A citric acid solution is an optimal test drink in the 
$^{13}$C-urea breath test for the diagnosis of 
*Helicobacter pylori* infection

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Abstract

**Background**—The $^{13}$C-urea breath test ($^{13}$C-UBT) is a simple, non-invasive and reliable test for the diagnosis of *Helicobacter pylori* infection. The duration of the test, the timing of breath sampling, and the accuracy of the method vary according to the test meal used.

**Aim**—To identify the optimal test meal or drink for rapid and accurate performance of the $^{13}$C-UBT for the detection of *H pylori* infection.

**Patients**—Eighty patients with dyspeptic symptoms were included. Of these, 48 patients had a positive *H pylori* status and 32 a negative one according to the results of the rapid urease test, histological examination, and culture.

**Methods**—A $^{13}$C-UBT was performed after an overnight fast, on three consecutive days. On each study day a different test meal or drink was given (0.1 N citric acid solution, a standard semiliquid meal, or a semiliquid fatty meal) 10 minutes before giving 75 mg $^{13}$C-urea. Breath samples were collected at 0, 15, 30, 45, and 60 minutes, and analysed by isotype ratio mass spectrometry. Results were expressed as delta ($\delta$) and considered as positive for *H pylori* if the highest $\delta$ (peak) was greater than 4.0.

**Results**—The $\delta$ peak obtained with the citric acid drink in *H pylori* positive subjects (mean $\delta$ (SEM) 14.1 (SEM 1.5)) was significantly higher than that obtained with any of the semiliquid meals (13.3 (SEM 1.1) and 17.1 (SEM 1.0) respectively, $p<0.001$). Furthermore, the $\delta$ peak was obtained earlier with the citric acid drink (30 (SEM 2) minutes) than with the other two meals tested (53 (SEM 2) min and 45 (SEM 2) min, $p<0.001$). The sensitivity of the $^{13}$C-UBT for the diagnosis of *H pylori* infection was 96–100% with all three test meals. This high sensitivity was, however, obtained from 15 minutes by giving citric acid as the test drink, from 45 minutes by giving a semiliquid fatty meal, and at 60 minutes by giving the semiliquid standard meal. The specificity was 100% for all test meals. Citric acid is inexpensive and palatable to patients.

**Conclusions**—The $^{13}$C-UBT procedure with citric acid as the test drink is superior to the previously proposed semiliquid test meals in terms of $^{13}$CO$_2$ recovery, time requirement, and cost. In routine clinical sampling, collection at times 0 and 30 minutes seems to be optimal and gives a high diagnostic accuracy.

**Keywords:** *Helicobacter pylori*, diagnosis, breath test.

The $^{13}$C-urea breath test ($^{13}$C-UBT) is a non-invasive and simple test that reflects the hydrolysis of $^{13}$C-labelled urea by *Helicobacter pylori* urease. The reliability of this test in diagnosing *H pylori* infection is very high, with a sensitivity ranging from 90 to 98% and a specificity from 92 to 100%. The $^{13}$C-UBT may become the preferred method of assessing the effect of therapeutic regimens. Furthermore, this test is adequate for epidemiological research, with an advantage over serological methods of detecting active infection.

Since the original description by Graham et al of the non-invasive and non-radioactive $^{13}$C-UBT to identify the presence of *H pylori* urease activity, several modifications have been published aiming at simplifying and optimising the test. Quantity of substrate given, type of test meal, number of samples, and timing of sample collection have been the key variables investigated.

A sufficiently high amount of substrate is required to avoid exhaustion by oral bacteria containing urease. However, the $^{13}$C-urea dose should be reduced as far as possible for cost reasons. The amount of substrate used varies widely among different authors from 75 mg to 350 mg (5 mg/kg body weight). A dose of 100 mg has been proposed in a standardised European protocol, but 75 mg $^{13}$C-urea has been shown to be equally effective. The number of samples to be taken depends basically on the aim of the test: clinical diagnosis or research. For clinical routines, in which the $^{13}$C-UBT is required as a "yes or no" test, collection of two samples (before and after $^{13}$C-urea ingestion) is sufficient to provide a high diagnostic accuracy. Both dose of $^{13}$C-urea and composition of the test meal influence the optimal timing for sample collection. A readily exhaustible amount of substrate will provide an early and short $^{13}$CO$_2$ recovery peak, limiting the time frame for sample collection. On the other hand, timing of sample collection does not seem to be critical for at least one hour if a high substrate dose is given. Besides the basal sample, a single sample at 30 minutes is usually recommended. The need for giving a test meal in the $^{13}$C-UBT has been shown in several studies.
because administration of the urea with no additional test meal leads to gastric emptying of the substrate before sufficient reaction with H. pylori urease can take place. To inhibit gastric emptying a semiliquid fatty test meal is recommended by several authors.1-3 Solutions of glucose polymers and standardised commercial semiliquid meals have also been used.3,4 Good results have been obtained by using a citric acid drink to inhibit gastric emptying.4,8 In the original paper by Graham et al the test meal composition was reported as being not important.1 However, the duration of the test, timing of breath sampling, and the accuracy of the method vary among different studies using different types of test meal.1-11 In the present study we aimed at identifying the optimal test meal or drink for rapid and accurate performance of the 13C-UBT for the detection of H. pylori infection. The accuracy of the optimised test was also calculated.

