Polar vectors as a method for evaluating the effectiveness of irritable bowel syndrome treatments: An analysis with pinaverium bromide 100 mg plus simethicone 300 mg po bid

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KEYWORDS
Irritable Bowel Syndrome; Clinical studies; Outcome measures; Polar vectors; Bristol Stool Scale; Pinaverium bromide plus Simethicone

Abstract
Background: Irritable Bowel Syndrome (IBS) is a disorder characterized by abdominal pain or discomfort associated with changes in bowel habit. Currently there are no objective outcome measures for evaluating the effectiveness of treatments for this disorder.
Aims: To determine the usefulness of a method of analysis that employs polar vectors to evaluate the effectiveness of IBS treatments.
Methods: Data from a Phase IV clinical study with 1677 active IBS-Rome III patients who received 100 mg of pinaverium bromide + 300 mg of simethicone (PB+S) po bid for a period of four weeks were used for the analysis. Using the Bristol Stool Scale as a reference, the consistency and frequency of each type of bowel movement were recorded weekly in a Bristol Matrix (BM) and the data were expressed as polar vectors.
Results: The analysis showed a differential response to the PB+S treatment among the IBS subtypes: in reference to the IBS with constipation subtype, the magnitude of the vector increased from 10.2 to 12.5, reaching maximum improvement at two weeks of treatment (p<0.05, Scheffé). In the IBS with diarrhea and mixed IBS subtypes, the magnitude of the vector decreased from 19 to 14 (p<0.05) and from 16.5 to 13 (p<0.05), respectively, with continuous improvement for a period of four weeks. There was no definable vectorial pattern in the unsubtyped IBS group.

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Conclusions: Analysis with polar vectors enables treatment response to be measured in different IBS subtypes. All the groups showed improvement with PB+S, but each one had its own characteristic response in relation to vector magnitude and direction. The proposed method can be implemented in clinical studies to evaluate the efficacy of IBS treatments.

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Figure 1  Improvement of abdominal pain and bloating in relation to the Bristol Matrix. A scale that corresponds to stool consistency measured by the Bristol Stool Scale is shown. For IBS-C, improvement should be from left to right, whereas for IBS-D and IBS-M it should be from right to left.

In contrast, in patients with IBS-D, clinical improvement implies bowel movements of a greater consistency, going from types 6 and 7 to type 5 or lower, with a reduction in evacuation frequency. Once the magnitude and direction of the vectorial movement is known (the degree of improvement), vectors can be transferred to a plane of polar coordinates that enables the treatment response in all the IBS subtypes to be observed at the same time. This method of analysis is based on the assumption that things are more easily understood if they can be visualized.

The aim of this study was to determine the usefulness of the method of analysis employing polar vectors for evaluating IBS pharmacologic treatment efficacy. The data used were obtained in an open Phase IV study conducted on patients with active IBS who received 100 mg of pinaverium bromide + 300 mg of simethicone (PB + S) po bid over a four-week period. The hypothesis stated that this method is adequate for evaluating the efficacy of any IBS treatment, by determining stool consistency and frequency improvement through a BM that allows for the treatment response in the various IBS subtypes (IBS-D, IBS-C, IBS-M, IBS-U) to be differentiated.

Methods

Population

The data used in this analysis were obtained from a Phase IV clinical study that included 1677 patients with active IBS in accordance with Rome III criteria. Patients were recruited by 1303 physicians with different specialties throughout the Mexican Republic. Selection criteria were:

(a) For inclusion – Patients of both sexes who were ≥18 and ≤50 years of age and who had a body mass index (BMI) <50. Clinical diagnosis of IBS according to the Rome III criteria: presenting with recurring abdominal pain or discomfort for at least three days a month in the last three months that was associated with 2 or more of the following characteristics: (1) improvement with defecation, (2) onset associated with a change in stool frequency, and (3) onset associated with a change in stool form (appearance). In addition, based on the Rome III classification, patients were categorized according to bowel habit as IBS-D, IBS-C, IBS-M, and IBS-U. IBS-D was identified by loose or liquid stools in at least 25% and hard or lumpy in less than 25% of bowel movements; IBS-C: hard or lumpy stools in at least 25% and loose or liquid in less than 25% of bowel movements; IBS-M: hard or lumpy stools in at least 25% and loose or watery stools in at least 25% of bowel movements; and IBS-U: modifications in stool consistency but insufficient to be classified as IBS-D, IBS-C, or IBS-M. Active IBS was defined as the presence of abdominal pain and/or discomfort at least twice a week during the previous seven days.

