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2016 ESC Guidelines for the Management of Atrial Fibrillation Developed in Collaboration With EACTS

ESC Scientific Document Group; Kirchhof, Paulus; Benussi, Stefano; Kotecha, Dipak; Ahlsson, Anders; Atar, Dan; Casadei, Barbara; Castellá, Manuel; Diener, Hans-Christoph; Heidbuchel, Hein; Hendriks, Jeroen; Hindricks, Gerhard; Manolis, Antonis S; Oldgren, Jonas; Alexandru Popescu, Bogdan; Schotten, Ulrich; Van Putte, Bart; Vardas, Panagiotis *DOI:*

10.1016/j.rec.2016.11.033

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Document Version Peer reviewed version

Citation for published version (Harvard):

ESC Scientífic Document Group, Kirchhof, P, Benussi, S, Kotecha, D, Ahlsson, A, Atar, D, Casadei, B, Castellá, M, Diener, H-C, Heidbuchel, H, Hendriks, J, Hindricks, G, Manolis, AS, Oldgren, J, Alexandru Popescu, B, Schotten, U, Van Putte, B & Vardas, P 2017, '2016 ESC Guidelines for the Management of Atrial Fibrillation Developed in Collaboration With EACTS', Revista española de cardiología (English ed.), vol. 70, no. 1, pp. 50-50. https://doi.org/10.1016/j.rec.2016.11.033

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10.1093/europace/euw295

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Document Version Peer reviewed version

Citation for published version (Harvard):

Kirchhof, P, Benussi, S, Kotecha, D, Ahlsson, A, Atar, D, Casadei, B, Castella, M, Diener, H-C, Heidbuchel, H, Hendriks, J, Hindricks, G, Manolis, AS, Oldgren, J, Popescu, BA, Schotten, U, Van Putte, B, Vardas, P & Authors/Task Force Members 2016, '2016 ESC Guidelines for the management of atrial fibrillation developed in collaboration with EACTS: The Task Force for the management of atrial fibrillation of the European Society of Cardiology (ESC). Developed with the special contribution of the European Heart Rhythm Association (EHRA) of the ESC. Endorsed by the European Stroke Organisation (ESO).' Europace. DOI: 10.1093/europace/euw295

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2016 ESC Guidelines for the management of atrial fibrillation developed in collaboration with EACTS

The Task Force for the management of atrial fibrillation of the European Society of Cardiology (ESC)

Developed with the special contribution of the European Heart Rhythm Association (EHRA) of the ESC

Endorsed by the European Stroke Organisation (ESO)

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The disclosure forms of all experts involved in the development of these guidelines are available on the ESC website <u>www.escardio.org/guidelines</u>

37 Keywords:

- 38 Guidelines Atrial fibrillation Anticoagulation Vitamin K antagonists Non vitamin-K-antagonist oral
- 39 anticoagulants Left atrial appendage occlusion Rate control Cardioversion Rhythm control -
- 40 Antiarrhythmic drugs Upstream therapy Catheter ablation AF surgery Valve repair Pulmonary vein 41 isolation - Left atrial ablation
- 42

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Working Groups: Cardiac Cellular Electrophysiology, Cardiovascular Pharmacotherapy

