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Impact of proton pump inhibitor treatment on gastrointestinal bleeding associated with non-steroidal anti-inflammatory drug use among post-myocardial infarction patients taking antithrombotics: nationwide study

Anne-Marie Schjerning Olsen,1,2 Jesper Lindhardsen,1 Gunnar H Gislason,1,3,4 Patricia McGettigan,2 Mark A Hlatky,5 Emil Fosbøl,6 Lars Køber,6 Christian Torp-Pedersen,7 Morten Lamberts8

ABSTRACT

STUDY QUESTION
What is the effect of proton pump inhibitors (PPIs) on the risk of gastrointestinal bleeding in post-myocardial infarction patients taking antithrombotics and treated with non-steroidal anti-inflammatory drugs (NSAIDs)?

METHODS
This was a nationwide cohort study based on linked administrative registry data from all hospitals in Denmark between 1997 and 2011. The study included patients aged 30 years and over who had a first myocardial infarction who survived at least 30 days after discharge. The association between PPIs and risk of gastrointestinal bleeding according to NSAID plus antithrombotic therapy was estimated using adjusted time dependent Cox regression models.

STUDY ANSWER AND LIMITATIONS
The use of PPIs was independently associated with decreased risk of gastrointestinal bleeding in post-myocardial infarction patients taking antithrombotics and treated with NSAIDs. Of 82,955 post-myocardial infarction patients (mean age 67.4 years, 64% (n=53,070) men), all of whom were taking single or dual antithrombotic therapy, 42.5% (n=35,233) filled at least one prescription for NSAIDs and 45.5% (n=37,771) received PPIs. Over a mean follow-up of 5.1 years, 3,229 gastrointestinal bleeds occurred. The crude incidence rates of bleeding (events/100 person years) on NSAID plus antithrombotic therapy were 1.8 for patients taking PPIs and 2.1 for those not taking PPIs.

The adjusted risk of bleeding was lower with PPI use (hazard ratio 0.72, 95% confidence interval 0.54 to 0.95) regardless of antithrombotic treatment regimen, type of NSAID, and type of PPI used. The main limitation of the study is its observational non-randomised design. The results suggest that PPI treatment probably has a beneficial effect regardless of underlying gastrointestinal risk and that when NSAIDs cannot be avoided in post-myocardial infarction patients, physicians might prescribe a PPI as well. The study does not clarify whether PPIs might be safely omitted in specific subgroups of patients with a low risk of gastrointestinal bleeding.

WHAT THIS STUDY ADDS
In post-myocardial infarction patients, bleeding complications have been associated with both antithrombotic and NSAID treatment. Concurrent use of PPIs was independently associated with a decreased risk of gastrointestinal bleeding in post-myocardial infarction patients taking antithrombotics and NSAID, regardless of antithrombotic treatment regimen, type of NSAID, and type of PPI used.

FUNDING, COMPETING INTERESTS, DATA SHARING
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WHAT IS ALREADY KNOWN ON THIS TOPIC
Although administration of non-steroidal anti-inflammatory drugs (NSAIDs) to post-myocardial infarction patients is discouraged owing to their cardiovascular risks, NSAIDs are still widely used because painful conditions are common. Among post-myocardial infarction patients, bleeding complications have been associated with both antithrombotic and NSAID treatment. The effect of proton pump inhibitors (PPIs) on the risk of gastrointestinal bleeding in post-myocardial infarction patients taking antithrombotics who are treated with NSAIDs is unknown.

WHAT THIS STUDY ADDS
Concurrent use of PPIs was independently associated with decreased risk of gastrointestinal bleeding in post-myocardial infarction patients taking antithrombotics who were treated with NSAIDs. This finding was independent of antithrombotic treatment regimen, type of NSAID, and type of PPI used. Post-myocardial infarction patients in whom NSAIDs are judged necessary might benefit from PPI treatment as well.
PPIs should be used with both antithrombotic and NSAID treatments in patients judged to be at high risk of gastrointestinal bleeding. In the United Kingdom, the National Institute for Health and Care Excellence (NICE) recommends routine PPI co-prescription with NSAIDs for everyone aged 45 years and older with osteoarthritis, rheumatoid arthritis, or chronic low back pain, as well as for people taking antithrombotics who are at high risk of gastrointestinal adverse effects including bleeding. However, the potential benefit of PPI therapy used with antithrombotic/NSAID combinations in post-myocardial infarction patients irrespective of gastrointestinal risk is unknown. We therefore investigated the effect of PPIs on the risk of bleeding in Danish post-myocardial infarction patients taking antithrombotics who were also treated with NSAIDs.

