14-day triple, 5-day concomitant, and 10-day sequential therapies for *Helicobacter pylori* infection in seven Latin American sites: a randomised trial


Summary

Background Evidence from Europe, Asia, and North America suggests that standard three-drug regimens of a proton-pump inhibitor plus amoxicillin and clarithromycin are significantly less effective for eradication of *Helicobacter pylori* infection than are 5-day concomitant and 10-day sequential four-drug regimens that include a nitroimidazole. These four-drug regimens also entail fewer antibiotic doses than do three-drug regimens and thus could be suitable for eradication programmes in low-resource settings. Few studies in Latin America have been done, where the burden of *H pylori*-associated diseases is high. We therefore did a randomised trial in Latin America comparing the effectiveness of four-drug regimens given concomitantly or sequentially with that of a standard 14-day regimen of triple therapy.

Methods Between September, 2009, and June, 2010, we did a randomised trial of empiric 14-day triple, 5-day concomitant, and 10-day sequential therapies for *H pylori* in seven Latin American sites: Chile, Colombia, Costa Rica, Honduras, Nicaragua, and Mexico (two sites). Participants aged 21–65 years who tested positive for *H pylori* by a urea breath test were randomly assigned by a central computer using a dynamic balancing procedure to: 14 days of lansoprazole, amoxicillin, clarithromycin, and metronidazole (concomitant therapy); or 5 days of lansoprazole and amoxicillin followed by 5 days of lansoprazole, clarithromycin, and metronidazole (sequential therapy). Eradication was assessed by urea breath test 6–8 weeks after randomisation. The trial was not masked. Our primary outcome was probability of *H pylori* eradication. Our analysis was by intention to treat. This trial is registered with ClinicalTrials.gov, registration number NCT01061437.

Findings 1463 participants aged 21–65 years were randomly allocated a treatment: 488 were treated with 14-day standard therapy, 489 with 5-day concomitant therapy, and 486 with 10-day sequential therapy. The probability of eradication with standard therapy was 82·2% (401 of 486), which was 8·6% higher (95% adjusted CI 2·6–14·5) than with concomitant therapy (73·6% [360 of 489]) and 5·6% higher (–0·04% to 11·6) than with sequential therapy (76·5% [372 of 486]). Neither four-drug regimen was significantly better than standard triple therapy in any of the seven sites.

Interpretation Standard 14-day triple-drug therapy is preferable to 5-day concomitant or 10-day sequential four-drug regimens as empiric therapy for *H pylori* infection in diverse Latin American populations.


Introduction *Helicobacter pylori* infects most of the world's adult population and is the principal cause of gastric cancer, accounting for an estimated 60% of cases.1,2 Gastric cancer is second only to lung cancer as a cause of cancer death worldwide, and almost all of the nearly 1 million cases and 0·75 million deaths each year occur in east Asia and Latin America.3 Although gastric cancer death rates have fallen in recent decades, the number of deaths has actually increased as a consequence of ageing populations, and gastric cancer is projected to rank among the ten leading global causes of death by 2030.4,5 *H pylori* is also the main cause of peptic ulcer disease, which accounts for the loss of about 4·6 million disability-adjusted life-years every year worldwide, with most of the burden borne by populations in low-income and middle-income countries.5,6 Population-wide eradication programmes seem to offer the most direct approach to reducing the enormous human and economic consequences of *H pylori* infection; however, none has been implemented to date.5

Large programmes for *H pylori* eradication require a practical and inexpensive antibiotic regimen that is effective in the specific locale where it will be used. Standard antibiotic regimens for *H pylori* usually entail a proton-pump inhibitor, amoxicillin, and clarithromycin, taken together over 7–14 days.5,6 However, the effectiveness of these triple-therapy regimens seems to have diminished over time, largely as a result of emerging resistance of the organism to clarithromycin.5,6,7 Recent meta-analyses have shown that regimens that add a nitroimidazole (metronidazole or tinidazole) to triple

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therapy and are given either sequentially for 10 days or concomitantly for 5 days are significantly more successful at eradication of *H pylori* infection than are triple-therapy regimens.14–16 These regimens also require fewer doses of antibiotics and thus might be more affordable in low-resource settings. Almost all the evidence supporting these four-drug regimens comes from Europe and Asia; few data are available from Latin America, a region with some of the world’s highest rates of gastric cancer mortality.17 We therefore undertook a randomised trial in Latin America comparing the effectiveness of four-drug regimens given concomitantly or sequentially with that of a standard 14-day regimen of triple therapy. The trial also provided insights into the feasibility of community-based programmes of *H pylori* eradication.

**Methods**

**Study design and patients**

The trial (SWOG S0701) was a randomised trial of empiric 14-day triple, 5-day concomitant, and 10-day sequential therapies for *H pylori* infection in seven Latin American sites: Chile (Santiago), Colombia (Tiquerres), Costa Rica (Guancaste), Honduras (Santa rosa de Copán), Mexico (Ciudad Obregón and Tapachula), and Nicaragua (León). Between September, 2009, and June, 2010, study research staff recruited potential participants from the general population of adult men and women aged 21–65 years and explained the purpose and eligibility requirements of the study to them. Staff in Colombia, Costa Rica, and Nicaragua selected individuals from a census of households, in Chile they selected potential participants from a list of individuals served by a large public primary care clinic, and in Honduras and Mexico (two sites) they recruited participants by walking house-to-house within the local community or through announcements at primary care clinics.