Methods

Patients

Eighty patients (mean age 40, range 16–82 years, 41 men, 39 women) presenting for routine gastroscopy with dyspeptic symptoms gave their informed written consent for the study. The H. pylori status was investigated in all patients by: (a) the rapid urease test (HUT-Test, Astra GmbH, Germany)12 on one biopsy sample from the antrum and one from the gastric body, (b) histology on two biopsy samples from the antrum and two from the body, and (c) culture on two biopsy samples from the antrum. Biopsy specimens for histological examination were fixed in formalin, embedded in paraffin wax, and stained with Giemsa stain for detection of H. pylori. Bacteria were cultured at 37°C in 5% O2 for three days on Wilkins-Chalgren agar supplemented according to Skirrow (Anaerocult® C, Merck, Darmstadt, Germany). Diagnosis of H. pylori infection was based on either a positive culture or on a positive result in both the rapid urease test and histology. Patients had not received a compound containing bismuth, antibiotics, or a proton pump inhibitor in the four weeks before the study. None of the patients had a history of gastric surgery. The protocol was approved by the ethics committee of the University of Bonn.

Methods

13C-UBT was performed in all patients after an overnight fast on three consecutive days. On each study day a different test meal was given in a randomised order: (1) 200 ml 0-1 N citric acid solution, with the addition of 25 mg saccharin as sweetener; (2) 250 ml of a standard semiliquid meal (Meritene®; Wander Pharma; 237 kcal; 5 g fat, 20 g protein, and 28 g carbohydrate); and (3) a semiliquid fatty meal consisting of 50 ml Ensure® (Abbott) and 50 ml Calogen® (SHS Pharma; 275 kcal; 26.7 g fat, 2 g protein, 6-7 g carbohydrate). The price of each of these meals per test was 0-25 DM for the citric acid solution and about 6 DM for either of the two semiliquid meals. Ten minutes after ingestion of the test meal a baseline exhaled breath sample was collected in a CO2 storage capsule (CEDIOX-System®,Topic AG, Lichtenstein) and thereafter 75 mg 13C-urea dissolved in 50 ml water was given orally (T0). Further breath samples were taken at 15, 30, 45, and 60 minutes using the same CO2 storage capsules. To facilitate the simultaneous study of several patients and to simplify the test, turning the patients on their sides was avoided and they stayed seated over the whole study period. Collected samples were analysed by isotope ratio mass spectrometry (CEDIOX®, Topic AG, Lichtenstein), with a quadrupole mass spectrometer (Balzers AG, Asilier, Germany). Once all three tests were done, patients were asked which was the most pleasant test meal or drink.

Results were expressed as delta (δ), defined as the ratio (r–r0)/r0 where r = 13CO2/12CO2 (0 = basal sample; i = 15, 30, 45, 60 min). The result of the 13C-UBT is considered positive for H. pylori if maximal δ was greater than 4·0.

The within subject variability of the 13C-UBT in subjects with H. pylori infection was (mean (SEM)) 12 (1-2·4)% (unpublished data obtained by the daily performance of the 13C-UBT with citric acid as the test drink over three consecutive days in 20 H. pylori positive subjects).

Analysis of Data

The maximal δ value (peak) provided the end point of the test. A curve was defined by spline interpolation of the δ values at the different sampling times, and the area under the curve was calculated. The time to the maximal value of this curve (Tmax) was also measured. Results are expressed as means (SEM). Comparison between the three different test procedures was performed by repeated measures analysis of variance (ANOVA) with Bonferroni’s correction for multiple comparisons. The sensitivity and specificity of the 13C-UBT according to the test meal used for the diagnosis of H. pylori infection was calculated.
considered as the result of the test (as usually done routinely), all 
H pylori positive patients had a positive test with citric acid (sensitivity 
100%), three patients had a false negative result with Ensure'-Calogen' (sensitivity 94%), and 13 patients had a false negative result with Merite" (sensitivity 73%). All H 
pylori negative subjects had a negative result in the 
1C-UBT with all three test meals (specificity 100%).

Discussion

Our study shows that the 
1C-UBT performed with a solution of 0.1 N citric acid as test drink provides higher 
1CO2 recovery values and peaks recorded at earlier times than both a 
standard commercialised semiliquid meal (Merite') and a semiliquid fatty meal (Ensure'-Calogen') in patients with 
H pylori infection. As a result the duration of the test may be reduced without loss of sensitivity and specificity by using citric acid as the test drink. Furthermore, citric acid is inexpensive and palatable to patients.

Since the description of the 
1C-UBT for the diagnosis of 
H pylori infection,5 several modifications have been reported with the aim of simplifying and optimising the test.1,10-12 Most of them have evaluated different substrate doses as well as different times of sample collection. On the one hand, the dose of 
1C-urea to be given must be high enough to prevent its rapid exhaustion; on the other, cost is a limiting factor. Doses up to 350 mg (5 mg/ kg body weight) of substrate have been given and 100 mg is the recent recommended dose by a European Working party.12 However, doses as low as 75 mg are as reliable as higher doses.5 Based on this finding 75 mg of 
1C-urea was used in the present study.

