Updated overview of evidence on optimal antithrombotic therapy in patients with atrial fibrillation undergoing percutaneous coronary intervention

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Abstract

Selection of the optimal peri- and postprocedural antithrombotic regimen in patients with atrial fibrillation (AF) undergoing percutaneous coronary intervention (PCI) is a common clinical problem which may pose a challenge to medical practitioners. This systematic review summarizes the updated evidence on this topic. Non-vitamin K oral anticoagulants (NOACs) at standard doses are the preferred option in most of post PCI patients with AF, except those few with a clear indication for a vitamin K antagonist (VKA). Reduced NOAC doses should be considered in dabigatran- or rivaroxaban-treated patients with a high bleeding risk, which prevail over concerns about stent thrombosis or ischemic stroke. There is insufficient evidence to favor one NOAC over another in this setting. In the early post stenting period, triple therapy comprising a NOAC, clopidogrel and aspirin is recommended. Timing of post PCI aspirin cessation should be based on a careful analysis of the bleeding and ischemic risk. There is only low quality evidence regarding the optimal approach to elective or urgent/emergency PCI procedures in patients requiring oral anticoagulation. It is suggested that there is no need of interruption of VKA and PCI procedure should be performed via radial artery access with a lower dose of unfractionated heparin. On the other hand, NOACs are usually stopped before elective PCIs, while urgent/emergency procedures may be performed with the addition of low-dose parental anticoagulation.

Key words: atrial fibrillation, percutaneous coronary intervention, non-vitamin K oral anticoagulants, vitamin K antagonist, anticoagulation.

Introduction

Optimal anticoagulant and antiplatelet treatment of patients with atrial fibrillation (AF) undergoing percutaneous coronary intervention (PCI) still poses a great challenge to medical practitioners. The number of patients in need of such a treatment is ever-growing. AF affects approximately 33 million people worldwide, out of which 5–15% will undergo PCI at some point of their lives [1–3]. Clinicians need to face the problem of combining anticoagulant therapy, which is recommended for the prevention of ischemic stroke and systemic embolism in patients with AF, and dual antiplatelet therapy, which reduces the risk of stent thrombosis and subsequent coronary events. On the other hand, triple antithrombotic therapy, comprising anticoagulant and dual antiplatelet therapy, is associated with a high risk of bleeding. Over the last few years, numerous clinical trials have been conducted with the aim of developing the best treatment strategies. Based on the results of the after-mentioned trials, new guidelines on how to manage such complex cases have been developed. The aim of the present review is to provide an updated overview of evidence regarding the optimal peri- and postprocedural antithrombotic regimen in AF patients undergoing PCI.

Search strategy

A systematic investigation of all published literature including oral presentations was conducted to minimize...
the risk of bias. A database search including PubMed, CENTRAL and Google Scholar databases, covering the period up to April 2020 was conducted by two independent investigators (M.K. and J.R.B.). We paid particular attention to randomized clinical trials, European and American guidelines and expert consensus documents. Proceedings of the Scientific Sessions of the American College of Cardiology (http://www.acc.org), American Heart Association (http://www.heart.org), European Society of Cardiology – ESC (http://www.escardio.org), Transcatheter Cardiovascular Therapeutics (http://www.tctmd.com) and EuroPCR (http://www.europcr.com) were also considered. The following keywords were applied: ‘PCI’, ‘stenting’, ‘antiocoagulant’, ‘anticoagulation’, ‘NOAC’, ‘DOAC’ and ‘warfarin’. References of retrieved studies, guidelines and position statements were searched manually for additional relevant publications.

**Periprocedural management during elective procedures**

According to the ESC guidelines on chronic coronary syndromes, for the patient treated with vitamin K antagonist (VKA) and admitted due to elective or emergency procedures, there is no need of VKA interruption. Radial artery access is preferred along with intraprocedural unfractionated heparin at a lower dose of 30–50 U/kg [4]. The uninterrupted strategy is supported by various studies. In the WOEST study, where 573 patients on VKA treatment underwent PCI, in 43% the procedure was carried out without VKA interruption and in 57% with VKA interruption plus low-molecular weight heparin bridging. At 30 days and at 1-year follow-up, the occurrence of bleeding events and major adverse cardiovascular events (MACE), including death, myocardial infarction, stroke, target vessel revascularization, and stent thrombosis was comparable in both groups [5].

