Objectives

The primary objective of this study was to assess the extent to which Philips Sonicare TongueCare+ improves breath odor compared to manual tongue brushing alone, and to Listerine Cool Mint Rinse alone, based on organoleptic assessment.

The secondary objectives of the study included safety evaluation, and the effects of the study products on hydrogen sulfide levels in exhaled breath and tongue microbes (aerobes, anaerobes).

Methodology

One hundred sixty-eight healthy adults meeting study requirements were enrolled in this IRB-approved, parallel, examiner-blinded, three-arm clinical trial. There were 91 female and 77 male participants (mean age, 38.9 years) with 56 subjects per treatment group. Eligible subjects were non-smokers aged 18–70 years with a minimum organoleptic (OL) score of 2.7–4.5 following 12–18 hours of oral hygiene abstention. The OL score was an average, computed following breath assessment by three calibrated, blinded organoleptic assessors. Eligible participants agreed to comply with study procedures related to oral hygiene, and food/drink restrictions.

The breath hygiene study groups were:
- Philips Sonicare TongueCare+ (TC+): powered tongue brush attachment with BreathRx tongue spray (three sprays followed by 20 sec tongue brushing, performed three times)
- Listerine Cool Mint (LCM) Antiseptic Rinse (20 ml for 30 sec)
- Manual Toothbrush (MTB) tongue brushing (ADA-reference manual toothbrush)

There were two study visits (Day 1 and Day 8) with assessments occurring at each visit at the following intervals: upon arrival, immediately, and four and eight hours following breath hygiene product use. Organoleptic assessments, as well as microbial at H,S samples were performed at each of these time points. In addition to the assigned breath hygiene regimen, all subjects used a standardized home toothbrushing regimen (Philips Sonicare power toothbrush and Crest Cool Mint Gel dentifrice) twice daily, during the seven-day home-use period.

Results*

Organoleptic evaluation

On Day 1 immediately post treatment, the LS Mean (95% CI) percent reduction from pre-treatment for OL was 53.04% (49.37%, 56.71%) for TC+, 49.59% (45.90%, 53.28%) for LCM and 47.1% (43.40%, 50.79%) for MTB. The pair-wise comparison between TC+ and MTB was statistically significant (p-value = 0.0330).

At four hours post-treatment, the percent reduction in OL was 44.06% (39.89%, 48.23%) for TC+, 34.22% (30.03%, 38.41%) for LCM, and 38.37% (34.16%, 42.57%) for MTB. The pair-wise comparison between TC+ and LCM was statistically significant (p-value = 0.0062).

At eight hours post-treatment, the percent reduction in OL was 46.67% (42.18%, 51.57%) for TC+, 22.83% (18.31%, 27.35%) for LCM and 26.19% (21.66%, 30.72%) for MTB. The differences between TC+ and MTB, and TC+ and LCM, were significant (p-value < 0.0001, for each pair-wise comparison).

For OL outcomes on Day 8, the TC+ treatment group exhibited the lowest value throughout the study visit. Statistically significant differences were observed between TC+ and LCM, and TC+ and MTB, at each time point: immediately, four hours and eight hours following product use.

Values are included below.

The reported safety events were mild in severity and were unrelated to test product use.

Conclusions:

The Philips Sonicare TongueCare+ breath hygiene regimen was effective at improving oral malodor. Improvements, based on organoleptic assessment, were evident immediately following, and up to eight hours following product use.

The Philips Sonicare TongueCare+ breath hygiene regimen improved oral malodor significantly better than either Listerine Cool Mint Antiseptic Rinse, or tongue brushing with an ADA-reference manual toothbrush, up to eight hours following product use.
Day 1
LS Mean Percent Reduction
Organoleptic Score

<table>
<thead>
<tr>
<th></th>
<th>Immediately Post</th>
<th>4 hour</th>
<th>8 hour</th>
</tr>
</thead>
<tbody>
<tr>
<td>TC+</td>
<td>53.04</td>
<td>49.59</td>
<td>46.67</td>
</tr>
<tr>
<td>LCM</td>
<td>47.1</td>
<td>34.22</td>
<td>22.83</td>
</tr>
<tr>
<td>MTB</td>
<td>44.06</td>
<td>38.37</td>
<td>26.19</td>
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</table>

Day 8
LS Mean Percent Reduction
Organoleptic Score

<table>
<thead>
<tr>
<th></th>
<th>Immediately Post</th>
<th>4 hour</th>
<th>8 hour</th>
</tr>
</thead>
<tbody>
<tr>
<td>TC+</td>
<td>49.96</td>
<td>39.25</td>
<td>34.54</td>
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<tr>
<td>LCM</td>
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<td>21.05</td>
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<tr>
<td>MTB</td>
<td>28.6</td>
<td>18.34</td>
<td>12.07</td>
</tr>
</tbody>
</table>

*For brevity, the outcomes for the secondary efficacy endpoints, H₂S and microbes, are not reported here. Please refer to the full publication for these details.