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180		s and acronyms
181	ABC	age, biomarkers, clinical history
182	ACE	angiotensin-converting enzyme
183	ACS	acute coronary syndromes
184	AF	atrial fibrillation
185	AFFIRM	Atrial Fibrillation Follow-up Investigation of Rhythm Management
186	AFNET	German Competence NETwork on Atrial Fibrillation
187	AHRE	atrial high rate episodes
188	ARB	angiotensin receptor blocker
189	ARISTOTLE	Apixaban for Reduction in Stroke and Other Thromboembolic Events in Atrial Fibrillation
190	ARNI	angiotensin receptor neprilysin inhibition
191	ATRIA	AnTicoagulation and Risk factors In Atrial fibrillation
192	AXAFA	Anticoagulation using the direct factor Xa inhibitor apixaban during Atrial Fibrillation
193		catheter Ablation: Comparison to vitamin K antagonist therapy
194	BAFTA	Birmingham Atrial Fibrillation Treatment of the Aged Study
195	BMI	body mass index
196	bpm	beats per minute
197	CABANA	Catheter Ablation versus Antiarrhythmic Drug Therapy for Atrial Fibrillation Trial
198	CAD	coronary artery disease
199	CHA ₂ DS ₂ -VASc	Congestive Heart failure, hypertension, Age \geq 75 (doubled), Diabetes, Stroke (doubled),
200		Vascular disease, Age 65–74, and Sex (female)
201	$CHADS_2$	Cardiac failure, Hypertension, Age, Diabetes, Stroke (Doubled)
202	CI	confidence interval
203	CKD	chronic kidney disease
204	CrCl	creatinine clearance
205	CT	computed tomography
206	DIG	Digitalis Investigation Group
207	EACTS	European Association for Cardio-Thoracic Surgery
208	EAST	Early treatment of Atrial fibrillation for Stroke prevention Trial
209	ECG	electrocardiogram/electrocardiography
210	EHRA	European Heart Rhythm Association
211	ENGAGE AF-TIM	
212		Thrombolysis in Myocardial Infarction 48
213	EORP	EURObservational Research Programme
214	FAST	Atrial Fibrillation Catheter Ablation vs Surgical Ablation Treatment
215	FEV1	forced expiratory volume in 1 second
216	GDF-15	growth differentiation factor 15
217	GFR	glomerular filtration rate
218	GFR	glomerular filtration rate
219	GUCH	grown up congenital heart disease
220	HARMONY	A Study to Evaluate the Effect of Ranolazine and Dronedarone When Given Alone and in
$\frac{1}{221}$		Combination in Patients With Paroxysmal Atrial Fibrillation
222	HAS-BLED	hypertension, abnormal renal/liver function (1 point each), stroke, bleeding history or
$\frac{1}{223}$		predisposition, labile INR, elderly (>65 years), drugs/alcohol concomitantly (1 point each)
224	HFmrEF	heart failure with mid-range ejection fraction
225	HFpEF	heart failure with preserved ejection fraction
226	HFrEF	heart failure with reduced ejection fraction
227	HR	hazard ratio
228	INR	international normalized ratio
229	LA	left atrium/atrial
230	LAA	left atrial appendage
231	LAAOS	Left Atrial Appendage Occlusion Study
232	LV	left ventricular
232	LVEF	left ventricular ejection fraction
233	LVH	left ventricular hypertrophy
235	MANTRA-PAF	Medical ANtiarrhythmic Treatment or Radiofrequency Ablation in Paroxysmal Atrial
235	1111 11 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	Fibrillation
230	MERLIN	Metabolic Efficiency With Ranolazine for Less Ischemia in Non ST-Elevation Acute
237		Coronary Syndrome
238	MRI	magnetic resonance imaging
	171111	

240	NOAC	non-vitamin K antagonist oral anticoagulant
241	NYHA	New York Heart Association
242	OAC	oral anticoagulation/oral anticoagulant
243	OR	odds ratio
244	ORBIT	Outcomes Registry for Better Informed Treatment of Atrial Fibrillation
245	PCI	percutaneous coronary intervention
246	PREVAIL	Prospective Randomized Evaluation of the Watchman LAA Closure Device In Patients
247		with AF Versus Long Term Warfarin Therapy trial
248	PROTECT AF	Watchman Left Atrial Appendage System for Embolic Protection in Patients With AF trial
249	PVI	pulmonary vein isolation
250	RACE	Rate Control Efficacy in Permanent Atrial Fibrillation
251	RATE-AF	Rate Control Therapy Evaluation in Permanent Atrial Fibrillation
252	RCT	randomized controlled trial
253	RE-CIRCUIT	Randomized Evaluation of dabigatran etexilate Compared to warfarin in pulmonaRy vein
254		ablation: assessment of different peri-proCedUral anticoagulation sTrategies
255	RE-LY	Randomized Evaluation of Long-Term Anticoagulation Therapy
256	ROCKET-AF	Rivaroxaban Once Daily Oral Direct Factor Xa Inhibition Compared with Vitamin K
257		Antagonism for Prevention of Stroke and Embolism Trial in Atrial Fibrillation
258	RR	risk ratio
259	SD	standard deviation
260	SPAF	Stroke Prevention in Atrial Fibrillation
261	TIA	transient ischaemic attack
262	TIMI	Thrombolysis In Myocardial Infarction
263	TOE	transoesophageal echocardiography
264	TTR	time in therapeutic range
265	UFH	unfractionated heparin
266	US	United States
267	VKA	vitamin K antagonist
268	WOEST	What is the Optimal antiplatElet and anticoagulant therapy in patients with oral
269		anticoagulation and coronary StenTing
270	WPW	Wolff-Parkinson-White syndrome
271		
272		

Preamble 1 273

274 Guidelines summarize and evaluate all available evidence on a particular issue at the time of the writing process, 275 with the aim of assisting health professionals in selecting the best management strategies for an individual 276 patient with a given condition, taking into account the impact on outcome, as well as the risk-benefit ratio of 277 particular diagnostic or therapeutic means. Guidelines and recommendations should help health professionals to 278 make decisions in their daily practice. However, the final decisions concerning an individual patient must be 279 made by the responsible health professional(s) in consultation with the patient and caregiver as appropriate.

