**Guideline**

2017 consensus of the Asia Pacific Heart Rhythm Society on stroke prevention in atrial fibrillation

Chern-En Chiang, MD, PhD\(^a\), Ken Okumura, MD, PhD\(^b\), Shu Zhang, MD\(^c\), Tze-Fan Chao, MD, PhD\(^d,e\), Chung-Wah Siu, MD\(^f\), Toon Wei Lim, MD\(^g\), Anil Saxena, MD\(^h\), Yoshihide Takahashi, MD\(^i\), Wee Siong Teo, MD\(^j\)

\(^a\) General Clinical Research Center and Division of Cardiology, Taipei Veterans General Hospital and National Yang-Ming University, Taipei, Taiwan
\(^b\) Division of Cardiology, Saiseikai Kumamoto Hospital Cardiovascular Center, Kumamoto, Japan
\(^c\) State Key Laboratory of Cardiovascular Disease, Fuwai Hospital; National Center for Cardiovascular Diseases, Chinese Academy of Medical Sciences and Peking Union Medical College, Beijing, People’s Republic of China
\(^d\) Division of Cardiology, Department of Medicine, Taipei Veterans General Hospital, Taipei, Taiwan
\(^e\) Institute of Clinical Medicine, and Cardiovascular Research Center, National Yang-Ming University, Taipei, Taiwan
\(^f\) Cardiology Division, Department of Medicine, The University of Hong Kong, Hong Kong, China
\(^g\) National University Heart Centre, National University Hospital, Singapore
\(^h\) Department of Cardiovascular Medicine, Tokyo Medical and Dental University, Tokyo, Japan
\(^i\) Department of Cardiovascular Medicine, Fortis Escorts Heart Institute, New Delhi, India
\(^j\) National Heart Centre, Singapore

**Article info**

Article history:
Received 3 April 2017
Received in revised form 29 April 2017
Accepted 16 May 2017
Available online 27 June 2017

Keywords:
Anticoagulation
Atrial fibrillation
Non-vitamin K antagonist oral anticoagulants
Vitamin K antagonist
Stroke

**Abstract**

Atrial fibrillation (AF) is the most common sustained arrhythmia, causing a 2-fold increase in mortality and a 5-fold increase in stroke. The Asian population is rapidly aging, and in 2050, the estimated population with AF will reach 72 million, of whom 2.9 million may suffer from AF-associated stroke. Therefore, stroke prevention in AF is an urgent issue in Asia. Many innovative advances in the management of AF-associated stroke have emerged recently, including new scoring systems for predicting stroke and bleeding risks, the development of non-vitamin K antagonist oral anticoagulants (NOACs), knowledge of their special benefits in Asians, and new techniques. The Asia Pacific Heart Rhythm Society (APHRS) aimed to update the available information, and appointed the Practice Guideline sub-committee to write a consensus statement regarding stroke prevention in AF. The Practice Guidelines sub-committee members comprehensively reviewed updated information on stroke prevention in AF, emphasizing data on NOACs from the Asia Pacific region, and summarized them in this 2017 Consensus of the Asia Pacific Heart Rhythm Society on Stroke Prevention in AF. This consensus includes details of the updated recommendations, along with their background and rationale, focusing on data from the Asia Pacific region. We hope this consensus can be a practical tool for cardiologists, neurologists, geriatricians, and general practitioners in this region. We fully realize that there are gaps, unaddressed questions, and many areas of uncertainty and debate in the current knowledge of AF, and the physician’s decision remains the most important factor in the management of AF.

© 2017 Japanese Heart Rhythm Society. Published by Elsevier B.V. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

**1. Epidemiology of atrial fibrillation-associated stroke in Asia**

Stroke and systemic thromboembolism are the most clinically important complications observed in patients with atrial fibrillation (AF) [1]. Stroke caused by AF is defined as cardioembolic stroke, and once it occurs, it often results in death (up to 20%) or...
disability (approximately 60%) [2–4]. Therefore, appropriate thromboprophylaxis is mandatory.

Cardioembolic stroke occurs most commonly in the elderly, especially the oldest-old AF patients [5,6]. The Asian population is rapidly aging, and in 2050, the estimated population with AF will reach 72 million, of whom 2.9 million may suffer from AF-associated stroke [7]. Thus, stroke prevention in AF is an urgent healthcare and public-health concern in Asia.

The incidence of AF-associated stroke has been extensively investigated worldwide. Overall, the incidence of stroke in patients with AF is 4–5-fold higher than that in patients without AF [8,9]. Importantly, the incidence varies significantly with patients’ clinical characteristics and risk factors, the more common ones being included in risk scores such as the CHA2DS2-VASc (Congestive heart failure, Hypertension, Age ≥ 75 [doubled], Diabetes, Stroke [doubled]-Vascular disease, Age 65–74, Sex category [female]) score [10]. Other risk factors may include reduced renal function or chronic kidney disease (CKD) [11,12], and low body weight [13].

In addition, the incidence of stroke is also significantly affected by the use of oral anticoagulation therapy (OAC) and the quality of anticoagulation control. The overall annual incidences of ischemic stroke reported in Asia ranged from 1.3% in Japanese AF patients (n = 3588; mean age, 68.1 ± 13.5 years; mean CHA2DS2-VASc score, 2.4) from three prospective registries (Shinken Database, J-RHYTHM Registry, and Fushimi AF Registry) [14] to 10.4% in hospitalized Chinese AF patients (n = 3333; mean age, 79.5 ± 9.2 years; mean CHA2DS2-VASc score, 3.8) from the Queen Mary Hospital, Hong Kong [15]. Of the 186,570 AF patients without OAC selected from the National Health Insurance Research Database (NHIRD) in Taiwan, 23,723 (12.7%) experienced ischemic stroke during the follow-up period of 3.4 years (3.7%/year) [16]. Even higher incidences of AF-associated stroke, ranging from 13.0% to 15.4% for 1-to-3-year periods, were reported in the Far East and Southeast Asia [17]. The reported incidence of AF-associated stroke varies substantially due to clinical setting (hospitalized versus community), unrecorded use of OAC at follow-up, different methods of analysis, and diverse clinical characteristics of AF patients in different regions.

Is the prevalence of AF-associated stroke in Asia higher or lower when compared with that reported from Western countries? The annual incidence of ischemic stroke in non-anticoagulated AF patients in the United States was 2.1%, with the incidence increasing from 0.57% in patients < 65 years, to 1.41% in those between 65 and 74 years, to 2.58% in those between 75 and 84 years, and even further to 4.42% in those > 85 years of age [18]. Although no direct comparisons have been made in non-anticoagulated AF patients, recent global clinical trials may offer some insights [19–22]. For example, a sub-analysis of the RE-LY trial [23] comparing rates of ischemic and hemorrhagic stroke events between Asians and non-Asians demonstrated the absolute rate of ischemic stroke was numerically higher in Asians than in non-Asians in all treatment groups (2.05%/year versus 1.14%/year in the dabigatran 110 mg group, 1.12%/year versus 0.81%/year in the dabigatran 150 mg group, and 2.02%/year versus 0.98%/year in the warfarin group) [23]. The rates of hemorrhagic stroke in the warfarin group were significantly higher in Asians than in non-Asians (hazard ratio, 2.4; 95% confidence interval, 1.3–4.7) [23]. Similarly, the sub-analysis of the ROCKET AF trial [24] comparing event rates between East Asians (not including Japan) and non-East-Asians revealed the absolute rate of ischemic stroke was numerically higher in East Asia compared to non-East Asia (2.24/100 patient-years versus 1.60/100 patient-years in the warfarin group), and the rate of hemorrhagic stroke similarly higher in East Asia (1.24/100 patient-years versus 0.39/100 patient-years) [24]. The sub-analysis of the ENGAGE AF trial [25] comparing East Asians and non-East-Asians revealed the absolute rates of ischemic stroke were numerically higher in East Asia than in non-East Asia (1.31/100 patient-years versus 0.89/100 patient-years in the warfarin group), with the rate of hemorrhagic stroke also higher in East Asia (1.23/100 patient-years versus 0.41/100 patient-years) [25]. Similar trends were also demonstrated in the ARISTOTLE trial [26]. Thus, Asian AF patients are more prone to suffer from ischemic stroke compared with non-Asians, even with anticoagulation. In addition, Asian patients are more prone to hemorrhagic stroke, as previously reported [27].

2. Stroke risk scores

2.1. CHADS2 versus CHA2DS2-VASc scores

The risk of AF-associated stroke is not homogeneous and depends on patients’ ages and comorbidities, which have been used to formulate clinical scores to aid risk stratification. The CHADS2 (congestive heart failure, hypertension, age ≥ 75 [doubled], diabetes mellitus, and prior stroke or transient ischemic attack [TIA]) score has been commonly used to guide antithrombotic therapies for AF patients since its original validation in 2001 (Table 1) [28]. In 2010, the CHA2DS2-VASc was developed [10], and has been confirmed to be superior to the CHADS2 score in identifying truly low-risk patients (Table 1) [29–31]. The CHA2DS2-VASc score is recommended by the European Society of Cardiology (ESC) [32], American College of Cardiology/American Heart Association (ACC/AHA) [33], and the National Institute for Health and Care Excellence (NICE) for stroke risk stratification in AF (http://guidance.nice.org.uk/CG/Wave0/638).

The diagnostic accuracy of CHADS2 and CHA2DS2-VASc scores was compared among 186,570 AF patients in Taiwan, who did not receive anti-platelet agents or OAC [34]. The CHA2DS2-VASc score outperformed CHADS2 score in predicting ischemic stroke. More importantly, the stroke risk in patients with a CHADS2 score of 0 was not low; the annual stroke rate ranged from 1.15% (CHA2DS2-VASc score = 0) to 4.47% (CHA2DS2-VASc score = 3). This is consistent with data from the Danish nationwide cohort study, where patients with a CHADS2 score of 0 had a stroke rate as high as 3.2%/year when further stratified by the CHA2DS2-VASc score [35]. The annual risk of ischemic stroke for Asian AF patients stratified by CHADS2 and CHA2DS2-VASc scores is shown in Table 2. Additionally, the CHA2DS2-VASc score has been demonstrated to be better than the ATRIA (anticoagulation and risk factors in atrial fibrillation) score for the prediction of ischemic stroke for Asian AF patients [16]. Based on the current evidence in Asians,

Table 1: Calculations of the CHADS2 and CHA2DS2-VASc score.

|               | CHADS2 | CHA2DS2-VASc |
|---------------|--------|--------------|
| Congestive heart failure | 1 | 1 |
| Hypertension | 1 | 1 |
| Age ≥ 75 y | 1 | 2 |
| Diabetes mellitus | 1 | 1 |
| Previous Stroke/TIA | 2 | 2 |
| Vascular disease (prior MI, PAD, or aortic plaque) | – | 1 |
| Age 65–74 y | – | 1 |
| Sex category (i.e., female sex) | – | 1 |
| Maximum score | 6 | 9 |

CHADS2 congestive heart failure, hypertension, age ≥ 75 [doubled], diabetes mellitus, and prior stroke or transient ischemic attack; CHA2DS2-VASc congestive heart failure, hypertension, age ≥ 75 [doubled], diabetes, stroke [doubled]-vascular disease, age 65–74, sex category [female]; MI, myocardial infarction; PAD, peripheral artery disease; TIA, transient ischemic attack.
use of the CHA2DS2-VASc score is recommended for stroke risk stratification in Asian AF patients.

2.2. Should Asian AF patients with one stroke risk factor be treated?

In a recent registry study from Taiwan which enrolled 12,935 AF males with a CHA2DS2-VASc score of 1 and 7900 AF females with a CHA2DS2-VASc score of 2 (i.e., one non-sex stroke risk factor) [36], AF males with a CHA2DS2-VASc score of 1 had an annual stroke rate ranging between 1.96% and 3.50%, depending on the specific covariate composing the score. For AF female patients with one additional stroke risk factor (CHA2DS2-VASc score of 2), the annual stroke rate ranged from 1.91% to 3.34%. The annual risk of ischemic stroke for these patients, left untreated, exceeds the treatment threshold for the initiation of OAC (1.7%/year for warfarin and 0.9%/year for non-vitamin K antagonist oral anticoagulants [NOACs]) [37]. Therefore, we recommend that OAC should be considered for Asian AF patients with one additional risk factor beyond sex: i.e., CHA2DS2-VASc score of 1 for males and 2 for females. This recommendation is similar to the 2016 ESC AF guidelines [38].

Given that the CHA2DS2-VASc score is best at identifying “low-risk” patients, and the benefits of stroke prevention are evident with ≥ 1 non-sex stroke risk factors, the initial step should be to identify low-risk patients (i.e., CHA2DS2-VASc score 0 in males, 1 in females) who do not need antithrombotic therapy – rather than focus on identifying high-risk patients. Thus, the default should be to offer stroke prevention (i.e., OAC) to all patients with AF, unless they can be categorized as “low-risk.”

2.3. Do Asian AF patients have a lower age threshold for stroke?

The risk of ischemic stroke for Asian AF patients is higher than that of non-Asians [39]. Previous studies of NOACs showed that Asian AF patients treated with NOACs had a higher risk of ischemic stroke than non-Asians, despite similar CHADS2 and CHA2DS2-VASc scores [7]. Although the detailed mechanism(s) behind this remained unknown, a recent study from Taiwan has demonstrated that the risk of ischemic stroke may start to rise from age 50 upwards [40]. For these Chinese patients aged 50–64 years, the annual stroke risk was 1.78%, which exceeds the treatment threshold for OAC use for stroke prevention [40]. A similar age threshold (i.e., 50 years) for an increased risk of ischemic stroke was also observed in a study from Hong Kong [41].

A modified CHA2DS2-VASc score, mCHA2DS2-VASc, which assigned one point for patients aged 50–74 years, outperformed CHA2DS2-VASc score for stroke risk stratification for Chinese AF patients, with a higher C-index (0.71 versus 0.69, DeLong test P < 0.0001) and an improved net reclassification index [42]. Most importantly, for patients with a CHA2DS2-VASc score of 0 (males) or 1 (females), having a mCHA2DS2-VASc score of 1 (males) or 2 (females) due to the resetting of the age threshold, the use of warfarin was associated with a positive net clinical benefit when balancing the benefit of ischemic stroke reduction against the risk of intracranial hemorrhage (ICH) [42]. Whether the mCHA2DS2-VASc score could be used to guide stroke-prevention strategies for Asian AF patients needs to be confirmed via further studies.

3. Bleeding risk assessment

The stroke risk reduction with OAC should be balanced against the increased risk of bleeding, especially ICH. Several scoring systems have been proposed to estimate the risk of bleeding in AF, such as the HEMORRHAGES, HAS-BLED, ATRIA, ORBIT, and ABC-bleeding scores [43–47]. Bleeding scores have been subject to inappropriate use – they should be used to “flag up” the patients at risk for bleeding for more regular review and follow-up, and importantly, to address reversible bleeding-risk factors [48].

The HAS-BLED (Hypertension, Abnormal renal/liver function, Stroke, Bleeding history or predisposition, Labile International Normalized Ratio (INR), Elderly, Drugs/alcohol concomitantly) score has been proposed as a simple clinical score to predict clinically relevant bleeding in AF patients (Table 3). A HAS-BLED score ≥ 3 indicates a high risk of bleeding, and previous studies have demonstrated that the HAS-BLED score performed better than other bleeding scores [49,50]. In warfarin users, HAS-BLED would significantly outperform the ATRIA and ORBIT scores that do not consider “labile INR” as a risk factor [51,52]. Also, the HAS-BLED score has been validated in AF patients on no antithrombotic therapy, aspirin, warfarin and non-warfarin anticoagulants (and thus, is applicable to every step of the AF patient treatment pathway), as well as being validated in Asian AF patients [53]. A high HAS-BLED score should not be used to exclude patients from OAC therapy but allows clinicians to address the correctable risk factors for bleeding, such as uncontrolled hypertension, labile INRs (for a warfarin user) and concomitant use of aspirin, NSAIDs or alcohol excess/abuse.

| CHADS2 score | Incidence (per 100 person-years) |
|--------------|---------------------------------|
| 0            | 1.80                            |
| 1            | 3.08                            |
| 2            | 4.49                            |
| 3            | 5.33                            |
| 4            | 4.86                            |
| 5            | 5.80                            |
| 6            | 7.10                            |

| CHA2DS2-VASc score | Incidence (per 100 person-years) |
|-------------------|---------------------------------|
| 0                 | 1.15                            |
| 1                 | 2.11                            |
| 2                 | 3.39                            |
| 3                 | 3.89                            |
| 4                 | 4.61                            |
| 5                 | 5.12                            |
| 6                 | 5.18                            |
| 7                 | 6.22                            |
| 8                 | 7.98                            |
| 9                 | 10.50                           |

CHA2DS2, congestive heart failure, hypertension, age ≥ 75 [doubled], diabetes mellitus, and prior stroke or transient ischemic attack; CHA2DS2-VASc, congestive heart failure, hypertension, age ≥ 75 [doubled], diabetes, stroke [doubled]-vascular disease, age 65–74, sex category [female].
Table 3
Calculation of the HAS-BLED score.

| Clinical characteristics | Definition | Score |
|--------------------------|------------|-------|
| Hypertension             | SBP > 160 mmHg | 1 |
| Abnormal renal and liver function (1 score each) | Renal: dialysis, transplantation, or creatinine ≥ 2.3 mg/dL; Liver: chronic hepatits, cirrhosis, bilirubin > 2 ULN, with ALT > 3 ULN | 1 or 2 |
| Stroke                   | Previous history, particularly lacunar | 1 |
| Bleeding tendency or predisposition | Recent bleed, anemia, etc. | 1 |
| Labile INRs              | Unstable/high INR, or TTR < 60% | 1 |
| Elderly                  | Age > 65 y, extreme frailty | 1 |
| Drugs or alcohol (1 score each) | Drugs: concomitant antiplatelet, or NSAID use; Alcohol excess | 1 or 2 |

Maximum score: 9

ALT: alanine transaminase; Cr: creatinine; INR: international normalized ratio; NSAID: non-steroidal anti-inflammatory drugs; TTR: time in therapeutic range; ULN: upper limit of normal

Recommendations

- The HAS-BLED score is recommended for the prediction of bleeding risk in Asian patients with non-valvular AF.
- A HAS-BLED score ≥ 3 suggests a high risk of bleeding, but does not preclude the use of OAT. Such patients should have regular review and follow-up of the modifiable bleeding-risk factors (uncontrolled hypertension, labile INRs [for a warfarin user] and concomitant use of aspirin, NSAIDs or alcohol excess/abuse).