In the AFCAS registry of 663 patients with AF who underwent PCI (for stable coronary artery disease (CAD) in approximately 50%), at 3-month follow-up, both unadjusted major bleeding and MACE rates were significantly higher in the low-molecular-weight heparin (LMWH)-bridging group. After propensity score matching, and subsequent comparison of patients, in whom frequency of femoral access was similar, major bleeding was more frequent in the LMWH-bridging group at both 3 and 12 months, whereas MACE rates were comparable [6]. In a meta-analysis, uninterrupted oral anticoagulation (OAC) was found to be at least as safe as interrupted OAC, and safer than interrupted OAC with bridging strategy [7]. Although, it was shown that the incidence of both early (24-h) and late (30-day) radial artery occlusion was significantly higher in the ongoing warfarin group comparing to standard, intra-procedural unfractionated heparin (UFH) administration [8]. However, doses as low as 30–50 U/kg have been shown to be effective in preventing ischemic complications related to PCI [9].

Limited data are available for AF patients undergoing PCI while on NOAC. There is an inconsistency of the results reported with the two different NOACs (dabigatran and rivaroxaban). In the phase IIa, multicenter clinical trial of 50 patients undergoing elective PCI, dabigatran appeared to provide insufficient anticoagulation and clinical outcomes tended to be higher in the dabigatran group [10]. Different results were obtained in another phase IIa, multicenter trial [11]. In the X-PLORER study, patients undergoing elective PCI and on dual antiplatelet therapy were randomized to a single dose of either rivaroxaban 10 mg, rivaroxaban 20 mg, rivaroxaban 10 mg plus intravenous bolus of UFH, or standard UFH. In all groups receiving rivaroxaban, coagulation was effectively suppressed, comparably to standard UFH. No patients in the three rivaroxaban arms required bail-out antithrombotic medication and/or had clinical signs of catheter-related thrombosis. No significant bleeding was observed in either group up to 30 days after PCI.

According to the ESC guidelines on chronic coronary syndromes (CCS), it is recommended that NOAC therapy is stopped for 12–48 h before elective PCI, depending on renal function and the particular NOAC regimen [4]. Radial artery access is preferred along with intraprocedural unfractionated heparin at a standard dose (70–100 U/kg). Pre-treatment with aspirin 75–100 mg daily is recommended, and clopidogrel (300–600 mg loading dose if not on long-term maintenance therapy) is recommended in preference to prasugrel or ticagrelor. VKA-treated patients receiving aspirin and clopidogrel post-PCI should have a target international normalized ratio in the range of 2.0–2.5, aiming for high time in therapeutic range (> 70%).

**Periprocedural management during urgent and emergency procedures**

No randomized trials have specifically studied periprocedural management in anticoagulated patients developing an acute coronary syndrome (ACS) and undergoing acute angiography with or without PCI. According to “2018 Joint European consensus document on the management of antithrombotic therapy in atrial fibrillation”, patients with AF taking OAC presenting with ACS and/or undergoing percutaneous cardiovascular interventions should receive aspirin immediately (150–300 mg oral loading dose) [12, 13]. Anticoagulated patients undergoing primary PCI should receive additional low-dose parenteral anticoagulation regardless of the timing of the last dose of OAC (VKA or NOAC) (e.g. enoxaparin 0.5 mg/kg i.v. or UFH 60 IU/kg). In patients on VKA and international normalized ratio (INR) value > 2.5, it is suggested not to administer additional intraprocedural UFH [14].

To reduce the risk of bleeding there is a possibility to postpone the administration of P2Y12 inhibitors to
the time of PCI, when the anatomy of coronary artery is known [14]. The P2Y12 inhibitor of choice is clopidogrel with 600 mg loading dose [14]. Prasugrel and ticagrelor are not routinely recommended in anticoagulated patients due to their increased bleeding risk [14, 15]. The radial approach is strongly recommended [16].

It is worth mentioning that a big discussion concerning dual therapy over triple therapy does not concern periprocedural management during coronary procedures as the majority of them were done on two antiplatelet agents.

Post PCI therapy with VKAs

Trials with VKAs which compared double versus triple therapy in AF patients undergoing PCI are the ISAR-TRIPLE trial (Triple Therapy in Patients on Oral Anticoagulation After Drug Eluting Stent Implantation) [17] and the WOEST trial (What Is the Optimal Antiplatelet & Anticoagulant Therapy in Patients With Oral Anticoagulation and Coronary Stenting) [5].

In the ISAR-TRIPLE trial, patients treated with OAC were randomized to either 6 weeks or 6 months of DAPT (OAC plus clopidogrel) after PCI and DES (drug-eluting stent) implantation (one-third with an ACS) [17]. The main finding is that there was no difference in major bleeding events between patients treated with a 6-week or 6-month duration of clopidogrel. The authors explained that it might be due to the fact that approximately one-half of all bleeding events occurred in the first 6 weeks after PCI, when both groups received the same therapy consisting of aspirin, clopidogrel, and OAC. There was no extra risk of the composite ischemic endpoint with shorter duration therapy and the rate of stroke was comparable in both groups.