280 A great number of Guidelines have been issued in recent years by the European Society of Cardiology (ESC) 281 and by the European Association for Cardio-Thoracic Surgery (EACTS), as well as by other societies and 282 organisations. Because of the impact on clinical practice, quality criteria for the development of guidelines have 283 been established in order to make all decisions transparent to the user. The recommendations for formulating 284 and issuing ESC Guidelines can be found on the ESC website (http://www.escardio.org/Guidelines-&-285 Education/Clinical-Practice-Guidelines/Guidelines-development/Writing-ESC-Guidelines). ESC Guidelines 286 represent the official position of the ESC on a given topic and are regularly updated.

287 Members of this Task Force were selected by the ESC, including representation from the European Heart 288 Rhythm Association (EHRA), and EACTS as well as by the European Stroke Organisation (ESO) to represent 289 professionals involved with the medical care of patients with this pathology. Selected experts in the field 290 undertook a comprehensive review of the published evidence for management (including diagnosis, treatment, 291 prevention and rehabilitation) of a given condition according to ESC Committee for Practice Guidelines (CPG) 292 policy and approved by the EACTS and ESO. A critical evaluation of diagnostic and therapeutic procedures was 293 performed, including assessment of the risk-benefit ratio. Estimates of expected health outcomes for larger 294 populations were included, where data exist. The level of evidence and the strength of the recommendation of 295 particular management options were weighed and graded according to predefined scales, as outlined in Tables 1 296 and 2.

297 The experts of the writing and reviewing panels provided declaration of interest forms for all relationships that

298 might be perceived as real or potential sources of conflicts of interest. These forms were compiled into one file

299 and can be found on the ESC website (http://www.escardio.org/guidelines). Any changes in declarations of

300 interest that arise during the writing period must be notified to the ESC and EACTS and updated. The Task 301 Force received its entire financial support from the ESC and EACTS without any involvement from the

302 healthcare industry.

303 The ESC CPG supervises and coordinates the preparation of new Guidelines produced by task forces, expert 304 groups or consensus panels. The Committee is also responsible for the endorsement process of these Guidelines. 305 The ESC Guidelines undergo extensive review by the CPG and external experts, and in this case by EACTS and 306 ESO-appointed experts. After appropriate revisions the Guidelines are approved by all the experts involved in 307 the Task Force. The finalized document is approved by the CPG, EACTS and ESO for publication in the 308 European Heart Journal, Europace, and in the European Journal of Cardio-Thoracic Surgery as well as in the 309 International Journal of Stroke (TBC). The Guidelines were developed after careful consideration of the 310 scientific and medical knowledge and the evidence available at the time of their dating.

311 The task of developing ESC and EACTS Guidelines covers not only integration of the most recent research, but 312 also the creation of educational tools and implementation programmes for the recommendations. To implement 313

the guidelines, condensed pocket guideline versions, summary slides, booklets with essential messages,

314 summary cards for non-specialists and an electronic version for digital applications (smartphones, etc.) are 315

produced. These versions are abridged and thus, if needed, one should always refer to the full text version, 316 which is freely available on the ESC website. The National Societies of the ESC are encouraged to endorse,

317

translate and implement all ESC Guidelines. Implementation programmes are needed because it has been shown 318 that the outcome of disease may be favourably influenced by the thorough application of clinical

319 recommendations.

320 Surveys and registries are needed to verify that real-life daily practice is in keeping with what is recommended 321 in the guidelines, thus completing the loop between clinical research, writing of guidelines, disseminating them 322 and implementing them into clinical practice.

323 Health professionals are encouraged to take the ESC and EACTS Guidelines fully into account when exercising 324 their clinical judgment, as well as in the determination and the implementation of preventive, diagnostic or

325 therapeutic medical strategies. However, the ESC and EACTS Guidelines do not override in any way

326 whatsoever the individual responsibility of health professionals to make appropriate and accurate decisions in 327 consideration of each patient's health condition and in consultation with that patient and the patient's caregiver

328 where appropriate and/or necessary. It is also the health professional's responsibility to verify the rules and 329 regulations applicable to drugs and devices at the time of prescription.

