope of the
The present investigation has been limited to observing the effects in vitro on the clinical parameters of hydrogel lenses. Further investigation is needed to determine the significance of other clinical effects when lenses are treated by patients in vivo.

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