Functional dyspepsia and the satiety test: Its usefulness in clinical practice

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Abstract

Introduction: According to the Rome III Criteria, functional dyspepsia (FD) is classified as post-prandial distress syndrome (PDS) and epigastric pain syndrome (EPS). On the other hand, the satiety test (ST) has been used to evaluate gastric accommodation and emptying, distinguishing healthy individuals from those with dyspepsia.

Aims: To determine whether the ST can distinguish dyspeptic individuals from healthy ones and to evaluate its usefulness in differentiating the two FD subtypes.

Methods: Adults with FD were consecutively enrolled in a cross-sectional study within the timeframe of August 2011 and October 2012. Healthy subjects participated as controls. The ST consisted of the intake of a nutritional supplement (Fortisip®, Nutricia Bagó®) at a constant speed; satiety was graded at 5-minute intervals (1 to 5 points). Intake was suspended when the maximum score was reported. The total ingested volume and caloric intake was recorded and the Mann-Whitney U test was used in the statistical analysis.

Results: The study included 39 dyspeptic patients and 20 control individuals. The patients were predominantly women (84.6 vs. 25%; p < 0.0001) and they were similar in age (39.59 ± 13.53 vs. 34.70 ± 9.85 years) and BMI (24.32 ± 3.52 vs. 25.82 ± 3.34 kg/m²) with respect to the controls. The FD subtype percentages were PDS: 61%, EPS: 31%, and Mixed syndrome: 8%. There was a lower ingested volume and caloric intake on the part of the dyspeptic patients (185 vs. 300 ml and 277 vs. 520 Kcal, respectively. Both: P<.001). No differences in the ST were observed between the two pure dyspepsia subtypes.

Conclusions: There was a difference in the ST between healthy individuals and those with dyspepsia, but the ingested volume and caloric intake in the two FD subtypes were similar.

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Dyspepsia, whose etymologic meaning refers to poor digestion, is a condition with a symptomatology that is common in the general population. It represents 2-3% of the medical consultations in general practice and 40% of those to gastroenterologists. However, only one fourth of the patients with dyspepsia consult a physician. Epidemiologic studies based on clinical practice have a clear selection bias, and the most representative are the population studies. An epidemiologic study evaluating “non-studied” dyspepsia in a population sample in Argentina reported a prevalence of 29.6%.

There are 2 main pathophysiologic alterations in functional dyspepsia (FD): (a) motor abnormalities of the proximal digestive tract (motility alterations) and (b) sensory visceral dysfunction (sensory perception alterations).

Many patients with FD have an antral motor dysfunction or a delay in gastric emptying, but it has not been possible to show a correlation between symptoms and this abnormality.

A subgroup of patients that present with symptoms of early satiety and weight loss, has gastric accommodation alterations. This appears to be a very important manifestation, due to the fact that it finally demonstrates some kind of correlation between certain types of symptoms and an identified pathophysiologic alteration.

Visceral sensory alteration (the capacity to perceive or not perceive what is happening within the digestive tract) is another important pathophysiologic aspect because this pathophysiologic mechanism of hypersensitivity is linked to epigastric pain, burping, and weight loss. Stress and anxiety modify sensory perception and they cause subjects to present with high levels of anxiety, depression, hypervigilance, and multiorgan somatic complaints.

Since the early 1990s, digestive function pathology has been classified according to the Rome Criteria. The current guidelines are those of the Rome III Criteria that recommend differentiating between patients with epigastric pain and those with postprandial distress, identified as distinct pathophysiologic subgroups. This subclassification is based on the main symptoms of pain or intermittent burning sensation localized in the epigastrium of at least average intensity presenting for a minimum of once a week, and postprandial fullness that occurs after a normal-sized meal various times a week. The criteria must be met during the last 3 months and the symptoms must have begun a minimum of 6 months before diagnosis. The 2 subtypes can coexist.

The exact etiopathogenic mechanism that causes the symptoms in an individual patient is still difficult to identify. Therefore, up to the present, clinical manifestations are the most effective diagnostic method and they determine the treatment strategy. The so-called satiety test (ST) is a recently described functional exploration of the digestive tract that enables the objective measurement of early satiety, a symptom observed in an important number of patients with FD. This simple, non-invasive test appears to hold an important place among the techniques used in the study of the pathophysiology of the motor and functional disorders of the stomach. It was originally designed for

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**Introduction**

Dyspepsia is a condition that can present with various symptoms such as abdominal pain, bloating, early satiety, and nausea. It is often associated with poor digestion and can have a significant impact on an individual's quality of life. Dyspepsia can be functional or organic, with functional dyspepsia (FD) being the most common form.