The WOEST trial, despite the small sample size (573 patients), provided randomized evidence that the early discontinuation of aspirin reduces bleeding in comparison to triple therapy [5]. The trial found a 64% relative decrease in minor bleeding episodes without increase in ischemic events. The WOEST trial contributed to initiation of multiple trials of aspirin-free strategies in PCI. What is more, in light of the results of the WOEST trial, the 1-year duration of triple therapy has been questioned, and the standard duration of triple therapy has become shorter.

Post PCI therapy with NOACs

The PIONEER AF PCI trial (Open-Label, Randomized, Controlled, Multicenter Study Exploring Two Treatment Strategies of Rivaroxaban and a Dose-Adjusted Oral Vitamin K Antagonist Treatment Strategy in Subjects with Atrial Fibrillation who Undergo Percutaneous Coronary Intervention) compared 3 treatment strategies after PCI in 2,124 patients with AF: a low-dose rivaroxaban (15 mg once daily – OD) plus clopidogrel, a very low-dose rivaroxaban (2.5 mg (BID) twice daily) plus aspirin and clopidogrel, and control triple therapy group with a VKA plus aspirin and clopidogrel [18]. It is important to realize that doses of rivaroxaban used in the PIONEER AF PCI trial do not represent the approved doses for stroke prevention in AF. They were chosen based on dose-finding investigations. The clinically significant bleeding (bleeding requiring medical attention) at 12 months was reduced by both rivaroxaban-based strategies compared with the control group. The efficacy in the prevention of cardiovascular events was similar for all study groups, but the power was low for ischemic endpoints. In the PIONEER AF PCI trial, aspirin was dropped at the investigator’s discretion, most patients received triple therapy ≥ 6 months. Variable duration of dual therapy in the 2.5 mg BID (twice daily) rivaroxaban triple therapy group was 1, 6, or 12 months according to the physician’s choice.

Another trial, RE-DUAL PCI (Randomized Evaluation of Dual Antithrombotic Therapy with Dabigatran versus Triple Therapy with Warfarin in Patients with Nonvalvular Atrial Fibrillation Undergoing Percutaneous Coronary Intervention) was designed to compare the safety of using different doses of dabigatran (110 mg and 150 mg) in combination with clopidogrel and triple antithrombotic therapy with warfarin [19]. In the triple antithrombotic therapy group, aspirin was discontinued after 1 month in patients who received a bare-metal stent (17%) and after 3 months in patients who received a DES (83%). Dual therapy was continued for at least 12 months. Ticagrelor in the RE-DUAL PCI was used in 12%. The risk of major or clinically relevant non-major bleeding in the 110 mg and 150 mg dabigatran group was significantly lower than in the warfarin group. These results did not differ between stable CAD or ACS group or whether clopidogrel or ticagrelor was used. It should be noted, however, that it is impossible to state clearly if the reduction in the risk of bleeding among patients treated with dabigatran was related to the effect of the drug or the non-use of aspirin. There were no significant differences between the two dabigatran-based arms and the warfarin-based strategy when evaluating the risk of death, thromboembolic events and non-planned revascularization, both in ACS and elective PCI populations.

The AUGUSTUS trial (An Open-label, 2 Factorial, Randomized Controlled, Clinical Trial to Evaluate the Safety of Apixaban vs. Vitamin K Antagonist and Aspirin vs. Aspirin Placebo in Patients With Atrial Fibrillation and Acute Coronary Syndrome or Percutaneous Coronary Intervention) showed that the safest combination was apixaban and placebo in addition to P2Y12 inhibitor, most frequently clopidogrel. Apixaban 5 mg BID was associated with significantly less major or clinically relevant non-major bleeding than VKA [20]. The study also proved that aspirin, compared with placebo, was associated with significantly more bleedings. Apart from this the authors showed a reduced combined endpoint composed of
death and hospitalization. It is worth remembering that the used dose of 5 mg BID is a basic regular dose for stroke protection in AF patients. Nevertheless there was a non-significant increase in stent thrombosis events and myocardial infarction with placebo compared to the arm with aspirin. However, the trial was not powered to assess differences in these events between the investigated groups. One of the limitation of the study is that randomization was allowed within 14 days after ACS/PCI and in practice the patients were enrolled at a median of 6 days from ACS and/or PCI, which suggests that most patients in the trial had at least short-term aspirin use before randomization. In the RE-DUAL PCI [19] and PIONEER AF [18] trials patients were enrolled within 120 and 72 h after PCI, respectively. There is one more significant difference between these three trials: in the AUGUSTUS trial, 23.9% of patients with ACS were medically managed, 37.3% with ACS were treated with PCI and 38.8% were enrolled after elective PCI. What does it mean for our patients? The RE-DUAL PCI and the PIONEER AF results describe PCI population, while the AUGUSTUS results are more applicable to medically managed ACS population. Besides, time in therapeutic range (TTR) of warfarin may play an important role in the comparison between warfarin and NOAC treated populations. Mean TTR in the AUGUSTUS trial was 59% [20], while in the RE-DUAL PCI and the PIONEER AF trials TTR was 64–65% [18, 19]. The higher quality of TTR the more robust safety results. Also the follow-up time in the AUGUSTUS trial was shorter than in the PIONEER-AF PCI, the RE-DUAL PCI and ENTRUST AF-PCI trials (6 months vs. 12 months) [18–21].

