INTRODUCTION

Gastroesophageal reflux disease (GERD) is defined by our medical community as: “a chronic condition resulting from the reverse flow of gastroduodenal contents into the esophagus and/or adjacent organs, causing a variable range of esophageal and/or extraesophageal signs and/or symptoms, with or without tissue damage”.

It is a condition of great medical and social importance due to its high and growing incidence, causing long-term symptoms, which considerably affect the patients’ quality of life. It is estimated that GERD affects approximately 20% of the adult population in the United States and Europe.

Upper gastrointestinal (GI) endoscopy and esophageal pH monitoring are the methods directly linked to GERD diagnosis. The first method identifies forms of the disease causing esophagitis, allowing material collection for histologic examination, while the second helps diagnosing pathological gastroesophageal reflux (GER).

Esophageal pH monitoring, developed in the 1960s, was introduced into clinical practice in the 70’s. Initially, a glass pH probe was used in inpatients. A development of this technique was introduced in the early 80’s, with the use of flexible catheters and portable esophageal pH recorders in an outpatient setting. In the late 80’s, another important feature, the symptom index, was added to this method.

Prolonged esophageal pH monitoring enabled a better understanding of GERD. It is indicated for a practical approach, providing details of gastroesophageal and pharyngolaryngeal acid reflux, while associating clinical complaints to reflux episodes. However, there are some limitations: substantial discomfort,
Restriction of routine activities, and non-identification of pathological reflux in a considerable portion of patients with clinical and endoscopic evidence, suggestive of reflux.

Because the conventional esophageal pH monitoring is associated with discomfort, patients tend to exhibit reduced food intake and behave differently during the monitored period\(^{(7, 19)}\). Normal values of esophageal pH monitoring, in patients with endoscopic esophagitis, ranges from 17% to 31.4%\(^{(3, 5, 14, 15, 17, 21, 22, 28)}\). As a result, a normal pH monitoring does not exclude GERD diagnosis\(^{(3)}\).

A wireless esophageal pH monitoring system was developed in an attempt to improve the diagnostic sensitivity of the method, since it is better tolerated and allows a longer period of monitoring. Bile and multichannel intraluminal impedance pH monitoring (MII-pH) analyze other forms of GERD-related symptoms not assessed by pH monitoring: bile reflux and “non-acid” reflux, respectively.

Although wireless esophageal pH monitoring may cause chest discomfort because of the capsule, it is believed to be better tolerated than the conventional esophageal pH monitoring. It is believed that wireless esophageal monitoring may help provide better diagnostic sensitivity of GER, as it does not limit patient daily activities and allowing for longer periods of monitoring. However, there are no local and only a few international publications available, when it comes to the simultaneous comparative study of the wireless esophageal monitoring system and the conventional esophageal pH monitoring.

**Objective**

The objective is to compare the first 24 hours of the pH monitoring results, with and without catheter, positioned 3 cm above the lower esophageal sphincter (LES), with regard to: occurrence of technical failures during monitoring; ability to detect gastroesophageal acid reflux; ability to diagnose pathological gastroesophageal reflux; and the ability to relate clinical complaints to reflux episodes.

**METHODS**

Patients referred to the Esophageal Functional Investigation Laboratory of the Digestive System Surgery Department (Hospital das Clínicas da Universidade de São Paulo, São Paulo, SP, Brazil) were prospectively screened for 7 consecutive months, for esophageal pH monitoring.

Inclusion criteria were: heartburn and/or regurgitation as the main clinical complaint; at least 18 years of age; recent upper GI endoscopy (within the last 2 months); interruption in the administration of proton pump inhibitors for 7 days preceding the pH monitoring; and signature of the free and informed consent form. Exclusion criteria were: esophageal diverticula, strictures and varices; hiatal hernia greater than or equal to 3 cm; erosive esophagitis Los Angeles grades C or D; Barrett’s esophagus; and neoplasms, obstructive diseases or previous surgery of the gastrointestinal tract.

All patients underwent: clinical interview, nasal and oral esophageal manometry, pH monitoring with and without catheter for 24 and 48 hours, respectively, with simultaneous initial period.

The following GERD complaints were investigated during clinical interview: typical (heartburn and regurgitation), atypical (chest pain and globus sensation), and extraesophageal (cough, asthma, dysphonia and hem).

All patients underwent upper GI endoscopy at the Gastrointestinal Endoscopy Department of the Hospital das Clínicas da Faculdade de Medicina da Universidade de São Paulo. The presence of erosive esophagitis and hiatal hernia were assessed. The Los Angeles grade system was used for the characterization of esophagitis; the protrusion of part of the stomach 2 or more centimeters into the diaphragm, during deep inspiration, was considered hiatal hernia.