4. Role of aspirin

There is no evidence for the effectiveness of aspirin in stroke prevention in AF in Asia. In a Japanese trial, aspirin was no better than placebo in low-risk patients [54]. In a recent Hong Kong cohort study, aspirin showed a non-significant reduction in ischemic strokes, compared with no therapy [55]. OAC is more effective than aspirin for stroke prevention in AF, and the net clinical benefit is positive for OAC versus no treatment or aspirin, but neutral or negative for aspirin versus no antithrombotic therapy, even with a single stroke risk factor [55–58].

The risks of ischemic stroke and ICH in a real-world cohort of Chinese AF patients were reported recently from Hong Kong [15]. The incidence of ischemic stroke on aspirin was higher than that on dabigatran (110 mg) (7.95%/year vs 2.24%/year). The incidence of ICH was lower in dabigatran (110 mg) users than in those on aspirin (0.32%/year vs 0.80%/year) [15]. In the AVERROES trial, the risk of stroke was significantly lower in the apixaban group than in aspirin group (relative risk reduction 45%, P < 0.001), with a similar risk of major bleeds [59]. The risk of ICH was numerically lower in the apixaban group [59]. The totality of these data suggest that there is no role for aspirin in stroke prevention in Asians.

Nonetheless, the use of aspirin is highly prevalent in many Asian countries [60,61]. In the Registry on cardiac rhythm disORDers (RecordAF-Asia Pacific [AP]) registry, a prospective observational study of the management of patients with recently diagnosed AF in eight Asian-Pacific countries, aspirin was more commonly used than VKAs (56–66% vs. 35–47%) [62]. A recent study using Taiwan’s NHIRD between 2001 and 2008 showed that the percentage of AF patients who received warfarin, aspirin, or no treatment in Taiwan was 16%, 62% and 22%, respectively [63]. In Phase I of the Global Registry on Long-Term Oral Antithrombotic Treatment in Patients with Atrial Fibrillation (GLORIA-AF) registry, 49.6% of Chinese AF patients received aspirin alone [64]. This is reaffirmed in recent real-life data from Hong Kong, where 61% of patients received aspirin [65].

This continued use of aspirin may be partly explained by the misconception that aspirin is associated with a lower risk of bleeding. In addition, there are several issues specific to the Asia-Pacific region. First, INR control is generally poor in the Asia-Pacific region [66]. In a recent study from Hong-Kong, for example, the median time in therapeutic range (TTR) was 38.8% [15]. This may be due to limited access to anticoagulation clinics or to the interaction of VKAs with food or herbal drugs, which are commonly used in this region. Second, Asians treated with VKAs are at higher risk of ICH [7,67], which may discourage physicians from prescribing VKAs. Third, the financial burden of using NOACs needs to be considered. Because there are no data showing benefit of aspirin, this consensus statement does not recommend the use of aspirin solely for stroke prevention in AF patients.

Recommendations

- Aspirin is not recommended solely for stroke prevention in AF.

5. Role of vitamin K antagonists (VKAs)

VKAs have been the mainstay of treatment for stroke prevention in AF for more than half a century. In a meta-analysis of 6 randomized control trials (RCTs) involving 2900 patients with non-valvular AF, VKA therapy with a target INR between 2.0 and 3.0 reduced the risk of stroke by 64% and mortality by 26%, compared with placebo or no therapy [68]. Although VKA therapy doubles the risk of ICH, the absolute risk increases by only 0.2%/year.

Despite this, VKA therapy has long been grossly underutilized in Asia-Pacific countries, with a utilization rate typically around 15–20% [69–71]. In addition to various limitations related to the narrow therapeutic window and the wide assortment of drug-food interactions, the higher baseline risk of ICH [72–74] and poorer TTR [75,76] observed in Asian populations both undermine the benefits of VKA therapy.

Recently, the NOACs, including dabigatran, rivaroxaban, apixaban, and edoxaban have been shown in large RCTs to be at least as effective as VKA in stroke prevention, but with a consistently lower risk of ICH [19–22]. NOACs are regarded as the preferred agents for stroke prevention in non-valvular AF [32,33,39], and have been used extensively over the past few years. However, VKAs would remain a viable option in certain clinical scenarios.

5.1. Role of VKAs in chronic kidney disease

AF and CKD commonly co-exist. The prevalence of AF increases with reduced glomerular filtration rate (GFR) [12,77–80]. Patients with CKD can be categorized according to GFR (Table 4). While long-term OAC can effectively reduce ischemic stroke risk in

Table 4
Staging of chronic kidney disease.

| CKD stage | GFR level (ml/min/1.73 m²) |
|-----------|--------------------------|
| Stage 1   | ≥ 90                     |
| Stage 2   | 60–89                    |
| Stage 3   | 30–59                    |
| Stage 4   | 15–29                    |
| Stage 5   | < 15                     |

CKD: chronic kidney disease; GFR: glomerular filtration rate.
general AF patients, whether this can be extended to severe CKD patients remains inconclusive. Among patients with AF and a GFR of 30 mL/min or above, i.e., mild to moderate renal impairment or Stage 1 to 3 CKD, both VKA [81] and all 4 of the NOACs have been shown to be as effective in stroke reduction as in the general AF population [32,33,58]. AF patients with GFR < 25–30 mL/min were excluded from all pivotal NOAC trials.

Previous observational studies of AF patients with end-stage CKD on hemodialysis therapy have reported conflicting results on the net clinical benefits [82] and harms [83–86]. For example, the Dialysis Outcomes and Practice Patterns Study (DOPPS), an international, observational study of hemodialysis practices and outcomes, demonstrated that hemodialysis patients with AF receiving VKA therapy had higher stroke risks compared to non-VKA users [87,88]. In a North American study consisting of 1671 hemodialysis patients with AF, VKA therapy was associated with a nearly 2-fold increase in stroke [83]. Furthermore, the risk of hemorrhagic stroke among hemodialysis patients with AF on VKA also increases substantially [84,89].

As a result, clinical guidelines from Kidney Disease: Improving Global Outcomes (KDIGO) no longer recommend warfarin therapy for stroke prevention in AF among dialysis patients [90]. It is plausible that frequent heparinization during hemodialysis, reduced levels of protein C, protein S and antithrombin III [91–95], as well as the fluctuations in blood pressure in hemodialysis patients with AF might diminish the overall benefit of warfarin due to a higher thrombotic and bleeding risk. Nonetheless, good-quality anticoagulation control with high TTR may mitigate risks [96].

In contrast to developed countries, peritoneal dialysis instead of hemodialysis is the primary mode of renal replacement therapy in many Asian countries or regions. Despite the paucity of clinical data, VKA therapy appears to have a net clinical benefit in terms of ischemic stroke, ICH, and mortality among AF patients on peritoneal dialysis [97,98]. Until further RCTs to evaluate the net clinical benefit of OAC in dialysis patients with AF become available, the choice of long-term OAC should be highly individualized.

5.2. Role of VKAs in valvular heart disease

The efficacy of VKA therapy in stroke prevention in patients with underlying valvular heart diseases, particularly chronic rheumatic heart disease and prosthetic heart valves, has long been established. Although in the pivotal studies of NOACs, patients with moderate or severe mitral stenosis or mechanical heart-valve prosthesis were excluded, patients with other mitral or aortic valvular disease were allowed to participate [99–102]. The safety and efficacy of NOACs do not appear to be different with respect to the valvular status of individual patients, including those with bioprosthetic valves [103]. In a recent meta-analysis of 4 phase III AF trials comprising 13,585 patients with and 58,098 without valvular heart diseases, high-dose NOACs provided similar efficacy and safety [104].

To date, there has been only one randomized controlled study comparing warfarin and NOAC in patients with mechanical heart-valve prosthesis, which was prematurely terminated because of excessive strokes and bleeding with dabigatran [105]. Mechanistically, mechanical heart-valve prosthesis induces sufficient thrombin generation via the intrinsic pathway, overwhelming the clinically relevant concentration of dabigatran [106].

Taken collectively, VKAs remain the only drugs for patients with moderate or severe mitral stenosis and patients with mechanical heart-valve prosthesis.

5.3. Identifying patients likely to do well on VKA with good anticoagulation control: the SAMe-TT2R2 score

Anticoagulation control with warfarin is influenced by many demographic and clinical factors. The more common factors have been used to formulate the SAMe-TT2R2 (Sex [female], Age [less than 60], Medical history [more than two comorbidities], Treatment [interacting medications, e.g., amiodarone], Tobacco use [doubled], Race [doubled]) score, which may help in predicting whether a patient is likely to have a good anticoagulation control if VKA is used [107,108]. A SAMe-TT2R2 score 0–2 predicts a good response to VKA (i.e., high TTR > 65%), while a SAMe-TT2R2 score > 2 suggests that the patient is less likely to achieve a good TTR on VKA, thus flagging up the patient for additional education/counseling or more regular INR checks and clinic reviews, or for use of a NOAC [107,108].

The SAMe-TT2R2 score has been validated in two Asian populations where a score of 0–2 predicted a TTR ≥ 70% (Hong Kong cohort) or a TTR ≥ 60% (Singapore cohort), and a score of ≥ 3 predicted a TTR < 70% (Hong Kong cohort) or a TTR < 60% (Singapore cohort) [109,110]. Therefore, the SAMe-TT2R2 score can be used to help predict the performance of VKA in Asians by identifying those patients likely to achieve good or poor TTR, and to assist in decision-making in the selection of OAC (i.e., VKA or NOAC).

Recommendations

- VKAs have a role in stroke prevention in patients with stage 4, and possibly stage 5, CKD.
- VKA remains the only drug for stroke prevention in patients with moderate or severe mitral stenosis and those with mechanical-valve prosthesis.
- The SAMe-TT2R2 score may help predict whether a patient is likely to have good anticoagulation control if a VKA is used. A SAMe-TT2R2 score 0–2 predicts a good response to VKA, while a SAMe-TT2R2 score > 2 flags up the patient for additional education/counseling or more regular INR checks and clinic reviews, or use of a NOAC.

6. Role of non-vitamin K antagonist oral anticoagulants (NOACs)

NOACs have revolutionized the approach to stroke prevention in AF [111,112]. There are now four NOACs available: one oral direct thrombin inhibitor (dabigatran); and three oral factor Xa inhibitors (rivaroxaban, apixaban, and edoxaban). The pharmacokinetic data of the four NOACs are shown in Table 5 [32,113–115].

Compared with warfarin, all NOACs have more predictable pharmacokinetics and fewer drug-drug interactions, allowing fixed dosing without the need for regular monitoring of anticoagulation status [116].

6.1. Major randomized clinical trials

The efficacy and safety of the four NOACs have been tested in four major RCTs: the RE-LY trial, the ROCKET AF trial, the ARISTOTLE trial, and the ENGAGE AF trial [19–22]. The primary efficacy endpoints were stroke plus systemic embolization events (SEEs). In general, NOACs showed non-inferiority in primary efficacy endpoints when compared with dose-adjusted warfarin with target INR of 2.0–3.0, except dabigatran 150 mg and apixaban 5 mg, which showed superiority to warfarin. Most NOACs showed a decreased risk of major bleeding compared with warfarin, except dabigatran 150 mg and rivaroxaban 20 mg. A pre-specified meta-analysis comprising these 4 major RCTs of NOACs also
demonstrated a favorable risk-benefit profile, with significant reductions in stroke, ICH, and mortality, and with similar major bleeding as for warfarin, but increased gastrointestinal bleeding [112].

One should be careful to integrate this information into patient care in Asia, as Asians are prone to bleeding with warfarin use. Therefore, a more detailed examination of the subsets of Asians from these RCTs is important.

6.2. Asian sub-analyses of major RCTs

Among 71,783 participants in the four major RCTs of NOACs [19–22], 7650 patients were from Asia, mostly East Asian countries. The Asian sub-analyses of all these RCTs have been published [23,24,26,117]. The efficacy endpoints (stroke/SEE, ischemic stroke, hemorrhagic stroke, myocardial infarction, all-cause mortality, and cardiovascular mortality) and safety endpoints (major bleeding, ICH, gastrointestinal bleeding, and bleeding due to any cause) of NOACs versus warfarin in the 4 RCTs are summarized in Table 6 [58]. These data suggest great advantages of using NOACs for stroke prevention in AF patients in Asia.

6.3. Meta-analysis of NOACs in Asia

In a recent meta-analysis, the differences in efficacy and safety outcomes of NOACs in Asian patients were compared with those in non-Asian patients [118]. The 5 RCTs included the studies RE-LY, ROCKET AF, J-ROCKET AF, ARISTOTLE, and ENGAGE AF, comprising 8928 Asian patients (5250 with NOACs and 3678 with VKAs) and 64,033 non-Asian patients (37,800 with NOACs and 26,233 with VKAs) [19–26,119,120]. There were 2 separate analyses: a meta-analysis for standard-dose NOACs (dabigatran 150 mg, edoxaban 60 mg, rivaroxaban 20 mg, and apixaban 5 mg); and a meta-analysis for low-dose NOACs (dabigatran 110 mg, edoxaban 30 mg, and rivaroxaban 15 mg) (Table 7) [118].

Standard-dose NOACs significantly reduced stroke/SEE in both Asian and non-Asian patients, and the effect size of this reduction was greater in Asian patients than in non-Asians (P interaction=0.045) (Table 7). All-cause mortality was also reduced in Asian and non-Asian patients, but heterogeneity was not significant (P interaction=0.219) [118]. In Asians, standard-dose NOACs significantly reduced major bleeding, ICH, and hemorrhagic stroke, compared with warfarin. There was no increase in GI bleeding. In non-Asians, standard-dose NOACs significantly reduced ICH and hemorrhagic stroke, with an increase in GI bleeding, compared with warfarin. Heterogeneity was evident in major bleeding, hemorrhagic stroke, and GI bleeding among Asians versus non-Asians [118].

The efficacy of low-dose NOACs in stroke/SEE, ischemic stroke, myocardial infarction and all-cause mortality, was generally similar to that of warfarin in both Asians and non-Asians, except for an increase in myocardial infarction and a decrease in all-cause mortality among NOAC users in non-Asians (Table 7). No significant heterogeneity could be found between Asians and non-Asians. The main benefits of low-dose NOACs were in the safety aspects, because major bleeding, ICH, and hemorrhagic stroke significantly reduced in Asians while ICH and hemorrhagic stroke significantly reduced in non-Asians, compared with warfarin. Both Asians and non-Asians showed a numerically lower risk of GI bleeding.

Table 5
Pharmacokinetic characteristics of NOACs.

| Drug       | Absorption with food | Intake with food recommended | Renal clearance | Bioavailability | CYP metabolism | Transporter | Hours to Cmax | Half-life, hours |
|------------|----------------------|-----------------------------|----------------|----------------|----------------|-------------|---------------|-----------------|
| Dabigatran | No effect            | No                          | 80%            | 6%             | None           | P-glycoprotein | 3             | 12–17           |
| Rivaroxaban| +39%                 | Mandatory                   | 35%            | 80%            | 66%            | P-glycoprotein | 2–4           | 5–13            |
| Apixaban   | No effect            | No                          | 27%            | 60%            | 15%            | P-glycoprotein | 3             | 9–14            |
| Edoxaban   | +6(–22%)             | No                          | 50%            | 62%            | <4%            | P-glycoprotein | 1–2           | 10–14           |

Cmax: maximal concentration; CYP: cytochrome P 450; NOACs: non-vitamin K antagonist oral anticoagulants.