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333 **Table 1 Classes of recommendations**

Table 1: Classes of Recommendations						
Classes of Recommendations	Definition	Suggested wording to use				
Class I	Evidence and/or general agreement that a given treatment or procedure is beneficial, useful, effective.	ls recommended/is indicated				
Class II	Conflicting evidence and/or a divergence of opinion about the usefulness/efficacy of the given treatment or procedure.					
Class lla	Weight of evidence/opinion is in favour of usefulness/efficacy.	Should be considered				
Class Ilb	Usefulness/efficacy is less well established by evidence/opinion.	May be considered				
Class III	Evidence or general agreement that the given treatment or procedure is not useful/effective, and in some cases may be harmful.	Is not recommended				

Table 2 Levels of evidence

Level of Evidence A	Data derived from multiple randomized clinical trials or meta-analyses.	
Level of Evidence B	Data derived from a single randomized clinical trial or large non-randomized studies.	
Level of Evidence C	Consensus of opinion of the experts and/or small studies, retrospective studies, registries.	

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340 **2 Introduction**

Despite good progress in the management of patients with atrial fibrillation (AF), this arrhythmia remains one of the major causes of stroke, heart failure, sudden death, and cardiovascular morbidity in the world. Furthermore, the number of patients with AF is predicted to rise steeply in the coming years. To meet the growing demand for effective care of patients with AF, new information is continually generated and published, and the last few years have seen substantial progress. It therefore seems timely to publish this 2nd edition of the ESC guidelines on AF.

Reflecting the multidisciplinary input into the management of patients with AF, the Task Force includes cardiologists with varying subspecialty expertise, cardiac surgeons, stroke neurologists, and specialist nurses amongst its members. Supplementing the evidence review as outlined in the preamble, this task force identified three PICOT questions on relevant topics for the guideline. The ESC commissioned external systematic reviews to answer these three questions. These reviews informed specific recommendations.

Further to adhering to the standards for generating recommendations that is common to all ESC guidelines (see preamble), this task force discussed each draft recommendation during web-based conference calls dedicated to specific chapters, followed by consensus modifications and an online vote on each recommendation. Only recommendations that were supported by at least 75% of the task force members were included in the guideline.

We hope that this guideline will help to deliver good care to all patients with AF based on the current state-ofthe-art evidence in 2016.

362 **3 Epidemiology and impact for patients**

363 3.1. Incidence and prevalence of atrial fibrillation

364 In 2010, the estimated numbers of men and women with atrial fibrillation (AF) worldwide were 20.9 million and 12.6 million, respectively, with higher incidence and prevalence rates in developed countries.^{1, 2} One in four 365 middle-aged adults in Europe and the United States (US) will develop AF.³⁻⁵ By 2030, 14–17 million AF 366 patients are anticipated in the European Union, with 120,000-215,000 newly diagnosed patients per year.^{2, 6, 7} 367 368 Estimates suggest an AF prevalence of approximately 3% in adults age 20 years or older.^{8,9} with more AF in elderly persons¹ and in patients with conditions such as hypertension, heart failure, coronary artery disease (CAD), valvular heart disease, diabetes mellitus, and chronic kidney disease (CKD).^{7, 10-15} The increase in AF 369 370 prevalence can be attributed to better detection of silent AF^{16-18} and increasing age and conditions predisposing 371 372 to AF.¹⁹

373

374 3.2. Morbidity, mortality, and healthcare burden of atrial fibrillation

AF is independently associated with a twofold increased risk of all-cause mortality in women and a 1.5-fold increase in men²⁰⁻²² (*Table 3*). Death due to stroke can largely be mitigated by anticoagulation, while other cardiovascular deaths, for example due to heart failure and sudden death, remain common even in AF patients treated according to the current evidence-base.²³ AF is also associated with increased morbidity, such as heart failure and stroke.^{21, 24, 25} Contemporary studies show that 20–30% of patients with an ischaemic stroke have AF diagnosed before, during, or after the initial event.^{17, 26, 27} White matter lesions in the brain, cognitive impairment,²⁸⁻³⁰ decreased quality of life,^{31, 32} and depressed mood³³ are common in AF patients, and between 10% and 40% of AF patients are hospitalized each year.^{23, 34, 35}

The direct costs of AF already amount to approximately 1% of total healthcare spending in the UK, and between \$6.0 and \$26.0 billion in the US for 2008,^{36,37} driven by AF-related complications (e.g. stroke) and AF-

between \$6.0 and \$26.0 billion in the US for 2008,^{36,37} driven by AF-related complications (e.g. stroke) and AFrelated treatment costs (e.g. hospitalizations). These costs will increase dramatically unless AF is prevented and treated in a timely and effective manner.