**Esophageal manometry**

Before pH monitoring, patients underwent two manometric examinations to identify the LES: nasal and oral.

Nasal esophageal manometry was performed conventionally, with nostrils local anesthesia using 2% lidocaine gel and a 4.5 mm (diameter) polyvinyl, flexible catheter, with eight recording channels perfused with 0.6 mL distilled water flow per minute. A complete esophageal manometry was performed; however, for the purpose of this study, only the location (distance from the nostril) of the LES was considered. The oral manometry was performed with local oropharynx anesthesia using 10% liquid aerosol lidocaine, with the same equipment used for the nasal manometry, for location (distance from the upper dental arch) of the LES.

**Esophageal pH monitoring**

After evaluating the distance of the LES in relation to the nostril and to the upper dental arch through manometry, a catheter of the conventional esophageal pH monitoring system was introduced, followed by the capsule of the wireless esophageal pH monitoring. Each patient underwent conventional pH monitoring for 24 hours, and wireless pH monitoring for 48 hours, with simultaneous monitoring recording starting time.

The equipment used for the conventional pH monitoring consisted of: portable recording device (Medtronic/Synetics, USA), calibration solutions and pH monitoring catheter (Alacer, Brazil). The 2.1 mm in diameter catheter displayed two antimony sensors (2 cm away from each other) for pH registration, and an external reference electrode. The distal sensor was positioned 3 cm above the superior border of the LES, identified through nasal esophageal manometry. By internationally accepted standards, the proximal sensor was positioned 5 cm above the superior border of the LES.

The wireless monitoring system equipment (Bravo, Medtronic/Synetics, USA) consisted of: portable pH monitoring recording device, calibration solutions, pH monitoring capsule, and capsule delivery device. The pH monitoring capsule used contained an antimony sensor, sensitive to pH changes, and an internal reference electrode. The capsule sensor was systematically calibrated before each test, using the same solutions at pH 7 and pH 1. The capsule, measu-
ring 6.0 x 6.3 x 26.0 mm was inserted through the mouth and positioned in the esophagus, 3 cm above the superior border of the LES, identified by oral manometry and at the same level as the conventional catheter’s distal sensor. The suction system was applied by a vacuum pump (510 mm Hg during 60 s) and the esophagus mucosa penetrated into the capsule compartment (4 mm in diameter). The pin was released, transfixing the suctioned mucosa, while attaching the capsule to the esophageal wall. The vacuum was turned off and the capsule released from the distal end of the delivery device, which was removed. The pH recording was started and transmitted by radio waves (telemetry) to the portable recording device.

Patients were advised to try to maintain their daily activities, to fill out the pH monitoring log, and to return to the laboratory after 24 hours (1st day) to remove the pH monitoring system, and again after another 24 hours (2nd day) to remove the external recording device of the wireless pH monitoring system.

It is important to note that the data recorded by both the conventional and the wireless pH monitoring system sensors positioned 3 cm above the superior border of the LES were compared. The data related to the conventional pH monitoring sensor positioned 5 cm above the superior border of the LES was used to complete the pH monitoring routine report, enabling the continuation of the patient’s usual treatment. However, in this study, we do not compare the results recorded 5 and 3 cm above the superior border of the LES.

To compare the two types of esophageal pH monitoring, the following parameters were considered: occurrence of relevant technical failures during the monitoring period; ability to detect gastroesophageal acid reflux; and ability to relate clinical symptoms with acid reflux episodes.

Any incident preventing or impairing the proper monitoring of reflux, such as early capsule drop, extended interference periods and absence of signal, were all considered relevant technical failures.

The data recorded over the first day of monitoring was used to evaluate the acid reflux detection ability; the following parameters were considered: percentage of total reflux time, percentage of reflux time in upright position, percentage of reflux time during supine position and characterization of the reflux pattern (physiological or pathological). It is noteworthy that parameters of normality for the characterization of pathological reflux were established by measuring reflux 5 cm above the LES and were only used in this study as reference values. The normal parameters used were: rate of total reflux time up to 4.5, rate of reflux time in an upright position up to 8.4, and rate of reflux time in a supine position up to 3.5. Please refer to Table 1 for specific values.

The patient was considered to be affected by pathological gastroesophageal reflux if: any of the three percentages of reflux time adopted were at levels higher than normal; or had quantitatively normal reflux, but with a significant relationship with the symptoms. The relationship between clinical complaint and gastroesophageal acid reflux was assessed by the symptom index and considered positive when equal or greater than 50%.