FD is characterized by symptoms such as epigastric pain, bloating, early satiety, and a feeling of fullness after eating a normal-sized meal. These symptoms are often intermittent and may not be associated with any underlying organic disease. The exact cause of FD is unknown, but it is believed to be linked to factors such as stress, anxiety, and dietary habits.

In recent years, research has focused on understanding the pathophysiology of FD, and several subtypes have been identified based on symptom characteristics. These subtypes include dyspeptic pain syndrome (DPS), dyspeptic early satiety syndrome (DES), and the mixed type.

The satiety test (ST) is a non-invasive and simple method used to differentiate between these subtypes. The ST involves measuring the volume and energy intake over a series of meals. Patients are instructed to eat until they feel full, and their eating behavior is observed.

The ST has been shown to be a useful tool in the clinical setting, as it can help to identify the type of FD a patient has. This information can then be used to tailor the treatment strategy, potentially leading to improved symptom relief and quality of life for patients with FD.
evaluating gastric symptoms during the intake of a liquid food, as a continuous and slow infusion, until reaching the level of maximum satiety. The test measured satiety by grading the volume of the ingested liquid on a Linkert scale of 0 to 5, with 5 being the highest score. A validation study conducted on a group of patients with FD suggested that the low caloric content test was correlated with the severity of early satiety symptoms and was potentially useful in assessing gastric accommodation.\textsuperscript{11}

Likewise, there is evidence that the ST enabled the differentiation between healthy individuals and those presenting with dyspepsia in European\textsuperscript{5,12} and North and Central American populations.\textsuperscript{13} Such a study has not been conducted in Mexico and therefore we wish to evaluate this distinction in the Mexican population.

Our working hypothesis was that the ST is a diagnostic method that would enable the differentiation between the 2 dyspepsia subtypes, postprandial distress syndrome (PDS) or epigastric pain syndrome (EPS), thus managing the therapy with greater precision.\textsuperscript{13}

Aims

1. To evaluate whether there was a difference in the ST (ingested volume and caloric intake) between healthy individuals and those presenting with dyspepsia.
2. To analyze the usefulness of the ST in differentiating between the 2 dyspepsia subtypes (PDS or EPS).

Methods

Study population

Adults of both sexes \( \geq 18 \) years of age diagnosed with FD according to the Rome III Criteria that signed statements of informed consent were included in the study. The exclusion criteria were: evidence of organic disease through laboratory tests, upper digestive and/or abdominal endoscopy, the use of prokinetic agents, calcium blockers, anti-depressives, opioid analgesics, nonsteroidal anti-inflammatory drugs or iron supplements, known systemic disease altering motility, hydroelectric disorders, a past history of upper digestive tract surgeries, pregnancy or lactation, dementia or inability to follow the protocol, and a history of alcohol and/or illegal drug abuse.

Controls

Twenty healthy volunteers that did not fit any of the Rome III Criteria for FD, that reported no personal history of digestive diseases, and that were not taking any concomitant medication for digestive disorders were recruited as controls.

Consecutive recruitment was carried out within the time frame of August 2011 to September 2012 at the Hospital de Gastroenterología in Buenos Aires, Argentina.

Study design

An exploratory, comparative, prospective, and cross-sectional study was conducted.

Procedures

Symptom questionnaire

Before the ST was performed, each patient was classified according to the dyspepsia subtype and the severity of the common symptoms through the reproducible dyspepsia questionnaire employed by Tack and Cuomo in 1998 and 2001, respectively\textsuperscript{13,14} (appendix 1).

The ST was subsequently carried out, recording the appearance and severity of symptoms before (baseline determination) and immediately after ingestion, and then 30 min after having completed the test (appendix 2).

Satiety Test

After a fasting period of 12 h, the subjects ingested a nutritional supplement (a semi-liquid foodstuff of a known and standardized composition: Fortisip, Nutricia Bagó\textsuperscript{®}). It was free of both fiber and lactose and had a balanced content of carbohydrates (49%), proteins (16%), and lipids (36%). It was ingested at a constant speed (15 ml/min) until reaching maximum satiety (the sensation of gastric fullness that inhibits the desire to continue drinking or eating). Kcal intake was determined when the patient reached maximum satiety, in other words, when the patient stated «I can’t drink another drop». This was registered in the questionnaire that was applied at 5-min intervals (appendix 3).