Methods

Data sources
Diagnostic data came from the Danish National Patient Registry, which using ICD-10 (international classification of diseases, 10th revision) to classify hospital admissions (supplementary table 1). Each hospital admission is registered with one main discharge diagnosis and one or more supplementary diagnoses if appropriate. Information on vital status (dead or alive) came from the civil registration system through Statistics Denmark. We obtained primary, secondary, and contributing causes of death recorded by a physician from the National Causes of Death Registry. The National Prescription Registry provided information on the date of dispensing, quantity dispensed, strength, and formulation of all drugs dispensed from Danish pharmacies and classified according to the Anatomical Therapeutic Chemical (ATC) system (supplementary table 2). The partial reimbursement of drug expenses by the Danish healthcare system requires all pharmacies to register each drug dispensed in the National Prescription Registry, ensuring complete registration.

In Denmark, every resident has a permanent unique civil registration number that enables linkage across administrative registries.

Study population and follow-up
We identified all patients aged 30 years and over in the National Patient Registry between 1997 and 2011 who had a primary diagnosis of acute myocardial infarction (ICD-10 code I21 to I22), received antithrombotics, and survived at least 30 days from discharge (date of inclusion). The 30 day restriction defined a quarantine period. The follow-up period started after the 30 day restriction defined a quarantine period to minimise risk of immortal time bias. To avoid selection bias, we used a new user design, excluding patients who collected a prescription for an NSAID during the quarantine period (n=5711). Patients were followed until one of the following events (which ever came first): event of interest, emigration, death, or end of study period (31 December 2011). The diagnosis of myocardial infarction has been validated with a specificity exceeding 90%.20

Antithrombotic, PPI, and NSAID treatment
We used claimed prescriptions of aspirin or clopidogrel to characterise patients as receiving either antithrombotic monotherapy (aspirin or clopidogrel) or dual therapy. We excluded post-myocardial infarction patients who received no antithrombotic (n=15353) or an oral anticoagulant only (n=1440). We identified all claimed prescriptions for NSAIDs (ATC M01A, excluding glucosamine (M01AX05)). We categorised rofecoxib and celecoxib as cyclo-oxygenase-2 selective (COX 2) inhibitors; ibuprofen, diclofenac, and naproxen as non-selective NSAIDs; and all other NSAIDs as “other” NSAIDs. The only NSAID available in Denmark without prescription is ibuprofen (since 2001). We categorised PPI treatment in one group (ATC A02BC); we also examined five individual PPIs—omeprazole, lansoprazole, pantoprazole, esomeprazole, and rabeprazole.

We calculated exposure periods for NSAIDs and PPIs for each patient by estimating a daily dose after comparing the accumulated dose and the elapsed time from consecutive prescriptions for the drug under investigation. We determined ongoing exposure by dividing the number of tablets/capsules dispensed by the estimated average dosage. If only one prescription was registered for an individual, we used a standard dosage, defined as the minimal recommended dosage, to estimate the daily dose. We used information on increasing or decreasing dosage only to continuously assess whether tablets were available. We defined exposure as having occurred when patients had drug available and discontinuation as when they had no more drug available. Methods for determining dose and treatment duration have been described previously.21 22 For most patients, treatment regimens changed during the study period, so we treated NSAID and PPI use in the analysis as time varying exposures—that is, patients changed exposure group according to claimed prescriptions. Each patient’s exposure group at inclusion defined baseline treatment, shown in the table with the covariate distributions (supplementary figure).

Comorbidity
We identified comorbidities from previous diagnoses and at discharge from the index myocardial infarction, as specified in the Ontario acute myocardial infarction mortality prediction rule, and potential risk factors for bleeding (previous bleeding, alcohol consumption, liver disease, and ulcers). The Ontario acute myocardial infarction mortality prediction rule is a logistic regression model that predicts 30 day and one year mortality by using 11 variables determinable from hospital discharge databases (age, sex, shock, diabetes with complications, congestive heart failure, cancer, cerebrovascular disease, pulmonary oedema, acute renal failure, chronic renal failure, cardiac dysrhythmia). In our analyses, we incorporated each variable as a covariate and permitted diagnoses up to one year previously (supplementary table 1). To account for accumulation of risk factors during follow-up, we did an analysis including all variables in the main analysis as time dependent (exempt for inclusion year and
percutaneous coronary intervention status (considered as related to myocardial infarction inclusion criteria) and previous peptic ulcer disease (considered an intermediate variable of the primary outcome during follow-up)). During the entire span of the database, patients could change status on the date of exposure.

