Proton pump inhibitors in rheumatic diseases: clinical practice, drug interactions, bone fractures and risk of infections

Inhibitori di pompa protonica in reumatologia: pratica clinica, interazioni farmacologiche, fratture e rischio infettivo

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INTRODUCTION

Patients affected by acute coronary syndrome (ACS) or by chronic inflammatory musculoskeletal and connective tissue diseases (i.e. systemic sclerosis), often need antiaggregant therapy (ASA or clopidogrel).

The concomitant use of proton pump inhibitors (PPIs) is suggested to reduce the risk of haemorrhage. Clopidogrel is a prodrug activated by cytochrome P 450. PPIs too have a CYP P450 metabolism, and a drug interaction has been observed between PPIs and clopidogrel. 25% of non-

responsiveness to clopidogrel is due to this drug interaction (1). Some studies have demonstrated that the use of PPIs is associated with an increased risk of bone fractures and Clostridium difficile infection.

CLOPIDOGREL PHARMACOGENOMICS AND RISK OF INEFFECTIVE ANTI-AGGREGATION

Many clinical trials have demonstrated that the administration of clopidogrel together with aspirin (ASA) versus aspirin alone re-
roduces the risk of death or of recurrent acute coronary syndrome.

In particular, clopidogrel has proven beneficial in patients undergoing percutaneous coronary intervention (PCI), such that patients receiving bare metal stents and drug-eluting stents are currently recommended to receive clopidogrel maintenance therapy for 1-2 months and at least 12 months, respectively (2).

Nevertheless, “clopidogrel non-responsiveness” occurs in approximately 25% of the population and it is associated with an increased risk of recurrent events (2); therefore the pharmacogenomic and pharmacodynamic knowledge of clopidogrel and the relative drug interactions became very important and the object of many studies.

Clopidogrel belongs to the thieno-pyridine class of chemical compounds. It is a prodrug and it is converted to its active metabolite by cytochrome P450 (CYP) enzymes. Clopidogrel is absorbed in the duodenum and approximately 85% of this dose is hydrolyzed by esterases to an inactive carboxylic acid derivative. The remaining 15% is metabolized by CYP into an active intermediate, after an oxidation of the thioephene ring.

After being released into the systemic circulation, the active metabolite’s thiol group forms an irreversible disulfide bond with the platelet ADP P2Y12 receptor (P2RY12) inhibiting ADP-mediated platelet activation.

There are many isoforms of CYP P450, but the most important for PPI metabolism is the CYP2C19. This enzyme is responsible for the metabolism and clearance of many other drugs, such as cyclophosphamide, diazepam and PPI.

The CYP2C19 gene is on chromosome 10 and it contains 9 exons and it is highly polymorphic; 25 CYP2C19 variant alleles have been identified (2).

It is well established that there is an efflux pump (P-gp, ABCB1) on the bowel mucous membranes that represents a barrier to clopidogrel absorption.

Many clinical and genomic factors are responsible for clopidogrel non-responsiveness (Table I).

Genetic variations in other CYP isoforms (different from CYP2C19) that contribute to metabolic conversion of clopidogrel to its active metabolite in vitro have not consistently proven to be associated with clopidogrel pharmacokinetics and pharmacodynamics in humans.

CYP2C19*2 variant allele carriers are at significantly higher risk of adverse cardiovascular events compared with wild-type individuals, including death, myocardial infarction and stent thrombosis.

Polymorphism ABCB1 3435 C>T seems to be related to a bigger outflow of the drug and then to a smaller intestinal absorption. Polymorphism of the receptor P2Y12 (P2RY2744C>T) also seems to be associated with a smaller response to anti-aggregant therapy.

Diabetic and overweight individuals (BMI>=25 kg/m²) are significantly more likely to exhibit impaired inhibition of platelet activation, and are at higher risk of morbidity and mortality.

It is known that diabetes is associated with a pro-inflammatory and pro-thrombotic state, which may be partially mediated by hyperglycemia-induced up-regulation of P2Y12 receptor expression and increased oxidative stress. Moreover, diabetes seems to interfere with the pharmacokinetics of clopidogrel (2).