In the ENTRUST AF-PCI trial, patients were randomly assigned from 4 h to 5 days after PCI to either edoxaban (60 mg OD) plus a P2Y12 inhibitor for 12 months or a vitamin K antagonist (VKA) in combination with a P2Y12 inhibitor and aspirin (100 mg OD, for 1–12 months). The edoxaban dose was reduced to 30 mg if at least one of the following factors (creatinine clearance ≤ 60 kg, or concomitant use of specified potent P-glycoprotein inhibitors) was present. The study showed that double therapy with edoxaban is non-inferior to combined therapy with VKA in terms of major and non-major clinically relevant bleeding events. Main efficacy outcome which was composed of cardiovascular death, stroke, systemic embolism, myocardial infarction and stent thrombosis did not differ between the strategy based on edoxaban and warfarin. Although the study was not designed to test either superiority or non-inferiority of edoxaban-based regimen vs VKA-based regimen in terms of the efficacy outcome [21].

Details regarding the design and results of landmark trials on the role of NOACs in AF patients treated with PCI are summarized in Table I. These randomized trials showed that double therapy with a P2Y12 inhibitor and NOAC is safer with respect to bleeding than triple therapy (including warfarin). However, none of these studies were sufficiently powered for efficacy and the populations studied were a mixture of stable CAD and ACS patients.

Results of recent meta-analyses

Two meta-analyses including the AUGUSTUS, ENTRUST AF-PCI, PIONEER AF-PCI, and RE-DUAL PCI studies found that dual when compared with triple antithrombotic therapy reduced the risk for major bleeding, without any impact on all-cause mortality and the risk of myocardial infarction [22, 23]. Importantly, one of these papers also demonstrated that this benefit was counterbalanced by a significant increase in the risk of stent thrombosis (relative risk (RR) 1.59, 95% confidence interval (CI): 1.01–2.50) [22].

Current recommendations

Anticoagulants in AF patients have demonstrated superiority over aspirin monotherapy or clopidogrel-based DAPT for stroke prevention, and are therefore recommended for this indication [24].

Current ESC recommendations on CCS from 2019 (Table II) showed that in patients who are eligible for a NOAC, it is recommended that apixaban 5 mg BID, dabigatran 150 mg BID, edoxaban 60 mg OD, or rivaroxaban 20 mg OD is used in preference to a VKA in combination with antiplatelet therapy (class of recommendation I, level of evidence A) [4]. In patients with a high bleeding risk, rivaroxaban 15 mg OD should be considered in preference to rivaroxaban 20 mg OD or dabigatran 110 mg BID should be considered in preference to dabigatran 150 mg BID for the duration of concomitant single or dual antiplatelet therapy (class of recommendation IIa, level of evidence B). Triple therapy with aspirin, clopidogrel and an OAC for ≥ 1 month should be considered when the risk of stent thrombosis outweighs the bleeding risk, with the total duration ≤ 6 months, decided according to assessment of these risks (class of recommendation IIa, level of evidence C) [4].

