Increased Risk of Spontaneous Bacterial Peritonitis in Cirrhotic Patients Using Proton Pump Inhibitors

Abdel-Naser Elzouki\textsuperscript{a,b}  Nadia Neffati\textsuperscript{a}  Fatma A. Rasoul\textsuperscript{a}  Ali Abdallah\textsuperscript{a}  Muftah Othman\textsuperscript{a}  Abdelkarim Waness\textsuperscript{a}

\textsuperscript{a}Department of Medicine, Hamad General Hospital, Hamad Medical Corporation, Doha, Qatar; \textsuperscript{b}Weill Cornell Medical College, Doha, Qatar

Abstract

**Background:** The association between bacterial infections and proton pump inhibitors (PPIs) has recently been studied with debatable results. **Aim:** The aim of this study was to investigate the relationship between PPIs and the development of spontaneous bacterial peritonitis (SBP) or other bacterial infections in cirrhotic patients. **Materials and Methods:** Consecutive cirrhotic patients hospitalized from 2007 through 2012 to Hamad General Hospital, Doha, Qatar, were enrolled and classified as PPI users or non-users according to PPI consumption in the 90 days prior to hospitalization. Cirrhosis was clinically diagnosed by a combination of physical, biochemical, radiological, and endoscopic findings, or by liver biopsy. **Results:** A total of 333 patients were included in this study, of whom 171 (51.4\%) used PPIs and 162 (48.6\%) did not use PPIs. PPI users were significantly older in age ($p = 0.001$). There was no statistical difference between the 2 groups in sex distribution and etiology of cirrhosis ($p > 0.05$ for both parameters). PPI users had a significantly higher incidence of overall bacterial infection (38\%) than non-PPI users (13.6\%), $p = 0.0001$. Statistical significance is observed specifically for SBP and chest infection ($p = 0.0006$ and $p = 0.01$, respectively). In multivariate analysis, older age ($>60$ years; OR = 1.246, 95\% CI 1.021–0.8486; $p = 0.02$), and PPI use (OR = 2.149, 95\% CI 1.124–0.6188; $p = 0.01$) were independent predicting factors for SBP and overall bacterial infection. **Conclusion:** The present study shows that PPI use, as well as older age ($>60$ years), was an independent predicting factor for the development of overall infection and SBP in hospitalized cirrhotic patients. Unless it is indicated, PPI therapy should be avoided in this group of patients, particularly in those older than 60 years of age.
Resumo

Introdução: A associação entre infecções bacterianas e os inibidores da bomba de prótons (IBPs) tem vindo a ser estudada com resultados discutíveis. **Objetivo:** O objetivo deste estudo foi investigar a relação entre IBPs e o desenvolvimento de peritonite bacteriana espontânea (PBE) ou outras infecções bacterianas em doentes cirróticos. **Material e Métodos:** Doentes consecutivos com cirrose hospitalizados entre 2007 e 2012 no Hamad General Hospital-Qatar foram selecionados e classificados como utilizadores ou não utilizadores de IBPs de acordo com o seu consumo nos 90 dias prévios ao internamento. A cirrose foi clinicamente diagnosticada por uma combinação de achados no exame físico, no estudo bioquímico, radiológico e endoscópico; ou por biopsia hepática. **Resultados:** Um total de 333 doentes foi incluído neste estudo, 171 (51.4%) medicados com IBPs e 162 não (48.6%). Os utilizadores de IBPs eram significativamente mais velhos (p = 0.001). Não se observaram diferenças estatísticas entre os dois grupos no que se refere ao sexo ou etiologia da cirrose (p > 0.05 para os dois parâmetros). A incidência global de infecções bacterianas foi significativamente superior nos utilizadores de IBPs (38%) do que nos não utilizadores (13.6%), p = 0.0001. O significado estatístico desta diferença foi observado especificamente para a PBE e para as infecções pulmonares (p = 0.0006 e p = 0.01, respectivamente). Na análise multivariada, a idade superior a 60 anos (OR = 1.246, 95% CI 1.021–0.8486; p = 0.02), e a utilização de IBPs (OR = 2.149, 95% CI 1.124–0.6188; p = 0.01) foram fatores preditivos independentes para PBE e para infecção bacteriana no global. **Conclusão:** Este estudo mostra que a utilização de IBPs, assim como a idade superior a 60 anos, são fatores preditivos independentes para o desenvolvimento de infecções bacterianas no global e para PBE nos doentes cirróticos hospitalizados. A não ser que esteja especificamente indicado a utilização de IBPs deve ser evitada neste grupo de doentes, particularmente naqueles com idade superior a 60 anos.