Modified from Camm et al.,[32] Heidbuchel et al.,[113] Eriksson et al.,[114] and Lip et al.[115]

Table 6
Efficacy and safety endpoints of different NOACs in Asians [23–26,119].

| NOAC | Stroke/SEE | Ischemic stroke | Hemorrhagic stroke | Myocardial infarction | All-cause death | CV death | Major bleeding | Intracranial hemorrhage | GI bleeding | Bleeding due to any cause |
|------|------------|-----------------|-------------------|----------------------|-----------------|---------|----------------|-------------------------|-------------|---------------------------|
|      |            |                 |                   |                      |                 |         |                |                         |             |                           |
| Dabigatran | V    | V               | V                 | NR                   | V               | V       | V              | V                       | V           |                           |
| 150 mg     | V    | V               | V                 | NR                   | V               | V       | V              | V                       | V           |                           |
| Dabigatran | V    | V               | V                 | NR                   | V               | V       | V              | V                       | V           |                           |
| 110 mg     | V    | V               | V                 | NR                   | V               | V       | V              | V                       | V           |                           |
| Rivaroxaban | V    | V               | V                 | NR                   | V               | V       | V              | V                       | V           |                           |
| 60 mg      | V    | V               | V                 | NR                   | V               | V       | V              | V                       | V           |                           |
| Edoxaban   | V    | V               | V                 | NR                   | V               | V       | V              | V                       | V           |                           |
| 30 mg      | V    | V               | V                 | NR                   | V               | V       | V              | V                       | V           |                           |

CV: cardiovascular; GI: gastrointestinal; NOACs: non-vitamin K antagonist oral anticoagulants; NR: not reported; SEE: systemic embolization events; V: P value less than 0.05 when compared with warfarin.

Modified from Lip et al.[58] with permission

* China, Japan, South Korea, Taiwan, Hong Kong, Philippines, Singapore, Malaysia, Thailand, India.
* China, South Korea, Taiwan, Hong Kong.
+ China, Japan, South Korea, Taiwan, Hong Kong, Philippines, Singapore, Malaysia.
+ China, Japan, South Korea, Taiwan.
bleeding, compared with warfarin. No heterogeneity was shown between Asians and non-Asians [118].

In another meta-analysis of 3155 Asian patients with NOACs in the RE-LY and ENGAGE AF trials, efficacy and safety with standard-dose versus low-dose NOACs were compared [121]. Risks of stroke/SEE and ischemic stroke significantly reduced with standard-dose versus low-dose NOACs (RR 0.62, 95% CI 0.45–0.85; and RR 0.55, 95% CI 0.38–0.79, respectively). Rates of major, intracranial, and life-threatening bleeding with the two dosing regimens were broadly similar (RR 1.31, 95% CI 0.74–2.33; RR 1.54, 95% CI 0.72–3.30; and RR 1.49, 95% CI 0.87–2.55, respectively). Therefore, standard-dose NOACs represent a more appealing therapeutic option than low-dose NOACs in Asians, with a significant reduction in ischemic stroke without an excess of major bleeding [121]. Nevertheless, label- or guideline-adherent NOAC dosing offers a balance of efficacy and safety outcomes, compared to warfarin [122].

**Recommendations**

- For Asian patients with non-valvular AF, standard-dose NOACs (dabigatran 150 mg bid, rivaroxaban 20 mg od, apixaban 5 mg bid, or edoxaban 60 mg od) are the default doses of choice for stroke prevention unless label guidance recommends low-dose regimens as follows:
  - For dabigatran, the 110mg bid dose is recommended in the elderly (age > 75), in patients with a high bleeding risk (HAS-BLED ≥ 3) or in patients receiving interacting drugs (e.g. verapamil).
  - For rivaroxaban, the 15 mg od dose is recommended where the Cockcroft-Gault creatinine clearance (CrCl) is 30–49 mL/min.
  - For apixaban, 2.5 mg is used in patients with two or more of the following criteria: age ≥ 80 years, body weight ≤ 60 kg, or serum creatinine ≥ 1.5 mg/dL.
  - For edoxaban, the 30-mg od dose is recommended in patients with any one of the following criteria: eGFR of 30 to 50 mL/min, a body weight ≤ 60 kg, or the concomitant use of verapamil or quinidine (potent P-glycoprotein [P-gp] inhibitors).
  - Low-dose NOACs (rivaroxaban 15 mg od, apixaban 2.5 mg bid, edoxaban 30 mg od) should be used with caution in severe CKD (CrCl 15–30 mL/min) according to drug labels.

- Dabigatran should not be used in patients with CrCl < 30 mL/min.

### 7. Role of left atrial appendage closure and excision

#### 7.1. Rationale and techniques for LAA closure

The left atrial appendage (LAA) is thought to be the most important site of thrombus formation leading to ischemic stroke in AF patients. Oclusion or excision of the LAA has been proposed as a way to reduce thromboembolic events [123]. However, not all strokes in AF patients are cardio-embolic or due to AF, and the LAA is probably not the only left atrial region where thrombi can potentially originate. Even after removal or closure of the LAA, antithrombotic therapy may still be needed [124].

Surgical excision or ligation of the LAA is often performed as a concomitant procedure during open heart surgery. Less invasive techniques using epicardial or trans-septal approaches have been developed to occlude the LAA [125–127]. These techniques are aimed at providing an alternative for AF patients at high risk for stroke but with contraindications for chronic OAC use.

Devices for trans-septal LAA occlusion include the WATCHMAN device (Boston Scientific, Natick, MA, USA) and the Amplatzer Cardiac Plug (St. Jude Medical, Minneapolis, MN, USA). The WATCHMAN device is deployed percutaneously via a transseptal puncture and has a permeable, polyester fabric membrane that covers a self-expanding nitinol cage with barbs to anchor the device in the LAA [128]. The Amplatzer Cardiac Plug consists of a proximal disc and a distal lobe with hooks to anchor the device in the LAA. It does not require anticoagulation [127].

An alternative strategy is to tie off the LAA using an epicardial suture device, referred to as the LARIAT device (SentreHEART, Redwood City, CA, USA) [129]. More recently, the US FDA alerted against its off-label use for LAA closure, as its safety and effectiveness for this indication has not been established. (http://www.fda.gov/MedicalDevices/Safety/AlertsandNotices/ucm454501.htm; date last accessed, June 26, 2016).

#### 7.2. Percutaneous closure of LAA

The most data exist for the WATCHMAN device and early studies suggest non-inferiority to warfarin for the composite
predicted by their CHADS2 scores (1.7% versus 7.3%) [128]. This was because the WATCHMAN device implanted had a lower rate of stroke andSEE, which are more limited. In the ASAP registry study, patients with a higher risk of stroke or SEE from AF who have significant contraindications for any OAC therapy.

For patients with a contraindication for warfarin, the data are more limited. In the ASAP registry study, patients with a WATCHMAN device implanted had a lower rate of stroke andSEE, and it was lower than would be expected for the level of risk as predicted by their CHADS2 scores (1.7% versus 7.3%) [128]. This was at the expense of an 8.7% rate of serious procedural or device-related complications, and in comparison with historical CHADS2 score event rates.

Less data are available for the Amplatzer devices, with earlier studies mostly being retrospective, nonrandomized case series that also included devices that were not dedicated to LAA occlusion [127,133,134]. In the larger series, procedure-related complications occurred in around 5% of patients, but the lack of a control group precludes any comparisons with pharmacological treatment.

While the use of percutaneous approaches for LAA closure is feasible, there are still sparse data for most. The exception is the WATCHMAN device, for which there is some trial evidence of its non-inferiority (and even superiority in one trial) to warfarin. This does come at the cost of peri-procedural and device-related complications and significant financial outlay. Of note, none of the trials included significant numbers of Asian patients, and thus its utility in our patient population is uncertain. Nonetheless, it may be reasonable to consider the use of this device in patients at high risk of stroke orSEE from AF who have significant contraindications for any OAC therapy.

### 7.3. Surgical occlusion or excision of LAA

There is no conclusive evidence that surgical LAA excision or occlusion reduces stroke risk in AF patients [124]. Retrospective or observational studies in different patient populations have shown inconsistent results for surgical LAA excision or occlusion [135]. This could be partly due to low rates of successful closure using current techniques, though this appears to improve with experience [136,137]. A large randomized trial of left atrial ligation in patients undergoing cardiac surgery (LAAOS III) is underway, and may provide some evidence for surgical closure of the LAA [138].

Randomized studies to date have been small and did not show any benefit of LAA occlusion at the time of coronary-artery bypass surgery in patients at risk of stroke. In the LAAOS study, 52 out of 77 patients were randomized to receive LAA occlusion; 2 of them suffered a perioperative thromboembolic event [137]. After 13 ± 7 months, no additional patients had stroke. Similarly, the LAAOS II study, which was conducted to assess the feasibility of a larger trial (i.e., LAAOS III) also found no significant reduction in embolic events in the LAA-occlusion arm but concluded that concomitant LAA occlusion was safe and feasible [139]. On follow-up, surgical excision resulted in a higher rate of LAA closure (73%) than either suture occlusion (23%) or stapler occlusion (0%) [136].

Surgical techniques and devices for occluding the LAA are still being developed. The most widely used device, the AtriClip (Atricure, West Chester, OH, USA), consists of 2 parallel, straight, rigid titanium tubes and 2 nitinol springs with a knit-braided polyester fabric [140]. Early non-randomized studies of its use during open heart surgery have demonstrated high success rates and good short-term durability (98.4%) based on imaging studies [141].

Surgical LAA closure may be conceptually reasonable, but there is a paucity of evidence for its efficacy and safety. In the absence of stronger clinical data, especially in the form of RCTs, recommendations can only be based on expert consensus. Importantly, there is no evidence for its use in place of OAC for patients who could otherwise be treated with the latter. It does appear safe in patients who are undergoing open heart surgery for other indications and may be considered in these patients.

### Recommendations

- Interventional percutaneous LAA closure with the WATCHMAN device may be considered in patients with non-valvular AF who have high risk of stroke, but major contraindications to OAC therapy.
- Surgical excision of the LAA may be considered in patients undergoing concomitant cardiac surgery.

### 8. Practical use of NOACs

#### 8.1. Drug-drug and drug-food interaction

The use of VKAs is complicated by their unpredictable and variable performance, which is due to several factors. These include numerous drug-food and drug-drug interactions [142–144]. Many herbs used as foods, and also medicinal supplements, can interact with VKAs [145]. There is evidence that
bleeding rates are higher on warfarin and NOACs seem preferable for stroke prevention in Asians [7,146]. One of the main advantages of NOACs has been a predictable onset and offset of action, and fewer drug-drug interactions as compared to warfarin [143]. As NOAC usage is increasing, various drug-drug interactions are coming to light (Table 8) [144,147].

The action of NOACs can be influenced at various stages in their absorption, metabolism, and elimination. These drugs have variable renal excretion, hepatic metabolism, and re-secretion into the gut via a P-gp transporter. The cytochrome P450 (CYP 450) enzyme system is responsible for hepatic clearance of NOACs. All these elimination pathways could be points of interaction with food or drugs.

Dabigatran is predominantly (80%) eliminated by renal excretion, and not affected by CYP 450 enzyme modulators. Clinicians prescribing anticoagulants should have accurate knowledge of various modes of elimination of each drug and their possible modification by various drugs. When a patient on NOACs develops a thrombotic or bleeding complication, co-medications should be carefully reviewed for a possibility of drug-drug interactions.

There is a significant re-secretion of NOACs into the intestine via the transporter P-gp, which may be involved to some extent in renal excretion [148]. Inhibitors of P-gp may therefore result in higher plasma levels and consequently, increased anticoagulant activity [149]. Common P-gp inhibitors used in AF patients include verapamil, drosdronarone, amiodarone, and quinidine.

Rivaroxaban and apixaban are mainly metabolized by CYP3A4 [150]. Any concomitant medication that modulates CYP3A4 may therefore affect plasma concentrations and effects, and should be evaluated [151].

8.1.1. Interaction with rate- and rhythm-control drugs

Patients with AF requiring anticoagulation are frequently co-administered various rate- or rhythm-controlling drugs. Several of these agents can interact with anticoagulants (Table 8). In particular, amiodarone and verapamil have been shown to increase the bioavailability of dabigatran [152]. In one study, a single 120-mg dose of verapamil 1 h before dabigatran increased its plasma concentration (AUC) and peak serum concentration (Cmax) by 143% and 179% respectively [153]. The effect was minimized if verapamil was given 2 h before dabigatran [153].

8.1.2. Interaction with antifungals and antibiotics

Antifungal agents like ketoconazole, itraconazole, and posaconazole are very strong inhibitors of P-gp, and are contraindicated for use with NOACs (Table 8) [150]. Among antifungals, fluconazole was found to have least effect on rivaroxaban and can be used with caution [150]. One study found a 2-fold increase in apixaban exposure with co-administration of ketoconazole [154].

Clarithromycin has been shown to increase the bioavailability of dabigatran from 6.5% to 10.1%, while the Cmax and AUC increased by 60.2% and 49.1%, respectively [155]. Being a strong inhibitor of both CYP3A4 and P-gp, clarithromycin can also increase plasma levels of rivaroxaban and lead to bleeding [156]. Erythromycin may significantly increase the activity of rivaroxaban by about 34% [150]. Ritonavir, an antiretroviral agent, increased the activity of rivaroxaban by 158% [150]. Rifampin has been shown to reduce exposure to edoxaban while increasing exposure to its active metabolites M4 and M6 without significantly changing aPTT [157].

8.1.3. Interaction with miscellaneous drugs

Several antiepileptic drugs including carbamazepine, phenytoin, valproic acid, levetiracetam, and topiramate are inducers of P-gp, and may affect the anticoagulant activity of NOACs. Naproxen has been shown to increase the activity of apixaban [158], and can increase bleeding time in patients on rivaroxaban [159] and edoxaban [160]. Co-administration of pantoprazole caused moderate reduction in dabigatran absorption [161]. Antacids have no effect on apixaban, rivaroxaban, and edoxaban, while they may reduce bioavailability of dabigatran by 12–28% without affecting its efficacy [152].

8.1.4. Interaction with foods

One of the main factors for unpopularity of VKAs has been their strong predilection for interaction with various foods rich in vitamin K. Unlike VKAs, NOACs are not significantly affected by food intake, with the exception of rivaroxaban, which has better absorption and near 100% bioavailability when taken with food. Therefore, it is strongly recommended that rivaroxaban is taken with food at dinner time. Apixaban and edoxaban are not affected by food intake [115,162].

**Recommendations**

- NOACs should not be combined with potent P-gp inducers, such as rifampin, carbamazepine, phenobarbital, and phenytoin.
- NOACs should not be combined with potent P-gp inhibitors, such as HIV protease inhibitors, itraconazole, and ketoconazole. The only exception is edoxaban: a 50% reduction of edoxaban dose can be combined with itraconazole and ketoconazole.
- When combined with amiodarone, a lower dose NOAC is considered in the elderly.
- A 50% reduction in dabigatran dose is needed when combined with verapamil; while data for other NOACs are scarce.
- Dabigatran should not be combined with drosdronarone.
- A 50% reduction in the edoxaban dose is needed when combined with drosdronarone, while data for rivaroxaban and apixaban are lacking.

8.2. How to switch?

Switching between oral anticoagulants is often required in the clinical setting. In both the ROCKET AF and ARISTOTLE trials, the risk of stroke increased after the end of the trials in patients in whom the study NOAC was switched to open warfarin [163,164]. This was explained by a delay in achieving a therapeutic INR after switching. The timing of the interruption and initiation of the drugs must be chosen to minimize any gap in the therapeutic anticoagulation status.

When a VKA is switched to a NOAC, the NOAC should be started once the INR is approximately 2.0 or less. When a NOAC is switched to a VKA, VKA should be combined with parenteral heparin or combined with a NOAC until the INR approaches 2.0. Since factor Xa inhibitors affect the INR, the INR should be measured 24 h after the last NOAC intake.

Recently, the Dresden NOAC registry from Germany reported the risk associated with switching from a VKA to a NOAC [165]. At 30 days of follow-up from switching, major cardiovascular events and major bleeding events occurred in 0.8% and 0.3%, respectively. In general, switching between oral anticoagulants can be performed safely with careful monitoring of the INR.

**Recommendations**

- When a VKA is switched to a NOAC, the NOAC should be started once the INR is approximately 2.0 or less.
- When a NOAC is switched to a VKA, VKA should be combined with parenteral heparin or combined with a NOAC until the INR is approximately 2.0.
8.3. Patients with chronic kidney disease

The prevalence of CKD, similar to that of AF, is increasing worldwide, including in the Asia-Pacific (AP) region. The staging of CKD is shown in Table 4. All individuals with a GFR < 60 mL/min for 3 months are classified as having CKD, irrespective of the presence or absence of kidney damage. End-stage renal disease (ESRD) is defined as a CrCl of less than 15 mL/minute/1.73 m².

