24 hour results from a US based, randomized, placebo-controlled, multi-centered trial, to evaluate a novel peppermint oil delivery system. Results from the IBSREST™ trial.

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Introduction

Irritable bowel syndrome (IBS) is a chronic disorder with periodic exacerbations of symptoms that can be severe. The incidence of the 3 major forms of IBS has been reported as IBS-M (61.0%), IBS-D (29.3%), and IBS-C (9.7%).1 While there are prescription options for IBS-C, there are no approved products for IBS-M and limited options, including restricted use alosetron, for IBS-D. The Irritable Bowel Syndrome Reduction Evaluation & Safety Trial (IBSREST) comprised of patients with IBS-M and IBS-D (in equal numbers) in order to address the needs of this large group of patients. Disturbance of small intestinal homeostasis, in particular transit time, has been implicated as a potential source of the clinical manifestations of IBS. IBgard® contains peppermint oil (PO), a proven ingredient to help normalize digestive and absorptive processes (including intestinal transit times), which are customarily disrupted by IBS. Along with the desired properties of PO’s principal component L-menthol, in promoting uptake of important nutrients, it has been shown to possess anti-spasmodic, anti-inflammatory, serotoninergic, and anti-bacterial properties. PO is available in the US as a medical food, as single-unit, liquid-filled, enteric-coated capsules. Such PO delivery methods are not specifically designed for targeted delivery to the small intestine, have a higher risk of dose-dumping, and generally reside longer in the stomach. As a result, patients receiving single-unit, enteric-coated PO can experience heartburn, nausea, and anal burning. IBgard® is a modern version of PO delivery consisting of triple-coated, sustained release microspheres designed to provide quick and reliable release of PO in the small intestine.

IBSREST® Trial Objectives

Evaluate the effectiveness and safety of IBgard® for the management of IBS.

• Confirm results of previous European clinical trials of PO in a U.S. population.
• Determine if PO with Site Specific Targeting (SST®) technology results in rapid action and improved tolerability of PO in patients with IBS-M and IBS-D.

* Irritable Bowel Syndrome Reduction Evaluation & Safety Trial

Methods

Randomized to 2 IBgard capsules (90 mg PO) or identical placebo TID 30-90 min. before meals

Rescue medications were not allowed during the study

Before enrollment, patients underwent testing to exclude organic disease and a 2-week washout period of all prohibited medications

Patients maintained a 2-week baseline diary with daily assessments of bowel movements and IBS symptoms

Prior to and 24 hours after randomization, measurements of IBS symptom frequency and intensity (both on a scale of 0-4) were obtained.

• IBS symptoms included abdominal pain or discomfort, abdominal bloating or distention, pain at evacuation, urgency of bowel movement, constipation, diarrhea, passage of mucus or gas, and a sense of incomplete evacuation.
• Methods and endpoints (TISS) approximated those of Cappello et al.2

Inclusion Criteria

• Females and males 18-60 years of age
• Meets Rome III IBS criteria for IBS-M or IBS-D
• Average daily IBS-related abdominal pain on a 0 to 4 scale each week of 2 week baseline period
• TISS ≥2.9 on a 0 to 4 scale
• Subject was not planning to change his/her usual diet and lifestyle during the course of the study

Exclusion Criteria

• Diagnosis of IBS-C or IBS-U
• Organic gastrointestinal disease
• Refusal to discontinue one or more prohibited medications for at least 7 days prior to beginning the baseline diary and throughout the remainder of the study

Baseline demographics were shown in Table 1.

Results

Baselines reported in all tables and figures were not significantly different between IBgard and placebo (P > 0.05).

Table 1. Demographic Characteristics

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<th>IBgard®</th>
<th>Placebo</th>
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<td>Weight (kg)</td>
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<tr>
<td>Height (cm)</td>
<td>0.6 (0.1)</td>
<td>0.6 (0.1)</td>
<td>0.87</td>
</tr>
</tbody>
</table>

Figure 1. Reduction in Total IBS Symptom Score at 24 Hours*

*Statistically significant vs. placebo (P<.05).

Figure 2. Reduction in All 8 Symptoms at 24 Hours†

†Statistically significant vs. placebo (P<.05).

Figure 3. Reduction in abdominal pain and discomfort at 24 hours

Figure 4. Reduction in Intensity of Bowel Movement Urgency at 24 Hours

Conclusions

In the IBSREST Trial, measurements of all 8 IBS symptoms (average of intensity and frequency) trended in favor of IBgard, as compared to placebo, within 24 hours of the first dose, and abdominal pain and discomfort reached statistical significance.

Subjects in the IBgard treated group experienced statistically significant decreases from baseline, compared to the placebo group, at 24 hours, in the following:

• TISS (total IBS symptom score)
• Abdominal pain and discomfort (average of frequency and intensity)
• Urgency of bowel movements (intensity)

IBgard was well tolerated and associated with no treatment emergent adverse events during the first 24 hours, which was comparable to placebo.

References

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