(b) For exclusion – The presence of alarm symptoms over the past 6 months, e.g.: involuntary weight loss, anorexia, inexplicable anemia, palpable lymph nodes or masses, fever, digestive tract bleeding, the suspicion or confirmed presence of malignant disease in any system or organ; women who were suspected of being pregnant, women who were pregnant, or who were breastfeeding; the suspicion or confirmed presence of rectoanal stricture; esophageal varices; a history of chronic non-specific ulcerative colitis, Crohn’s disease, or rectoanal ulcer; major upper or lower abdominal surgery; digestive tract malformation; or bowel obstruction.

Design

An open, prospective, descriptive, and multicenter study was conducted. Treatment was PB + S for a period of four weeks. Stool consistency and frequency and improvement in cardinal symptom intensity (abdominal pain and bloating) were evaluated on a weekly basis.

Data collection instrument

The intensity of abdominal pain and bloating were evaluated by the patients through a 10-cm-long VAS for each symptom. Stool consistency (type) was recorded according to the Bristol Stool Scale on a 7-day format that also took into consideration the frequency of all types of bowel movements (Fig. 2). This format, or BM, was used during the four weeks of treatment. With the BM data, a vectorial calculation was made (see the statistical analysis) to evaluate the changes obtained with PB + S treatment.

Statistical analysis

An omnibus variable was created that included two variables: consistency (type) and frequency of evacuations obtained from the BM. Given that the aim was to demonstrate the treatment changes through graphs, a two-dimensional configuration using polar vectors was decided upon. Polar vectors are dimensionless and can be evaluated by determining the treatment response in all the IBS subtypes to be observed at the same time. This method of analysis is based on the assumption that things are more easily understood if they can be visualized.

The vector calculation was made in the conventional manner:

\[ r = \sqrt{x^2 + y^2}, \]

where \( r \) is the magnitude of the vector and \( y \) is the equivalent of the hypotenuse of a right triangle and therefore is
equal to the square root of the sum of the weekly stool frequency (x), plus the total sum of the Bristol Stool Scale types the patient would have had (y), each one squared. The vectorial angle (which is equal to the direction of the vector) was calculated with the equation: \( \theta = \tan^{-1}(y/x) \), in which \( \tan^{-1} \) is the arc tangent of the coefficient of the numerator y (total sum of the Bristol types) and the denominator x (total weekly stool frequency). Thus, each patient had an r and \( \theta \) value that could be linedly combined with the other subjects in order to obtain a mean (the expected value) vector magnitude according to each IBS-Rome III subtype. The vector was obtained in such a way that it represented the sum of the variables. In short, the higher the value of the Bristol type and the frequency of the bowel movements were, the greater was the magnitude of the vector. The maximum value of x was the result of: \( \sum_{i=1}^{n} x_i = 28 \), in a patient who would have shown all the Bristol types during the week. In contrast, the frequency of bowel movements during one week determined the maximum value of y. The weekly stool frequency in IBS patients was in the first percentile (P1) of 2 defecations, the P10 of 3 defecations, the median (P50) of 8 defecations, the P90 of 15, the P95 of 19, and the P99 of 28 defecations. In this manner, certain mathematical equivalences are produced that, even though they are not exact, can be useful for clinical interpretation.

Data were expressed as mean and standard deviation (mean ± SD), and standard error (SE) was specified when used. The differences in the means of magnitude and angles were calculated with ANOVA for repeated measures and post hoc comparisons were run with the Scheffé test, obtaining homogeneous groups.

