Adding to the evidence or to the confusion: dual antithrombotic therapy in chronic coronary syndrome and atrial fibrillation

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Atrial fibrillation (AF) is the most common arrhythmia in the world with the lifetime risk estimated to be 1/3 in men and women over the age of 50 years.1 Cardiovascular disease, like misfortune, does not come singly in most cases. Coronary artery disease (CAD) with the chronic coronary syndrome (CCS) or acute coronary syndrome (ACS) is a common comorbid condition. Management of patients with AF with a comprehensive treatment of risk factors and concomitant diseases is the key to treat these patients. Yet, the devil is in the details as treatment becomes increasingly complex.

Patients with AF and CAD require antiplatelet therapy (APT) in addition to oral anticoagulation (OAC) after myocardial infarction or percutaneous coronary intervention (PCI) for a limited time period. Depending on ischaemic risk, bleeding risk and unplanned PCI, different treatment regimens are available with dual antithrombotic therapy with OAC and APT up to 12 months.2 Looking at each possible combination of drugs and therapy length, the cardiologist is left with 2.8 million possible combinations in the first 12 months only.3

PRECISION MEDICINE BUT TOO MANY OPTIONS WITH LOW EVIDENCE?

This subanalysis from the recently published randomised controlled ‘Atrial Fibrillation and Ischemic Events With Rivaroxaban in Patients With Stable Coronary Artery Disease Study’ (AFIRE) by Fukaya et al4 examines whether aspirin or P2Y12 inhibitors (clopidogrel in 94.6%) in addition to rivaroxaban impacts outcomes in patients with CCS. The main trial showed that dual antithrombotic therapy is associated with increased mortality compared with OAC monotherapy in patients with AF and CCS.

Patients in the dual antithrombotic arm were compared in regard to additional APT (P2Y12 inhibitors n=297 vs aspirin n=778) and a primary outcome of composite of stroke, systemic embolism, myocardial infarction, unstable angina requiring urgent revascularisation or death from any cause. No difference in the primary outcome was observed in patients receiving P2Y12 inhibitors or aspirin (6.76% vs 5.28%, p=0.178). However, mortality was higher in the P2Y12 inhibitors group (4.44% vs 2.85% per patient-year; p=0.041).

There is limited evidence in the comparison of aspirin with P2Y12 inhibitors in addition to OAC in patients with CCS. The AFIRE main trial showed that these patients do not benefit from additional APT,5 but doubts remain, and dual antithrombotic therapy is still common in clinical practice.6 The benefit seen with clopidogrel compared with aspirin in the monotherapy of patients undergoing PCI, recently shown with the ‘Harmonizing Optimal Strategy for Treatment of Coronary Artery Stenosis-EXTended Antiplatelet Monotherapy (HOST-EXAM)’ trial, is not evident in patients with AF in the analysis by Fukaya et al.4

Interpretation of this data is to be seen in the contexts of its many caveats. The AFIRE trial was randomised, but the comparison of P2Y12 inhibitors and aspirin was not, as seen with significant baseline differences between the groups. The design was open label, which certainly can introduce bias. Also, the generalisability of the cohorts to European/Western patients is limited. In particular, anticoagulant dosing is different here and significant numbers of patients received bare-metal stents (more than 30% in the aspirin group) which is not standard of care anymore in the guidelines. General assumptions on P2Y12 inhibitors should be avoided, since almost 95% of patients with P2Y12 inhibitors received clopidogrel instead of more potent antiplatelet drugs like prasugrel or ticagrelor. Prescribing physicians in the AFIRE trial were free to prescribe more potent agents but may have feared a higher bleeding risk with prasugrel or ticagrelor.

Interestingly, the authors report no difference in major bleeding between both cohorts (2.35% vs 2.95%, p=0.456) which again contrasts with the findings of HOST-EXAM where clopidogrel was associated not only with fewer thrombotic events (3.7% vs 5.5%, p=0.003) but also less bleeding (2.3% vs 3.3%, p=0.003).7 The pathophysiological background of aspirin-induced gastric bleeding is nicely explained by the inhibition of cyclo-oxygenase which not only leads to the desired inhibition of platelet-aggregator thromboxane A2 but also the synthesis of prostaglandins which are essential for gastric mucus secretion (figure 1).

One of the strengths in the work by Fukaya et al may be the utilisation of proton pump inhibitors (PPIs) in both cohorts (64.3% in the P2Y12 group and 62% in the aspirin group). In the ‘Rivaroxaban for the Prevention of Major Cardiovascular Events in Coronary or Peripheral Artery Disease (COMPASS)’ trial where rivaroxaban 2.5 mg daily in addition to aspirin in stable atherosclerotic disease was shown to reduce cardiovascular mortality, an interesting substudy was conducted. Patients who were not on PPI at baseline were randomised to PPI or placebo resulting in two groups over 8000 patients with a mean follow-up of 3 years. Therapy with PPI was shown to be very safe without increase in cardiovascular events, cancer or pneumonia or dementia but a twofold increase of Clostridium difficile infections.8 The COGENT trial showed that concomitant therapy with PPI (in this case omeprazole) in patients with APT reduces gastrointestinal bleeding without a rise in thrombotic events.9 However, the most recent European Society of Cardiology (ESC) guidelines for management of ACS without ST-segment elevation (NSTEMI) do not recommend omeprazole due to concerns of inhibition of CYP2C19 and thus efficacy of clopidogrel.2

The ESC guidelines on management of AF mention PPIs without giving a level of recommendation akin to the NSTEMI guidelines. With the abundance of data supporting the use of PPI in patients with antiplatelet or dual antithrombotic therapy, they should be considered more frequently and guidelines have to provide more powerful recommendation.
In conclusion, the authors shed some new light on the dual antithrombotic therapy of patients with AF and concomitant CCS. While not recommended without acute coronary events and certainly related to an increased risk of bleeding, it represents the clinical reality in many cases. Whereas the current study suggests that aspirin or a P2Y₁₂ inhibitor may be equivalent, not the current study suggests that aspirin or the clinical reality in many cases. Whereas

CONCLUSION

Figure 1 Overview of antithrombotic and anticoagulation mechanisms on haemostasis to prevent thrombotic events while increasing bleeding risk. In patients with AF and CCS, antithrombotic drugs can be either antithrombotic drugs or oral anticoagulation (OAC). Aspirin and P2Y₁₂ inhibitors lead to an inhibition of glycoprotein IIb/IIIa, a fibrinogen receptor that aids platelet activation. Aspirin blocks the formation of thromboxane A₂ (TxA₂) limiting platelet aggregation and also reduces gastric mucus secretion. OAC either inhibits factor Xa (FXa) or thrombin to reduce clotting formation. Both medications reduce thrombotic events such as ischaemic stroke (mainly OAC) and coronary ischaemic events (antplatelets) but increase bleeding risk, that is, gastric bleeding. Potential protection may arise from proton pump inhibitors (PPIs) and histamine H₂ receptor (H₂ blockers) may provide protection by reducing stomach acid production. Adapted from Collet et al. AF, atrial fibrillation; CCS, chronic coronary syndrome.

The personalised treatment of patients with concomitant AF and CCS remains complex and more evidence is required to make guideline recommendations.

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