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Introduction

Since the introduction of proton pump inhibitors (PPIs) to the biopharmaceutical market in the late 1980s, this class of drugs has had a tremendous impact on patient care, clinical practice, and research development. Compared to other antacid drugs, PPIs were initially thought to have an excellent safety profile, resulting in their widespread use in both inpatient and ambulatory settings [1, 2]. After more than 25 years of clinical post-marketing experience, improved reporting, and closer scrutiny, several serious adverse effects associated with the long-term use of PPIs have emerged. These include increased risks of falls and fractures in postmenopausal women [3], a reduction of renal and liver function, and an increased risk of community-acquired and nosocomial pneumonia [4, 5]. PPIs are also found to be an independent risk factor for Clostridium difficile infection even with a very short-term use of 2 days in the intensive care setting [6].

The possible association between PPI use and increased risk of spontaneous bacterial peritonitis (SBP) and overall bacterial infection in cirrhotic patients is a debatable subject in the medical literature. While some authors point to a straight causative effect, others are not so sure. Furthermore, to our knowledge, this issue has never been investigated in the State of Qatar or in any other Arab country. The aim of this study is to evaluate the possible relationship between the use of PPIs and the development of bacterial infections in cirrhotic patients and whether PPI use increases the risk of SBP development in such patients at the main tertiary care center in Qatar.

Material and Methods

Study Population and Setting

Consecutive cirrhotic patients >18 years of age hospitalized from 2007 through 2012 to Hamad General Hospital were enrolled in this study. Hamad General Hospital is a 620-bed tertiary center that covers all specialties and is affiliated to Hamad Medical Corporation in Qatar. Data were collected retrospectively from the medical records using a data collection form including age, gender, etiology of liver cirrhosis, presence of other comorbidities, and the cause of hospital admission. Other characteristics of cirrhotic patients who were admitted to the medical wards and/or medical intensive care unit of the hospital included acute cirrhotic complications such as hepatic encephalopathy, variceal bleeding, SBP or other infections, refractory ascites, hepatorenal syndrome, and first diagnosis of hepatocellular carcinoma. Cirrhotic patients with active gastrointestinal bleeding, disseminated malignancies, undergoing immunosuppressive therapy, or using antibiotics in the previous 2 weeks prior to hospitalization were excluded from this study.

Acid-Suppressive Medication Exposure

To determine drug exposure, medical records were reviewed for PPI use in the 90 days prior to hospitalization, and each subject was classified as PPI user versus non-PPI user. Exposure to acid-suppressive medication was defined as any order for a prescription of PPI or histamine-2 receptor antagonists (H2RAs). The PPIs included were either pure or compound medication containing omeprazole, lansoprazole, esomeprazole, pantoprazole, or rabe-
prazole. H2RAs included any drugs containing ranitidine, cimetidine, or famotidine. A cumulative treatment period for ≥7 days was required for inclusion. Unexposed status was defined as no prescription or cumulative treatment period of <7 days in the last 90 days prior to hospital admission.