CKD can affect up to 10% of the adult population, especially elderly people, and carries a high risk for AF. Up to 30% of patients with AF have some renal dysfunction; hence, it is important to screen for renal dysfunction in AF patients.

Higher incidence of AF has been reported even in patients with early renal dysfunction [78,166]. The prevalence of AF in patients with impaired kidney function or on dialysis is considerably higher than in the general population, with estimates that about one in 5–6 patients on hemodialysis has AF [80,87,167]. In an Asian study, the incidence rates of AF were 12.1, 7.3, and 5.0 per 1000 person-years for ESRD, CKD, and control patients, respectively [168]. Among patients with ESRD, age, hypertension, heart failure, coronary artery disease, peripheral arterial occlusive disease, and chronic obstructive pulmonary disease were significant risk factors for new-onset AF.

Further, AF and CKD have an unhealthy relationship: AF predisposes people to CKD and CKD increases the risk of AF [77,169]. CKD in adult patients with incident AF is independently associated with increased risk of developing ESRD [170].

The estimated CrCl (eCrCl) and the estimated GFR (eGFR) measure slightly different things. In the context of NOAC treatment, CrCl is best assessed by the Cockcroft-Gault (CG) method [171], as this method was used in the NOAC trials.

8.3.1. Stroke and bleeding in patients with CKD

CKD increases stroke risk in patients with AF, as well as the risk of major bleeds [172]. Renal impairment (CrCl < 60 mL/min) doubles the risk of stroke [173]. It has even been proposed to add renal function to the CHADS2 scoring system as a manner to improve prediction of stroke [11,174]. In patients on NOAC, impaired renal function was shown to be an independent risk factor for stroke in a sub-study of the ROCKET AF trial [11]. However, in other “real-world” studies, renal impairment was not an independent predictor of ischemic stroke or thromboembolism in AF and did not significantly improve the predictive ability of the CHADS2 or CHA2DS2-VASc scores [173,175].

In a Japanese study, as renal function declined below an eGFR of 60 mL/min, stroke risk increased regardless of whether AF was also present. The hazard ratios for stroke were 1.9 and 3.1 in patients with eGFR of 40–70 mL/min and < 40 mL/min, respectively, as compared to those with an eGFR > 70 mL/min [176].

The risk of ischemic stroke has also been reported to be higher in patients with CKD and ESRD [87,172,177]. Hemodialysis patients typically have multiple comorbidities, and AF would be expected to increase the risk of ischemic stroke at least as much as in patients without renal failure. New-onset AF in hemodialysis patients adversely affected the outcomes in a retrospective cohort study obtained from the Taiwanese NHIRD [178]. Compared to the control group, the patients with new-onset AF had higher risks of ischemic stroke (HR, 1.27), all-cause mortality (HR, 1.59), in-hospital cardiovascular death (HR, 1.83), myocardial infarction (HR, 1.33), and heart failure (HR, 1.9). After adjustment for in-hospital deaths, AF was associated with a higher risk of heart failure (HR, 1.56) and in-hospital cardiovascular death (HR, 1.65), but not stroke or myocardial infarction [178].

Renal dysfunction is a risk marker not only for stroke but also for death, myocardial infarction, and bleeding. Renal impairment (CrCl < 60 mL/min) especially ESRD, increases the risk of major bleeding by almost 60% in anticoagulated patients with AF [173]. In addition, warfarin may also promote vascular calcification in the CKD patient.

8.3.2. VKA in CKD patients

The extent to which CKD increases the risk of thromboembolism in patients with nonvalvular AF and the benefits of anticoagulation in this group remain unclear. Only warfarin was associated with a decreased risk of stroke or SEE among patients with CKD, whereas both warfarin and aspirin were associated with an increased risk of bleeding [172]. High-risk patients with AF (CHA2DS2-VASc ≥ 2) and renal failure still derive a net benefit from anticoagulation with warfarin, especially if quality of anticoagulation is good [96,179]. In a meta-analysis, the presence of CKD in patients with AF was associated with a 50% increase in risk of thromboembolism, which can be effectively decreased with appropriate antithrombotic therapy [180].

8.3.2.1. Patients with non-ESRD. There is evidence from certain AF studies suggesting the use of OAC in patients with mild to moderate CKD provides similar, or potentially even greater, benefit compared to its use in the general population. In a post hoc analysis of patients with CKD stage III in the Stroke Prevention in Atrial Fibrillation III trial, warfarin use markedly decreased ischemic stroke and/or systemic venous thromboembolism (VTE) by 76% (95% CI, 42–90; P < 0.001) compared to aspirin plus low, fixed doses of warfarin [81]. In patients with both CKD and AF one year after discharge from an acute myocardial infarction in the SWEDEHEART registry, Carrero et al. showed that warfarin was associated with lower risk of the composite endpoint including death, repeat myocardial infarction, or ischemic stroke [181]. Bleeding risk did not increase. The lower event rate for the primary outcome was observed across all strata of eGFR and primarily driven by mortality events, while the risk of bleeding was not significantly higher in patients treated with warfarin in any CKD stratum [181].

8.3.2.2. Patients with ESRD on dialysis. AF patients with severe renal impairment or on dialysis have been excluded from large RCTs evaluating antithrombotic therapy in AF. Therefore, the optimal approach to anticoagulation in patients with non-valvular AF who have severe renal disease or are on dialysis is controversial. Although warfarin is indicated to prevent ischemic strokes in most patients with AF, evidence supporting its use in hemodialysis patients is limited.

The systematic use of any oral anticoagulant or acetylsalicylic acid has not been demonstrated to be beneficial for AF patients who are hemodialysis-dependent. Several observational studies have raised concerns about the use of warfarin in dialysis patients with non-valvar AF [84,87,182]. Warfarin use in patients with AF and CKD was not associated with significant reductions in stroke risk or mortality in patients with AF on chronic hemodialysis, but might actually have contributed to greater bleeding risk. In a population-based, retrospective, cohort study of 1626 AF patients on dialysis from Montreal, Canada, no reduction in stroke risk was found with warfarin, even after adjusting for multiple factors (HR, 1.14; 95% CI, 0.78–1.67), but a significantly higher risk of bleeding on warfarin (HR, 1.44; 95% CI, 1.13–1.85) was observed [89]. Similar findings were reported by Winkelmayer et al. [84] In the Dialysis Outcomes and Practice Patterns Study (DOPPS), Wizemann et al. demonstrated that warfarin use in AF patients > 75 years of age (n=1107) was associated with a 2.2-fold higher risk for the composite stroke/death outcome, but in the two groups under age of 75, no difference with warfarin use was observed [87].

There is a possibility that the benefit of warfarin in these patients may be outweighed by its risks, and therefore, RCTs are
needed [183,184]. The current Canadian AF guidelines do not advocate any oral anticoagulation for stroke prevention and a high bleeding risk with warfarin [185,186]. In Europe, many centers routinely anticoagulate these patients with warfarin, but aiming for a TTR > 65–70%. Indeed, Olesen et al. demonstrated a favorable analysis for the use of warfarin in 901 patients with ESRD in a large observational study from the National Danish Registry from 1997–2008 evaluating patients with AF at hospital discharge; this showed a 56% reduction in risk for the composite stroke/death outcome compared to antithrombotic therapy (HR 0.44; 95% CI, 0.26–0.74) [172]. The use of warfarin in ESRD patients must thus be individualized, weighing the risks versus the benefits.

One meta-analysis on the use of warfarin for AF showed that it may have an unfavorable risk/benefit ratio in patients with ESRD but not in those with non-ESRD. Thirteen publications from 11 cohorts (6 retrospective and 5 prospective), including > 48,500 total patients with > 11,600 warfarin users, were included in the meta-analysis [187]. In patients with AF and non-ESRD, warfarin resulted in a lower risk of ischemic stroke/thromboembolism (HR, 0.70; 95% CI, 0.54–0.89; P = .004) and mortality (HR, 0.65; 95% CI, 0.59–0.72; P < .00001), but had no effect on major bleeding (HR, 1.15; 95% CI, 0.88–1.49; P = .31). In patients with AF and ESRD, warfarin had no effect on the risks of stroke (HR, 1.12; 95% CI, 0.69–1.82; P = .65) or mortality (HR, 0.96; 95% CI, 0.81–1.13; P = .60), but increased the risks of major bleeding (HR, 1.30; 95% CI, 1.08–1.56; P = .005) [187]. As mentioned, this would be highly dependent on the quality of anticoagulation control, and the net clinical benefit may still be positive where the TTR is > 70%.

8.3.3. NOACs

NOACs have a more stable dose response than warfarin but are dependent on renal clearance. Used correctly, NOACs are at least as safe as well-controlled warfarin. Similar to warfarin, they can cause bleeding, especially if used in excessive doses, in patients at higher risk for bleeds, and in patients with reduced kidney function.

8.3.3.1. Renal clearance of NOACs. Clearance of NOACs from the body is dependent on renal function, which should be assessed regularly (Table 5). Dabigatran is an oral direct thrombin inhibitor that is 80% renally cleared, and thus has a potential to cause more bleeding in patients with reduced renal function. Of the oral factor Xa drugs, rivaroxaban has 35% renal clearance while apixaban has 27% renal clearance and edoxaban has 50% renal excretion. Apixaban has the lowest renal clearance and is potentially safer in patients with renal impairment. The renal clearance of warfarin is < 1% and hence it may be safest pharmacokinetically for patients with severe CKD [186].

8.3.3.2. NOACs in patients with mild-to-moderate CKD. Subgroup analyses of RE-LY, ARISTOTLE, ROCKET AF, and ENGAGE AF have demonstrated that all four NOACs produced comparable results in the primary efficacy endpoints (stroke and SEE) and the primary safety endpoint (major bleeding) across different stages of renal function (Table 9) [188–191]. The only exception was found in the ARISTOTLE trial [189]. Apixaban seems to produce less bleeding compared to warfarin in patients with an eGFR ≤ 50 mL/min than in those with a higher eGFR (P for interaction 0.030).

| Subgroup | RE-LY | ARISTOTLE | ENGAGE AF |
|----------|-------|-----------|-----------|
| eGFR (mL/min) | CrCl (mL/min) | eGFR (mL/min) | CrCl (mL/min) |
| < 50 | ≥ 50 | < 50 | ≥ 50 |
| 50 to < 80 | 50 to < 80 | 50 to < 80 | 50 to < 80 |
| ≥ 80 | ≥ 80 | ≥ 80 | ≥ 80 |
| P (int) | P (int) | P (int) | P (int) |

Table 9

Efficacy and safety of NOACs in patients with CKD.

| Subgroup | RE-LY | ARISTOTLE | ENGAGE AF |
|----------|-------|-----------|-----------|
| eGFR (mL/min) | CrCl (mL/min) | eGFR (mL/min) | CrCl (mL/min) |
| < 50 | ≥ 50 | < 50 | ≥ 50 |
| 50 to < 80 | 50 to < 80 | 50 to < 80 | 50 to < 80 |
| ≥ 80 | ≥ 80 | ≥ 80 | ≥ 80 |
| P (int) | P (int) | P (int) | P (int) |

Apixaban seems to produce less bleeding compared to warfarin in patients with an eGFR ≤ 50 mL/min than in those with a higher eGFR (P for interaction 0.030).

8.3.3.3. NOACs in patients with mild-to-moderate CKD. Subgroup analyses of RE-LY, ARISTOTLE, ROCKET AF, and ENGAGE AF have demonstrated that all four NOACs produced comparable results in the primary efficacy endpoints (stroke and SEE) and the primary safety endpoint (major bleeding) across different stages of renal function (Table 9) [188–191]. The only exception was found in the ARISTOTLE trial [189]. Apixaban seems to produce less bleeding compared to warfarin in patients with an eGFR ≤ 50 mL/min than in those with a higher eGFR (P for interaction 0.030).
8.3.3.3. Meta-analysis of NOAC in CKD. There have been several meta-analyses addressing the efficacy of NOACs in relation to VKAs in patients with mild or moderate CKD [192,193]. The data are very consistent across studies, showing that in patients with mild or moderate CKD (eGFR 30–79 mL/min), all NOACs are associated with decreased rates of thromboembolism compared with warfarin. Among patients with mild CKD (defined as an eGFR of between 50 and 79 mL/min), major bleeding was also significantly reduced in patients receiving NOACs; however, in patients with moderate CKD (eGFR 30–49 mL/min) the overall bleeding rate was similar to that of warfarin [192,193].

8.3.3.4. Can we use NOACs in patients with CrCl < 30 mL/min? The major RCTs of NOACs have excluded patients with CrCl < 30 mL/min, except for an apixaban trial, which excluded patients with CrCl < 25 mL/min. European and American dosing recommendations state that apixaban and rivaroxaban can be administered to patients with an eCrCl > 15 mL/min. However, the evidence for this recommendation comes from pharmacokinetic studies in a limited number of patients. Because of limited experience with NOACs at this level of renal dysfunction, the Canadian guidelines have recommended that VKAs are generally the preferred agent for patients with an eCrCl of 15–30 mL/min [185].

The US Food and Drug Administration has approved dabigatran 75 mg bid for patients with an eGFR of 15–30 mL/min, and apixaban (5.0 mg bid) for ESRD with hemodialysis, based on their pharmacokinetic and pharmacodynamic modelling profiles. Dabigatran is considered a less-than-ideal choice because of its risk of increasing bleeding when CrCl drops below 50 mL/min. Drug levels may substantially fluctuate with dialysis treatment, particularly because dialysis clears 50% to 60% of the drug [194]. We do not recommend the use of NOACs in patients with ESRD with an eCrCl < 15 mL/min or in patients on dialysis, until there are clinical data to confirm their safety and efficacy [186].

**Recommendations**

- In the context of NOAC treatment, CrCl is best assessed by the Cockcroft-Gault method, as this was used in the NOAC trials.
- For patients with moderate CKD, i.e., eCrCl 30–49 mL/min, NOACs are preferred over VKA in stroke prevention in Asians, due to a lower risk of ICH.
- Standard-dose NOACs should not be used in patients with severe CKD, i.e., eCrCl < 30 mL/min (< 25 mL/min for apixaban).
- Low-dose NOACs (rivaroxaban 15 mg od, apixaban 2.5 mg bid, edoxaban 30 mg od) should be used with caution in severe CKD (CrCl 15–30 mL/min) according to drug labels.
- Dabigatran should not be used in patients with CrCl < 30 mL/min.
- In patients with ESRD or dialysis, NOACs are contraindicated. Although VKA with good-quality anticoagulation control (TTR > 70%) might be useful, the data are lacking.

8.4. Patients with coronary heart disease

Patients with AF may have concurrent coronary heart disease (CHD), either in a stable form or with acute coronary syndrome (ACS). The use of OAC and dual antiplatelet therapy (DAPT) may increase the bleeding risk. The use of glycoprotein IIb/IIIa inhibitors is not recommended due to a potentially higher risk of bleeding. In stabilized patients, OAC can be restarted after percutaneous anticoagulation is stopped. It is reasonable to restart the NOAC that the patient was taking before the ACS or elective procedure [113]. The same principle applies for AF patients after coronary bypass grafting. The use of ticagrelor or prasugrel as part of the triple therapy regimen is not recommended, given that their bleeding risk when associated with NOACs is unknown [195]. (Triple therapy stands for low-dose aspirin (75–100 mg/d), clopidogrel 75 mg/d, and an OAC; dual therapy means clopidogrel 75 mg/d and an OAC.)

For long-term management of AF patients after revascularization and/or ACS, a management algorithm has been suggested to reduce the risk of bleeding while protecting against coronary events (Fig. 1). The bleeding risk can be defined by the HAS-BLED score [44]. Those with a HAS-BLED score of 0–2 have a low bleeding risk; while a HAS-BLED score ≥ 3 suggests a high bleeding risk. There is no indication that the advantages of NOACs over VKAs are not preserved in AF patients with CHD, especially in Asians [39,113].

For all stable CHD patients with AF, the rule of thumb is to use anticoagulation as monotherapy and to discontinue any antiplatelet agents at 1 year after patient presentation with ACS, except for those with a very high risk of coronary events and an acceptably low bleeding risk [39,113].

In the recent PIONEER AF-PCI trial, 2124 participants with non-valvular AF who had undergone PCI with stenting were randomly assigned to receive low-dose rivaroxaban (15 mg od) plus a P2Y12 inhibitor for 12 months (Group 1), very-low-dose rivaroxaban (2.5 mg bid) plus DAPT for 1, 6, or 12 months (Group 2), or standard therapy with a dose-adjusted VKA (once daily) plus DAPT for 1, 6, or 12 months (Group 3) [196]. The primary safety outcome was clinically significant bleeding (a composite of major bleeding or minor bleeding according to Thrombolysis in Myocardial Infarction [TIMI] criteria, or bleeding requiring medical attention).