Outcome
We defined the primary outcome of gastrointestinal bleeding as hospital admission for or death from a bleeding gastrointestinal ulcer, haematemesis, melena, or unspecified gastrointestinal bleeding from the National Causes of Death Register and National Patient Register. Occurrence and type of bleeding as recorded in hospital databases have shown a positive predictive value of 89-99%.

Statistical methods
We calculated crude incidence rates as number of events per 100 person years according to the different treatment regimens. We estimated the effects of PPI treatment on gastrointestinal bleeding with adjusted Cox proportional hazards models in terms of hazard ratios and 95% confidence intervals for gastrointestinal bleeding with the drug exposure continuously updating—that is, as time varying exposure allocated according to treatment regimen. We considered patients to be at risk only when exposed to the drug (during active treatment). Each patient could have multiple treatment groups throughout follow-up. We calculated risk time (person years) only for the active treatment period. The timescale in the Cox model was days passed since inclusion. We adjusted all models for age, sex, year of index hospital admission, concomitant drugs, comorbidity, and percutaneous coronary intervention status. We did additional analyses to assess any association between PPI use and individual antithrombotic regimens and NSAIDs.

We did nine sensitivity analyses. (1) To take account of any effect of over the counter NSAID use, we ended follow-up in December 2010. (3) We examined cardiovascular death as a solo endpoint. (4) We stratified the cohort at 65 years to take account of guidelines recommending PPIs for people over 65 taking antithrombotic treatment. (5) We controlled for the variables included in the HAS-BLED score. (6) We did an analysis including all covariates as time dependent. (7) We stratified the population in two groups: high (previous bleeding) and low (no previous bleeding) risk of gastrointestinal bleeding. (8) Although the indication for NSAID use was not systematically available, we were able to do an analysis of patients with rheumatoid arthritis. (9) We examined the effect of duration of NSAID treatment (0-14 days and >14 days).

We confirmed the validity of the proportional hazard assumption, linearity of continuous variables, and lack of interaction and chose a significance level of 0.05. We used SAS 9.2 and Stata 11.0 for all statistical calculations.

Patient involvement
No patients were involved in setting the research question or the outcome measures; nor were they involved in the design and implementation of the study. There are no plans to involve patients in dissemination.

Results
A total of 128 751 patients were admitted with a first myocardial infarction in the period 1997 to 2011, of whom 82 955 (64.4%) were taking single or dual antithrombotic therapy and met the study inclusion criteria (fig 1). Baseline characteristics included 67.4 years mean age, 64% male, and 6.6% with a history of previous gastrointestinal bleeding (table). A total of 35 233 (42.5%) patients had one or more NSAID prescription claim after discharge; 37 711 (45.5%) received a PPI, and 10 613 (12.8%) had concurrent NSAID and PPI exposure (fig 2). The proportions receiving a PPI in the year after their first myocardial infarction increased from 15% in 1997 to 37% by 2010. The proportions receiving NSAIDs peaked at 20% in 2002 and subsequently declined, whereas the proportions receiving a PPI concurrently with an NSAID increased from less than 10% to more than 30% by 2010 (fig 2).

Gastrointestinal bleeding
We identified 3229 gastrointestinal bleeding events, 282 (8.7%) of which were fatal. A total of 327 events were registered during NSAID treatment. The overall crude incidence rate was 0.8 (95% confidence interval 0.7 to 0.8) events per 100 person years The crude incidence rate of gastrointestinal bleeding on NSAID treatment was 2.1 (1.8 to 2.4) events per 100 person years without concurrent PPI treatment and 1.8 (1.4 to 2.4) with concurrent PPI treatment (fig 3). After multivariable adjustment for baseline differences, use of PPIs concurrently with combined antithrombotic and NSAID treatment was associated with a significantly lower risk of gastrointestinal bleeding (hazard ratio 0.72, 95% confidence interval 0.54 to 0.95) compared with treatment without concurrent PPIs (fig 3). Compared with combined antithrombotic and NSAID treatment (without concurrent
Baseline characteristics of total study population and individual antithrombotic treatment groups. Values are numbers (percentages) unless stated otherwise

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Total population (n=82 955)</th>
<th>No PPI and no NSAID (n=68 044)</th>
<th>NSAID (n=2006)</th>
<th>PPI (n=12 334)</th>
<th>PPI and NSAID (n=57)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean (SD) age, years</td>
<td>67.4 (13.3)</td>
<td>66.7 (13.3)</td>
<td>67.7 (13.2)</td>
<td>71.4 (12.8)</td>
<td>68.7 (13.0)</td>
</tr>
<tr>
<td>Male sex</td>
<td>53 070 (64.0)</td>
<td>44 802 (65.8)</td>
<td>11 999 (59.8)</td>
<td>6 772 (54.9)</td>
<td>2 97 (52.0)</td>
</tr>
</tbody>
</table>