Study participants in Tapachula (Mexico), Nicaragua, and Chile were predominantly urban, and those in the other sites were from small, rural communities. Potential participants were deemed ineligible if they had been treated in the past for *H pylori* infection, had serious illnesses that might end their lives before completing the study, or had other disorders that required or precluded treatment with antibiotics or proton-pump inhibitors. They also had to agree to abstain from alcohol use for at least 2 weeks. Those who expressed an interest in participating and gave signed, informed consent then completed an interview regarding socioeconomic characteristics and health history and a detailed gastrointestinal-symptom-history assessment with the validated Spanish language version of the Rome III diagnostic questionnaire for the adult functional gastrointestinal disorders.18,19 The institutional review boards for each clinical centre and for the SWOG Statistical Center, Seattle, WA, USA approved the study protocol.

**Procedures**

Participants provided a urea breath test for *H pylori* infection by exhaling into foil balloons before, and 30 min after, consuming a 75 mg dose of ¹³C-labelled urea with water. Staff at each centre analysed the breath samples using an infrared mass spectrometry device (IRIS, Wagner Analysen Technik, Bremen, Germany) that produced a computer-generated result of positive (change relative to baseline ≥4·0%) or negative (change relative to baseline <2·5%); intermediate values were classified as inconclusive. If a participant reported use of an antibiotic or proton-pump inhibitor within the past 15 days, or if the result from the urea breath test was inconclusive, the test was rescheduled for a later date.

Study staff contacted participants at least once during treatment to encourage adherence and to remind them to return any unused doses at their follow-up visit, which was scheduled to occur 6–8 weeks after randomisation. During follow-up visits, participants completed another interview assessing adherence to therapy, their reasons for missing any doses of the regimens, and the occurrence of any new or worsened medical disorders that led them to seek medical attention. Study staff counted the number of drug doses returned and administered the follow-up urea breath test.

**Randomisation and masking**

Participants who had a positive urea breath test and met all other eligibility criteria were randomly assigned, in equal proportions, to one of three treatment groups: standard triple therapy of lansoprazole 30 mg, amoxicillin 1000 mg, and clarithromycin 500 mg taken twice a day for 14 days; 10-day sequential therapy of clarithromycin and amoxicillin for 5 days followed by lansoprazole and metronidazole for 5 days; and 14-day triple therapy of lansoprazole, amoxicillin, and clarithromycin. Participants were randomised in a 1:1:1 ratio to each treatment group using computer-generated random numbers. Randomisation code was made available to principal investigators and site coordinators but was unblinded until the end of the study. Patients and study staff were masked to treatment regimens within the study period and were unmasked only after the study was closed. Safety and adverse events were monitored through a standard case report form. Participants who completed the study were scheduled for a follow-up visit 6–8 weeks later, at which a repeat urea breath test was performed. Follow-up visits continued for participants who withdrew consent or were lost to follow-up, or participants whose test results indicated that they were negative for *H pylori* infection.

**Figure: Trial profile**

UBT=urea breath test.
14 days; concomitant therapy of lansoprazole 30 mg, amoxicillin 1000 mg, clarithromycin 500 mg, and metronidazole 500 mg taken twice a day for 5 days; or sequential therapy of lansoprazole 30 mg, clarithromycin 500 mg, and metronidazole 500 mg taken twice a day for 5 days. Each clinical centre purchased its own supply of the drugs, as generic (off-patent) preparations, from suppliers in country, except the centres in Honduras and Nicaragua, which both used the same supplier in Honduras. The drug suppliers provided quality-control data for the content and dissolution of each drug. Randomisation was implemented centrally through a dynamic balancing procedure at the SWOG Statistical Center to ensure balance within centre by age and sex across the three regimens. Staff at the clinical centres entered data for potentially-eligible, consented individuals into the SWOG Statistical Center computer using a web-based data entry system. If the participants met all eligibility requirements, the computer assigned them to a treatment group and immediately transmitted the treatment assignment to the clinical centre. All participants were randomly assigned in a 1:1:1 ratio within 2 weeks after a positive urea-breath-test result. The trial was not masked.

**Statistical analysis**

We designed the study to address two primary hypotheses. The first hypothesis was that 5-day concomitant therapy was not inferior to 14-day standard therapy, whereby inferiority was defined as a difference in eradication probability of 5% or greater in favour of standard therapy. We reasoned that the shorter duration and lower cost associated with concomitant therapy would make it clearly superior in eradication of *H pylori* infection. Our second hypothesis was that 10-day sequential therapy would be more effective than 14-day standard therapy, because a four-drug sequential regimen would be preferred over standard three-drug therapy only if it were clearly superior in eradication of *H pylori* infection. Assuming an eradication rate of 80% for standard therapy, 10% missing follow-up urea-breath-test results, and 210 randomised participants per centre (1470 total), each treatment group comparison would have 82% power to detect a difference of 8% or greater, based on a two-sided, 0.025-level test. Although a meta-analysis by Essa and colleagues suggested that concomitant therapy was about 10% better than sequential therapy, if the therapies were actually equivalent the trial would have 46% power to reject the inferiority of concomitant therapy, based on a one-sided, 0.025-level test.