Antacid intake on demand as rescue medication was permitted one week prior to the study.

Measurement results:

1. Satiety in healthy individuals and in those presenting with dyspepsia: the ingested volume/caloric intake at the appearance of the symptom of maximum satiety was recorded to evaluate whether there was a difference between the two groups. When the test was performed, the ingested volume and caloric intake were registered at the instant the symptom of “maximum satiety” appeared, along with any eventual concomitant symptoms.
2. Usefulness of the test for differentiating between the two types of dyspepsia: the existence or not, of differences in the ingested volume and caloric intake between the two dyspepsia subtypes was evaluated. Dyspepsia subtype diagnosis was made according to the type and severity of the predominant symptom after the test:
   - PDS:
     - Postprandial fullness: the sensation of fullness after a regular meal at least several times a week.
     - Early satiety: the sensation of fullness that impedes finishing a regular meal at least several times a week.
   - EPS:
     - Pain or burning sensation in the epigastrium of at least average intensity minimum of once a week.
   - Mixed Syndrome.

Statistical analysis

The information was processed using MedCalc version 11.2.1.0 and the data were reported as means ± standard deviation. The results of the different groups were compared using the
corresponding chi-square test or the Fisher exact test for the qualitative values and the corresponding Student’s t test or Mann-Whitney test for the quantitative variables, according to normal or non-parametric distribution, respectively. Statistical significance was considered when there was a \( p < 0.05 \).

Because the study was an exploratory one, sample size was established at a minimum of 30 patients presenting with dyspepsia.

**Ethical safeguards**  
The patients signed statements of informed consent and the study protocol was approved by the Bioethics Committee of the Hospital de Gastroenterología “Dr. Carlos Bonorino Udaondo”.

**Results**

**Description of the sample**

Thirty-nine patients and controls were enrolled in the study; the characteristics of both the dyspeptic patients and the controls are described in Table 1. The FD subtypes were categorized according to the Rome III Criteria and the most frequent subtypes, in descending order, were: PDS: 61% (24/39), EPS: 31% (12/39), and Mixed: 8% (3/39).

**Results**

1. The comparison of the ingested volume and caloric intake between the dyspeptic patients and the controls showed that there was a statistically significant lower volume and caloric intake in the ST of the patients with dyspepsia (\( p < 0.001 \)) (Table 2).

2. Usefulness of the test in differentiating between the two types of dyspepsia: the existence or not, of differences in the ST in patients with pure PDS and EPS was evaluated. The patients with mixed syndromes (n=3) were excluded from this analysis. No statistically significant differences were observed between the two dyspepsia subtypes (Figs. 1 and 2 and Table 3).

The most frequent symptoms accompanying ingestion suspension, according to the dyspepsia subtype, are described as follows:

- **PDS**: postprandial fullness 50% (12/24), early satiety and nausea —individually— 16% (4/24), bloating and epigastric pain 8.33% (2/24).
- **EPS**: postprandial fullness 50% (6/12), nausea 25% (3/12), epigastric pain and nausea —together— 25% (3/12).

**Discussion**

Different organic diseases can cause symptoms that are consistent with dyspepsia. This disorder is common and usually has a benign cause. Current recommendations suggest that young patients < 45 years of age, with no alarm symptoms or risk factors for gastric disease can be treated...
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empirically with anti-secretors or prokinetics for 4 weeks, after which the response is evaluated. The patients that continue to present with symptoms despite this strategy, or those > 45 years of age, should undergo endoscopic evaluation.\textsuperscript{15}

FD is defined by the Rome III Criteria and subclassified into EPS and PDS; symptoms should be present during the last 3 months and have begun at least 6 months prior to diagnosis. Symptoms are nonspecific and the pathophysiology is diverse, which is why an effective universal treatment is still the subject of research.\textsuperscript{16,17}

Dyspepsia evaluation includes upper digestive tract video-endoscopy, scintigraphy study of gastric emptying, breath test, ultrasound or nuclear magnetic resonance (NMR) imaging, and evaluation of a gastric accommodation through NMR, PET, or barostat.