![Flow chart for the long-term management of patients with atrial fibrillation and acute coronary syndrome/percutaneous intervention. ACS, acute coronary syndrome; M, month; OAC, oral anticoagulant; PCI, percutaneous intervention; Y, year.](image-url)
The study did not have enough power to examine the difference in major CV events (death from cardiovascular causes, myocardial infarction, or stroke) [197]. The rates of clinically significant bleeding were lower in the two groups receiving rivaroxaban than in the group receiving standard therapy, but there was no significant difference in major bleeding [196]. Compared with Group 3, the rates of cardiovascular death were 29% higher in Group 1 and 19% higher in Group 2, though these differences did not reach significance. One should interpret this trial cautiously in that the rates of ischemic stroke were numerically higher in Group 1 and Group 2, compared to Group 3 (Group 1 versus Group 3, HR 3.28, CI 0.68–15.78; Group 2 versus Group 3, HR 2.87, CI 0.58–14.23) [196], though these results, again, did not reach statistical significance. It is still too early to use only single antiplatelet therapy plus OAC to replace the conventional triple-therapy regimen (i.e., DAPT plus OAC) in this clinical setting (Fig. 1). Again, NOACs are preferred over VKAs in Asians [58], and no data suggest that one NOAC is better than another [39].

**Recommendations**

- The use of glycoprotein IIb/IIIa inhibitors is not recommended due to a potentially higher risk of bleeding.
- The use of ticagrelor or prasugrel as part of the triple-therapy regimen is not recommended, given that their bleeding risk when used with NOACs is unknown.
- For patients receiving elective PCI, triple therapy should be used for 1 month, followed by dual therapy up to 1 year (or up to 6 months for patients with high bleeding risk).
- For patients with ACS, triple therapy should be used for 6 months, and followed by dual therapy up to 1 year (or for 1 month, followed by dual therapy up to 1 year in patients with high bleeding risk).
- For all stable CHD patients with AF, the recommendation is to use anticoagulation as monotherapy and to discontinue any antiplatelet agents at 1 year after presentation with ACS, except for those with a very high risk of coronary events and an acceptably low bleeding risk.

8.5. Patients with a history of stroke

Prior stroke or transient ischemic attack (TIA) (history of stroke/TIA) is a powerful independent predictor of subsequent stroke, with a relative risk between 2.2 and 2.5 [198,199]. In a Japanese pooled analysis of three registries of 5,188 person-years, it was demonstrated via multivariate Cox regression analysis that the HR for subsequent recurrence was 3.25 (CI, 1.86–5.67) [14]. The prevalence of a history of stroke/TIA in AF patients was high in Asians (18.8% in China and 22.1% in Southeast Asia) compared with those in other regions (13.8% overall) [66]. Asian AF patients with prior stroke/TIA have strong indications for OAC, unless there are contraindications or inappropriate conditions for OAC. When prescribing OAC to patients with prior stroke/TIA, these patients are also at significantly higher risk for ICH during OAC than those without prior stroke/TIA [200–205].

8.5.1. Patients with a history of ischemic stroke

The efficacy and safety profiles of NOACs between the patient groups with and without prior stroke/TIA were consistent [200–203], indicating that NOACs can be used safely even in patients with prior stroke/TIA. There was no interaction in the efficacy and safety between patients with and without prior stroke [24]. Warfarin was shown to be associated with a numerically increased risk of major bleeding in the Asian patients compared with the non-Asians [24–26], and most importantly, it increased the incidence of ICH by 1.5–3.9 times in Asian patients [67].

Use of NOACs is associated with significantly lower risk of major bleeding than warfarin in Asian or East Asian AF patients, except for rivaroxaban [24–26], dabigatran 150 mg reduced stroke and see more effectively than warfarin [23], while other NOAC regimens, including dabigatran 110 mg, rivaroxaban 20 mg, apixaban 5 mg, and edoxaban 60 mg, had similar efficacy compared with warfarin [24–26]. Thus, NOACs seem the best option for stroke prevention when treating Asian patients with prior stroke/TIA.

The NOAC trials generally excluded patients within first 7–14 days after acute ischemic stroke. However, the risk of recurrent stroke is highest in the early phase after the first stroke/TIA [206]. The 2016 ESC Guidelines for the Management of Atrial Fibrillation proposed to initiate OAC between 1 and 12 days after an ischemic stroke [38], depending on stroke severity, using the so-called “1–3–6–12 day rule” as follows: (i) in patients with TIA, NOAC can be initiated at day 1; (ii) after mild stroke (National Institute of Health Stroke Scale (NIHSS < 8), NOAC can be initiated after 3 days, or after ICH is excluded by imaging modality (CT or MRI); (iii) in moderate stroke (NIHSS 8–16), NOAC can be initiated after 5–7 days; and (iv) in severe stroke (NIHSS > 16) after 12–14 days [113]. This is based on no trial evidence, and is simply expert opinion.

8.5.2. Patients with a history of intracranial hemorrhage

Patients with a history of ICH have higher recurrence rates of ICH when OAC is taken. Several studies have shown that in patients with a history of ICH, OAC treatment was associated with a significant reduction in ischemic stroke/all-cause mortality rates in comparison with no treatment [207,208].

In a recent report from the Taiwan NHIRD, warfarin use was found to be possibly beneficial for AF patients with prior ICH having a CHA2DS2-VASc score ≥ 6 [209]. Whether the use of NOACs can lower the threshold for treatment deserves further study [209]. One should know that, in all the RCTs of NOACs, a history of spontaneous ICH was an exclusion criteria. Therefore, the decision to use OAC should be individualized. In patients with high cardioembolic risk, CHA2DS2-VASc score ≥ 6 for instance, and a low ICH risk, OAC can be started after 4–8 weeks; the CHA2DS2-VASc score threshold may be lower for NOACs. For patients with low cardioembolic risk and high ICH risk, the use of a NOAC might be omitted [38].

**Recommendations**

- NOACs are preferred over VKA in patients with a history of ischemic stroke/TIA.
- After an acute episode of ischemic stroke/TIA, NOACs can be initiated based on the “1–3–6–12 day rule”": In patients with TIA, NOAC can be initiated at day 1. In patients with mild stroke (NIHSS < 8), NOAC can be initiated after 3 days. In patients with moderate stroke (NIHSS 8–16), NOAC can be initiated after 5–7 days, and in severe stroke (NIHSS > 16) after 12–14 days.
- For patients with a history of ICH, the decision to use OAC should be individualized. In patients with high cardioembolic risk, CHA2DS2-VASc score ≥ 6 for instance, and a low ICH risk, NOAC can be started after 4–8 weeks. For patients with low cardioembolic risk and high ICH risk, the use of a NOAC might be omitted.

8.6. Peri-operative use of NOACs

Surgical interventions or invasive procedures require careful planning and temporary discontinuation of anticoagulants. Both patient characteristics (kidney function, age, history of bleeding
complications, concomitant medication) and surgical factors should be taken into account when deciding the timing of discontinuing and restarting the anticoagulants [113,210–213]. Clinicians must always weigh the risks of thrombosis associated with discontinuing antithrombotic medications against the risks of potential bleeding inherent to the contemplated procedure [214].

8.6.1. Assessment of bleeding and stroke risk

Assessment of bleeding risk should include an understanding of the patient’s comorbidities and the invasive nature of contemplated procedure. Active cancer, thrombocytopenia, and a history of bleeding are associated with an increased risk of perioperative bleeding [215]. Formal bleeding-risk assessment is essential, and can be done by using the well-validated HAS-BLED score [44].

All patients with AF receiving anticoagulants should be assessed using the CHA2DS2-VASC score before surgery, and patients with higher scores should be considered for minimal duration of anticoagulation discontinuation, especially if they are on NOACs [211,216]. The duration of preoperative discontinuation and re-initiation of NOAC therapy should be determined on the basis of the pharmacokinetic properties of each agent [217–222].

8.6.2. Do we need a bridging strategy?

Since VKAs have a slower onset and offset of action, bridging with low molecular weight heparin (LMWH) or unfractionated heparin (UFH) has often been used in AF patients with higher thromboembolic risk [223]. However, bridging therapy is not always necessary in NOAC-treated patients since the predictable waning of the anticoagulation effect allows properly timed short-term cessation and re-initiation of NOAC therapy.

The BRIDGE trial has shown that in VKA-treated patients, bridging with LMWH has no benefit regarding thromboembolism but is instead an inferior strategy, since it can lead to a higher incidence of major bleeding [224]. In a meta-analysis of AF patients with intermediate CHADS2 scores who were anticoagulated with warfarin and required temporary interruption of warfarin for an elective surgery or procedure, periprocedural bridging with UFH or LMWH was associated with a higher rate of major bleeding with no significant difference in mortality or stroke [225].

Heparin bridging is generally not necessary for NOACs, because their half-lives are similar to those of LMWH. Bleeding and thromboembolic outcomes in the periprocedural period using NOACs versus warfarin have been investigated in the RE-LY, ROCKET AF, and ARISTOTLE trials [226–228]. In these studies, NOACs and warfarin were generally interrupted. There were no statistically significant differences between the dabigatran, rivaroxaban, or apixaban groups and their respective warfarin groups with respect to bleeding or thromboembolic complications.

In the Perioperative Dabigatran Study, a Canadian multicenter prospective study of perioperative management, 541 adult patients receiving dabigatran for any indication (AF, 97%) underwent an invasive procedure requiring NOAC interruption [229]. The outcomes of the study included major and minor bleeding, thromboembolism, and death, and the results suggested that interruption of dabigatran without bridging is safe [229]. Observational analyses from the Dresden NOAC Registry (76% rivaroxaban, 24% dabigatran) suggested no difference in bleeding or thromboembolic complications in the periprocedural period between rivaroxaban and dabigatran [230]. Heparin bridging did not reduce cardiovascular events but led to significantly higher rates of major bleeding [230].

8.6.3. Peri-operative strategy

When the intervention carries minimal bleeding risk and/or when adequate local hemostasis is possible, as with some dental procedures or interventions for cataract or glaucoma, the procedure can be scheduled 24 h after the last dose, and then, the drug can be restarted 6 h later. Stoppage of NOACs depending on renal function for patients undergoing procedures with minor or major bleeding risk is shown in Table 10.

Whenever a patient on anticoagulation needs surgery, the anticoagulant should be discontinued. If surgery cannot be delayed, reversal of the anticoagulant may be considered. Idarucizumab, a specific antibody for dabigatran, was tested in a phase III, (REVERSE-AD) trial in patients with acute life-threatening bleeding and those requiring urgent surgical intervention [231]. Idarucizumab showed a rapid and near-maximal reversal of the anticoagulant effects of dabigatran [231]. No reversal agents are currently available for clinical use to reverse the Xa inhibitors. Therefore, in emergency situations, off-label administration of prothrombin complex concentrate (PCC) or recombinant factor VII can be used in circumstances of life-threatening bleeding [232,233].

There are no data suggesting how to resume NOACs after surgical procedures. For procedures with immediate and complete hemostasis, the NOACs can be resumed 6–8 h after the intervention. In general, NOACs can be restarted 24 h after procedures with low bleeding risk, and may be restarted 48–72 h after procedures with high bleeding risk [113,222,234].

**Recommendations**

- Bridging with LMWH or UFH is not necessary for VKA-treated patients undergoing planned surgical intervention.
- Bridging with LMWH or UFH is not necessary for NOAC-treated patients undergoing planned surgical intervention.
- Stoppage of NOACs depends on renal function and the risk of bleeding of different surgical procedures.
- For patients receiving rivaroxaban, apixaban, and edoxaban, NOACs should be withheld for 24 h for procedures with low bleeding risk, or for 48 h for procedures with high bleeding risk, irrespective of renal function.
- For patients with an eCrCl ≥ 50 mL/min receiving dabigatran, the drug should be withheld for 24 h for procedures with low bleeding risk, or for 48 h for procedures with high bleeding risk.
- For patients with an eCrCl = 30–49 mL/min receiving dabigatran, the drug should be withheld for 48 h for procedures with low bleeding risk.

**Table 10**

| Dabigatran   | Rivaroxaban, Apixaban, Edoxaban |
|--------------|---------------------------------|
| Low bleeding risk | High bleeding risk               |
| CrCl ≥ 80 mL/min | ≥ 24 h                                  |
| CrCl 30–79 mL/min | ≥ 24 h                                  |
| CrCl 30–49 mL/min | ≥ 24 h                                  |

| Low bleeding risk | High bleeding risk |
|-------------------|-------------------|
| ≥ 24 h            | ≥ 48 h            |
| ≥ 24 h            | ≥ 48 h            |
| ≥ 24 h            | ≥ 48 h            |

CrCl: creatinine clearance; NOAC: non-vitamin K antagonist oral anticoagulants

Adapted from Heidbuchel et al., [113] Schulman et al., [229] and Chiang et al. [39] with permission.
low bleeding risk, or for 96 h for procedures with high bleeding risk.

- NOACs can be restarted 24 h after procedures with low bleeding risk, and may be restarted 48–72 h after procedures with high bleeding risk.

8.7. Patients undergoing cardioversion

Thromboembolism after cardioversion is relatively uncommon overall (<1% within 30 days) but is a potentially devastating complication [235,236]. Observational studies suggest that thromboembolic risk after cardioversion is highest in the first 72 h and that the majority of events occur within 10 days [235,237]. These thromboembolic events are thought to be due to the embolization of pre-existing thrombi at the time of cardioversion or from the formation and subsequent migration of thrombi that formed while atrial function was still depressed after cardioversion.

8.7.1. AF duration ≥ 48 h or of unknown duration

Anticoagulation with warfarin for at least 3 weeks before and continuing for at least 4 weeks after cardioversion has been shown to be associated with a low risk of thromboembolism [238,239]. Furthermore, the addition of trans-esophageal echocardiography (TEE) screening before cardioversion in patients who had adequate anticoagulation prior to cardioversion did not appear to reduce that risk; however, it did find left atrial thrombus in 7.7%, despite anticoagulation [239]. As for NOACs, subgroup studies of patients on dabigatran (RE-LY), rivaroxaban (ROCKET AF), and apixaban (ARISTOTLE) demonstrated comparable event rates to patients on warfarin, lending support to their use in patients with non-valvular AF [240–242]. While these studies all showed low event rates after cardioversion, they were not sufficiently powered to demonstrate superiority over warfarin.

A recent randomized study of rivaroxaban versus VKA also showed similar event rates between the 2 treatment arms, but a reduction in time to cardioversion in the rivaroxaban arm [243]. The ENSURE-AF trial was the largest prospective RCT of anticoagulation for cardioversion of patients with non-valvular AF [244]. A total of 2199 patients were randomly assigned to receive edoxaban or enoxaparin-warfarin. Both the primary efficacy endpoint (stroke/SEE, myocardial infarction, and cardiovascular mortality) and the primary safety endpoint (major and clinically relevant non-major bleeding) did not differ significantly between the 2 treatment groups [244].

In patients who have not had 3 weeks of anticoagulation before cardioversion, TEE guidance can be used to reduce the risk of thromboembolism [245,246]. If no thrombus is observed (including in the LAA), cardioversion can proceed with UFH or LMWH coverage and anticoagulation should then be continued for at least 4 weeks. This expedited anticoagulation strategy appears to be safe in observational studies [247]. If a thrombus is observed on TEE, the cardioversion should be deferred and the patient commenced on at least 4 weeks of anticoagulation. A repeat TEE to ensure thrombus resolution should be considered before another cardioversion attempt, as thrombi may not always resolve with anticoagulation [248]. If a thrombus persists on the repeat TEE, then an alternative strategy such as rate control with appropriate anticoagulation should be considered. The same principles can be applied to NOACs.

8.7.2. AF duration < 48 h

For patients who have been in AF for less than 48 h, it is common practice to perform cardioversion without TEE or antecedent anticoagulation, but there are no RCTs to support this. These recommendations are based on observational studies that found a low risk of thromboembolic events in patients who either spontaneously reverted or were cardioverted without having been anticoagulated [235,249]. It is recommended that these patients may be cardioverted early with UFH or LMWH coverage. However, it should be noted that some patients, such as those with diabetes, heart failure or significant valvular disease, may be at higher risk [235].

8.7.3. Emergent cardioversion

For patients with AF who require emergency cardioversion because of hemodynamic instability (angina, myocardial infarction, shock, or pulmonary edema), initiation of anticoagulation should not delay interventions to stabilize the patient. Again, there are no RCTs that have tested optimal anticoagulation strategies in such patients, but it may be reasonable to administer UFH or LMWH prior to cardioversion unless contraindicated. If the AF or atrial flutter has been present for more than 48 h or if its duration is unclear, oral anticoagulation is recommended for at least 4 weeks after emergency cardioversion. If warfarin is used, bridging with UFH or LMWH is indicated until the INR is therapeutic. For patients with AF and thromboembolic risk factors (CHA2DS2-VASc ≥ 1 for males, or ≥ 2 for females), long-term anticoagulation with an oral anticoagulant is recommended.

8.7.4. Atrial flutter

There is less data on the thromboembolic risk of cardioverting atrial flutter, but it is probably similar to AF and has been shown to be associated with thrombi and episodes of AF [238]. It is recommended that the anticoagulation management strategy for cardioversion of atrial flutter should hence be the same as for AF.