The evaluation of spontaneous detachment of the wireless system capsule from the esophageal wall was performed by a lateral chest X-ray, on the 30th day after its insertion.

This study was approved by the Ethics Committee for Analysis of Research Projects of the Clinical Hospital of the São Paulo University Medical School (number 1079/06). For the statistical study, conducted at the Laboratory of Statistics and Epidemiology, Department of Gastroenterology, São Paulo University Medical School, the following tests were used: Fisher’s exact test, Wilcoxon ratio, paired $t$-test and bilateral test. The rejection level for the null hypothesis was set at 0.05.

**RESULTS**

Twenty-five patients were included, 21 (84%) of which were female. Ages ranged from 34 to 73 years (average 52.4). All patients had as predominant symptom the typical GERD complaints. Sixteen (64%) patients had atypical complaints, and 10 (76%) had associated extraesophageal complaints. Upper GI endoscopy revealed erosive esophagitis in 8 (32%) patients and hiatal hernia in 11 (44%).

There was no significant difference between the two types of pH monitoring concerning technical failure during examination ($P = 0.463$). An early capsule drop occurred in one (4%) patient during the wireless method exam; there was no relevant technical failure in the group monitored with a catheter.

Regarding the detection capacity of gastroesophageal acid reflux (Table 1), there was significant difference between the two types of pH monitoring, with a higher reflux detection rate in patients being monitored with the wireless system (percentage of total reflux, $P = 0.001$; reflux in upright position, $P = 0.020$, and reflux in supine position, $P = 0.023$).

With regard to the reflux pattern, the conventional method detected pathological gastroesophageal reflux in 16 (64%) patients, while the wireless method in 19 (76%) patients. However, such difference did not reach statistical significance ($P = 0.355$).

**TABLE 1. Comparison of reflux time rate between the two methods of esophageal pH monitoring, positioned 3 cm above the lower esophageal sphincter, in 25 patients.**

<table>
<thead>
<tr>
<th>pH monitoring (day 1)</th>
<th>Min</th>
<th>Max</th>
<th>Median</th>
<th>Average</th>
<th>$P$</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Total reflux time rate</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conventional</td>
<td>0.1</td>
<td>13.8</td>
<td>4.1</td>
<td>5.0</td>
<td>0.001*</td>
</tr>
<tr>
<td>Wireless</td>
<td>0.1</td>
<td>21.4</td>
<td>6.1</td>
<td>7.0</td>
<td></td>
</tr>
<tr>
<td><strong>Reflux time rate in upright position</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conventional</td>
<td>0.5</td>
<td>13.6</td>
<td>6.2</td>
<td>6.3</td>
<td>0.020**</td>
</tr>
<tr>
<td>Wireless</td>
<td>0.1</td>
<td>19.2</td>
<td>7.4</td>
<td>7.8</td>
<td></td>
</tr>
<tr>
<td><strong>Reflux time rate in supine position</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Conventional</td>
<td>0.0</td>
<td>23.4</td>
<td>0.10</td>
<td>3.5</td>
<td>0.023*</td>
</tr>
<tr>
<td>Wireless</td>
<td>0.0</td>
<td>31.9</td>
<td>1.30</td>
<td>5.8</td>
<td></td>
</tr>
</tbody>
</table>

* Wilcoxon test; ** Paired t-test; Min = minimum; Max = maximum; Conventional = conventional esophageal pH monitoring; Wireless = wireless esophageal pH monitoring
As for the ability to relate clinical symptoms with reflux, out of the 25 patients studied, 20 (80%) exhibited symptoms during monitoring. The symptom index could be calculated in those patients with symptoms. There was no significant difference between the two types of pH monitoring systems, when it comes to the positivity of the symptom index ($P = 0.777$) (Table 2).

**TABLE 2.** Comparison of symptom index positivity between the two methods of esophageal pH monitoring, positioned 3 cm above the lower esophageal sphincter.

<table>
<thead>
<tr>
<th>Symptom index</th>
<th>Conventional (day 1)</th>
<th>Wireless (day 1)</th>
<th>$P$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$n$</td>
<td>%</td>
<td>$n$</td>
</tr>
<tr>
<td>Negative</td>
<td>8</td>
<td>32%</td>
<td>7</td>
</tr>
<tr>
<td>Positive</td>
<td>12</td>
<td>48%</td>
<td>13</td>
</tr>
<tr>
<td>No symptom</td>
<td>5</td>
<td>20%</td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
<td>25</td>
<td>100%</td>
<td>25</td>
</tr>
</tbody>
</table>

1 Bilateral proportion test ($H_0: p_1=p_2$, $H_1: p_1\neq p_2$). Conventional = conventional esophageal pH monitoring; Wireless = wireless esophageal pH monitoring.