Other more specific studies that evaluate motor function alterations include antro-duodenal-jejunal manometry that is used to record the myoelectric activity of the stomach; the barostat and satiety test are used for evaluating visceral sensitivity.\textsuperscript{18}

The pathophysiologic mechanisms described in FD include motor abnormalities such as emptying and/or gastric accommodation disorders and visceral sensitivity dysfunction of unexplained clinical significance.\textsuperscript{19} Likewise, the lack of sensitivity and specificity of defined symptomatic patterns for identifying the respective pathophysiologic alteration, together with the fact that the tests for evaluating these disorders are invasive, laborious, and occasionally associated with radiation exposure, make the management of some of these patients difficult.\textsuperscript{20}

The ST has been developed as a non-invasive method for evaluating gastric accommodation in healthy individuals and in those presenting with dyspepsia.\textsuperscript{11,21} These tests are easily performed without specific instruments and are well-tolerated; however, their sensitivity and specificity vary depending on the ingested substance used (water or nutritional supplement), the method of comparison employed for greater diagnostic performance (ultrasonography or scintigraphy), and the technique used for carrying out the test.\textsuperscript{14,22,23}

Studies conducted on volunteers without dyspepsia that correlated the ST with the barostat showed the former to be useful in evaluating accommodation and the threshold for inducing severe discomfort. Nevertheless, the correlation between the ingested volume and the accommodation volume measured by the barostat is controversial.\textsuperscript{24-26}

Tack et al. observed that patients with FD tolerated fewer volumes than healthy individuals and that the ingested volume, at different caloric densities, was the major determinant of gastric accommodation to a meal.\textsuperscript{10} Cuomo et al. analyzed symptom severity through a reproducible questionnaire that correlated the ST parameters with gastric emptying and dyspeptic symptoms and concluded that the ST was able to objectively measure satiety and relate it to symptom severity. The study also showed an indirect correlation between early satiety severity measured by the ST and gastric emptying, as well as an association between flatulence and delayed gastric emptying.\textsuperscript{9}

Boecxkstaens et al. evaluated dyspeptic symptoms with the ST employing water and a nutritional supplement (Nutridrink) and observed that the FD patients reported more severe and persistent symptoms during the ST than the controls. In addition, the patients underwent the barostat study and were pathophysiologically classified as individuals with normal physiology, visceral hypersensitivity, and accommodation alterations. The symptoms triggered by the ST were not influenced by the results of the barostat study.\textsuperscript{27}

Our results coincided with those previously reported, confirming the fact that healthy individuals can be

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**Table 3** ST in FD subtypes.

<table>
<thead>
<tr>
<th>Satiety test</th>
<th>EPS</th>
<th>PDS</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients</td>
<td>12</td>
<td>24</td>
<td>—</td>
</tr>
<tr>
<td>Age (years), mean (range)</td>
<td>34.2 (19-62)</td>
<td>41.4 (21-68)</td>
<td>0.14</td>
</tr>
<tr>
<td>Sex (W/M)</td>
<td>8/4</td>
<td>22/2</td>
<td>0.40</td>
</tr>
<tr>
<td>BMI kg/m(^2), mean ± SD</td>
<td>23.1 ± 0.7</td>
<td>24.6 ± 0.8</td>
<td>0.28</td>
</tr>
<tr>
<td>Volume (ml), median (range)</td>
<td>177.5 (75-450)</td>
<td>197.5 (60-650)</td>
<td>0.48</td>
</tr>
<tr>
<td>Kcal, median (range)</td>
<td>261 (112-675)</td>
<td>296 (90-810)</td>
<td>0.50</td>
</tr>
</tbody>
</table>


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**Figure 2** ST results expressed in Kcal in patients presenting with dyspepsia subtypes: PDS and EPS.
distinguished from those with dyspepsia through the ST. To the best of our knowledge, there are currently no studies having evaluated the ST and the FD subtypes. In 2010, Kato et al. conducted a study utilizing the ST with water in order to differentiate distinct pathophysiologic mechanisms, but they did not include the clinical subtypes. Therefore, our study is the first to distinguish the clinical subtypes of FD through the ST.

Conclusions

According to our results, we can conclude that the ST was different between healthy individuals and patients presenting with dyspepsia, even though the ingested volume and caloric intake was similar in patients with PDS and EPS.