Comorbidities:
- Cardiac arrhythmias: 7624 (9.2) vs 5821 (8.6), P=0.001
- Peripheral vascular disease: 7112 (9.0) vs 5324 (7.9), P=0.001
- Cerebral vascular disease: 3716 (4.5) vs 2 748 (4.0), P=0.001
- Diabetes with complications: 1340 (0.8) vs 89 (0.4), P=0.001
- Acute renal failure: 628 (0.8) vs 339 (0.5), P=0.001
- Chronic renal failure: 1027 (1.2) vs 600 (0.8), P=0.001
- Malignancy: 1796 (2.2) vs 1 296 (1.9), P=0.001
- Shock: 216 (0.3) vs 143 (0.2), P=0.001
- Chronic obstructive pulmonary disease: 690 (0.8) vs 522 (0.8), P=0.001
- Previous bleeding: 5311 (6.6) vs 3 466 (5.1), P=0.001
- Liver disease: 954 (1.2) vs 657 (1.0), P=0.001
- Peptic ulcer: 3481 (4.2) vs 1 706 (2.5), P=0.001
- Alcohol: 3255 (3.9) vs 2 415 (3.6), P=0.001
- Percutaneous coronary intervention: 32 376 (39.0) vs 26 764 (33.9), P=0.001

Concomitant drugs:
- β blockers: 60 789 (73.3) vs 50 541 (74.3), P=0.001
- Angiotensin converting enzyme inhibitors: 31 748 (38.3) vs 25 821 (38.0), P=0.001
- Statins: 48 829 (58.9) vs 40 214 (59.1), P=0.001
- Spironolactone: 4771 (5.8) vs 3560 (5.2), P=0.001
- Loop diuretics: 22 020 (26.5) vs 16 685 (24.5), P=0.001
- Glucose lowering drugs: 6388 (7.7) vs 4860 (71), P=0.001

NSAID=non-steroidal anti-inflammatory drug; PPI=proton pump inhibitor.

PPI, taking antithrombotics only (without NSAID or PPI) or antithrombotics with concurrent PPI treatment (without NSAID) were each associated with a lower risk of gastrointestinal bleeding. The bleeding risk associated with concurrent PPI and NSAID treatment was similar for each antithrombotic regimen (supplementary table 3). PPIs taken concurrently with NSAID and dual antithrombotic therapy were associated with a reduced risk of gastrointestinal bleeding (incidence rate 2.5 (1.3 to 4.8) events per 100 person years) compared with NSAID and dual antithrombotic therapy without concurrent PPIs (5.2 (3.9 to 6.8) events per 100 person years) (hazard ratio 0.41, 0.20 to 0.84).

Different NSAID groups (selective COX 2 inhibitors, non-selective NSAIDs, other NSAIDs) and PPIs were associated with similar adjusted hazard ratios and crude incidence rates as for overall NSAID treatment (fig 3, top). Individual PPIs (omeprazole, pantoprazole, lansoprazole, and esomeprazole) were each associated with reduced bleeding risks (fig 3, bottom). For individual NSAIDs (rofecoxib, celecoxib, diclofenac, ibuprofen, naproxen), hazard ratios for bleeding were lower with concurrent PPI use than without, but not significantly so owing to small numbers of events individually (supplementary table 4).

Sensitivity analyses
None of the sensitivity analyses changed the results appreciably (web appendix).

Discussion
This nationwide study suggests that PPIs reduce the risk of gastrointestinal bleeding associated with NSAID use among post-myocardial infarction patients taking antithrombotics. Four PPIs (omeprazole, pantoprazole, lansoprazole, and esomeprazole) were individually associated with a lower risk of bleeding compared with non-use for the antithrombotic regimens examined and irrespective of whether selective COX 2 inhibitors or non-selective NSAIDs were prescribed.
Adverse cardiovascular effects associated with NSAIDs have led to discouragement of their use in patients with cardiovascular disease. NSAIDs have also been associated with a substantial independent risk of bleeding in patients already taking antithrombotics. Nevertheless, they continue to be prescribed quite extensively for patients with cardiovascular risks. Previous studies have reported that around 40% of Danish patients with myocardial infarction are exposed to NSAIDs, consistent with the 42.5% found here. PPIs reduce the risks of gastrointestinal complications associated with antithrombotics and with NSAIDs, including among patients at high risk treated with selective or non-selective NSAIDs. Our results support these findings in showing that concurrent PPIs were associated with a reduced risk of gastrointestinal bleeding among post-myocardial infarction patients taking both NSAIDs and antithrombotics.