387 388

Event	Association with AF		
Death	Increased mortality, especially cardiovascular mortality due to sudden death, heart		
	failure, or stroke		
Stroke	20–30% of all strokes are due to AF. A growing number of patients with stroke are		
	diagnosed with 'silent', paroxysmal AF		
Hospitalizations	10-40% of AF patients are hospitalized every year		
Quality of life	Quality of life is impaired in AF patients independent of other cardiovascular		
	conditions		
LV dysfunction and	LV dysfunction is found in 20-30% of all AF patients. AF causes or aggravates LV		
heart failure	dysfunction in many AF patients, while others have completely preserved LV		
	function despite long-standing AF		
Cognitive decline and	Cognitive decline and vascular dementia increase even in anticoagulated patients.		
vascular dementia	Brain white matter lesions are more common in AF patients than in patients without		
	AF		

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AF = atrial fibrillation; LV = left ventricular.

390

391 3.3. Impact of evidence-based management on outcomes in atrial fibrillation 392 patients

Figure 1 depicts the major milestones in the management of AF. Despite these advances, substantial morbidity
 remains. Oral anticoagulation (OAC) with vitamin K antagonists (VKAs) or non-VKA oral anticoagulants

395 (NOACs) markedly reduces stroke and mortality in AF patients.^{38, 39} Other interventions such as rhythm control

and rate control improve AF-related symptoms and may preserve cardiac function, but have not demonstrated a $\frac{40}{41}$

397 reduction in long-term morbidity or mortality.^{40, 41}

First maze surgery for 000 AF treatment VKA superior to aspirin published for stroke prevention in ΔF PVI can suppress AF ACEi/ARBs prevent RF based maze AF in heart failure maintains SR after Rate control not inferior to rhythm control cardiovascular VKA reduces stroke in AF surgerv by 2/3 ARBs prevent AF in **PVI** maintains SR hypertension & LVH better than 2005 Ximelagatran as effective antiarrhythmic drugs as VKA Amiodarone not superior to rate ARBs do not prevent Dronedarone improves Dabigatran at least as control in heart outcomes in non-AF or adverse effective as VKA in AF failure outcomes in patients permanent AF without hypertension Lenient rate control AF ablation acceptable improves QoL Rixaroxaban and PUFA do not Apixaban at least as prevent AF Dronedarone harms effective as VKA in AF in permanent AF First-line PVI MRA prevent AF in Bipolar RF more maintains SR better HFrEF patients pre-Edoxaban at least as than antiarrhythmic effective than treated with ACEi/ conventional RF for effective as VKA in AF drugs beta-blockers stand-alone AF ACEi/ARBs prevent surgerv AF in hypertension Meta-analysis and healthcare PVI alone as Beta-blockers without effective as databases: NOACs safer and prognostic benefit in Beta-blockers complex ablation in Concomitant maze slightly more effective AF patients with prevent AF in HFrEF persistent AF surgery maintains SR 2015 compared to VKA HFrEF patients pre-treated but increases risk of Cryoenergy as with ACEi permanent pacemaker effective as RF for PVI

398 399 400 401 402 403 404 405

Figure 1 Timeline of major landmarks in AF management, including treatment of concomitant conditions and prevention (green), anticoagulation (blue), rate and rhythm control (orange and red), and surgical therapy (purple).
ACEi = angiotensin-converting enzyme inhibitor; AF = atrial fibrillation; ARB = angiotensin receptor blocker;

HF = heart failure; HFrEF = heart failure with reduced ejection fraction; LVH = left ventricular hypertrophy;
NOAC = non-vitamin K antagonist oral anticoagulant; PUFA = polyunsaturated fatty acid; PVI = pulmonary
vein isolation; QoL = quality of life; RACE = Rate Control Efficacy in Permanent Atrial Fibrillation; RF =
radiofrequency; SR = sinus rhythm; VKA = vitamin K antagonist.