**Case Diagnosis and Definitions**

The diagnosis of liver cirrhosis was established by a combination of physical, biochemical, radiological and endoscopic findings, or by liver biopsy. The decision of liver biopsy was based on the clinical needs as well as the patient’s choice and consent. Infection was defined according to standard criteria [7, 8]: (1) SBP: ascitic fluid polymorphonuclear cells more than 250 cells/mm³; (2) spontaneous bacteremia: positive blood cultures without a known source; (3) chest infection: new pulmonary infiltrate in the presence of (a) at least 1 respiratory symptom (cough, sputum production, dyspnea, pleuritic chest pain) with (b) at least 1 finding on auscultation (rales or crepitation) or signs of infection (i.e., with at least 2 of the following: (a) a core body temperature >38 °C or <36 °C, (b) tachycardia with a heart rate above 90 beats per minute at rest, a respiratory rate of more than 20 breaths per minute, and/or shivering or a leukocyte count >10,000/mm³ or <4,000/mm³) [9] in the absence of antibiotics; and (4) urinary tract infection: urine white blood cell >15/high power field with either a positive urine gram stain or culture.

**Statistical Analysis**

Analysis used were χ² for categorical variables expressed as number (percent) and the independent t test for continuous variables (e.g., age) expressed as the mean ± standard deviation. Multivariate logistic regression analysis was used to produce the prediction model for infections in cirrhotic patients.

**Results**

**Study Population and Use of PPIs**

A total of 333 cirrhotic patients were included in this study, and 78.1% of them were male. Hepatitis C infection (35.7%), alcohol abuse (20.1%), and hepatitis B infection (16.5%) were the main etiologies of cirrhosis. The presence of SBP was detected in 61 (18.3%) patients, and the severity of liver disease according to the Child-Pugh score was significantly associated with the risk of SBP development (HR = 1.9, 95% CI: 1.21–3.80, p = 0.01 in Child-Pugh B patients and HR = 4.10, 95% CI: 1.87–7.86, p = 0.001 in Child-Pugh C patients). There were 171 (51.4%) patients using PPIs and 162 (48.6%) not using PPIs. Only 4 patients in the PPI-user group had a history of a prior consumption of H2RAs. Specific indication for PPI use was not documented in 143 (43%) of our patients. Table 1 shows the comparison of demographic and clinical data between the 2 groups. PPI users were significantly older

<table>
<thead>
<tr>
<th>Independent variables</th>
<th>PPI users (n = 171)</th>
<th>Non-PPI users (n = 162)</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Demographics</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean age ± SD, years</td>
<td>55.3±11.7</td>
<td>50.2±11.8</td>
<td>0.0001</td>
</tr>
<tr>
<td>Males</td>
<td>126 (73.7)</td>
<td>134 (82.7)</td>
<td>0.46</td>
</tr>
<tr>
<td><strong>Comorbidities</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Smoker</td>
<td>55 (32.2)</td>
<td>48 (29.6)</td>
<td>0.617</td>
</tr>
<tr>
<td>Diabetes mellitus</td>
<td>94 (55)</td>
<td>59 (36.4)</td>
<td>0.001</td>
</tr>
<tr>
<td>Hypertension</td>
<td>66 (38.6)</td>
<td>35 (21.6)</td>
<td>0.001</td>
</tr>
<tr>
<td>Chronic kidney disease</td>
<td>9 (5.3)</td>
<td>3 (1.9)</td>
<td>0.097</td>
</tr>
<tr>
<td><strong>Etiology of cirrhosis</strong></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Chronic hepatitis B</td>
<td>27 (15.8)</td>
<td>28 (17.3)</td>
<td>0.714</td>
</tr>
<tr>
<td>Chronic hepatitis C</td>
<td>64 (37.4)</td>
<td>55 (34)</td>
<td>0.508</td>
</tr>
<tr>
<td>Alcoholic liver disease</td>
<td>45 (26.3)</td>
<td>49 (30.2)</td>
<td>0.426</td>
</tr>
<tr>
<td>Autoimmune hepatitis</td>
<td>5 (2.9)</td>
<td>3 (1.9)</td>
<td>0.523</td>
</tr>
<tr>
<td>Primary biliary cirrhosis</td>
<td>3 (1.8)</td>
<td>3 (1.9)</td>
<td>0.947</td>
</tr>
<tr>
<td>Cardiac cirrhosis</td>
<td>1 (0.6)</td>
<td>0 (0)</td>
<td>0.330</td>
</tr>
<tr>
<td>Cryptogenic cirrhosis</td>
<td>38 (22.2)</td>
<td>29 (17.9)</td>
<td>0.326</td>
</tr>
<tr>
<td>Other causes</td>
<td>5 (2.9)</td>
<td>7 (4.3)</td>
<td>0.494</td>
</tr>
<tr>
<td><strong>Child-Pugh score</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>A</td>
<td>64 (37.4)</td>
<td>71 (43.8)</td>
<td>0.194</td>
</tr>
<tr>
<td>B</td>
<td>65 (38.0)</td>
<td>56 (34.6)</td>
<td>0.514</td>
</tr>
<tr>
<td>C</td>
<td>42 (24.6)</td>
<td>35 (21.6)</td>
<td>0.522</td>
</tr>
</tbody>
</table>