### PPI AND DRUG INTERACTION

PPI can have different mechanisms through which they interfere with many drugs:

1) An increase in gastric pH causes a bigger absorption of weak acids (digoxin,
furosemide, aspirin) and a smaller absorption of weak bases (ketoconazole).
2) Interference with kidney excretion.
3) Interference with CYP P450 system.
PPI and clopidogrel interaction is still an object of discussion. It is not very clear, moreover, if the clinical meaning of their interaction is more pronounced in some subsets of population (eg: specific CY-
P2C19 allelic variant carriers, diabetics or those overweight (2).
Different studies show the existence of drug interaction between PPI and clopi-
dogrel (3-6).
In the OCLA study of Martine Gilard et al, a double-blind placebo-controlled trial, all consecutive patients undergoing coronary artery stent implantation received aspirin (75 mg/day) and clopidogrel (loading dose, followed by 75 mg/day) and were randomized to receive either associated omeprazole (20 mg/day) or placebo for 7 days (3).
Clopidogrel effect was tested on days 1 and 7 in both groups by measuring plate-
let phosphorylated -VASP expressed as a platelet reactivity index (PRI). The main end point compared PRI value at the 7-day treatment period in the 2 groups. Data for 124 patients were analyzed. On day 1, mean PRI was 83.2% and 83.9%, respectively, in the placebo and omeprazole groups (p=NS), and on day 7, 39.8% and 51.4%, respectively (p<0.0001) (7).
In a case–control study conducted by Da-
vid N. Juurlink et al, among 13,636 patients
prescribed clopidogrel following acute
myocardial infarction, 734 cases were re-
admitted with myocardial infarction. After extensive multivariables adjustment, cur-
rent use of PPI was associated with an in-
creased risk of re-infarction (adjusted odds ratio [OR] 1.27, 95% confidence interval [CI] 1.03-1.57).
In a stratified analysis, pantoprazole, which does not inhibit CYP P450 2C19, had no association with readmission for myocardial infarction (adjusted OR 1.02, 95% CI 0.70-1.47) (8). However, a meta-
alysis has been published on PPI effect on cardiovascular events in patients who take clopidogrel (9). 23 studies have been found on Medline, Embase, Cochrane in October 2009. No significant increase in mortality has been found in patients who take PPI together with clopidogrel.

### HOW TO PREVENT COMPETITIVE INHIBITION OF CYP2C19 BY PPI
Although each single PPI has similar ef-
ficacy in many cases, differences between them should be considered when choosing a treatment regimen. Our literature analy-
sis shows that different PPIs are not all the same. Juurlink et al show that pantoprazole, which does not inhibit CYP P450 2C19, had no drug interaction with clopi-
dogrel (8).
These data suggest, where needed, the pre-
scription of pantoprazole.
As PPIs and clopidogrel are each admin-
istered once daily and their presence in the bloodstream is short-lived, separating them by 12-15 h should in theory prevent any competitive inhibition of CYP metabol-
olism and any clinical effect. In addition, PPIs are most effective when taken before meals. Therefore, it is suggested that PPIs be given before breakfast and clopidogrel at bedtime, or, to minimize concern about poor CYP2C19 metabolizers, PPIs may be taken before dinner and clopidogrel at lunchtime (1).

### PPI AND BONE FRACTURES
Patients affected by chronic systemic in-
flamatory or connective tissue diseases i.e. systemic sclerosis, have high frequency of oesophageal mucosal abnormalities and should take long-term PPI therapy.
It has been suggested that an acidic envi-
ronment in the stomach and upper small bowel is required to free ingested calcium from the food matrix making it available for absorption. Impaired calcium absorp-
tion would lead to compensatory physi-
ologic responses, including secondary hy-
perparathyroidism.
Secondary hyperparathyroidism refers to
the increase in circulating levels of parathyroid hormone when serum levels of ionized calcium drop (as would occur with reduced efficiency in intestinal calcium absorption). Over time, parathyroid hormone would lead to an increase in the rate of skeletal turnover and a reduction in bone mass, both of which increase the risk of fracture (10).

If normal gastric acid production is required for calcium ionization and subsequent absorption, then the millions of individuals using PPIs may be at increased risk for calcium malabsorption, negative calcium balance, and potential bone loss. Unfortunately, there are no long-term studies on the effects of PPIs on calcium absorption (11).

Today, we have only short-term studies examining the effect of gastric acid suppression on calcium absorption and bone turnover (12-16).

A MEDLINE search was conducted to identify relevant articles regarding PPIs and fractures (10).