In contemporary, well-controlled, randomized clinical trials in AF, the average annual stroke rate is about 1.5%
and the annualized death rate is around 3%.⁴⁰ In real life, the annual mortality can be different (both higher and lower).⁴² A minority of these deaths are related to stroke, while sudden cardiac death and death from progressive heart failure are more frequent, emphasizing the need for interventions beyond anticoagulation.^{43, 44}
Furthermore, AF is also associated with high rates of hospitalization, commonly for AF management, but often

414 also for heart failure, myocardial infarction, and treatment-associated bleeding.^{34, 45}

415

416 **3.4.** Gender

417 In both developed and developing countries, the age-adjusted incidence and prevalence of AF are lower in women, while the risk of death in women with AF is similar to or higher than that in men with AF.^{1, 46, 47} Female 418 419 AF patients who have additional stroke risk factors (particularly older age) are also at greater risk than men of having a stroke, 48,49 even those anticoagulated with warfarin⁵⁰ (see Chapter 8 for details). Women with 420 diagnosed AF can be more symptomatic than men and are typically older with more comorbidities.^{51, 52} Bleeding risk on anticoagulation is similar in both sexes,^{49, 50, 53} but women appear less likely to receive 421 422 specialist care and rhythm control therapy,⁵⁴ while the outcomes of catheter ablation or AF surgery are comparable to those in men.^{55, 56} These observations highlight the need to offer effective diagnostic tools and 423 424 425 therapeutic management equally in women and men. 426

427 **Recommendations relating to gender**

Recommendations	Class ^a	Level ^b	Refs ^c
AF clinicians must offer effective diagnostic tools and therapeutic management to women and men equally to prevent stroke and death	I	A	39, 46, 57
Catheter or surgical ablation techniques should be regarded as equally effective in women and men	lla	В	55, 56

428 AF = atrial fibrillation 429

^aClass of recommendation.

430 ^bLevel of evidence. ^cReference(s) supporting recommendations.

431 432

433 Pathophysiological and genetic aspects that guide management 4

434 4.1. Genetic predisposition

435 AF, especially early-onset AF, has a strong heritable component, independent of concomitant cardiovascular 436 conditions.^{58, 59} A few young AF patients suffer from inherited cardiomyopathies or channelopathies mediated 437 by disease-causing mutations. These monogenic diseases also convey a risk for sudden death (see Chapter 5). 438 Up to one-third of AF patients carry common genetic variants that predispose to AF, albeit with a relatively low 439 added risk. At least 14 of these common variants, often single nucleotide polymorphisms, are known to increase the risk of prevalent AF in populations.⁶⁰⁻⁶² The most important variants are located close to the paired-like 440 homeodomain transcription factor 2 gene on chromosome 4q25.^{63, 64} These variants modify the risk of AF up to 441 442 sevenfold.⁶⁴ Several of the AF risk variants are also associated with cardioembolic or ischaemic stroke, possibly due to silent AF (see section 4.1).^{62, 65, 66} Changes in atrial action potential characteristics,⁶⁷⁻⁷⁰ atrial remodelling, 443 444 and modified penetration of rare gene defects⁶¹ have been suggested as potential mechanisms mediating increased AF risk in carriers of common gene variants. Genetic variants could in the future become useful for patient selection of rhythm control strategies,⁷¹⁻⁷³ but it is currently unknown whether common gene variants 445 446 447 differentially affect the efficacy of antiarrhythmic drugs or rate control medication.⁷⁴ While genomic analysis may provide an opportunity to improve diagnosis and management of AF in the future,^{75, 76} routine genetic 448 449 testing for common gene variants associated with AF cannot be recommended at present.⁷⁷

450

451 4.2. Mechanisms leading to atrial fibrillation

452 4.2.1. Remodelling of atrial structure and ion channel function

453 External stressors such as structural heart disease, hypertension, possibly diabetes, but also AF itself induce a 454 slow but progressive process of structural remodelling in the atria (Figure 2). Activation of fibroblasts, 455 enhanced connective tissue deposition, and fibrosis are the hallmarks of this process.⁷⁸⁻⁸⁰ In addition, atrial fatty infiltration, inflammatory infiltrates, myocyte hypertrophy, necrosis, and amyloidosis are found in AF patients with concomitant conditions predisposing to AF.⁸¹⁻⁸⁴ Structural remodelling results in electrical dissociation 456 457 between muscle bundles and local conduction heterogeneities,⁸⁵ favouring reentry and perpetuation of the 458 459 arrhythmia.⁸⁶ In many patients, the structural remodelling process occurs before the onset of AF.⁷⁸ As some of 460 the structural remodelling will be irreversible, early initiation of treatment seems desirable.⁸⁷ Table 4 gives an 461 overview of the most relevant pathophysiological alterations in atrial tissue associated with AF, and lists 462 corresponding clinical conditions that can contribute to these changes.