The spontaneous detachment of the pH monitoring capsule from the esophageal wall was confirmed in all patients in the study by a lateral chest X-ray, on the 30th day after the capsule insertion. No patient experienced severe chest pain or any other symptom requiring endoscopic removal of the capsule.

**DISCUSSION**

The comparison between wireless and conventional esophageal pH monitoring was conducted at a level below the traditional level, by positioning the capsule and catheter sensors 3 cm above the superior border of the LES. Reasons for this choice included: changes in the mucosa due to GERD commonly occur next to the esophagogastric junction; feasibility study of inserting the capsule closest to this transition zone; and compare, at this level, GERD detection between the two types of pH monitoring sensors (capsule versus catheter).

The percentage of total reflux time considered normal for wireless pH monitoring (48h) varies from 4.4% to 5.3% according to the literature (24, 33). In this study, the same parameters of the conventional method (5 cm) were used; it is important to note that the main objective was to compare the detection of reflux, using both methods, at 3 cm above the LES.

Contraindications of the wireless pH monitoring system are limitations to the method and include: severe esophagitis, esophageal varices, bleeding, stenosis and obstruction of the gastrointestinal tract, the use of cardiac pacemakers and defibrillators. It should be noted that such conditions do not represent contraindications to the pH monitoring with a catheter. Magnetic resonance (MRI) is not recommended for 30 days after insertion of the capsule because of the risk of perforation, in case the capsule has not yet been completely eliminated. This restriction is also not applicable to pH monitoring with a catheter.

Complaints associated with heartburn and/or regurgitation was significant: 64% of patients had atypical complaints, and 76% had associated extraesophageal complaints. Nasi et al. (23) observed that there was a prevalence of typical complaints in 49.7% of patients referred for the conventional pH monitoring, and prevalence of atypical and/or extraesophageal complaints in 50.3%. The conventional pH monitoring study with two pH sensors (one in the distal esophagus and the other in the upper esophageal sphincter, or just above it), can be performed in patients whose main symptom is extraesophageal or globus sensation. Thus, a considerable group of patients (about half of the cases) would have restrictions to the use of the wireless pH monitoring system, since it is not possible to insert a second capsule in the upper esophageal sphincter or in the pharynx.

The finding of 4% of relevant technical failures during wireless pH monitoring supports literature’s current data, that indicate failures in 4.1% to 5% of cases (2, 9, 26). However, there was an improvement of this rate when compared to earlier studies that indicate failures from 11% to 13.3% (24, 31, 33), due to changes in equipment manufacture and increased time of vacuum suction of the mucosa, which enabled better fixation of the capsule. Early drop of the capsule occurred in the only patient who experienced diffuse esophageal spasm in the manometric study. The relationship between the two events is, however, debatable.

The conventional pH monitoring method has been considered the best diagnostic method of gastroesophageal acid reflux, offering sensitivity ranging from 79% to 96%, specificity from 85% to 100%, and 98% accuracy (24, 31, 33). The authors of the first simultaneous study also observed a greater rate of supine reflux time in the conventional method, stating that the wireless method has lower sensitivity when compared to the conventional method (4). However, in these studies the capsule was positioned with the aid of an endoscopic parameter at 5 or 6 cm above the squamocolumnar transitional zone (4, 10, 25). In this study, the capsule and the catheter were positioned with the aid of a manometric parameter at 3 cm above the upper border of the LES, showing different results: the three rates of reflux time (total, upright and supine) were significantly higher in the wireless pH monitoring. Perhaps the difference in the placement method of the pH sensors may have influenced the difference between our results and those reported in the literature.

There are possible explanations for the reflux detection differences between the methods: the logging interval of pH samples is different in each method (the wireless method registers samples every 6 seconds, while the conventional method does it every 4 seconds); the capsule positioning is...
fixed, while the catheter’s varies in relation to the LES during swallowing; and the wireless method detects fewer reflux episodes, especially those of short duration, when compared to the conventional pH monitoring.\(^{24, 28}\)

When simultaneously comparing the first 24 h of the conventional method with those of the wireless method, there was an increase in the diagnosis of pathological GERD in 12% of cases (64.0% vs 76.0%). Despite the fact that this difference did not reach statistical significance levels, the trend observed suggests that the wireless method may exhibit a greater diagnostic sensitivity when compared to the conventional pH monitoring.