**Ethical aspects**

The Phase IV clinical study protocol that produced the data of this analysis was approved by the Ethics Committee of the *Universidad de Guanajuato* and the General Hospital of Naucalpan Dr. Maximiliano Ruiz Castañeda. The study was conducted in accordance with the Declaration of Helsinki, and the Good Clinical Practice, and the local regulations concerning clinical research. All participating patients signed informed consents.

**Results**

**Demographic characteristics of the population**

The 1677 patients with active IBS included in the clinical study were classified according to IBS subtype as: IBS-C, 42.9%; IBS-D, 5.6%; IBS-M, 48.6%; and IBS-U, 2.9%. The total sample mean age was 36.9 ± 8.8 years, BMI was 26.3 ± 4.8, and 76.8% were women. The percentage of patients who stated that their IBS progression was longer than one year was 36.4%. Of the patient total, 1369 (81%) completed the four weeks of treatment with PB + S and attended their final visit. Table 1 shows the demographic and clinical characteristics of this subset, according to the IBS subtypes.

**Intensity of abdominal pain and bloating**

There was a notable reduction in the intensity of abdominal pain and bloating with the PB + S treatment, regardless of the IBS subtype. A plateau was not reached at four weeks, but there was maximum improvement in that time period (Fig. 3).

**Vectorial analysis (stool consistency and frequency)**

Polar vector analysis made it possible to show that there was considerable improvement in the IBS-C patients within the first two weeks of treatment and that it remained steady during the final two weeks. In this group there was an increase in vector magnitude from 10.2 to 12.5, reaching maximum improvement at two weeks of treatment (\( p < 0.05 \), Scheffé). A vector magnitude of 12.5 is apparently equal to type 4 on the Bristol Stool Scale.

The magnitude of the vector was reduced in the patients with IBS-D and IBS-M. In the IBS-D patients the vector began at a magnitude of 19.0 and it went toward a magnitude of 14.0 (\( p < 0.05 \)), whereas the IBS-M group shared the same direction as the IBS-D group, starting at a magnitude of 16.5 and gradually moving toward a magnitude of 13.0 (\( p < 0.05 \)). There was important improvement in these two IBS subtypes during the second treatment week, and unlike the IBS-C group, it continued during weeks 3 and 4. The IBS-U group showed erratic behavior with no definite pattern in relation to vector magnitude and direction (Fig. 4).

**Discussion**

This study demonstrates that the transformation into polar vectors of the combination of the type of stool consistency according to Bristol Stool Scale, and the frequency of each of those stool types evaluated in a BM, is a useful method for evaluating IBS pharmacologic treatments, as it was the
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This has led to continuous meetings between the representatives of those agencies, experts on the subject matter, and representatives of the pharmaceutical industry in an effort to define the reported outcome measures for the patients, so-called PRO or Patient Reported Outcomes. This can be difficult when it entails validated outcome measures that objectively record their efficacy in clinical studies; the fact that there are no biological markers for the diagnosis of IBS also has to be taken into account. In the last few years, the most widely used primary outcome measures in IBS clinical studies have been global variables of binary response: adequate improvement of abdominal pain and discomfort or satisfactory relief of IBS symptoms, that followed the Rome recommendations. Nevertheless, certain psychometric aspects of these variables have not been adequately validated and therefore are not presently accepted by the regulatory agencies. This has led to continuous meetings between the representatives of those agencies, experts on the subject matter, and representatives of the pharmaceutical industry in an effort to define the reported outcome measures for the patients, so-called PRO or Patient Reported Outcomes. This can be difficult when it entails validated outcome measures that objectively record their efficacy in clinical studies; the fact that there are no biological markers for the diagnosis of IBS also has to be taken into account.

Table 1 Patient clinical characteristics according to IBS subtype.