463 The functional and structural changes in atrial myocardium and stasis of blood, especially in the left 464 atrial appendage (LAA), generate a prothrombotic milieu. Furthermore, even short episodes of AF lead to 465 myocardial damage and expression of prothrombotic factors on the atrial endothelial surface, and activation of platelets and inflammatory cells, and contribute to a generalized prothrombotic state.^{88, 89} The atrial and 466 467 systemic activation of the coagulation system can partially explain why short episodes of AF convey a long-468 term stroke risk.

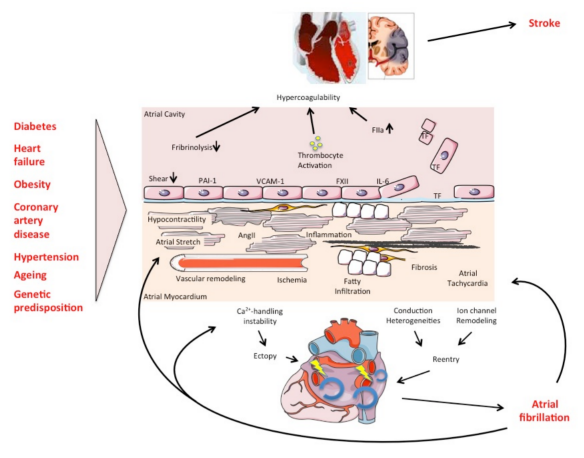


Figure 2 Major mechanisms causing AF that can be considered when guiding therapy. The various aetiological factors (left) cause a complex array of pathophysiological changes in the atria, including stretch-induced atrial fibrosis, hypocontractility, fatty infiltration, inflammation, vascular remodelling, ischaemia, ion channel dysfunction, and Ca²⁺-instability. These changes enhance both ectopy and conduction disturbances, increasing the propensity of the atria to develop or maintain AF. At the same time, some of these alterations are involved in the occurrence of the hypercoagulable state associated with AF. For example, hypocontractility reduces local endothelial shear stress, which increases PAI-1 expression, and ischaemia-induced inflammation enhances the expression of endothelial adhesion molecules or promotes shedding of endothelial cells, resulting in tissue factor exposure to the blood stream. These changes contribute to the thrombogenic milieu in the atria of AF patients. AF in itself can aggravate many of the mechanisms shown, which may explain the progressive nature of the arrhythmia. AngII = angiotensin II; TF = tissue factor; FXII = factor XII; IL-6 = interleukin 6; PAI-1 = plasminogen

activator inhibitor 1; VCAM-1 = vascular cell adhesion molecule 1.

Table 4 Pathophysiological alterations in atrial tissue associated with AF and clinical conditions that could contribute to such alterations

Pathophysiological alteration	Clinical conditions contributing to the alteration	Proarrhythmic mechanism/ functional consequence	References
Changes of the extract	ellular matrix, fibroblast function, an	d fat cells	
Interstitial and replacement fibrosis	AF (especially forms with a high AF burden), hypertension, heart failure, valvular heart disease (via pressure and volume overload)	Electrical dissociation, conduction block, enhanced AF complexity	78, 79, 90, 91
Inflammatory infiltration		Profibrotic responses, enhanced AF complexity	81
Fatty infiltration	Obesity (fatty infiltration)	Profibrotic/proinflammatory responses, localized conduction	82, 92

Amyloid deposition	Ageing, heart failure, CAD (via atrial scarring), genetic factors	block Conduction disturbances	83, 93
Ion channel alterations			
Ion channel remodelling	AF (especially forms with a high AF burden), genetic predisposition to AF	AF cycle shortening (if due to atrial tachycardia), AF cycle length prolongation (if due to heart failure), enhanced heterogeneity of atrial repolarization	94-96
Ca ²⁺ handling instability	AF (especially forms with a high AF burden), possibly heart failure and hypertension (possibly through increased sympathetic activation)	Enhanced propensity to ectopy	97, 98
Gap-junction redistribution	AF	Conduction disturbances	99
Myocyte alterations			
Apoptosis and necrosis	CAD, heart failure (through cardiomyocyte death and atrial scarring)	May induce replacement fibrosis	100
Myocyte hypertrophy	Atrial dilatation, AF	Aggravates conduction disturbances	84, 101
Endothelial and vascul	ar alterations		
Microvascular changes	Atherosclerosis, CAD and peripheral artery disease, possibly	Aggravation of atrial ischaemia, heterogeneity of electrical	102
Endocardial remodelling	AF	function, structural remodelling Enhanced risk for thrombus formation	103, 104
Changes of the autonor	nic nervous system		
Sympathetic hyperinnervation	Heart failure, hypertension	Enhanced propensity to ectopy	80, 105