Values are expressed as n (%), unless otherwise indicated. PPI, proton pump inhibitor; SD, standard deviation.
in age \( p = 0.0001 \), and there was no statistical difference between the 2 groups in gender distribution, etiology of cirrhosis, and Child-Pugh score \( p > 0.05 \) for all parameters. As shown in Table 2, PPI users had a significantly higher overall bacterial infection rate (38%) than non-PPI users (13.6%; \( p = 0.0001 \)). For SBP, the rate in PPI users was also higher (32.8%) than in non-PPI users (6.9%; \( p = 0.0014 \)) (Table 3).

**Multivariate Analysis**

On the multivariate analysis, age >60 years (OR = 1.246, 95% CI 1.021–0.848; \( p = 0.02 \)) and PPI use (OR = 2.149, 95% CI 1.124–6.188; \( p = 0.012 \)) were independent predicting factors for SBP and overall bacterial infection (Table 4).

**Discussion**

The present study has shown that the use of PPIs in patients with liver cirrhosis is an independent risk factor for overall infection and SBP development, hence support an association between PPI use and SBP. PPIs have been widely used worldwide since their introduction around 3 decades ago. A growing list of their adverse effects is being compiled. In an early study, acid-activated omeprazole was documented to inhibit in vitro human neutrophil phagocytosis and phagolysosome acidification [10].
Another small study (blood samples from 10 healthy subjects) documented the impairment of reactive oxygen intermediates by human neutrophils after omeprazole administration. This finding possibly alludes to the bacterial activity of reduced neutrophils [11].

Patients with liver cirrhosis usually suffer from a poor synthetic function with a decreasing albumin production. There is a growing body of evidence that they suffer from a complex and not fully understood cirrhosis-associated immune dysfunction syndrome [12]. The evolving explanation for this impaired immunity includes: a decrease in T-helper cells and phagocytic potential of both monocytes and neutrophils [13], an increased level of cyclooxygenase-derived eicosanoid prostaglandin E2 [14], and a decreased HLA-DR expression on monocytic cells defined as immune paralysis [15].

The use of PPIs in patients with cirrhosis may further degrade their already compromised immune system. Garcia-Martinez and coworkers have recently demonstrated that PPIs significantly decrease granulocyte and monocyte cellular oxidative burst in patients afflicted with cirrhosis; a pathogenic finding that possibly explains the reported high rates of bacterial infections in these patients [16]. The most common infection is SBP [17]. The mechanism of development of SBP in cirrhotic patients is not fully explicated; however, increased intestinal permeability, altered intestinal motility, and increased small intestinal bacterial overgrowth may play an important role in facilitating bacterial translocation [18, 19]. PPIs have been identified as an important factor for increased small intestinal bacterial overgrowth [20].