Our primary statistical analyses adhered to the intention-to-treat principle and included all randomised eligible participants, with those without a definitive follow-up urea breath test judged to be treatment failures (urea-breath-test positive). To compare concomitant versus standard therapy, we used a two-sample Z test of the null hypothesis, in which the difference in the estimated probabilities of eradication was 5% or greater

<table>
<thead>
<tr>
<th>Centre</th>
<th>14-day standard therapy (N=488)</th>
<th>5-day concomitant therapy (N=489)</th>
<th>10-day sequential therapy (N=486)</th>
<th>Total (N=1463)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Santiago (Chile)</td>
<td>69 (14%)</td>
<td>70 (14%)</td>
<td>70 (14%)</td>
<td>209 (14%)</td>
</tr>
<tr>
<td>Túquerres (Colombia)</td>
<td>71 (15%)</td>
<td>69 (15%)</td>
<td>70 (14%)</td>
<td>212 (15%)</td>
</tr>
<tr>
<td>Guanacaste (Costa Rica)</td>
<td>70 (14%)</td>
<td>70 (14%)</td>
<td>70 (14%)</td>
<td>210 (14%)</td>
</tr>
<tr>
<td>Copán (Honduras)</td>
<td>70 (14%)</td>
<td>70 (14%)</td>
<td>70 (14%)</td>
<td>210 (14%)</td>
</tr>
<tr>
<td>Tapachula (México)</td>
<td>71 (15%)</td>
<td>70 (14%)</td>
<td>70 (14%)</td>
<td>210 (14%)</td>
</tr>
<tr>
<td>Obregón (México)</td>
<td>70 (14%)</td>
<td>69 (14%)</td>
<td>70 (14%)</td>
<td>210 (14%)</td>
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<tr>
<td>León (Nicaragua)</td>
<td>67 (14%)</td>
<td>67 (14%)</td>
<td>65 (12%)</td>
<td>199 (14%)</td>
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</table>

<table>
<thead>
<tr>
<th>Sex</th>
<th></th>
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<th></th>
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</thead>
<tbody>
<tr>
<td>Women</td>
<td>287 (59%)</td>
<td>288 (60%)</td>
<td>286 (59%)</td>
<td>861 (59%)</td>
</tr>
<tr>
<td>Men</td>
<td>201 (41%)</td>
<td>201 (41%)</td>
<td>200 (41%)</td>
<td>602 (41%)</td>
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</table>

<table>
<thead>
<tr>
<th>Age (years)</th>
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<th></th>
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<tbody>
<tr>
<td>20–29</td>
<td>66 (14%)</td>
<td>66 (14%)</td>
<td>91 (19%)</td>
<td>223 (15%)</td>
</tr>
<tr>
<td>30–39</td>
<td>137 (28%)</td>
<td>139 (28%)</td>
<td>117 (24%)</td>
<td>333 (23%)</td>
</tr>
<tr>
<td>40–49</td>
<td>142 (29%)</td>
<td>117 (24%)</td>
<td>127 (26%)</td>
<td>386 (26%)</td>
</tr>
<tr>
<td>≥50</td>
<td>143 (29%)</td>
<td>167 (34%)</td>
<td>151 (31%)</td>
<td>461 (32%)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Years of education</th>
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<tbody>
<tr>
<td>≤4</td>
<td>88 (18%)</td>
<td>77 (16%)</td>
<td>80 (17%)</td>
<td>245 (17%)</td>
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<tr>
<td>5–8</td>
<td>135 (28%)</td>
<td>166 (34%)</td>
<td>143 (29%)</td>
<td>444 (30%)</td>
</tr>
<tr>
<td>9–12</td>
<td>146 (30%)</td>
<td>135 (28%)</td>
<td>136 (28%)</td>
<td>417 (29%)</td>
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<tr>
<td>≥13</td>
<td>71 (15%)</td>
<td>71 (15%)</td>
<td>70 (14%)</td>
<td>212 (14%)</td>
</tr>
<tr>
<td>Not reported</td>
<td>48 (10%)</td>
<td>40 (8%)</td>
<td>52 (12%)</td>
<td>140 (10%)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Chronic dyspeptic symptoms</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Present</td>
<td>125 (26%)</td>
<td>121 (25%)</td>
<td>127 (26%)</td>
<td>373 (26%)</td>
</tr>
<tr>
<td>Absent</td>
<td>363 (74%)</td>
<td>368 (75%)</td>
<td>359 (74%)</td>
<td>1090 (75%)</td>
</tr>
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</table>