According to “2018 Joint European consensus document on the management of antithrombotic therapy in atrial fibrillation patients presenting with acute coronary syndrome and/or undergoing percutaneous cardiovascular interventions”, triple therapy can be shortened to one month and then followed by dual therapy with OAC plus clopidogrel up to 12 months in patients with a high bleeding risk (estimated using the HAS-BLED score) [12]. Patients with a very high bleeding risk (e.g. recent bleeding event) can be treated with dual therapy (OAC plus clopidogrel) up to 3–6 months without a one-month triple therapy period. Time frames remain similar for AF patients presenting with elective PCI or ACS undergoing PCI. However, in patients with non-ST-segment elevation acute coronary syndrome (NSTEMI) triple therapy should last 3 to 6 months, while in ST-segment elevation
| Table I. Summary RCTs on NOACs in patients with atrial fibrillation undergoing PCI |
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| **Acronym** | **Primary end point** | **Efficacy end point** | **Study design/Investigated population/compared treatment regimens** | **Results: primary end point** | **Results: efficacy end point** | **Conclusions** |
| AUGUSTUS [20] (apixaban) | ISTH major or clinically relevant non-major bleeding | Death or hospitalization and a composite of ischemic events | • RCT with factorial design and 6-month follow-up, anticoagulation comparison open-label, comparison between aspirin and placebo double-blind • 4614 patients treated with a P2Y12 inhibitor who suffered from ACS or underwent PCI within 14 days (median time from the index event to randomization 6 days) First randomization: • Apixaban 5 mg BID • VKA Second randomization: • Aspirin • Placebo Clopidogrel was used in 92.6% of participants. A total of 229 of 2290 patients (10%) who had been assigned to apixaban received the dose of 2.5 mg BID | Anticoagulation regimens: • Apixaban (N = 2290) – 241 (10.5%) • VKA (N = 2259) – 332 (14.7%) Comparison: p for non-inferiority < 0.001; p for superiority < 0.001 Antiplatelet regimens: • Aspirin (N = 2277) – 367 (6.1%) • Placebo (N = 2279) – 204 (9.0%) Comparison: p for non-inferiority < 0.001; p for superiority < 0.001 | Death or hospitalization Anticoagulation regimen: • Apixaban (N = 2306) – 541 (23.5%) • VKA (N = 2308) – 632 (27.4%) Comparison: p for superiority = 0.002 Antiplatelet regimen: • Aspirin (N = 2307) – 604 (26.2%) • Placebo (N = 2307) – 569 (24.7%) Comparison: p for superiority = NS Death or ischemic event Anticoagulation regimens: • Apixaban (N = 2306) – 154 (6.7%) • VKA (N = 2308) – 163 (7.1%) Comparison: p for superiority = NS Antiplatelet regimen: • Aspirin (N = 2307) – 149 (6.5%) • Placebo (N = 2307) – 168 (7.3%) Comparison: p for superiority = nt | Patients with AF after PCI who were treated with a P2Y12 inhibitor in the apixaban study arm, with no aspirin, were at a lower risk of bleeding and hospitalization than patients receiving VKA, aspirin or both these drugs |
| Acronym (investigated NOAC) | Primary end point | Efficacy end point | Study design/Investigated population/compared treatment regimens | Results: primary end point | Results: efficacy end point | Conclusions |
|-----------------------------|-------------------|--------------------|----------------------------------------------------------------|---------------------------|---------------------------|-------------|
| ENTRUST AF-PCI [21] (edoxaban) | Major or clinically relevant non-major bleeding | Cardiovascular death, stroke, systemic embolic events, myocardial infarction, definite stent thrombosis | • Open-label RCT with parallel-group design and 12-month follow-up 1506 patients who underwent 4 h – 5 days ago successful PCI due to stable CAD or ACS (median time from PCI to randomization 45 h) Study arms:  • Edoxaban 60 mg daily plus a P2Y12 inhibitor (clopidogrel 93%, ticagrelor 7%) for 12 months  • VKA plus a P2Y12 inhibitor (clopidogrel 92%, ticagrelor 8%) and aspirin 100 mg for 1–12 months | • Edoxaban regimen (N = 753) – 128 (17%) • VKA regimen (N = 755) – 152 (20%) Comparison: \( p \) for non-inferiority = 0.001; \( p \) for superiority = NS | • Edoxaban regimen (N = 753) – 49 (7%) • VKA regimen (N = 755) – 46 (6%) Comparison: \( p \) for superiority = NS | Edoxaban-based therapy was non-inferior for bleeding when compared to VKA. No significant differences were found in terms of ischemic events |
| PIONEER-AF [18] (rivaroxaban) | Clinically significant bleeding (a composite of major bleeding or minor bleeding events according to TIMI criteria or bleeding requiring medical attention) | Major adverse cardiovascular event (a composite of death from cardiovascular causes, myocardial infarction, stroke) | • Open-label RCT with parallel-group design and 12-month follow-up 2124 patients who underwent PCI with stent placement due to stable CAD or ACS (randomization occurred within 72 h after sheath removal) Study arms:  • Low-dose rivaroxaban (15 mg OD) plus a P2Y12 inhibitor (clopidogrel 93%, prasugrel 2%, ticagrelor 5%) for 12 months  • Very-low-dose rivaroxaban (2.5 mg BID) plus DAPT (clopidogrel 94%, prasugrel 1%, ticagrelor 5%) for 1, 6 or 12 months  • Triple therapy with VKA (INR 2.0–3.0) plus DAPT (clopidogrel 96%, prasugrel 1%, ticagrelor 3%) for 1, 6 or 12 months | • Low-dose rivaroxaban plus a P2Y12 inhibitor (N = 696) – 109 (16.8%) • Very-low-dose rivaroxaban plus DAPT (N = 706) – 117 (18%) • Triple therapy with VKA plus DAPT (N = 697) – 167 (26.7%) Comparison between the low-dose rivaroxaban group and the triple therapy group: \( p \) for superiority < 0.001 Comparison between the very-low-dose rivaroxaban group and the triple therapy group: \( p \) for superiority < 0.001 | • Low-dose rivaroxaban plus a P2Y12 inhibitor (N = 696) – 41 (6.5%) • Very-low-dose rivaroxaban plus DAPT (N = 706) – 36 (5.6%) • Triple therapy with VKA plus DAPT (N = 697) – 36 (6%) Comparison between the low-dose rivaroxaban group and the triple therapy group: \( p \) for superiority = NS Comparison between the very-low-dose rivaroxaban group and the triple therapy group: \( p \) for superiority = NS | The use of both low-dose rivaroxaban (15 mg) with a P2Y12 inhibitor and very-low-dose rivaroxaban (2.5 mg) with DAPT was associated with a significantly lower risk of bleeding than standard triple therapy with warfarin and DAPT. Efficacy rates were similar for all three groups, but wide confidence intervals reduce the degree of certainty with which conclusions about the efficacy of the treatment can be drawn |
Table I. Cont.