In the clinical setting, the role of PPIs in the pathogenesis of SBP in cirrhotic patients has been hotly debated for the last decade. When it comes to PPI use and SBP association, we can clearly identify 2 opposite groups. The first one is made of authors who concluded that there is no causative effect. An example is the 2008 case-control study by Campbell et al. [21] that included 116 consecutive cirrhotic patients. A second retrospective study by Mandorfer et al. [22] that enrolled a larger number of subjects (607 consecutive patients) reported no association between PPI use and SBP or other infections or mortality. Another large prospective study involved 770 decompensated cirrhotic consecutive patients in 23 hospitals in Argentina and failed to demonstrate an association between PPI therapy and an increased risk of SBP [23]. In a more recent retrospective cohort study that enrolled 307 cirrhotic patients with a previous SBP in Korea, the incidence of second SBP did not statistically differ between PPI users and non-PPI users who were followed up for 5 years [24]. Furthermore, a more recent cohort study conducted in southern Brazil that enrolled 258 cirrhotic patients with ascites did not confirm the high risk of SBP development in PPI users compared to PPI-non users [25].

The second group of authors claims the opposite view. They advocate a direct correlation between PPI use and the increased risk of developing SBP. Their number has been increasing recently, and their evidence is growing stronger. We identified many studies in this respect, including 3 meta-analyses. The first one, conducted in 2011, included 4 studies involving 772 patients. It found a significant association between PPI use and the development of SBP (OR 2.77, 95% CI 1.82–4.23) [26]. In 2013, a second meta-analysis that reviewed 8 studies was conducted. It examined the association between acid-suppressive therapy (PPI and H2RAs) in cirrhotic patients and the risk of developing SBP. This risk was greater in subjects on PPI treatment (n = 3,815; OR 3.15, 95% CI 2.09–4.74) compared to those taking H2RAs (n = 562; OR 1.71, 95% CI 0.97–3.01). Authors concluded that patients with cirrhosis receiving PPIs have approximately 3 times the risk of developing SBP compared with those not taking these medications [27]. More recently, in 2015, a third meta-analysis reviewed 17 studies published between 2008 and 2014 (12 journal articles and 5 conference abstracts) involving 8,204 patients. These studies were conducted with North American (8 studies), European (4 studies), South East Asian (4 studies), and South American (1 study) populations; none of them examined the Arabic population. The result showed that PPI use in cirrhotic patients was significantly associated with an increased risk of SBP (OR 2.17, 95% CI 1.46–3.23) and an overall risk of bacterial infection (OR 1.98, 95% CI 1.36–2.87) [28]. Many other investigations in the medical literature confirm this association. A large Korean study (1,140 patients) by Kwon et al. [29] confirmed that PPI use (within 30 days) in cirrhotic patients having ascites increased their risk of developing SBP, especially in the elderly and in patients with a high model for end-stage liver disease score on admission. Further, the mortality risk was also found to be higher in this group of patients. Another recent Taiwanese case-control study published in 2015 identified a total of 947 patients among 86,418 patients with advanced liver cirrhosis using acid suppression. It further confirmed the risk of developing SBP with higher cumulative days of gastric acid suppression being associated with a higher risk of infection (p < 0.0001) [19]. In a more recent study, the use of PPIs has been found to be a risk factor for developing hepatic...
en cephalapathy and SBP in cirrhotic patients with ascites [30].

The present study included 333 cirrhotic patients between 2007 and 2012. It further solidified the argument that PPI use and older age (>60 years) are independent risk factors for the development of SBP in these patients. We side with all prior authors that further trials (especially prospective comparative ones) will probably shed more light on all potential adverse effects of PPIs in patients with liver cirrhosis. We also recommend that adequate precautions are taken to utilize such treatment in cirrhotic patients only when benefit may outweigh potential harm.

In conclusion, the evidence that PPIs pose health risks in certain patient populations is expanding. Cirrhotic patients receiving this form of therapy seem to have a higher risk of developing SBP. The present study shows that PPI use, as well as older age (>60 years), was an independent predicting factor for the development of bacterial infection including SBP in hospitalized cirrhotic patients. Further studies are needed to settle the current debate between supporters and opponents of these conclusions. We recommend that unless it is clearly indicated, PPI therapy should be avoided in patients using PPIs, in particular those who are older than 60 years of age.

**Statement of Ethics**

The ethical approval for this study was obtained from the Institutional Research Review Board at Hamad Medical Corporation, Qatar (research approval No. 13246/2013).

**Disclosure Statement**

The authors have no conflicts of interest to declare.

**References**