**Table 2:** Adherence to treatment of patients that returned for 6-week follow-up, by treatment group

<table>
<thead>
<tr>
<th>Amount of drugs taken</th>
<th>Standard therapy (N=475)</th>
<th>Concomitant therapy (N=471)</th>
<th>Sequential therapy (N=470)</th>
<th>Total (N=1416)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>All (100%)</td>
<td>427 (90%)</td>
<td>442 (94%)</td>
<td>437 (93%)</td>
<td>1306 (92%)</td>
</tr>
<tr>
<td>Nearly all (&gt;80%)</td>
<td>7 (2%)</td>
<td>0 (0)</td>
<td>2 (&lt;1%)</td>
<td>9 (&lt;1%)</td>
</tr>
<tr>
<td>Most (50–80%)</td>
<td>24 (5%)</td>
<td>14 (3%)</td>
<td>21 (4%)</td>
<td>59 (4%)</td>
</tr>
<tr>
<td>Less than half (&lt;50%)</td>
<td>10 (2%)</td>
<td>8 (2%)</td>
<td>5 (1%)</td>
<td>23 (2%)</td>
</tr>
<tr>
<td>Undetermined (but not all)</td>
<td>7 (2%)</td>
<td>5 (1%)</td>
<td>5 (1%)</td>
<td>17 (1%)</td>
</tr>
<tr>
<td>None</td>
<td>0 (0)</td>
<td>2 (&lt;1%)</td>
<td>0 (0)</td>
<td>2 (&lt;1%)</td>
</tr>
</tbody>
</table>

**Reasons for not taking all drugs**

<table>
<thead>
<tr>
<th>Reason</th>
<th>Standard therapy (N=475)</th>
<th>Concomitant therapy (N=471)</th>
<th>Sequential therapy (N=470)</th>
<th>Total (N=1416)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Concern about having or developing side-effects</td>
<td>41 (9%)</td>
<td>28 (6%)</td>
<td>32 (7%)</td>
<td>102 (7%)</td>
</tr>
<tr>
<td>Unrelated illness or injury</td>
<td>3 (1%)</td>
<td>1 (&lt;1%)</td>
<td>5 (1%)</td>
<td>9 (&lt;1%)</td>
</tr>
<tr>
<td>Forgot or inconvenient</td>
<td>36 (8%)</td>
<td>25 (5%)</td>
<td>37 (8%)</td>
<td>98 (7%)</td>
</tr>
<tr>
<td>Reason not given</td>
<td>0 (0)</td>
<td>3 (&lt;1%)</td>
<td>2 (&lt;1%)</td>
<td>5 (&lt;1%)</td>
</tr>
</tbody>
</table>

Data are n (%). *Includes 1414 participants with a valid follow-up urea breath test and two (from the sequential therapy group) whose urea breath test results were inconclusive. †Based on count of returned drugs and self-report among participants who returned for a follow-up urea breath test. Multiple responses allowed.

Data are n (%). *Includes 1414 participants with a valid follow-up urea breath test and two (from the sequential therapy group) whose urea breath test results were inconclusive. †Based on count of returned drugs and self-report among participants who returned for a follow-up urea breath test. Multiple responses allowed.
in favour of the standard regimen, based on a one-sided test of non-inferiority. Comparison of sequential versus standard therapies was based on a two-sample Z test for no difference between eradication probabilities. Sensitivity to missing data assumptions was examined by excluding data from participants without a conclusive follow-up urea breath test. To establish how poor adherence could have affected our conclusions, we further restricted the analyses to participants with a definitive urea breath test who had taken at least 80% of their assigned study drugs.

Secondary analyses assessed variability in treatment outcome by sex, age, presence of chronic dyspeptic symptoms, and clinical centre. Tests of interaction were calculated as deviance tests comparing logistic regression models with the treatment group indicators, the designated covariate, and their interaction terms, with those without the interaction terms.

All analyses were done with SAS version 9.2 and R version 2.12.2 statistical software. Bonferroni-adjusted 95% CIs were used to account for the two primary comparisons, and p values less than 0.025 were classed as statistically significant. No corrections for multiplicity were applied to secondary analyses, because they were considered to be exploratory. All p values were two-sided except the test of non-inferiority.

This trial is registered with ClinicalTrials.gov, registration number NCT01061437.

**Role of the funding source**

The sponsors of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report. All authors had full access to the data in the study and participated in the decision to submit for publication.

**Results**

1859 potentially-eligible adults agreed to participate in the study and completed the screening interview and intake questionnaire, but seven withdrew before the urea breath test (figure). The test was positive for 1471 (79%) of 1852 tested participants. Eight patients with a positive test were not randomly assigned because they opted not to be treated, could not be randomly assigned within 2 weeks of their urea breath test, or had a disqualifying factor (eg, pregnancy). Six participants with a negative urea breath test, who were randomly assigned incorrectly because of data entry errors, were withdrawn from the study before receiving treatment, and their data were not included in any of the following analyses. Of the eligible participants who were randomised, 861 (59%) of 1463 were women, 847 (58%) of 1463 were age 40 years or older, and 373 (25%) of 1463 had chronic dyspeptic symptoms as classified by the Rome III criteria. Participant characteristics were balanced between the three treatment groups (table 1). 47 participants did not return for their follow-up urea breath test, and two participants had follow-up tests that were inconclusive after two repeat tests. Thus, we obtained definitive follow-up urea-breath-test results for 1414 (97%) of 1463 of the randomised eligible participants (figure).