| Acronym (investigated NOAC) | Primary end point | Efficacy end point | Study design/Investigated population/compared treatment regimens | Results: primary end point | Results: efficacy end point | Conclusions |
|-----------------------------|-------------------|-------------------|----------------------------------------------------------------|--------------------------|--------------------------|-------------|
| RE-DUAL AF-PCI [19] (dabigatran) | ISTH major or clinically relevant non-major bleeding | Death, myocardial infarction, systemic embolism, unplanned revascularization | ▪ Open-label RCT with parallel-group design, mean duration of treatment with the trial anticoagulant 12.3 months, mean duration of follow-up 14 months  ▪ 2725 patients who underwent successful PCI due to stable CAD or ACS within the previous 120 h Study arms:  ▪ Dabigatran 110 mg BID plus a P2Y12 inhibitor (clopidogrel, ticagrelor)  ▪ Dabigatran 150 mg BID plus a P2Y12 inhibitor (clopidogrel, ticagrelor)  ▪ Warfarin (INR 2.0–3.0) plus a P2Y12 inhibitor (clopidogrel, ticagrelor) plus aspirin All patients in the United States and nonelderly (< 80 years of age) patients outside the United States were assigned to one of the above listed three arms. Elderly patients outside the United States were assigned to either the smaller dabigatran dose group or the triple therapy with warfarin group Overall use of ticagrelor was 12%. Aspirin was discontinued after 1 and 3 months in patients treated with BMS and DES, respectively | • Dual therapy with dabigatran 110 mg (N = 981) – 151 (15.4%)  ▪ Triple therapy with warfarin (N = 981) – 264 (26.9%)  ▪ Dual therapy with dabigatran 150 mg (N = 763) – 154 (20.2%)  ▪ Corresponding triple therapy with warfarin (N = 764)* – 196 (25.7%) Comparison between dual therapy with dabigatran 110 mg and triple therapy with warfarin: p for non-inferiority < 0.001; p for superiority < 0.001 Comparison between dual therapy with dabigatran 150 mg and corresponding triple therapy with warfarin: p for non-inferiority < 0.001; p for superiority = 0.002 | • Dual therapy with dabigatran 110 mg (N = 981) – 149 (15.2%)  ▪ Triple therapy with warfarin (N = 981) – 131 (13.4%)  ▪ Dual therapy with dabigatran 150 mg (N = 763) – 90 (11.8%)  ▪ Corresponding triple therapy with warfarin (N = 764)* – 98 (12.8%)  ▪ Combined dual therapy groups (N = 1744) – 239 (13.7%) Comparison between combined dual therapy groups and triple therapy with warfarin: p for non-inferiority = 0.005; p for superiority = 0.74 Comparison between dual therapy with dabigatran 110 mg and triple therapy with warfarin: p for superiority = 0.30 Comparison between dual therapy with dabigatran 150 mg and corresponding triple therapy with warfarin: p for superiority = 0.44 | Dual therapy with dabigatran (110 or 150 mg) in combination with clopidogrel had a significantly lower risk of major or CRNM bleeding than triple therapy with warfarin, with a non-inferior efficacy outcome composed of thromboembolic events, death or unplanned revascularization |