<table>
<thead>
<tr>
<th></th>
<th>IBS-C (N = 723)</th>
<th>IBS-D (N = 94)</th>
<th>IBS-M (N = 812)</th>
<th>IBS-U (N = 48)</th>
<th>Total (N = 1677)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Women/men (%)</td>
<td>81.7/18.3</td>
<td>56.4/43.6</td>
<td>74.8/25.2</td>
<td>77.1/22.9</td>
<td>76.8/23.2</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Age in years</td>
<td>36.3 (9.0)</td>
<td>37.8 (8.7)</td>
<td>37.3 (8.5)</td>
<td>36.8 (9.0)</td>
<td>36.9 (8.8)</td>
<td>0.13</td>
</tr>
<tr>
<td>BMI</td>
<td>26.1 (4.8)</td>
<td>26.9 (4.4)</td>
<td>26.4 (4.8)</td>
<td>26.0 (4.4)</td>
<td>26.3 (4.8)</td>
<td>0.39</td>
</tr>
<tr>
<td>Heart rate × min</td>
<td>75.8 (7.4)</td>
<td>76.6 (7.7)</td>
<td>76.3 (7.8)</td>
<td>76.3 (7.6)</td>
<td>76.3 (7.6)</td>
<td>0.19</td>
</tr>
<tr>
<td>Respiratory rate × min</td>
<td>18.8 (3.0)</td>
<td>19 (3.0)</td>
<td>19 (3.0)</td>
<td>18.2 (3.0)</td>
<td>18.9 (3.0)</td>
<td>0.29</td>
</tr>
<tr>
<td>Systolic BP (mmHg)</td>
<td>115 (10.0)</td>
<td>118 (13.0)</td>
<td>116 (11.0)</td>
<td>117 (12.0)</td>
<td>116 (11.0)</td>
<td>0.08</td>
</tr>
<tr>
<td>Diastolic BP (mmHg)</td>
<td>74 (8.0)</td>
<td>76 (7.0)</td>
<td>74 (7.0)</td>
<td>74 (8.0)</td>
<td>74 (8.0)</td>
<td>0.34</td>
</tr>
</tbody>
</table>

BMI: body mass index; min: minutes; BP: blood pressure. Data expressed as mean (standard deviation).

* Contrast carried out with $\chi^2 = 33.8$, df = 8. The remaining contrasts were carried out through one-way ANOVA.

Figure 3 Changes in the intensity of abdominal pain and bloating. The intensity of abdominal pain and bloating was evaluated by the patients through a 10 cm VAS. The changes in time were significant ($p < 0.001$, repeated measures MANOVA) for pain (A) and bloating (B). The contrast between subtypes showed no significant difference. Blue: IBS-C, red: IBS-D, green: IBS-M, and pink: IBS-U. The figure shows means ± SE.

A current limitation in the development of new IBS therapeutic options is the lack of validated outcome measures that objectively record their efficacy in clinical studies; the fact that there are no biological markers for the diagnosis of IBS also has to be taken into account. In the last few years, the most widely used primary outcome measures in IBS clinical studies have been global variables of binary response: adequate improvement of abdominal pain and discomfort or satisfactory relief of IBS symptoms, that followed the Rome recommendations. Nevertheless, certain psychometric aspects of these variables have not been adequately validated and therefore are not presently accepted by the regulatory agencies. This has led to continuous meetings between the representatives of those agencies, experts on the subject matter, and representatives of the pharmaceutical industry in an effort to define the reported outcome measures for the patients, so-called PRO or Patient Reported Outcomes. This can be difficult when it entails validating the results in patients with FGIDs that do not have an organic or structural lesion and whose evaluation is made through symptom-based criteria.