Table 2 shows adherence to treatment of patients that returned for 6-week follow-up. 1313 (92%) of

### Table 3: Helicobacter pylori eradication by treatment group for three definitions of analysis population

<table>
<thead>
<tr>
<th>N</th>
<th>Helicobacter pylori eradication</th>
<th>Difference from standard group (adjusted 95% CI for difference)</th>
<th>p value for interaction*</th>
</tr>
</thead>
</table>
|### Intention to treat (N=1463)###
| 14-day standard therapy | 488 | 401 (82.2% [78.5 to 85.5]) | – |
| 5-day concomitant therapy | 349 | 360 (73.6% [69.5 to 77.5]) | 8.6% (2.6 to 14.5) |
| 10-day sequential therapy | 486 | 372 (76.5% [72.5 to 80.2]) | 5.6% (-0.4 to 11.6) |
|### Definitive 6-week UBT (N=1414)###
| 14-day standard therapy | 475 | 401 (84.4% [80.8 to 87.6]) | – |
| 5-day concomitant therapy | 347 | 360 (76.4% [72.3 to 80.2]) | 8.0% (2.2 to 13.7) |
| 10-day sequential therapy | 468 | 372 (79.4% [75.5 to 83.1]) | 4.9% (-0.9 to 10.8) |
|### Adherent to therapy (N=1314)###
| 14-day standard therapy | 347 | 372 (76.5% [72.5 to 80.2]) | 5.6% (-0.4 to 11.6) |
| 5-day concomitant therapy | 344 | 360 (73.6% [69.5 to 77.5]) | 8.6% (2.6 to 13.7) |
| 10-day sequential therapy | 348 | 355 (81.1% [77.1 to 84.6]) | 6.0% (0.3 to 11.8) |

Data are number (%) unless otherwise indicated. UBT=urea breath test.
1416 participants who returned for their follow-up visit had taken at least 80% of their assigned drugs, as assessed by pill count and self-report (table 2). Five participants had new, therapy-related symptoms (stomach discomfort, nausea or vomiting, or fatigue or weakness) that led them to seek medical attention; one in the standard treatment group and two each in the concomitant and sequential treatment groups. In intention-to-treat analyses, the estimated probability of Helicobacter pylori eradication was higher with 14-day standard triple therapy than with 5-day concomitant therapy and 10-day sequential therapy (table 3). The null hypothesis of inferiority of concomitant therapy to standard therapy could not be rejected (one-sided \( p=0.91 \)); thus, the data are consistent with 5-day concomitant therapy being inferior to 14-day standard therapy. The difference between 10-day sequential therapy and standard therapy was not significant (\( p=0.04 \)) at the \( p=0.025 \) level, but the near statistical significance of this two-sided test suggests that sequential therapy was not as effective as standard therapy. Sensitivity analyses that excluded participants without a definitive follow-up urea breath test and those that were confined to participants who had adhered to the prescribed regimens showed similar patterns of treatment contrasts as the intention-to-treat analysis, but with somewhat higher estimated eradication probabilities (table 3). Secondary analyses examining potential interactions showed no significant interactions between treatment and sex, age, presence of chronic dyspeptic symptoms at baseline, or study centre (table 4). Differences between study sites occurred in the overall probability of eradication in both the intention-to-treat populations (table 4) and the adherent-to-therapy population (webappendix), but nowhere was either four-drug regimen clearly better than standard triple therapy.

### Discussion

Our principal outcome measure, the probability of Helicobacter pylori eradication, was higher for 14-day standard triple therapy than for both four-drug regimens, and these results did not vary significantly by age, sex, study site, or history of chronic dyspeptic symptoms. The prevalence of Helicobacter pylori infection in screened participants was high, and nearly all individuals who had a positive urea breath test were randomly assigned into the trial. Participants tended to adhere closely to the protocol by returning for their follow-up urea breath test and taking their prescribed tablets. In the three treatment groups, the difference in adherence and serious side-effects was small.

Helicobacter pylori presents a major global health challenge, which can probably best be addressed through practical, inexpensive, and population-based interventions in the resource-limited countries where it is most prevalent. From this perspective the size of our trial, technical simplicity, broad geographic coverage within Latin America, community-based population, use of locally sourced generic drugs, and statistically robust findings across important subgroups reflect the realities of a region where diseases associated with Helicobacter pylori are especially burdensome. Our results are important because they challenge those of meta-analyses showing that four-drug regimens (triple therapy plus a nitroimidazole) given concomitantly or sequentially were clearly better than triple therapy, and they suggest that findings based primarily on data from Europe and other high-income regions might not be readily generalisable to lower-income countries (panel).