Des Varannes et al.\(^{10}\) found a lower positivity in wireless pH monitoring compared to the conventional method (22.6% vs 29.0%), among clinical complaints and GER in the first 24 h using the symptom association probability. However, using the symptom index, we detected a higher positivity in the wireless method compared to the conventional method (52% vs 48%). But none of the studies reached statistical significance.

As for radiological control, Des Varannes et al.\(^{10}\) and Remes-Troche et al.\(^{26}\) observed spontaneous capsule detachment in all patients, during the first 14 days. While using the wireless method in 245 patients, with radiological control on the 14th day, Lin et al.\(^{10}\) observed that the capsule remained in 1% of cases. Considering this and the recommendation not to perform MRI in the first 30 days, radiological control was carried out in this study after 1 month of capsule insertion. After this time, if the capsule was still in place, an endoscopy withdrawal would be scheduled.

There was no need for capsule removal in any patient studied by Remes-Troche et al.\(^{28}\), which was also the case of the present study. However, there are reports of endoscopic capsule removal in 1.4% to 3.5% of cases, in large samples (90 to 245 patients).\(^{16, 23}\) The most common reason for withdrawal was severe chest pain.\(^{28}\) Other complications of the wireless pH monitoring reported in the literature are: esophageal perforation during insertion, esophageal ulcer, capsule migration to the nasopharynx after cough, and capsule retention in a colon diverticulum.\(^{16, 32}\) Because of this complication, we believe a simple radiography of the abdomen should be required for complete evaluation of the capsule elimination.

In a review article\(^{11}\) comparing GER monitoring methods (bile, pH and MII-pH monitoring), the wireless method is described as having better tolerability and greater sensitivity with regard to the conventional method. MII-pH monitoring significantly contributes to the understanding of the GERD pathogenesis, however it still has a very limited availability in social clinical care.

Finally, it is important to emphasize that a higher reflux detection rate by wireless pH monitoring was observed in this study, when compared to the conventional method. There was a slight advantage of the wireless method over the conventional method in the diagnosis of pathological GER, and in the ability to relate clinical complaints with GER, although without statistical significance levels.

**CONCLUSIONS**

Considering the conditions of this study, we may conclude that:

1. There is no significant difference between esophageal pH monitoring with and without catheter in terms of occurrence of relevant technical failures during monitoring;
2. The wireless pH monitoring detects reflux at significantly higher levels than conventional pH monitoring, when it comes to the three variables considered: total reflux time rate, reflux rate in upright and in supine positions;
3. There is no significant difference between the two methods of pH monitoring on the ability of diagnosing pathological gastroesophageal reflux;
4. The conventional esophageal pH monitoring and the wireless pH monitoring have similar capabilities of relating clinical complaints with gastroesophageal reflux.


**Objetivo** - Comparar as primeiras 24 horas das pHmetrias convencional e sem cateter, posicionadas a 3 cm acima do esfincter inferior do esôfago, em relação à: ocorrência de falhas técnicas relevantes, capacidade de detecção do refluxo e capacidade de relacionar as queixas clínicas com o refluxo.

**Métodos** - Foram estudados, de modo prospectivo, 25 pacientes encaminhados para pHmetria esofágica, com sintomas típicos da doença do refluxo gastroesofágico, submetidos a entrevista clínica, endoscopia digestiva, manometria esofágica e realização, com período inicial simultâneo, de pHmetrias com cateter por 24 horas e com cápsula por 48 horas.

**Resultados** - Houve queda precoce da cápsula em um paciente (4%) e nenhuma falha técnica na pHmetria com cateter (P = 0,463). As percentagens de tempo de refluxo (total, ortostático e supino) foram mais elevadas na pHmetria sem cateter (P<0,05). Refluxo gastroesofágico patológico foi diagnosticado em 16 (64,0%) pacientes com o cateter e em 19 (76,0%) com a cápsula (P = 0,355). O índice de sintomas foi positivo em 12 (48%) pacientes na pHmetria com cateter e em 13 (52%) na pHmetria sem cateter (P = 0,777).

**Conclusões** - 1) Não há diferença significante entre as duas modalidades de pHmetria (cápula vs cateter), em relação à ocorrência de falhas técnicas relevantes durante o exame; 2) A pHmetria sem cateter detecta refluxo em percentagens superiores às detectadas pela pHmetria convencional; 3) Os dois métodos de pHmetria têm capacidades semelhantes de diagnóstico de refluxo gastroesofágico patológico e capacidades semelhantes de relacionar as queixas clínicas com o refluxo gastroesofágico.

REFERENCES