The method of analysis used in this study has taken the current tendencies and recommendations into account, such as including diagnosed patients through the Rome III criteria and evaluating the changes in stool consistency and frequency. In addition, it was based on the collection of symptoms through diaries and the weekly evaluation of treatment response. It also contemplated the variables in a multidimensional context and evaluated the individual response in the different IBS subtypes.
Figure 4  Polar vectors showing the changes in magnitude (r) and direction, during four weeks of treatment with PB + S, according to each IBS subtype. Notice that the vectorial magnitude axes were adjusted according to the space covered during the four weeks and therefore, at its maximum improvement (r < 12.5), IBS-C did not reach the lower end of IBS-M (r > 13); in addition, considering the magnitude of the vector, there is overlapping in the IBS-D, IBS-M, and the IBS-U spaces. The letters correspond to homogeneous Scheffé groups in the post hoc analysis. See the statistical analysis section for details.

A first limitation of the suggested method is the fact that the physician must graph the weekly vector results himself however, it is possible to develop computerized programs that can facilitate the follow-up of changes that occur in the patients and that can show the results of different treatments. A second limitation is that this method is applicable exclusively to medications that have an effect on stool consistency and frequency, and as a result, a third limitation is the fact that polar vectors do not evaluate the effect of treatment on abdominal pain or discomfort (abdominal bloating) that are key symptoms in IBS. However, it is worth noting that a recent review of the literature on questionnaires used in IBS to create a framework by which PRO can be developed for this disorder, established that abdominal pain and discomfort are two different symptoms. Therefore questions about discomfort response should be avoided, because it is very nonspecific. In the future, evaluations will have to be made on whether improvement in stool consistency and frequency not only reduces abdominal pain, but also other symptoms such as associated anxiety, and also whether the effect on pain and anxiety can be analyzed through vectors. A fourth limitation of this study is that the clinical significance of the change in vector magnitude and direction has not been determined. Regulatory agencies have emphasized the importance of identifying objective endpoints that would allow clinically significant improvement to be established, and thus report the percentage of patients that manage to go beyond that endpoint. Finally, the vectorial method was studied in patients who were seen in private medical practices in Mexico and it is necessary not only to validate the results in studies on open populations, but also on subjects from other cultures and in other languages.

In conclusion, polar vectors can be useful in clinical trials for the integral evaluation of the changes in stool consistency and frequency in response to an IBS treatment. Vectorial analysis shows that the combined therapy of PB + S, administered for at least four weeks, differentially modifies the frequency and consistency of bowel movements in all IBS subtypes, but with differences in the magnitude and velocity of the change between them. It should be pointed out that, even though the polar vectors analyzed in this study did not evaluate the response to pain and/or subjective abdominal bloating, the improvement of these symptoms analyzed through VAS was also significant for all the IBS subtypes. It is necessary to apply this method of analysis in placebo-controlled studies in order to confirm the results observed in this study.

Financial disclosure

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Conflict of interests

Dr. Juan Carlos López-Alvarenga is currently a Biometric Scientist for Clinical Trials of Takeda México SA de CV. Has received research grants by Silanes, Roche, Servier. Has been a speaker for Takeda, Roemmers, Bayer, Roche and Novartis.

Dr. Sergio Sobrino-Cossío has been a speaker for Takeda México SA de CV, Astra, Ferrer, Olympus, Boston Scientific, Trasmédica, Wilson Cook.

Dr. José-María Remes-Troche is a member of the Advisory Board of Takeda Pharmaceuticals, Alfa-Wasserman and Janssen. Has been a speaker for Nycomed-Takeda, Advance Medical, Astra-Zeneca and Bristol-Myers-Squibb. Has received a research grant from Sanofi-Pasteur.

Dr. Jazmín Chiu-Ugalde and Dr. José Antonio Vargas-Romero, are employees of the Medical Direction of Takeda México SA de CV.

Dr. Max Schmulson has served as a consultant for Procter and Gamble, Novartis, Schering-Plough, Alfa-Wasserman, Janssen, Nestle Ltd and Almirall. Has been a speaker for Takeda México SA de CV, Schering-Plough, Mayoli-Spindler, Alfa-Wasserman, Janssen and Novartis. Has received research grants from Takeda México SA de CV and Nestle Ltd.

Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.rgmx.2012.10.003.

References