We reported probabilities of Helicobacter pylori eradication of less than 80% with 5-day concomitant and 10-day sequential regimens by the intention-to-treat analysis, whereas meta-analyses had reported probabilities greater than 90% for both. Investigators of a trial from Taiwan comparing 10-day regimens of concomitant versus sequential four-drug therapies also reported that both

<table>
<thead>
<tr>
<th>Study site</th>
<th>N</th>
<th>Helicobacter pylori eradication</th>
<th>Difference from standard group (adjusted 95% CI)</th>
<th>p value for interaction*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Santiago (Chile)</td>
<td>209</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
</tr>
<tr>
<td>14-day standard</td>
<td>69</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
</tr>
<tr>
<td>5-day concomitant</td>
<td>70</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
</tr>
<tr>
<td>10-day sequential</td>
<td>70</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
</tr>
<tr>
<td>Tíqueres (Colombia)</td>
<td>212</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
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<tr>
<td>14-day standard</td>
<td>71</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
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<tr>
<td>5-day concomitant</td>
<td>72</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
</tr>
<tr>
<td>10-day sequential</td>
<td>69</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
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<tr>
<td>Guanacaste (Costa Rica)</td>
<td>210</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
</tr>
<tr>
<td>14-day standard</td>
<td>70</td>
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<td>0.28</td>
</tr>
<tr>
<td>Copán (Honduras)</td>
<td>213</td>
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<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
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<tr>
<td>Tapachula (México)</td>
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<tr>
<td>Ciudad Obregón (México)</td>
<td>210</td>
<td>0.85 (5%)</td>
<td>-1.9% (-10.6 to 7.8)</td>
<td>0.28</td>
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<tr>
<td>León (Nicaragua)</td>
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Articles

regimens were more than 90% successful.26 The estimated effectiveness of our three-drug regimen (82%) in the intention-to-treat analysis was only modestly greater than that reported in the meta-analyses (77–79%).14–16 Thus, the divergence between our findings and those of the meta-analyses represents the substantially worse performance of the four-drug regimens.

Geographical variations in the pattern of H pylori resistance to antibiotics might account for some of the discrepancies between the results. In clinical series from several countries included in our trial, clarithromycin resistance in H pylori isolates has been reported to be less prevalent than in Europe, and metronidazole resistance substantially more prevalent (as high as 80%).21–24 Clarithromycin resistance strongly diminishes the effectiveness of triple therapy, so a better outcome with triple therapy would be expected if the prevalence of resistance in our trial population was actually low.13,21 A high prevalence of resistance to metronidazole in our study population is a plausible, but less certain, explanation for the worse-than-expected success of the four-drug regimens.25 Results according to antibiotic resistance are available from only two trials of sequential therapy versus triple therapy; the combined data showed that 68 of 71 (96%) patients with metronidazole-resistant organisms were treated successfully with sequential therapy compared with 46 of 59 (78%) treated with triple therapy.26,28 No comparable data from trials of concomitant versus triple therapy are available. The presence of organisms resistant to both antibiotics would likely cause treatment to fail for all three studied regimens, but data for this topic from Latin America are scarce.

We used a 14-day regimen of triple therapy in our trial, whereas previous trials of the four-drug regimens generally compared them to 7 days or 10 days of triple therapy. In one meta-analysis 14-day triple therapy regimens were slightly, but significantly, superior to those of 7 days or 10 days; thus, longer duration conceivably enhanced the performance of triple therapy in our trial.27 Success with the four-drug regimens perhaps would also improve with longer duration,13,20,28 however, this would increase their cost.

We recruited our participants from the general adult population in the community, whereas previous trials studied patients with gastrointestinal symptoms.14–16 The success of H pylori treatment with triple therapy or sequential therapy has not been shown to be affected by a diagnosis of peptic-ulcer disease or dyspepsia, and the superiority of triple therapy over the four-drug regimens in our trial did not vary significantly according to history of chronic dyspeptic symptoms.24

Our trial was undertaken in the general population in community settings where it was not feasible to mask the study or to obtain bacterial specimens for antibiotic sensitivity testing. We also obtained generic drugs from a variety of sources and did not have a completely objective way of determining adherence to therapy. Each of these factors is a potential limitation of the trial, but we doubt that any of them represents a serious threat to the validity of our results. Despite the absence of masking, our principal outcome measure (urea breath test) was objectively measured and seems unlikely to be biased. The absence of antibiotic sensitivity data also should not affect study validity, but it does make it more difficult to generalise our results to other regions where sensitivity patterns could be different. Inaccuracies in patient reports of adherence to therapy could have lowered our estimates of treatment effectiveness in the subgroup of participants classified as adherent, but these inaccuracies would not compromise the validity of the main results from our intention-to-treat analyses comparing the relative effectiveness of the three regimens. Lastly, although we cannot guarantee that the locally available generic drugs were of uniform efficacy, differences in treatment response between centres cannot be easily attributed to variable drug quality, since some of the greatest differences were seen between Nicaragua and Honduras, where investigators obtained drugs from the same source.