Three case-control studies assessed fractures and PPI use. A study of all subjects with fracture in Denmark in 2000 revealed adjusted OR=1.18 (1.12-1.43) for PPI use within the last year (hip fracture OR=1.45, 1.28-1.65); no dose-response relationship was identified (17).

A study of hip fractures in UK patients ≥50 years found adjusted OR=1.44 (1.30-1.59) for >1 year of PPIs; duration and average daily dose were significantly associated with fracture risk: adjusted OR for >1.75 times average daily dose for >1 year was 2.65 (1.80-3.90) (18).

A study of vertebral, wrist, and hip fractures in Manitoba patients ≥50 years found significant ≥7 years (OR=1.92, 1.16-3.18) (19) risk factor.

### PPI AND RISK OF CLOSTRIDIUM DIFFICILE INFECTION (CDI)

Clostridium difficile (CD) is a Gram-positive, spore-forming, anaerobic bacillus. The PPI’s use increases the risk of CDI. In a case-control study performed on hospitalized patients, PPI use was associated with C. difficile diarrhoea (OR 2.7, 95% confidence interval (CI) 1.4–5.2) and with relapse (OR 5.2, 95% CI 1.1–24.6) (20).

Another study showed that hospitalized patients using PPIs were 4.2-fold more likely to have recurrent disease. This risk was not observed in patients using H2 blockers (21).

Probably, the chronic use of PPIs, reducing gastric acid environment, reduces a natural defense of our organism.

Also in the community PPI use was associated with an increase in CDI (OR 2.9, 95% CI 2.4-3.4) (22).

Recently a retrospective cohort study has been published. It shows a risk of recurrent CDI higher in patients taking PPIs versus patients not exposed to this class of drugs (25.2% vs 18.5%) with a 42% increased risk of recurrence (23).

CDI has been demonstrated to be rarely associated with reactive arthritis, a condition that may develop in response to a gastrointestinal infections usually due to Salmonella, Shigella, Yersinia or Campylobacter (24).

Recently 46 cases in adults have been reported. The sex ratio was close to 1 and the patients were older than those with other causes of reactive arthritis (25).

### CONCLUSIONS

In several long term therapeutic programs PPI should be adopted as a protection to the stomach or to avoid gastroesophageal reflux disease (GERD).

It appears very clear that the choice of PPI in several cases should be made on a carefully assessed risk/benefit ratio assessment.

In particular a thoughtful analysis should be performed in all patients needing antiplatelet therapy and in all patients with osteoporosis, either primary or secondary.
Platelet activation and aggregation are key elements of the pathogenesis of acute coronary syndromes, of endothelial damage in chronic inflammatory and connective tissue disease (i.e. systemic sclerosis-SSc). Patients affected by chronic inflammatory diseases as well as by connective tissue diseases such as systemic sclerosis, often have the need to take anti-platelet therapy (e.g. ASA or clopidogrel). Current consensus recommendations state that patients prescribed clopidogrel plus aspirin should receive a proton pump inhibitor (PPI) to reduce gastrointestinal bleeding. Although each single PPI has similar efficacy in many cases, differences between them should be considered when choosing a treatment regimen. Many studies show PPI and clopidogrel drug interaction, with clopidogrel non-responsiveness in about 25% of the population. Only pantoprazole, which does not inhibit CYP P450 2C19, doesn’t seem to have interaction with clopidogrel or other drugs. Patients affected by systemic sclerosis have high frequency of oesophageal mucosal abnormalities and should take long-term PPI therapy. When addressing long-term therapy safety data are clearly needed. Two recent studies have reported increased hip fracture rates with long-term PPI use, raising concerns about adverse effects of this class of drugs on mineral metabolism. The use of PPIs is also associated with an increase in the risk of development of Clostridium difficile infection (CDI) and the use of PPIs during CDI treatment is associated with an increased risk of recurrence.

In order to achieve the desired results and, as with all medications, PPIs should always be used appropriately taking care never to exceed correct dosage and duration. When necessary use of pantoprazole arises as one of the best possible choices.

Parole chiave: Inibitori di pompa, fratture, sindrome coronarica acuta, interazioni farmacologiche.

Key words: Proton pump inhibitor; cytochrome P450; clostridium difficile; bone fractures; drug interactions; acute coronary syndrome; systemic sclerosis.

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