*The corresponding triple therapy with warfarin group includes only patients who had been eligible to the 150 mg dual therapy group (i.e. did not include elderly patients outside the United States). ACS – acute coronary syndrome, AF – atrial fibrillation, BID – bis in die (twice daily), BMS – bare-metal stent, CAD – coronary artery disease, CRNM – clinically relevant non-major bleeding, DAPT – dual antiplatelet therapy, DES – drug-eluting stent, INR – international normalized ratio, ISTH – International Society on Thrombosis and Haemostasis, N – number, NOAC – non-vitamin K oral anticoagulant, ns – non-significant, nt – not tested, OD – once daily, PCI – percutaneous coronary intervention, RCT – randomized controlled trial, TIMI – Thrombolysis in Myocardial Infarction, VKA – vitamin K antagonist.
myocardial infarction (STEMI) patients, 6 months [12]. A different point of view is presented in the American consensus [25]. Duration of the triple therapy is recommended only for peri-PCI time, or 1 month in patients at high thrombotic risk and low bleeding risk.

Summarizing, a North American perspective suggests that the triple therapy should be used only during in-hospital stay followed by OAC and clopidogrel for 6 to 12 months depending on the bleeding risk and this should be the way of treatment of the vast majority of all patients treated with PCI, including ACS patients [25]. So the American consensus goes along with the RE-DUAL PCI, PIONEER AF-PCI and AUGUSTUS trials results [18–20]. While the European perspective is still more conservative and suggests that triple therapy should be stopped at discharge, 1 month, or 3 to 6 months depend-

### Table II. The ESC recommendations on antithrombotic therapy in patients with CCS and AF, including post-PCI recommendations [4]

| Recommendation | Class of recommendation | Level of evidence |
|----------------|------------------------|------------------|
| When oral anticoagulation is initiated in a patient with AF who is eligible for a NOAC, a NOAC is recommended in preference to a VKA | I | A |
| Long-term OAC therapy (NOAC or VKA with time in therapeutic range > 70%) is recommended in patients with AF and a CHA₂DS₂-VASc score ≥ 2 in males and ≥ 3 in females | I | A |
| Long-term OAC therapy (NOAC or VKA with time in therapeutic range > 70%) should be considered in patients with AF and a CHA₂DS₂-VASc score of 1 in males and 2 in females | IIa | B |
| Aspirin 75–100 mg daily (or clopidogrel 75 mg daily) may be considered in addition to long-term OAC therapy in patients with AF history of MI, and at high risk of recurrent ischemic events who do not have a high bleeding risk | IIb | B |
| It is recommended that peri-procedural aspirin and clopidogrel are administered to patients undergoing coronary stent implantation | I | C |
| In patients who are eligible for a NOAC, it is recommended that a NOAC (apixaban 5 mg BID, dabigatran 150 mg Bid, edoxaban 60 mg OD, or rivaroxaban 20 mg OD) is used in preference to a VKA in combination with antiplatelet therapy | I | A |
| When rivaroxaban is used and concerns about the high bleeding risk prevail over concerns about stent thrombosis or ischemic stroke, rivaroxaban 15 mg OD should be considered in preference to rivaroxaban 20 mg OD for the duration of concomitant single or dual antiplatelet therapy | IIa | B |
| When dabigatran is used and concerns about the high bleeding risk prevail over concerns about stent thrombosis or ischemic stroke, dabigatran 110 mg BID should be considered in preference to dabigatran 150 mg b.i.d. for the duration of concomitant single or dual antiplatelet therapy | IIa | B |
| After uncomplicated PCI, early cessation (≤ 1 week) of aspirin and continuation of dual therapy with an OAC and clopidogrel should be considered if the risk of stent thrombosis is low, or if concerns about the bleeding risk prevail over concerns about the risk of stent thrombosis irrespective of the type of stent used | IIa | B |
| Triple therapy with aspirin, clopidogrel, and an OAC for ≥ 1 month should be considered when the risk of stent thrombosis outweighs the bleeding risk, with the total duration (≤ 6 months) decided according to assessment of these risks and clearly specified at hospital discharge | IIa | C |
| In patients with an indication for a VKA in combination with aspirin and/or clopidogrel, the dose intensity of the VKA should be carefully regulated with a target international normalized ratio in the range of 2.0–2.5 and with time in therapeutic range > 70% | IIa | B |
| Dual therapy with an OAC and either ticagrelor or prasugrel may be considered as an alternative to triple therapy with an OAC, aspirin, and clopidogrel in patients with a moderate or high risk of stent thrombosis irrespective of the type of stent used | IIb | C |
| The use of ticagrelor or prasugrel is not recommended as part of triple antithrombotic therapy with aspirin and an OAC | III | C |
| Concomitant use of a proton pump inhibitor is recommended in patients receiving aspirin monotherapy, DAPT, or OAC monotherapy who are at high risk of gastrointestinal bleeding | I | A |
In the long-term treatment, aspirin 75–100 mg daily (or clopidogrel 75 mg daily) may be considered in addition to OAC therapy in patients with AF, history of myocardial infarction, and at high risk of recurrent ischemic events. This strategy is recommended in patients who do not have a high bleeding risk (class of recommendation IIb, level of evidence B). Dual therapy with an OAC and either ticagrelor or prasugrel may be considered as an alternative to triple therapy with an OAC, aspirin, and clopidogrel in patients with a moderate or high risk of stent thrombosis, irrespective of the type of stent used (class of recommendation IIb, level of evidence C). The use of ticagrelor or prasugrel is not recommended as part of triple antithrombotic therapy with aspirin and an OAC (class III, level of evidence C) because of the high bleeding risk [4].