For individuals with H pylori infection in much of Latin America, 14 days of triple therapy is probably the preferred empiric treatment. Nevertheless, the 87% eradication

Panel: Research in context

Systematic review

Consensus groups that represent both global and Latin American perspectives have designated triple-drug regimens of a proton-pump inhibitor plus amoxicillin and clarithromycin taken for 7–14 days as a standard approach for eradicating Helicobacter pylori.10,11 However, the effectiveness of these regimens seems to have diminished to unacceptably low levels over time.12,13 and recent meta-analyses of clinical trials from Europe and Asia show that four-drug regimens that add metronidazole or tinidazole to triple therapy achieve superior results.14,15 The reported meta-analyses have not included any trials that were undertaken in Latin America, an area where H pylori-associated diseases are common and where wide-scale eradication programmes might be needed. We did a Medline search of all publications up to 2010, with no language restrictions, using the search terms “Helicobacter” in combination with the term for each country in Latin America. No publications of clinical trials that compared 14-day triple, 5-day concomitant, and 10-day sequential therapies for Helicobacter pylori infection were identified.

Interpretation

Our findings showed that in the Latin American populations we studied, by contrast with those for European and some Asian populations, 14-day standard triple therapy is more effective than 5-day concomitant or 10-day sequential four-drug regimens that include metronidazole for eradication of H pylori. Thus, effectiveness of H pylori eradication regimens in one area might not be as equally effective elsewhere.
success in participants who adhered to the regimen is
suboptimum in the clinical setting, and its effectiveness
might decrease over time as clarithromycin resistance
increases. Improved H. pylori treatment regimens for
Latin American populations could conceivably be
designed on the basis of local antibiotic resistance data.
However, the small number of published reports of
antibiotic resistance in this region generally pertain to
H. pylori isolates obtained from symptomatic patients
undergoing endoscopy in urban academic centres, and
thus the results might not be applicable to broader
populations.57 In view of the paucity of representative
data for resistance, and the daunting technical and
financial challenges of obtaining these data, treatment
guidelines for Latin America and other regions with
limited resources might have to rely primarily on the
results of large, simple clinical trials of empiric therapies
in the specific populations to which the guidelines
would apply. These data could be supplemented by subsequent
monitoring of effectiveness in practice over time and by
the results of antibiotic-resistance testing in selected
patients, when feasible.

We designed our study as a preliminary step towards
implementation of programmes of gastric cancer
prevention in Latin America. H. pylori-associated gastric
cancer results from a long progression from normal
mucosa to invasive cancer.3,29 The most promising
preventive approach seems to be eradication of H. pylori
infection before cancer develops, and several randomised
trials have assessed this strategy. The results show that
eradication of this infection slows or reverses progression
of premalignant histological lesions, but no trial has been
large enough to show a definitive cancer-preventive effect.30,31
Nevertheless, analyses have suggested that H. pylori eradication programmes would be cost effective
over the long term if they prevented only 10% of gastric-
cancer deaths; over the short term they would reduce
costs of care for peptic ulcers and dyspepsia symptoms.32–34
Eradication programmes are potentially even more cost
effective in regions such as Latin America, where the
burden of H. pylori-associated diseases is high.

Our results suggest that population-wide clinical trials
or public health programmes of H. pylori eradication are feasible in Latin America. Individuals with a positive
urea breath test readily agreed to be randomly assigned
to antibiotic treatment, and all three regimens of generic
drugs resulted in probabilities of eradication comparable
to those reported in previous prevention programmes.11
The 14-day triple-drug regimen had superior results, but
the lower cost of the shorter duration regimens might
make them acceptable for use in prevention programmes,
where resources are particularly scarce. Other consider-
ations, including risks of recrudescence and reinfection
after eradication, will also be important.

Contributors
All the authors participated in the design and oversight of the trial and
in the interpretation and reporting of results, and have seen and
approved the final report. LEB, RLD, CF, RH, MMM-M, RP, and ES-M
directed the clinical activities at the study centres. GLA had principal
responsibility for the statistical analyses of the data.

Conflicts of interest
DRM has submitted a patent application through the University of
North Carolina for a technique using molecular endoscopy to detect
cancer in the gastrointestinal tract, and has received funding from AstraZeneca
for his participation in a speakers’ bureau; he has also received a
research grant from AstraZeneca, for a proton-pump inhibitor study in
Hispanic populations in the USA, and from Given Imaging, for ongoing
efficacy studies of colon endocapsule efficacy. All other authors declare
that they have no conflicts of interest.

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Paris Heidt; Universidad del Valle, Cali, Colombia—Luz Stella García,
Yolanda Mora; Hospital Regional de Occidente, Santa Rosa de Copan,
Honduras—Jean Paul Higuero, Glenda Jeanette Euceda Wood,
Lesly Maritza Castellanos; Pontificia Universidad Católica de Chile,
Santiago, Chile—María Paz Cook, Paul Harris, Antonio Rollán; Fundación
INCIENSA, San José, Costa Rica—Silvia Jiménez, Paula González,
Ana Cecilia Rodríguez, Lidiana Morera, Blanca Cruz Reyes; Instituto
Nacional de Salud Pública, Cuernavaca, Mexico—Rogerio Danis,
Erika Marlen Hurtado Salgado, María del Pilar Hernández Neváres;
Instituto Tecnológico de Sonora, Ciudad Obregón, Mexico—Myriam Bringas,
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Yesenia Zapata; and also Charles A Colman, David S Alberts, and
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Mass eradication of *Helicobacter pylori*: feasible and advisable? 