Recommendations

- For patients with AF or atrial flutter of 48 h duration or longer, or when the duration of AF is unknown, anticoagulation with warfarin (INR 2.0 to 3.0) or NOAC is recommended for at least 3 weeks before and 4 weeks after cardioversion, regardless of the CHA2DS2-VASc score and the method (electrical or pharmacological) used.
- For patients with AF or atrial flutter of 48 h duration or longer, or when the duration of AF is unknown, and who have not been anticoagulated for the preceding 3 weeks, it is reasonable to perform TEE before cardioversion and to proceed with cardioversion if no LA thrombus is observed, including in the LAA, provided that anticoagulation is achieved before TEE with therapeutic warfarin (INR 2.0–3.0) or NOAC, and maintained after cardioversion for at least 4 weeks.
- For patients undergoing a TEE-guided strategy in whom a thrombus is identified, VKA (INR 2.0–3.0) or NOAC is recommended for at least 4 weeks, followed by a repeat TEE to ensure thrombus resolution.
- If thrombus resolution is evident on repeat TEE, cardioversion should be performed, and OAC should be considered for at least 4 weeks, or lifelong if risk factors are present (i.e., CHA2DS2-VASc ≥ 1 in males or ≥ 2 in females).
- For patients with AF or atrial flutter of more than 48 h duration or unknown duration that requires immediate cardioversion for hemodynamic instability, anticoagulation with UFH or LMWH should be initiated as soon as possible and anticoagulation should be continued for at least 4 weeks after cardioversion unless contraindicated.
- For patients with AF or atrial flutter of less than 48 h duration and with high risk of stroke, anticoagulation with UFH or LMWH is recommended as soon as possible before cardioversion, followed by long-term anticoagulation therapy.
- Following cardioversion for AF or flutter of any duration, long-term anticoagulation therapy should be considered in patients with a CHA2DS2-VASc ≥ 1 in males or ≥ 2 in females.
8.8. Peri-ablation procedure

Catheter ablation of AF is associated with a potential risk of SEE not only due to the introduction and manipulation of catheters and long sheaths within the atrium, but also the endocardial damage produced by ablations [250]. Recent studies show that the incidence of SEE with AF ablation was approximately 1–5% [251,252].

Several studies have assessed the safety and efficacy of NOACs compared to warfarin during the periprocedural period of AF ablation. The results of these studies have demonstrated that NOACs, such as dabigatran, could be a safe and effective alternative to warfarin during the periprocedural period of AF ablation [253,254]. Antithrombotic strategies for the prevention of SEE should be specified for 3 different stages, that is, pre-ablation, during-ablation, and post-ablation stages [250].

8.8.1. Pre-Ablation

For patients who have had AF for more than 48 h, or for an unknown duration, systemic anticoagulation with NOACs or VKAs for at least 3 weeks is recommended. For patients being treated with VKAs, the ablation should be performed without interruption of VKAs.

Several non-randomized studies have shown that uninterrupted VKA use is associated with a lower risk of thromboembolic and hemorrhagic complications compared to interrupted VKAs [251,255,256]. The VENTURE AF trial showed similar event rates in patients on uninterrupted rivaroxaban compared with uninterrupted VKA [257]. In the recent RE-CIRCUIT trial, 704 patients who were scheduled for catheter ablation of AF were randomly assigned to receive either dabigatran (150 mg bid) or warfarin (target INR, 2.0 to 3.0) [258]. Ablation was performed after 4 to 8 weeks of uninterrupted anticoagulation, which was continued during ablation and for 8 weeks afterwards. The incidence of major bleeding events (primary endpoint) during and up to 8 weeks after ablation was significantly lower with dabigatran than with warfarin (1.6% vs. 6.9%, absolute risk difference, −5.3 percentage points; 95% CI −8.4 to −2.2; P < 0.001). The relative risk reduction versus warfarin was 77.2% (HR 0.22, 95% CI, 0.08–0.59). Dabigatran was associated with fewer periprocedural pericardial tamponades and groin hematoma than warfarin. One thromboembolic event occurred in the warfarin group. These findings confirmed that uninterrupted dabigatran was associated with fewer bleeding complications than uninterrupted warfarin, while maintaining antithrombotic efficacy [258].

A TEE should be performed for patients who do not receive appropriate systemic anticoagulants for 3 weeks or for whom the duration of treatment is uncertain before the intervention [259]. Several studies suggested that even adequate anticoagulation could not exclude the possibility of the presence of left atrial thrombi or sludge in the LAA [260–262]. TEE can be performed on the day of, or up to 24 h before, the ablation to rule out the presence of atrial thrombi. Once a left atrial thrombus is detected, ablation should be delayed; it may be reconsidered after a 3-month treatment of systemic anticoagulants. However, TEE may not be necessary in patients with paroxysmal AF and full anticoagulation for > 3 weeks and with a low risk of thromboembolism such as CHA2DS2-VASc 0 for males and 1 for females.

For patients in sinus rhythm with a CHA2DS2-VASc score of 0 for males or 1 for females, NOACs can be initiated on the day of the ablation procedure and continued post-ablation [250]. Although prospective and randomized controlled trials are lacking, there is a trend toward the initiation of the antithrombotic treatments before ablation, even for patients who present with sinus rhythm.

8.8.2. During Ablation

Heparin should be administered prior to or immediately after trans-septal puncture, and the dose should be adjusted to achieve and maintain an activated clotting time (ACT) of 300 to 400 seconds [259]. An intravenous loading dose of 5,000–15,000 units (or 90–200 U/kg) of heparin should be administered at the beginning of the procedure [250]. Previous studies showed that patients on NOAC required a larger dose of heparin and took longer time to reach the target ACT level than those on a VKA [257,263]. After the loading dose of heparin, continuous heparin infusion at an initial rate of 1000–1500 U/kg/h can be started depending on the levels of ACT. The target ACT level should be achieved and maintained by administering heparin ranging between 2,500 and 7,500 U intermittently. The ACT level should be checked every 10–15 min before the therapeutic anticoagulation is achieved and, then every 15–30 min for the rest of the procedure [259]. Uninterrupted VKAs or NOACs influence the ACT and the time needed to reach the target ACT level [250,254,264].

All sheaths should be continuously flushed with heparinized saline solution, with a suggested dose of 2000 units per 250 mL [265,266]. Heparin infusion can be discontinued once all catheters are removed from the left atrium. To decrease the risk of bleeding of the puncture site, ACT should be less than 250 s before the sheath is removed, otherwise protamine should be used to reverse the anticoagulation effect of heparin.

8.8.3. Post-ablation

The atria are vulnerable to the formation of thrombi after ablation, and therefore adequate post-ablation anticoagulation is crucial to prevent SEE. Systemic oral anticoagulants should be continued for at least 2 months after ablation [259]. Recently, a cohort study demonstrated that most thromboembolic events occurred within 4 weeks post-ablation, and the thromboembolic risk beyond 3 months after ablation was relatively low compared with a matched non-ablated AF cohort [267]. For patients who discontinue VKAs or with a suboptimal INR at the time of ablation, LMWH should be administered 4–6 h after AF ablation, along with the re-initiation of VKAs once hemostasis has been achieved. The LMWH should be maintained until a therapeutic INR level (2.0–3.0) has been achieved [259]. Because of the increased risk of bleeding on full-dose LMWH (1 mg/kg bid), a dosage reduction of LMWH (0.5 mg/kg) should be considered [259]. In patients who continue VKAs, the use of LMWH may be avoided and the INR should be maintained between 2.0 and 3.0 [259].

In patients on NOACs, the next dose should be resumed within 3–4 h once hemostasis is achieved [268]. Decisions regarding the use of systemic anticoagulants for longer than two months following ablation should be based on the patient’s risk for stroke [259]. Long-term anticoagulation is recommended for patients at a high risk of stroke (CHA2DS2-VASc score ≥ 1 in males or ≥ 2 in females). Continuous ECG monitoring should be considered to detect asymptomatic AF in patients who discontinue systemic anticoagulants [259]. Recently, several studies have demonstrated that noninvasive ambulatory ECG monitoring can significantly improve the detection of AF [269,270]. In patients who are at an increased risk of stroke, noninvasive ambulatory ECG monitoring may be necessary.

Recommendations

- NOACs can be safe and effective alternatives to warfarin during the periprocedural period of AF ablation.
- For patients who have had AF for more than 48 h or an unknown duration, systemic anticoagulation with NOACs or VKAs for at least 3 weeks before any ablation procedure is recommended.
• For patients treated with VKAs or NOACs, the ablation should be performed without interruption of VKAs or NOACs.
• A TEE should be performed for patients who do not receive appropriate systemic anticoagulants for 3 weeks, or for whom the duration of treatment is uncertain before the intervention.
• TEE can be performed on the day of, or up to 24 h before, the ablation to rule out the presence of atrial thrombi.
• TEE may not be necessary in patients with paroxysmal AF and full anticoagulation for > 3 weeks and with a low risk of thromboembolism such as when CHA2DS2-VASc = 0 for males and = 1 for females.
• For patients in sinus rhythm with a CHA2DS2-VASc score = 0 for males or = 1 for females, NOACs can be initiated on the day of the ablation procedure and continued post-ablation.
• Heparin should be administered prior to, or immediately after, trans-septal puncture and the dose should be adjusted to achieve and maintain an ACT of 300 to 400 seconds.
• Systemic oral anticoagulants should be continued for at least 2 months after ablation, and for longer in the presence of risk factors, irrespective of the apparent success of rhythm control.
• For patients who discontinue VKAs or who have a suboptimal INR at the time of ablation, LMWH should be administered 4–6 h after AF ablation along with the re-initiation of VKAs once hemostasis has been achieved.
• In patients on NOACs, the next dose should be resumed within 3–4 h once hemostasis is achieved.
• Long-term anticoagulation is recommended for patients at a high risk of stroke (CHA2DS2-VASc score ≥ 1 in males or ≥ 2 in females).

8.9. Management of bleeding complications

The management strategy for bleeding in patients on OAC therapy depends primarily on the severity of bleeding, and the types of OAC used. Generally, bleeding events are broadly classified into mild, moderate, and severe bleeding. Mild bleeding events include epistaxis, small bruises, and bleeding after minor trauma; moderate bleeding events include gross hematuria, spontaneous large bruises, and any bleeding requiring transfusion but no hemodynamic compromise; and severe bleeding events refer to potentially life-threatening bleeding, such as bleeding into critical sites (ICH and retroperitoneal bleeding), or bleeding leading to hemodynamic instability.

General measures for active bleeding in patients on OAC include: mechanical compression if the bleeding site is accessible; hemodynamic assessment; clarification of the types and last dose of OAC; and blood tests for complete blood count, liver and kidney function, and basic coagulation tests. In cases of mild bleeding which stops spontaneously, no intervention is needed after ascertaining the dosage and timing of drug for patients taking NOACs.

In case of mild bleeding which stops spontaneously, no intervention is needed after ascertaining the dosage and timing of drug for patients taking NOACs.

In case of moderate bleeding, OAC should be withheld immediately and standard supportive measurements must be started.

In the setting of moderate bleeding, OAC should be withheld immediately and standard supportive measurements must be started. These include fluid replacement to maintain urine output, blood transfusion, and additional hemodynamic support. For patients on VKA, intravenous vitamin K (1–10 mg) may be considered [271]. Activated charcoal can reduce further absorption of NOAC if the drug has been administered within 2–4 h. Regardless of the types of OAC, specific diagnostic procedures and therapeutic interventions to control bleeding, such as endoscopy, should be considered.

In case of severe or life-threatening bleeding, in addition to the aforementioned measures, treatments to reverse anticoagulation effects should be considered. For patients on VKA, fresh frozen plasma may restore coagulation more rapidly than vitamin K.

For patients on dabigatran, idarucizumab should be considered [231]. When idarucizumab is not available, hemodialysis accelerates the clearance of dabigatran from the body. On the contrary, andexanet alfa, a recombinant modified human factor Xa decay protein, has also been demonstrated to be effective in reversing the anticoagulant effect of various factor Xa inhibitors; but it has yet to be approved by regulatory bodies. Unlike dabigatran, hemodialysis is in general not effective in the removal of factor Xa inhibitors out of the body. For patients on factor Xa inhibitors, PCC or activated prothrombin complex (aPCC) concentrates can be considered in severe or life-threatening bleeding with a starting dose of 25 U/kg and can be repeated if clinically indicated.

Recommendations

• In case of mild bleeding which stops spontaneously, no intervention is needed after ascertaining the dosage and timing of drug for patients taking NOACs.
• In the setting of moderate bleeding, OAC should be withheld immediately and standard supportive measurements must be started.
• In case of severe or life-threatening bleeding, treatments to reverse anticoagulation effects should be considered, such as idarucizumab in patients receiving dabigatran.
• When idarucizumab is not available, hemodialysis accelerates the clearance of dabigatran from the body.
• For patients on factor Xa inhibitors, PCC or aPCC concentrates can be considered in severe or life-threatening bleeding.

8.10. Reversal agents

Specific reversal agents for NOACs are now available. Idarucizumab is a monoclonal antibody fragment and binds dabigatran with an affinity that is 350 times as high as that observed with thrombin [272]. In the recent RE-VERSE AD study, the efficacy and safety of idarucizumab was tested in dabigatran-treated patients who had serious bleeding or required urgent procedures [231]. In the recent interim analysis of the first 90 patients, idarucizumab completely reversed the anticoagulant effect of dabigatran within minutes [231]. Immediately after the administration of idarucizumab, the concentration of unbound dabigatran was reduced to a level at or near the lower limit of quantification in all but 1 patient [231]. The US FDA has granted accelerated approval to idarucizumab (Praxbind®) to rapidly reverse the effects of dabigatran.

Andexanet alfa (andexanet) is a specific reversal agent for both direct and indirect factor Xa inhibitors [273]. Andexanet is a recombinant modified human factor Xa decay protein that is catalytically inactive but that retains the ability to bind factor Xa inhibitors in the active site with high affinity and a 1:1 stoichiometric ratio. In a recently published clinical trial, andexanet reversed the anticoagulant activity of apixaban and rivaroxaban in healthy older participants within minutes after administration and for the duration of infusion, without clinical evidence of toxic effects [274]. In the ANNEXA-4 trial, an initial bolus and subsequent 2-h infusion of andexanet substantially reduced anti-factor Xa activity in patients with acute major bleeding associated with factor Xa inhibitors, with effective hemostasis occurring in 79% [275]. However, the US FDA has delayed approval of andexanet. Aripazine (ciraparantag, PER 977) is a small molecule that interacts with anticoagulants through non-covalent hydrogen bonding and electrostatic interactions. This agent appears to inhibit nearly all anticoagulants with the exception of vitamin K.
antagonists and argatroban [276]. Clinical trials are awaited to confirm its efficacy and safety in AF patients.

**Recommendations**

- Idarucizumab, a specific reversal agent for dabigatran, is indicated in patients with serious bleeding or requiring urgent procedures.

**9. Management algorithm**

CHA2DS2-VASc score has outperformed other scoring systems in predicting AF-associated stroke in Asians [34,36]; therefore, the APHRS consensus on stroke prevention in AF recommends the use of CHA2DS2-VASc scores in the prediction of stroke risk. A management algorithm is shown in Fig. 2.

The first step is to identify those patients with low risk (i.e. CHA2DS2-VASc score 0 in males, 1 in females); no antithrombotic agent is recommended for them. The second step is offer stroke prevention to those with ≥ 1 additional stroke risk factors. The third step is to use the SAME-TT2R2 score to identify patients who have a possibility of doing well with VKA (SAME-TT2R2 score, 0–2) or those patients who are unlikely to achieve a good TTR by taking VKA (SAME-TT2R2 score ≥ 3), so a NOAC should be used initially, without subjecting the patient to a "trial of warfarin" period.

No head-to-head RCT has tested the superiority of one NOAC versus another, and therefore, one can choose any NOAC, based on available evidence.

**Conflict of interest**

Chern-En Chiang has been on the speakers' bureau for AstraZeneca, Bayer, Boehringer Ingelheim, Chugai, Daiichi-Sankyo, MSD, Novartis, Pfizer, Roche, Sanofi-aventis, Servier, Tanabe, Takeda, and TTY.

Ken Okumura has received remuneration from Boehringer Ingelheim, Daiichi-Sankyo, Medtronic and Johnson & Johnson.

Shu Zhang has been an advisory board member of Boston Scientific, an investigator for Boston Scientific, and an investigator for Medtronic.

Tze-Fan Chao has declared no conflict of interest related to this paper.

Chung-Wah Siu has declared no conflict of interest related to this paper.

Toon Wei Lim has received research funding from Bayer, Biotronik, Boehringer Ingelheim, Boston Scientific, Medtronic, and Pfizer. He has been on the advisory board of Bayer, Boehringer Ingelheim, and Pfizer. He has received travel support & honoraria from Bayer, Biotronik, Boehringer Ingelheim, Boston Scientific, Medtronic, Pfizer, and St. Jude Medical.

Anil Saxena worked as consultant for and attended advisory board meetings of Boehringer-Ingelheim, Bayer Pharma, and Pfizer.