The main goal for giving a test meal is to slow the gastric emptying of the substrate and thus allow a longer reaction time between 
1C-urea and 
H pylori urease. A lower gastric emptying is achieved by the semiliquid meals through their fatty content, whereas with the citric acid solution this effect is obtained by its low pH (3.0–3.5). We have recently found that the intraduodenal infusion of the citric acid solution used in the present study induces an
Sensitivity (% 95% CI) of the \(^{13}\)C-UBT for the diagnosis of \(H\) pylori infection according to the test meal used and the sampling time

<table>
<thead>
<tr>
<th>Test meal</th>
<th>15</th>
<th>30</th>
<th>45</th>
<th>60</th>
</tr>
</thead>
<tbody>
<tr>
<td>Citric acid (CA)</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>(92-6-100)</td>
<td>(92-6-100)</td>
<td>(92-6-100)</td>
<td>(92-6-100)</td>
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</tr>
<tr>
<td>Mereine</td>
<td>56</td>
<td>73</td>
<td>88</td>
<td>96</td>
</tr>
<tr>
<td>(42-3-69-3)</td>
<td>(59-0-83-4)</td>
<td>(75-3-94-1)</td>
<td>(86-0-98-8)</td>
<td></td>
</tr>
<tr>
<td>Ensure/Calogen (EC)</td>
<td>90</td>
<td>94</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>(77-9-95-5)</td>
<td>(83-2-97-8)</td>
<td>(92-6-100)</td>
<td>(92-6-100)</td>
<td></td>
</tr>
<tr>
<td>Statistical comparison</td>
<td>CA v M: p&lt;0.001</td>
<td>CA v M: p&lt;0.001</td>
<td>CA v M: p&lt;0.05</td>
<td>CA v M: NS</td>
</tr>
<tr>
<td></td>
<td>CA v EC: p&lt;0.05</td>
<td>CA v EC: NS</td>
<td>CA v EC: NS</td>
<td>CA v EC: NS</td>
</tr>
<tr>
<td>(binomial test)</td>
<td>EC v M: p&lt;0.001</td>
<td>EC v M: p&lt;0.05</td>
<td>EC v M: p&lt;0.05</td>
<td>EC v M: NS</td>
</tr>
</tbody>
</table>

inhibition of antral motility and a relaxation of the gastric fundus similar to that achieved by the intraduodenal perfusion of a fatty solution. 

The selection of the optimal sampling time not only depends on the quantity of substrate given, but also on the time needed for the hydrolysis of \(^{13}\)C-urea by contact with the \(H \) pylori urease. The test meal chosen for this purpose does, therefore, plays an important part.

A single sample at 30 or 60 minutes\(^3\) \(^6\) or multiple sample collections up to 90 minutes\(^3\) \(^5\) \(^11\) have been recommended by performing the test with different semi-liquid meals. Three studies have been reported with citric acid solution as the test drink; and the collection of a single sample at 20 minutes\(^3\) or at 30 minutes\(^8\) \(^9\) has been recommended in these cases. In the present study we found that the time required to reach the maximal \(^{13}\)C-exhalation was 30 (2) minutes when citric acid was given, 45 (2) minutes when a semi-liquid fatty meal was used, and 53 (2) minutes when a standard commercial semi-liquid meal was administered. Collection of a single sample at an earlier time (for example, 30 minutes) leads to a lower sensitivity. Therefore, a single postprandial breath sample is adequate only if the optimal sampling time is identified. With the citric acid solution this time point was at 30 minutes, although all patients had already shown a positive result at 15 minutes, compared with 45 minutes in the best case when a semiliquid test meal was used.

Some authors recommend turning the patients on to each side, head down for several minutes after ingestion of the test meal and \(^{13}\)C-urea to guarantee the contact of the substrate with the bacterial urease in the stomach.\(^1\) \(^2\) This seems to be unnecessary from the high diagnostic accuracy obtained in our study, in which we did not do this. This simplification of the test allows the simultaneous study of several patients.

The greater and earlier \(^{13}\)CO\(_2\) recovery obtained with citric acid as the test drink may indicate an enhanced contact between \(^{13}\)C-urea and \(H \) pylori urease mediated by this solution. Other mechanisms such as alterations in the metabolism of the organism by citric acid cannot however be excluded.

The citric acid solution is about 24 times cheaper than any of the semiliquid meals. Furthermore, and contrary to the semiliquid fatty meal, the citric acid solution can easily be stored at 4°C for weeks.

In conclusion, citric acid as the test drink in the \(^{13}\)C-UBT seems to be practical and accurate for detecting \(H \) pylori infection. This test procedure is superior to the previously proposed semiliquid test meals in terms of \(^{13}\)CO\(_2\) recovery, time requirement, and cost. Citric acid is, furthermore, palatable for patients. In clinical routine, sampling collection at times 0 and 30 minutes seems to be optimal and gives a high diagnostic accuracy.

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