In *The Lancet*, E Robert Greenberg and colleagues present an Article about the efficacy of three different therapeutic regimens for the eradication of *Helicobacter pylori* in several Latin American countries. A large sample of adults with symptomatic and asymptomatic disease was studied. Besides aiming to identify the best therapeutic regimen, the study also served as a preliminary step to support future programmes of gastric cancer prevention in the Latin American population. Gastric cancer is the fifth most common cause of cancer in this region (figure) and is the leading cause of cancer mortality among males in many Latin American countries. The study showed that the 14-day standard therapy with lansoprazole, amoxicillin, and clarithromycin was superior to both the 5-day concomitant and sequential four-drug regimens (lansoprazole, amoxicillin, clarithromycin, and metronidazole).

Generic (off-patent) preparations of the study drugs were used in this trial. The use of low-cost generic drugs is an important factor for public health strategies in Latin America. These preparations were validated in the USA and the results from the study done by Greenberg and colleagues show that good quality generic drugs were used. However, if these drugs are used in large populations for *H pylori* eradication, quality control of these drugs will be crucial. Recent reports have brought attention to problems related to the quality of generic drugs, including antibiotics.

Importantly, Greenberg and colleagues’ study confirms that the standard therapy, which includes generic drugs, remains effective in Latin America, by contrast with developed countries where resistance to clarithromycin has been identified. We believe however, that costs could be reduced and the potential for development of resistant bacterial strains could be decreased, if the standard therapy could be prescribed over a short period of time. Findings from two studies done 10 years apart in Brazil using standard therapy over a 10-day period showed *H pylori* eradication rates of about 90%.

*H pylori* affects around 50% of men, with prevalence much higher in developing countries than in developed countries (ie, prevalence in North America is 30%, in Central America 70–90%, and in South America 70–90%). In 1994, the International Agency for Research on Cancer defined *H pylori* as a human carcinogen for gastric adenocarcinoma, raising concerns about how to treat people with asymptomatic infection.

A disturbing issue presented by Greenberg and colleagues is the suggestion of developing a mass eradication programme for *H pylori* in Latin America, aimed at long-term prevention of at least 10% of gastric cancer cases, and short-term gain of saving money by reducing care costs associated with peptic ulcer and dyspepsia symptoms.

Although the role of *H pylori* in gastric carcinogenesis is well defined, no definitive evidence shows that mass eradication could reduce incidence of gastric cancer. Wong and colleagues showed no benefit in the prevention of gastric cancer with the eradication of *H pylori*. By contrast, a recent meta-analysis suggested that eradication could indeed reduce the risk of gastric cancer.

Policies aimed at population-wide *H pylori* eradication could have individual and social repercussions. Amoxicillin could cause fatal anaphylactic reactions, while clarithromycin has been associated with increased mortality in patients with ischaemic heart disease. Furthermore, almost all antibiotics can cause *Clostridium difficile* infection, which could eventually become life threatening. Even if infrequent, these complications could become important when eradicating *H pylori* at a population level. Another

### Figure: Mortality of gastric cancer in men and women in Latin America (2008)

(A) Number of deaths in men per 100 000 people-12.8. (B) Number of deaths in women per 100 000 people-6.9. Modified from GLOBOCAN 2008, International Agency for Research on Cancer.
key point to address is the probability of re-infection. Annual recurrence of \textit{H pylori} infection is about 12% in developing countries.\textsuperscript{24} However, the most serious issue relates to development of bacterial resistance for not only \textit{H pylori} but potentially many other bacteria. Thus, we believe that although any public policy aimed at decreasing the prevalence of \textit{H pylori} infection could lead to reductions in gastric cancer and peptic ulcer, the risks associated with antibiotic use might represent an important limitation to such a strategy. Furthermore, restricted financial resources in Latin America represent a major public health challenge. Therefore, appropriate studies are needed that compare the effect of investment in sanitation, hygiene, and education with the effect of mass eradication of \textit{H pylori}, to assess which options will be most beneficial to the Latin American population. Another possible approach is the development of effective immunisation as a safer method of decreasing rate of \textit{H pylori} infection.\textsuperscript{55} On the one hand, we are aware that sanitation and immunisation will not reduce the risk of \textit{H pylori}-associated disease for people who are already infected. On the other hand, doubts exist about the reversibility of already established precursor lesions in the carcinogenesis cascade in infected patients. Wong and colleagues\textsuperscript{22} showed that eradication significantly decreased the development of gastric cancer only in the subgroup of \textit{H pylori} carriers without precancerous lesions.

Mass eradication is a major issue requiring further investigation. We agree with Greenberg and colleagues that future properly designed studies addressing this issue are essential to assess the balance between positive and negative effects of each strategy. Local epidemiological data, among other factors, will dictate allocation of public health resources: countries with high mortality rates secondary to gastric cancer, such as Colombia, Chile, and Guatemala, should discuss whether development of a programme for mass eradication of \textit{H pylori} is worthwhile.

In conclusion, mass eradication of \textit{H pylori} is potentially feasible but in view of the differing socioeconomic realities of Latin American countries, doubts remain about the advisability of such a policy.

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