Yoshhide Takahashi has received speaker fees from Biosense Webster.

Wee Siong Teo has received research grants from Pfizer.

**References**

[1] Freedman B, Potpara TS, Lip GYH. Stroke prevention in atrial fibrillation. Lancet 2016;388:806–17.

[2] Gladstone DJ, Bui E, Fang J, et al. Potentially preventable strokes in high-risk patients with atrial fibrillation who are not adequately anticoagulated. Stroke 2009;40:235–40.

[3] Tomita H, Hagii J, Metoki N, et al. Impact of sex difference on severity and functional outcome in patients with cardioembolic stroke. J Stroke Cerebrovasc 2015;24:2631–8.

[4] Perera KS, Vannaische T, Bosch J, et al. Global survey of the frequency of atrial fibrillation–associated stroke. Embolic stroke of undetermined source global registry. Stroke 2016;47:2197–202.

[5] Yamashita Y, Hamatani Y, Esato M, et al. Clinical characteristics and outcomes in extreme elderly (age ≥ 85 Years) Japanese patients with atrial fibrillation: the Fushimi AF registry. Chest 2016;149:401–12.

[6] Leyden JM, Kleining TJ, Newbury J, et al. Adelaide stroke incidence study: declining stroke rates but many preventable cardioembolic strokes. Stroke 2013;44:1226–31.

[7] Chiang CE, Wang KL, Lip GYH. Stroke prevention in atrial fibrillation: an Asian perspective. Thromb Haemost 2014;111:789–97.

[8] Wolf PA, Abboud RB, Kannel WB. Atrial fibrillation as an independent risk factor for stroke: the Framingham study. Stroke 1991;22:2983–8.

[9] Chien KL, Su TC, Hsu HC, et al. Atrial fibrillation prevalence, incidence and risk of stroke and all-cause death among Chinese. Int J Cardiol 2010;139:173–80.

[10] Lip GY, Nieuwlaat R, Pisters R, et al. Refining clinical risk stratification for predicting stroke and thromboembolism in atrial fibrillation using a novel risk factor-based approach: the euro heart survey on atrial fibrillation. Circulation 2011;123:263–72.

[11] Piccini JP, Stevens SR, Chang Y, et al. Renal dysfunction as a predictor of stroke and systemic embolism in patients with nonvalvular atrial fibrillation: validation of the R(2)CHA2DS(2) index in the ROCKET AF (rivaroxaban once-daily, oral, direct factor Xa inhibition compared with vitamin K antagonist for prevention of stroke and embolism trial in atrial fibrillation) and ATRIA (Anticoagulation and risk factors in atrial fibrillation) study cohorts. Circulation 2013;127:224–32.

[12] Lau YC, Proietti M, Guiducci E, et al. Atrial fibrillation and thromboembolism in patients with chronic kidney disease. J Am Coll Cardiol 2016;68:1452–64.

[13] Hamatani Y, Ogawa H, Uozumi R, et al. Low body weight is associated with the incidence of stroke in atrial fibrillation patients—insight from the Fushimi AF registry. Circ J 2015;79:1009–17.

[14] Suzuki S, Yamashita T, Okumura K, et al. Incidence of ischemic stroke in Japanese patients with atrial fibrillation not receiving anticoagulation therapy-pooled analysis of the shinken database, J-HYTHEM registry, and Fushimi AF registry. Circ J 2015;79:432–8.
[5] Ho C-W, Ho M-H, Chan P-H, et al. Ischemic stroke and intracranial hemorrhage with aspirin, dabigatran, and warfarin: impact of quality of anticoagulation control. Stroke 2015;46:23–30.

[6] Chao TF, Liu CJ, Wang KL, et al. Using the CHA2DS2-VASc score for refining stroke risk stratification in low-risk Asian patients with atrial fibrillation. Am J Cardiol 2014;64:1658–65.

[7] Guo Y, Lip GY, Apostolakis S. The unmet need of stroke prevention in atrial fibrillation in the far East and South East Asia. Malays J Med Sci 2012;19:1–7.

[8] Singer DE, Chang Y, Bönschky LH, et al. A new risk scheme to predict ischemic stroke and other thromboembolism in atrial fibrillation: the ATRIA study stroke risk score. J Am Heart Assoc 2013;2:e000250.

[9] Connolly SJ, Eikelboom J, Granger CB, et al. Oral anticoagulants for atrial fibrillation in patients with cardiovascular disease. N Engl J Med 2009;361:1191–9.

[10] Patel MR, Mahaffey KW, Garg J, et al. Rivaroxaban versus warfarin in nonvalvular atrial fibrillation. N Engl J Med 2011;365:981–92.

[11] Giugliano RP, Ruff CT, Braunwald E, et al. Edoxaban versus warfarin in atrial fibrillation: a derivation and validation study. Lancet 2016;387:2302–11.

[12] Haji Z, Öldjerg J, Lindbäck J, et al. The novel biomarker-based ABC (age, biomarkers, clinical history)-bleeding risk score for patients with atrial fibrillation: a derivation and validation study. J Am Coll Cardiol 2015;63:3258–64.

[13] O’Brien EC, Simon DN, Thomas LE, et al. The ORBIT bleeding score: a simple bedside score to assess bleeding risk in atrial fibrillation. Eur J Heart Fail 2015;17:1711–14.

[14] Friberg L, Rosenqvist M, Lip GYH. Evaluation of risk stratifi cation schemes for ischaemic stroke and bleeding in 182 678 patients with atrial fibrillation: the Swedish Atrial Fibrillation cohort study. Eur Heart J 2012;33:1500–10.

[15] Apostolakis S, Lane DA, et al. Performance of the HEMORR2HAGES, ATRIA, and HAS-BLED bleeding risk prediction scores in patients with atrial fibrillation undergoing anticoagulation: the MADEUSE (Evaluating the use of SR34006 compared to apixaban or acenocumarol in patients with atrial fibrillation) Study. J Am Coll Cardiol 2012;60:861–7.

[16] Proietti M, Senoo K, Lane DA, et al. Major bleeding in patients with nonvalvular atrial fibrillation: impact of time in therapeutic range on concomitant regular antiplatelet therapy. Lancet 2015;385:1639–46.

[17] Senoo K, Proietti M, Lane DA, et al. Evaluation of the HAS-BLED, ATRIA, and ORBIT bleeding risk scores in patients with atrial fibrillation taking warfarin. Am J Med 2016;129:600–7.

[18] Naganuma M, Shio K, Tanaka Y, et al. Clinical outcome in Japanese elderly patients with non-valvular atrial fibrillation taking warfarin: a single-center observational study. Thromb Res 2012;130:21–6.

[19] Sato H, Ishikawa K, Kitabatake A, et al. Low-dose aspirin for prevention of stroke in low-risk patients with atrial fibrillation: Japan atrial fibrillation stroke trial. Stroke 2006;37:447–51.

[20] Siu CW, Lip GY, Lam KT, et al. Risk of stroke and intracranial hemorrhage in 9772 Chinese patients with atrial fibrillation in Hong Kong. Heart Rhythm 2013;10:1401–8.

[21] Lip GYH, Skirka F, Rasmussen JH, et al. Net clinical benefit for oral anticoagulation, aspirin, or no therapy in nonvalvular atrial fibrillation patients with 1 additional risk factor of the CHA2DS2-VASc score (Beyond Sex). J Am Coll Cardiol 2015;66:488–90.

[22] Senoo K, Lau YC, Dzeshika M, et al. Efficacy and safety of non-vitamin K antagonist oral anticoagulants vs. warfarin in Japanese patients with atrial fibrillation–meta-analysis. Circ J 2015;79:339–45.

[23] Lip GY, Wang KL, Chiang CE. Non-vitamin K antagonist oral anticoagulants (NOACs) for stroke prevention in Asian patients with atrial fibrillation: time for a reappraisal. Int J Cardiol 2015;180:246–54.

[24] Connolly SJ, Eikelboom J, Joyner C, et al. Apixaban in patients with atrial fibrillation. N Engl J Med 2011;364:806–17.

[25] Guo Y, Pisters R, Apostolakis S, et al. Stroke risk and suboptimal thromboembolism in Chinese patients with atrial fibrillation: would the novel oral anticoagulants have an impact? Int J Cardiol 2013;168:315–22.

[26] Gama H, Murin J, Chiang CE, et al. Use of antithrombotics in atrial fibrillation in Africa, Europe, and South America: insights from the International RealiseAF Survey. Arch Cardiovasc Dis 2014;107:77–87.

[27] Amerena J, Chen SA, Sirinatanasathorn C, et al. Insights into management of atrial fibrillation in Asia Pacific gained from baseline data from REgistry on cardiac rhythm disORDers (ReCORDAsia–Pacific [AP]i) registry. Am J Cardiol 2012;109:378–82.

[28] Chang CH, Yang YH, Chen JH, et al. Cost-effectiveness of dabigatran etexilate for the prevention of stroke and systemic embolism in atrial fibrillation in Taiwan. Thromb Res 2014;133:782–9.

[29] Huissman MV, Ma CS, Diener H-C, et al. Antithrombotic therapy use in patients with atrial fibrillation before the era of non-vitamin K antagonist oral anticoagulants: the global registry on long-term oral antithrombotic treatment in patients with atrial fibrillation (GLORIA-AF) phase I cohort. EuroIntervention 2016;10:1388–1405.

[30] Chan EW, Lau WCY, Siu CW, et al. Effect of suboptimal anticoagulation treatment with antiplatelet therapy and warfarin on clinical outcomes in patients with nonvalvular atrial fibrillation: a population-wide cohort study. Heart Rhythm 2016;13:1581–8.

[31] Oldgren J, Healey JS, Ezekowitz M, et al. Variations in cause and management of atrial fibrillation in Asia Pacific. Eur Heart J 2015;36:3258–64.

[32] Hart RG, Pearce LA, Aguilar ML. Meta-analysis: antithrombotic therapy to prevent stroke in patients who have nonvalvular atrial fibrillation. Ann Intern Med 2007;146:857–67.
Compliance with antithrombotic prescribing guidelines for patients with atrial fibrillation—A nationwide descriptive study in Taiwan. Clin Ther 2008;30:1726–36.

Su CW, Tse HF. Net clinical benefit of warfarin therapy in elderly Chinese patients with atrial fibrillation. Circ Arrhythm Electrophysiol 2014;7:300–6.

Li SY, Zhao XQ, Wang CX, et al. One-year clinical prediction in Chinese ischemic stroke patients using the CHADS2 and CHA2DS2-VASC scores: the China National Stroke Registry. CNS Neurosci Ther 2012;18:988–93.

Zhang LF, Yang J. Proportion of different subtypes of stroke in China. Stroke 2003;34:2091–6.

Huang CY, Chan FY, Yu YL, et al. Cerebrovascular disease in Hong Kong Chinese. Stroke 1999;30:735–50.

Yang QB, Niu Q, Zhou YH, et al. Incidence of cerebral hemorrhage in the Changsha community. Cerebrovas Dis 2004;17:303–11.

Wallentin L, Yusuf S, Ezekowitz MD, et al. Efficacy and safety of dabigatran compared with warfarin at different levels of international normalised ratio confirmed by strong prediction: an analysis of the RE-LY trial. Lancet 2010;376:975–83.

Singer DE, Heikkamp AS, Pipicino JP, et al. Impact of global geographic region on time in therapeutic range on warfarin anticoagulant therapy: data from the ROCKET AF clinical trial. J Am Heart Assoc 2013;2:e000067.

Baber U, Howard VJ, Halperin JL, et al. Association of chronic kidney disease with atrial fibrillation among adults in the United States: reasons for geographic and racial differences in stroke (REGARDS) study. Circ Arrhythm Electrophysiol 2011;4:26–12.

Alonso A, Lopez FL, Matsushita K, et al. Chronic kidney disease is associated with the incidence of atrial fibrillation: the atherosclerosis risk in community (ARIC) study. Circulation 2011;123:3946–54.

Ananthapurnayut N, Watan N, Sudolph EH, et al. Prevalence of atrial fibrillation and its predictors in nondialysis patients with chronic kidney disease. Clin J Am Soc Nephrol 2010;5:173–81.

Fujimura EZ, Prineas RS, et al. Chronic kidney disease and prevalent atrial fibrillation: the chronic renal insufficiency cohort (CRIC). Arch Med Res 2010;19:1022–7.

Hart RG, Pearce LA, Asinger RW, et al. Warfarin in atrial fibrillation patients with moderate chronic kidney disease. Clin J Am Soc Nephrol 2011;6:2599–604.

Abbott KC, Trespalacios FC, Taylor AJ, et al. Atrial fibrillation in chronic dialysis patients in the United States: risk factors for hospitalization and mortality. BMC Nephrol 2003;4:1.

Chan KE, Lazarus JM, Thadhani R, et al. Warfarin use associates with increased risk for stroke in hemodialysis patients with atrial fibrillation. J Am Soc Nephrol 2009;20:2223–33.

Winkelmaier WC, Liu J, Setoguchi S, et al. Effectiveness and safety of warfarin substitution in older nondialysis patients with incident atrial fibrillation. Clin J Am Soc Nephrol 2011;6:2662–8.

Elliott MJ, Zimmerman D, Holden RM. Warfarin anticoagulation in hemodialysis patients: a systematic review of bleeding rates. Am J Kidney Dis 2007;50:431–40.

Biggs JA, Remmers Jr AR, Glassford DM, et al. The risk of anticoagulation in hemodialysis patients. Nephron 1977;18:109–13.

Wizemann V, Tong L, Satayathum S, et al. Atrial fibrillation in hemodialysis patients: clinical features and associations with anticoagulant therapy. Kidney Int 2010;77:1098–106.

Tobe SW, Clase CM, Gao P, et al. Cardiovascular and renal outcomes with felodipin, ramipril, or both in patients at high renal risk: results from the RENAAL study. Circulation 2011;123:1058–107.

Shah M, Avgil Tsadok M, Jackevicius CA, et al. Warfarin use and the risk for stroke in surgical patients with atrial fibrillation on time in therapeutic range on warfarin anticoagulant therapy: data from the ROCKET AF clinical trial. J Am Heart Assoc 2013;2:e000067.

Baber U, Howard VJ, Halperin JL, et al. Association of chronic kidney disease with atrial fibrillation among adults in the United States: reasons for geographic and racial differences in stroke (REGARDS) study. Circ Arrhythm Electrophysiol 2011;4:26–12.

Larsen TB, Lip GYH. Warfarin or novel oral anticoagulants for atrial fibrillation and valvular heart disease: the DANISH-AF trial. J Thromb Haemost 2011;9:1272–82.

Davies AI, Raskin F, Giugliano RP, et al. Non-Vitamin K antagonist oral anticoagulants in patients with atrial fibrillation and valvular heart disease. J Am Coll Cardiol 2017;69:1363–71.

Eikelboom JW, Connolly SJ, Bruereckmann M, et al. Dabigatran versus warfarin in patients with mechanical heart valves. N Engl J Med 2013;369:1266–14.

Jaffer BI, Stafford AR, Frederburgh JC, et al. Dabigatran is less effective than warfarin at attenuating mechanical heart valve-induced thrombin generation. J Am Heart Assoc 2015;4:e002322.

Apostolakis S, Sullivan RM, Oldhams B, et al. Factors affecting quality of anticoagulation control among patients with atrial fibrillation on warfarin: the SAME–TTZ2R score. Chest 2013;144:1555–63.

Larsen TB, Lip GYH. Novel or oral anticoagulants for atrial fibrillation. Circulation 2014;130:1538–52.

Chan PH, Hui J, Chan EW, et al. Use of the SAME–TTZ2R score to predict good anticoagulation control with warfarin in Chinese patients with atrial fibrillation: relationship to ischemic stroke incidence. PLoS One 2016;11:e0163474.

Bennants N, Ching CK, Chan L, et al. The sex, age, medical history, treatment, tobacco use, race (SAME–TTZ2R) score predicts warfarin control in a Singaporean population. J Stroke Cerebrovasc Dis 2017;26:64–9.

Husted S, De Caterina R, Andreotti F, et al. Non-vitamin K antagonist oral anticoagulants (NOACs): no longer new or novel. Thromb Haemost 2014;111:781–2.

Ruff CT, Giugliano RP, Braunwald E, et al. Comparison of the efficacy and safety of new oral anticoagulants with warfarin in patients with atrial fibrillation: a meta-analysis of randomised trials. Lancet 2014;383:955–62.

Heidbuchel H, Verhamme P, Alings M, et al. Updated European Heart Rhythm Association practical guide on the use of non-vitamin K antagonist anticoagulants in patients with non-valvular atrial fibrillation. Europace 2014;16:1467–507.

Ericsson BI, Quinlan DJ, Weitz JJ. Comparative pharmacodynamics and pharmacoekinetiks of oral direct thrombin and factor Xa inhibitors in development. Clin Pharmacokinet 2009;48:1–22.

Lip GYH, Agnelli G, Edoxaban: a focused review of its clinical pharmacology. Eur Heart J 2013;35:1844–55.