In Figure 1, we propose an algorithm which combines both European and American perspectives.

### Gaps in the evidence

There are several gaps in the evidence that warrant exploration. Firstly, optimal duration of periprocedural antithrombotic therapy with aspirin in PCI-treated AF patients remains controversial and the risk of ischemic coronary events (e.g. stent thrombosis) related to early aspirin cessation cannot be neglected. Secondly, optimal management of patients on NOAC treatment who require urgent or emergency PCI needs to be investigated. Finally, safety and efficacy of dual therapy with a NOAC and ticagrelor or prasugrel should be studied.

**Antithrombotic management**

**OAC:**
- A NOAC should be preferred in most patients and used at established stroke prevention doses; lower doses are not recommended unless specifically tested
- If VKA is chosen, maintain INR at the lower end of the therapeutic range (e.g. 2.0–2.5)
- Maintain OAC life-long

**APT:**
- Aspirin in the peri-PCI phase and continued through hospital discharge
- Clopidogrel is the P2Y12 inhibitor of choice; ticagrelor may be a reasonable alternative in high ischemic and low bleeding risk patients; avoid prasugrel
- Discontinue SAPT at 1 year in most patients; consider earlier SAPT discontinuation (e.g. 6 months) in patients at low ischemic or high bleeding risks and prolonging SAPT (> 1 year) for select patients with high ischemic and low bleeding risks

**Strategy (double vs. triple therapy):**
- A double-therapy regimen (OAC plus P2Y12 inhibitor) early after hospital discharge for most patients
- Consider triple-therapy by extending aspirin use for a limited period of time (e.g. 3–6 months) only in patients at high ischemic and low bleeding risks

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**Figure 1.** Algorithm for the management of patients with atrial fibrillation requiring oral anticoagulation undergoing PCI. Modified based on Angiolillo et al. [25]

*APT – antiplatelet therapy, DES – drug-eluting stent, FFR – fractional flow reserve, INR – international normalized ratio, IVUS – intravascular ultrasound, NOAC – non-vitamin K antagonist oral anticoagulant, NSAID – nonsteroidal anti-inflammatory drug, OCT – optical coherence tomography, PCI – percutaneous coronary intervention, PPI – proton pump inhibitor, SAPT – single antiplatelet therapy, VKA – vitamin K antagonist.*
Conclusions

NOACs at standard doses are the preferred option in most of post PCI patients with AF, except those with a clear indication for VKAs (i.e. patients with prosthetic heart valve or with mitral stenosis) [27]. Reduced NOAC doses should be considered in dabigatran- or rivaroxaban-treated patients with a high bleeding risk, which prevail over concerns about stent thrombosis or ischemic stroke. There is insufficient evidence to favor one NOAC over another in this setting. In the early post stenting period, triple therapy comprising a NOAC, clopidogrel and aspirin is recommended. Timing of post PCI aspirin cessation should be based on a careful analysis of the bleeding and ischemic risk. Depending on the result, all very early (≤ 1 week), intermediate (at 1–3 months) and relatively late (at 6 months) aspirin discontinuation may be considered. There is only low quality evidence regarding the optimal approach to elective or urgent/emergency PCI procedures in patients requiring OAC. Undoubtedly, all AF patients undergoing coronary stenting should be treated with a new generation DES. It is suggested that there is no need of interruption of VKA and PCI procedure should be performed via radial artery access with a lower dose of unfractionated heparin. On the other hand, NOACs are stopped before elective PCIs, while emergency procedures may be performed with the addition of low-dose parenteral anticoagulation.

Conflict of interest

Marek Koziński received speaker fees from Bayer, Boehringer-Ingelheim and Pfizer. Jolanta Maria Siller-Matula received speaker fees from Bayer, Bristol Myers Squibb and Daiichi Sankyo. Anna Tomaszuk-Kazberuk received speaker and consultation fees from Boehringer-Ingelheim and consultation fee from Bayer. Other authors disclosed that they have no significant relationships with or financial interests in any commercial companies related to this review.

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