Concern has been expressed that some PPIs might diminish the antiplatelet effect of clopidogrel, most likely through inhibition of CYP2C19 mediated conversion of clopidogrel into its active form, but studies have been contradictory and definitive data are awaited. A similar situation exists regarding concerns that PPIs might impair the cardiovascular protective efficacy of aspirin. Consequently, PPIs are recommended for aspirin treated patients at high risk of gastrointestinal bleeding. American specialty guidelines recommend that in patients at high risk of gastrointestinal bleeding, a PPI should be used concomitantly with antithrombotics and with NSAID treatment. Our data suggest that PPI treatment probably has a beneficial effect regardless of underlying gastrointestinal risk and that when NSAIDs cannot be avoided in post-myocardial infarction patients, physicians might prescribe a PPI as well. Our study does not clarify whether PPIs might be safely omitted in specific subgroups of patients with a low risk of gastrointestinal bleeding.

Limitations of study
The main limitation of this study is its observational non-randomised design. Furthermore, we do not have information on whether patients stop taking antithrombotics when treated with NSAIDs; however, given post-myocardial infarction treatment guidelines, we think that this is unlikely. We cannot exclude a possible effect of unmeasured confounders. However, if an unmeasured confounder or a combination of confounders were present in 20% of the NSAID treated cohort, our calculations suggest that the confounder would have to increase the risk by a factor of 1.8 to 3.5 to explain the increased risk observed. Existence of such a confounder or combination of confounders is unlikely, but not impossible, as we had no information on other risk factors such as smoking, lipid concentrations, or body mass index. Moreover, we did an analysis with continuous assessment of co-variables to account for accumulation of risk factors during follow-up to further minimise the effect of potential confounders; the results were unaffected. We did not have information about the indication for NSAID treatment, but this was probably for non-cardiac disease, as NSAIDs are not used to treat ischaemic heart disease. Having users treated with the drug(s) under investigation before inclusion could result in confounding (healthy user effect). Our estimates when we excluded prevalent NSAID users remained the same, so it is unlikely that confounding by indication alone could drive the observed results. The only NSAID available over the counter in Denmark is ibuprofen (since 2001) in low doses (200 mg) and in limited quantity. Restricting the analyses to 1997-2000 did not change our results. Aspirin is also available without prescription, but because of partial reimbursement of drug expenses, chronic users, in particular those with a history of myocardial infarction, are most likely to be issued with a prescription for aspirin as thrombo-prophylaxis. For these reasons, we think unrecorded over the counter drug use is unlikely to have had a major effect on the study results. Several estimates of a beneficial effect of concomitant specific PPI and specific NSAID treatment on risk of gastrointestinal bleeding were non-significant. This was potentially owing to a low number of events. Nevertheless, all estimates were comparable to our main analysis.

Conclusion
This study of a real life cohort of post-myocardial infarction patients taking antithrombotics suggests that use of a PPI diminishes the risk of gastrointestinal bleeding associated with NSAID treatment, regardless of the type of NSAID or PPI prescribed. Consequently, when NSAIDs cannot be avoided in post-myocardial infarction patients, physicians might consider prescribing a PPI as well.

Contributors: AMSO and ML made primary contributions to study design, data collection, data analysis, interpretation of results, and writing the manuscript. ML helped to write the first draft. JI, GHS, and CT-P contributed to the study design. All authors contributed to the interpretation of results and critical revision of the manuscript; all approved the final manuscript. AMSO is the guarantor.
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Competing interests: All authors have completed the ICMJE uniform disclosure form available at www.icmje.org/coi_disclosure.pdf (available on request from the corresponding author) and declare: no support from any organisation for the submitted work; no financial relationships with any organisations that might have an interest in the submitted work in the previous three years, no other relationships or activities that could appear to have influenced the submitted work.

Ethical approval: The Danish Data Protection Agency approved this study (No. 2007-58-0015; internal ref: GEH-2014-074-I Suite 02732), and data at the individual level were made available to us such that specific patients could not be identified. Retrospective register studies do not by law require ethical approval in Denmark.

Transparency declaration: The lead author (the manuscript’s guarantor) affirms that this manuscript is an honest, accurate, and transparent account of the study being reported; that no important aspects of the study have been omitted; and that any discrepancies from the study as planned (and, if relevant, registered) have been explained.

Data sharing: No additional data available.

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