De Caterina R, Husted S, Wallentin L, et al. New oral anticoagulants in atrial fibrillation and acute coronary syndromes: e sc working group on thrombosis-risk factor interaction in patients with heart disease position paper. J Am Coll Cardiol 2012;59:1413–25.

Shimada YJ, Yamashita T, Koreszny E, et al. Effects of regional differences in asia on efficacy and safety of edoxaban compared with warfarin-insights from the ENTEGRATE AF-TIMI 48 study. Eur Heart J 2015;36:3211–20.

Saito K, Yamagata Y, Kurosawa Y, et al. Edoxaban for the prevention of cardioembolic stroke in patients with nonvalvular atrial fibrillation: a meta-analysis. Circulation 2014;130:1511–15.

Wang KL, Lip GY, Lin SJ, et al. Non-Vitamin K antagonist oral anticoagulants for stroke prevention in Asian patients with nonvalvular atrial fibrillation: meta-analysis. Stroke 2015;46:2555–61.

Koreszny E, Yamagata Y, Yang Y, et al. Edoxaban versus warfarin in East-Asian (Including Japanese) patients with atrial fibrillation—an ENTEGRATE AF-TIMI 48 Sub-analysis. Circulation 2014;130:848–48.

Hori M, Matsumoto M, Tanahashi N, et al. Rivaroxaban vs. warfarin in Japanese patients with atrial fibrillation—the J-ROCKET AF study. Circ J 2012;76:1024–11.

Wang KL, Giugliano RP, Goto S, et al. Standard dose versus low dose non-vitamin K antagonist oral anticoagulants in Asian patients with atrial fibrillation: a meta-analysis of contemporary randomized controlled trials. Heart Rhythm 2016;13:2340–7.

Lip GYH, Clemens A, Noack H, et al. Patients outcomes using the European label for dabigatran. A post-hoc analysis from the RE-LY database. Thromb Haemost 2014;111:533–40.

Blackshear JL, Odell JA. Appendage obliteration to reduce stroke in cardiac surgical patients with atrial fibrillation. Ann Thorac Surg 1996;61:755–9.

Whitlock RP, Healey JS, Connolly SJ. Left atrial appendage occlusion does not eliminate the need for warfarin. Circulation 2009;120:1927–32.

Bayard YL, Omran H, Neuzil P, et al. PLATeTO (Percutaneous left atrial appendage transcather occlusion) for prevention of cardioembolic stroke in non-anticoagulation eligible atrial fibrillation patients: results from the PERMA Belgian trial. Eur Heart J 2015;36:225–6.

Holmes DR, Reddy VY, Turi ZG, et al. Percutaneous closure of the left atrial appendage versus warfarin therapy for prevention of stroke in patients with atrial fibrillation: a randomised non-inferiority trial. Lancet 2009;374:534–42.
Macle L, Cairns JA, Andrade JG, et al. The 2014 atrial fibrillation guidelines companion: a practical approach to the use of the Canadian cardiovascular society guidelines. Can J Cardiol 2015;31:1207–18.

Witt CT, Healey JS. Oral anticoagulant use in patients with chronic kidney disease: how to do it right, and the importance of empiric human data. Can J Cardiol 2014;30:853–4.

Dahal K, Kunwar S, Rijal J, et al. Stroke, major bleeding, and mortality outcomes in warfarin users with atrial fibrillation and chronic kidney disease: a meta-analysis of observational studies. Chest 2016;149:951–60.

Hijazi Z, Hohnloser SH, Oldgren J, et al. Efficacy and safety of dabigatran compared with warfarin in relation to baseline renal function in patients with atrial fibrillation: a RE-LY (Randomized evaluation of long-term anticoagulation therapy) analysis. Circulation 2014;129:591–70.

Hohnloser SH, Hijazi Z, Thomas L, et al. Efficacy of apixaban when compared with warfarin in relation to renal function in patients with atrial fibrillation: insights from the ARISTOTLE trial. Eur Heart J 2012;33:2821–30.

Fox KAA, Piccini JP, Wojdyla D, et al. Prevention of stroke and systemic embolism with rivaroxaban compared with warfarin in patients with nonvalvular atrial fibrillation and moderate renal impairment. Eur Heart J 2011;32:2387–94.

Bhatt DLO. PIONEERs! the beginning of the end of full-dose triple therapy in atrial fibrillation. Circulation 2017;135:334–366.

Dahal K, Kunwar S, Rijal J, et al. Stroke, major bleeding, and mortality outcomes in warfarin users with atrial fibrillation and chronic kidney disease: a meta-analysis of observational studies. Chest 2016;149:951–60.

Hijazi Z, Hohnloser SH, Oldgren J, et al. Efficacy and safety of dabigatran compared with warfarin in relation to baseline renal function in patients with atrial fibrillation: a RE-LY (Randomized evaluation of long-term anticoagulation therapy) analysis. Circulation 2014;129:591–70.

Hohnloser SH, Hijazi Z, Thomas L, et al. Efficacy of apixaban when compared with warfarin in relation to renal function in patients with atrial fibrillation: insights from the ARISTOTLE trial. Eur Heart J 2012;33:2821–30.

Fox KAA, Piccini JP, Wojdyla D, et al. Prevention of stroke and systemic embolism with rivaroxaban compared with warfarin in patients with nonvalvular atrial fibrillation and moderate renal impairment. Eur Heart J 2011;32:2387–94.

Bhatt DLO. PIONEERs! the beginning of the end of full-dose triple therapy in atrial fibrillation. Circulation 2017;135:334–366.

Dahal K, Kunwar S, Rijal J, et al. Stroke, major bleeding, and mortality outcomes in warfarin users with atrial fibrillation and chronic kidney disease: a meta-analysis of observational studies. Chest 2016;149:951–60.

Hijazi Z, Hohnloser SH, Oldgren J, et al. Efficacy and safety of dabigatran compared with warfarin in relation to baseline renal function in patients with atrial fibrillation: a RE-LY (Randomized evaluation of long-term anticoagulation therapy) analysis. Circulation 2014;129:591–70.

Hohnloser SH, Hijazi Z, Thomas L, et al. Efficacy of apixaban when compared with warfarin in relation to renal function in patients with atrial fibrillation: insights from the ARISTOTLE trial. Eur Heart J 2012;33:2821–30.

Fox KAA, Piccini JP, Wojdyla D, et al. Prevention of stroke and systemic embolism with rivaroxaban compared with warfarin in patients with nonvalvular atrial fibrillation and moderate renal impairment. Eur Heart J 2011;32:2387–94.

Bhatt DLO. PIONEERs! the beginning of the end of full-dose triple therapy in atrial fibrillation. Circulation 2017;135:334–366.

Dahal K, Kunwar S, Rijal J, et al. Stroke, major bleeding, and mortality outcomes in warfarin users with atrial fibrillation and chronic kidney disease: a meta-analysis of observational studies. Chest 2016;149:951–60.

Hijazi Z, Hohnloser SH, Oldgren J, et al. Efficacy and safety of dabigatran compared with warfarin in relation to baseline renal function in patients with atrial fibrillation: a RE-LY (Randomized evaluation of long-term anticoagulation therapy) analysis. Circulation 2014;129:591–70.

Hohnloser SH, Hijazi Z, Thomas L, et al. Efficacy of apixaban when compared with warfarin in relation to renal function in patients with atrial fibrillation: insights from the ARISTOTLE trial. Eur Heart J 2012;33:2821–30.

Fox KAA, Piccini JP, Wojdyla D, et al. Prevention of stroke and systemic embolism with rivaroxaban compared with warfarin in patients with nonvalvular atrial fibrillation and moderate renal impairment. Eur Heart J 2011;32:2387–94.

Bhatt DLO. PIONEERs! the beginning of the end of full-dose triple therapy in atrial fibrillation. Circulation 2017;135:334–366.
Seidl K, Rameken M, Drögemüller A, et al. Embolic events in patients with atrial fibrillation and effective anticoagulation: value of transesophageal echocardiography to guide direct-current cardioversion: final results of the Ludwigsafen observational cardioversion study. J Am Coll Cardiol 2002;39:1436–42.

Nagarakanti R, Ezekowitz MD, Oldgren J, et al. Dabigatran versus warfarin in patients with atrial fibrillation: an analysis of patients undergoing cardioversion. Circulation 2011;123:131–6.

Piccini JP, Stevens SR, Lohynchina Y, et al. Outcomes after cardioversion and atrial fibrillation ablation in patients treated with rivaroxaban and warfarin in the ROCKET AF trial. J Am Coll Cardiol 2013;61:1998–2006.

Flaker G, Lopes RD, Al-Khatib SM, et al. Efficacy and safety of apixaban in patients after cardioversion for atrial fibrillation: insights from the ARISTOTLE trial (apixaban for reduction in stroke and other thromboembolic events in atrial fibrillation). J Am Coll Cardiol 2014;63:1082–7.

Caputo R, Ezekowitz MD, Klein AL, et al. Rivaroxaban vs. vitamin K antagonists for cardioversion in atrial fibrillation. Eur Heart J 2014;35:3346–55.

Goette A, Merino JL, Ezekowitz MD, et al. Edoxaban versus enoxaparin–warfarin in patients undergoing cardioversion of atrial fibrillation (ENSURE–AF): a randomised, open-label, phase 3b trial. Lancet 2016;388:1995–2003.

Klein AL, Grimm RA, Murray RD, et al. Use of transesophageal echocardiography to guide cardioversion in patients with atrial fibrillation. N Engl J Med 2001;344:1411–20.

Weigner MJ, Thomas LR, Patel U, et al. Early cardioversion of atrial fibrillation facilitated by transesophageal echocardiography: short-term safety and impact on maintenance of sinus rhythm at 1 year. Am J Med 2001;110:694–702.

Wu LA, Chandrasekaran K, Friedman PA, et al. Safety of expedited anticoagulation in patients undergoing transesophageal echocardiographic-guided cardioversion. Am J Med 2006;119:142–6.

Jaber WA, Prior DL, Thamilarasan M, et al. Efficacy of anticoagulation in resolving left atrial and left atrial appendage thrombus: a transesophageal echocardiographic study. Am Heart J 2000;140:150–6.

Weigner MJ, Caulfield TA, Darias PG, et al. Risk for clinical thromboembolism associated with conversion to sinus rhythm in patients with atrial fibrillation lasting less than 48 hours. Ann Intern Med 1997;126:615–20.

Sticherling C, Marin BJ, Binnie D, et al. Antithrombotic management in patients undergoing electrophysiological procedures: a European Heart Rhythm Association (EHRA) position document endorsed by the ESC working group thrombosis, heart rhythm society (HRS), and Asia Pacific heart rhythm society (APHRS). Europace 2015;17:197–214.

Di Biase L, Burkhardt JD, Mohanty P, et al. Periprocedural stroke and management of major bleeding complications in patients undergoing catheter ablation of atrial fibrillation. Impact Periprocedural Ther Int Norm Ratio Circ 2010;121:2550–6.

Santangeli P, Di Biase L, Horton R, et al. Ablation of atrial fibrillation under therapeutic warfarin reduces periprocedural complications: evidence from a meta-analysis. Circ Arrhythm Electrophysiol 2012;5:302–11.

Kim JS, She F, Jongsarangsin K, et al. Dabigatran vs warfarin for radiofrequency catheter ablation of atrial fibrillation. Heart Rhythm 2013;10:483–9.

Bassiony M, Saliba W, Rickard J, et al. Use of dabigatran for periprocedural anticoagulation in patients undergoing catheter ablation for atrial fibrillation. Circ Arrhythm Electrophysiol 2013;6:460–6.

Wazni OM, Beheiry S, Fahmy T, et al. Atrial fibrillation ablation in patients with therapeutic international normalized ratio. Comp Strateg Anticoagulation Manag Periprocedural Period Circ 2007;116:2531.

Schmitt M, Segerson NM, Marschang H, et al. Atrial fibrillation ablation in patients with therapeutic international normalized ratio. Pacing Clin Electrophysiol 2009;32:995–9.

Caputo R, Marchlinski FE, Hohnloser SH, et al. Uninterrupted rivaroxaban vs. uninterrupted vitamin K antagonists for catheter ablation in non-valvular atrial fibrillation. Eur Heart J 2015;36:1805–11.

Calkins H, Willems EF, Linz W, et al. Uninterrupted dabigatran versus warfarin for ablation in atrial fibrillation. N Engl J Med 2017;376:1627–36.

Calkins H, Kuck KH, Caputo R, et al. 2012 I/EHRA/ECAS expert consensus statement on catheter and surgical ablation of atrial fibrillation: recommendations for patient selection, procedural techniques, patient management and follow-up, definitions, endpoints, and research trial design: a report of the heart rhythm society (HRS) task force on catheter and surgical ablation of atrial fibrillation. Developed in partnership with the European Heart Rhythm Association (EHRA), a registered branch of the European Society of Cardiology (ESC) and the European Cardiac Arrhythmia Society (ECAS); and in collaboration with the American College of Cardiology (ACC), American Heart Association (AHA), the Asia Pacific Heart Rhythm Society (APHRS), and the Society of Thoracic Surgeons (STS). Endorsed by the governing bodies of the American College of Cardiology Foundation, the American Heart Association, the European Cardiac Arrhythmia Society, the European Heart Rhythm Association, the Society for Thoracic Surgeons, the Asia Pacific Heart Rhythm Society, and the Heart Rhythm Society. Heart Rhythm 2012;9:632–96.

Fridh M, Eitel C, Bollmann A, et al. Should transesophageal echocardiography be done in all patients who underwent catheter ablation of atrial fibrillation? A case report and review of the literature Clin Res Cardiol 2010;99:125–8.

Puwavanant S, Varr BC, Shrestha K, et al. Role of the CHADS2Score in the evaluation of thromboembolic risk in patients with atrial fibrillation undergoing transesophageal echocardiography before pulmonary vein isolation. J Am Coll Cardiol 2009;54:2032–9.

Scherr D, Sharma K, Dalal D, et al. Incidence and predictors of periprocedural cerebrovascular accident in patients undergoing catheter ablation of atrial fibrillation. Ann Pharmacother 2015;49:278–84.

May MA, Gruel Y, Faucher L, Letter by May et al regarding article, “Use of dabigatran for periprocedural anticoagulation in patients undergoing catheter ablation for atrial fibrillation” by Bassiony et al. Circ Arrhythm Electrophysiol 2013;6:e55.

Di Biase L, Gaita F, Toso E, et al. Does periprocedural anticoagulation management of atrial fibrillation affect the prevalence of silent thromboembolic lesion detected by diffusion cerebral magnetic resonance imaging in patients undergoing radiofrequency atrial fibrillation ablation with open irrigated catheters? Results from a prospective multicenter study Heart Rhythm 2014;11:791–8.

Caschera F, Extramiana F, Caschera S, et al. High-flow perfusion of sheaths for prevention of thromboembolic complications during complex catheter ablation in the left atrium. J Cardiovasc Electrophysiol 2004;15:276–83.

Karasyo D, Gislason GH, Hansen J, et al. Oral anticoagulation therapy after radiofrequency ablation of atrial fibrillation and the risk of thromboembolism and serious bleeding: long-term follow-up in nationwide cohort of Denmark. Eur Heart J 2015;36:307–15.

Lakkireddy D, Reddy YM, Di Biase L, et al. Feasibility and safety of dabigatran versus warfarin for periprocedural anticoagulation in patients undergoing radiofrequency ablation for atrial fibrillation: results from a multicenter prospective registry. J Am Coll Cardiol 2012;59:1168–74.

Gladstone DJ, Sprig M, Doran P, et al. Atrial fibrillation in patients with mechanical heart valves: assessment of bleeding risk during oral anticoagulation by direct and indirect inhibitors of coagulation factor Xa. Nat Med 2003;9:2003.

Armbruster HL, Lindsley JP, Moranville MP, et al. Safety of novel oral anticoagulants compared with uninterrupted warfarin for catheter ablation of atrial fibrillation. Ann Pharmacother 2015;49:278–84.

Schnabel C, Extramiana F, Caschera S, et al. High-flow perfusion of sheaths for prevention of thromboembolic complications during complex catheter ablation in the left atrium. J Cardiovasc Electrophysiol 2004;15:276–83.

Karasoy D, Gislason GH, Hansen J, et al. Oral anticoagulation therapy after radiofrequency ablation of atrial fibrillation and the risk of thromboembolism and serious bleeding: long-term follow-up in nationwide cohort of Denmark. Eur Heart J 2015;36:307–15.

Lakkireddy D, Reddy YM, Di Biase L, et al. Feasibility and safety of dabigatran versus warfarin for periprocedural anticoagulation in patients undergoing radiofrequency ablation for atrial fibrillation: results from a multicenter prospective registry. J Am Coll Cardiol 2012;59:1168–74.

Glodstone DJ, Sprig M, Doran P, et al. Atrial fibrillation in patients with mechanical heart valves: assessment of bleeding risk during oral anticoagulation by direct and indirect inhibitors of coagulation factor Xa. Nat Med 2003;9:2003.