There is presently no scientific evidence at hand for recommending different alimentary instruction for these patients, and so clinical parameters continue to be the available tools for selecting the best-tolerated foods.

The reproducibility of this study with a greater number of patients would be useful for confirming our results.

Financial disclosure

No financial support was received in relation to this article.

Conflict of interests

The authors declare that there is no conflict of interest.

Appendix 1. Dyspepsia questionnaire.

Patient common symptom register

1. Postprandial fullness: sensation of fullness after a regular meal at least several times a week.

   0: absent
   1: mild, but does not interfere with customary activities
   2: relevant, bothersome, but does not interfere with customary activities
   3: severe, interferes with customary activities

2. Bloating: an excess of gas in the bowel that causes intestinal spasms and abdominal distension (the abdomen swells).

   0: absent
   1: mild, but does not interfere with customary activities
   2: relevant, bothersome, but does not interfere with customary activities
   3: severe, interferes with customary activities

3. Epigastric pain: pain or burning sensation located in the epigastrium of at least average intensity and presenting at least once a week.

   0: absent
   1: mild, but does not interfere with customary activities
   2: relevant, bothersome, but does not interfere with customary activities
   3: severe, interferes with customary activities

4. Early satiety: a sensation of fullness that impedes finishing a regular meal at least several times a week.

   0: absent
   1: mild, but does not interfere with customary activities
   2: relevant, bothersome, but does not interfere with customary activities
   3: severe, interferes with customary activities

5. Nausea:

   0: absent
   1: mild, but does not interfere with customary activities
   2: relevant, bothersome, but does not interfere with customary activities
   3: severe, interferes with customary activities

6. Vomiting:

   0: absent
   1: mild, but does not interfere with customary activities
   2: relevant, bothersome, but does not interfere with customary activities
   3: severe, interferes with customary activities

7. Burping:

   0: absent
   1: mild but does not interfere with customary activities
   2: relevant, bothersome, but does not interfere with customary activities
   3: severe, interferes with customary activities

8. Epigastric burning: A burning sensation in the epigastrium (mouth of the stomach) of at least average intensity and presenting for a minimum of one time a week.

   0: absent
   1: mild but does not interfere with customary activities
   2: relevant, bothersome, but does not interfere with customary activities
   3: severe, interferes with customary activities

Appendix 2. Register of symptoms before ingestion (baseline), immediately after ingestion, and 30 minutes after having completed the Satiety Test

Please mark an “X” on the scale below corresponding to the intensity of the symptom described:

0  Absence of symptoms
10 Maximum pain
1) Postprandial fullness
2) Bloating
3) Epigastric pain
4) Early satiety
5) Nausea
6) Epigastric burning

Appendix 3. Satiety Test

The level of satiety of the patient will be measured in 5-minute intervals through a visual analog scale from 1 to 5:

1: absence of symptoms
2: minimum satiety
3: moderate satiety
4: intense satiety
5: maximum satiety (inability to continue ingesting)

Additionally, the ingested volume, caloric intake, and the symptom identified as that motivating the ingestion suspension will be registered.

<table>
<thead>
<tr>
<th>Satiety register</th>
</tr>
</thead>
<tbody>
<tr>
<td>At 5 min:</td>
</tr>
<tr>
<td>Ingested volume:</td>
</tr>
<tr>
<td>Caloric intake:</td>
</tr>
<tr>
<td>1</td>
</tr>
<tr>
<td>At 10 min:</td>
</tr>
<tr>
<td>Ingested volume:</td>
</tr>
<tr>
<td>Caloric intake:</td>
</tr>
<tr>
<td>1</td>
</tr>
<tr>
<td>At 15 min:</td>
</tr>
<tr>
<td>Ingested volume:</td>
</tr>
<tr>
<td>Caloric intake:</td>
</tr>
<tr>
<td>1</td>
</tr>
<tr>
<td>At 30 min:</td>
</tr>
<tr>
<td>Ingested volume:</td>
</tr>
<tr>
<td>Caloric intake:</td>
</tr>
<tr>
<td>1</td>
</tr>
</tbody>
</table>

Mark with an “X” the symptom that motivated the ingestion suspension:

1. Postprandial fullness
2. Bloating
3. Epigastric pain
4. Early satiety
5. Nausea
6. Epigastric burning

References