488 AF = atrial fibrillation; CAD = coronary artery disease.

489

490 3.2.1. Electrophysiological mechanisms of atrial fibrillation

491 AF provokes a shortening of the atrial refractory period and AF cycle length during the first days of the 492 arrhythmia, largely due to downregulation of the Ca²⁺-inward current and upregulation of inward rectifier K⁺ 493 currents.^{94, 95} Structural heart disease, in contrast, tends to prolong the atrial refractory period, illustrating the heterogeneous nature of mechanisms that cause AF in different patients.⁹⁶ Hyperphosphorylation of various 494 Ca^{2+} handling proteins may contribute to enhanced spontaneous Ca^{2+} release events and triggered activity, ^{97,98} 495 thus causing ectopy and promoting AF. Although the concept of Ca^{2+} handling instability has been challenged 496 recently, ^{106, 107} it may mediate AF in structurally remodelled atria and explain how altered autonomic tone can generate AF.^{80, 105} 497 498

499

500 Focal initiation and maintenance of AF: The seminal observation by Haissaguerre et al¹⁰⁸ was that a focal 501 source in the pulmonary veins can trigger AF, and ablation of this source can extinguish the arrhythmia. The mechanism of focal activity might involve both triggered activity and localized reentry.^{109, 110} Hierarchic 502 organization of AF with rapidly activated areas driving the arrhythmia has been documented in patients with 503 paroxysmal AF,^{111, 112} but is more challenging in patients with persistent AF.¹¹³ 504 505

506 The multiple wavelet hypothesis and rotors as sources of AF: Moe and Abildskov¹¹⁴ proposed that AF can be 507 perpetuated by continuous conduction of several independent wavelets propagating through the atrial 508 musculature in a seemingly chaotic manner. As long as the number of wavefronts does not decline below a 509 critical level, they will be capable of sustaining the arrhythmia. Numerous experimental and clinical

observations can be reconciled with the multiple wavelet hypothesis.¹¹⁵ All localized sources of AF (ectopic 510

foci, rotors, or other stable reentry circuits) cause fibrillatory conduction remote from the source, which is
 difficult to distinguish from propagation sustaining AF by multiple wavelets, and either of these phenomena
 may generate 'rotors' picked up by intracardiac^{116, 117} or body surface¹¹⁷ recordings.

514 515

516 **5 Diagnosis and timely detection of atrial fibrillation**

517 5.1. Overt and silent atrial fibrillation

The diagnosis of AF requires rhythm documentation using an electrocardiogram (ECG), with the typical pattern
of AF. ECG-documented AF was the entry criterion in trials forming the evidence for these guidelines. By
accepted convention, an episode lasting at least 30 seconds is diagnostic. Individuals with AF may be
symptomatic or asymptomatic ('silent AF'). Many AF patients have both symptomatic and asymptomatic
episodes of AF.¹¹⁸⁻¹²¹

523 Silent, undetected AF is common,^{120, 122} with severe consequences such as stroke and death.¹²³⁻¹²⁵ 524 Prompt recording of an ECG is an effective and cost-effective method to document chronic forms of AF.¹²⁶ The 525 technology to detect paroxysmal, self-terminating AF episodes is rapidly evolving (see Chapter 5 for a 526 definition of AF patterns). There is good evidence that prolonged ECG monitoring enhances the detection of 527 undiagnosed AF, for 72 hours after a stroke,^{27, 127} for even longer periods,^{18, 128} or by daily short-term ECG 528 recording in patients over 75 years of age¹²⁹ (*Web Addenda Figure 1*). Ongoing studies will determine whether 529 such early detection alters management (e.g. initiation of anticoagulation) and improves outcomes.

Once the ECG diagnosis of AF has been established, further ECG monitoring can inform management
in the context of: (1) a change in symptoms or new symptoms; (2) suspected progression of AF; (3) monitoring
of drug effects on ventricular rate; and (4) ECG monitoring of antiarrhythmic drug effects or catheter ablation
for rhythm control.

535 **5.2.** Screening for silent atrial fibrillation

536 5.2.1. Screening for atrial fibrillation by electrocardiogram in the community

537 Undiagnosed AF is common, especially in older populations and in patients with heart failure.¹³⁰ Opportunistic screening for silent AF seems cost-effective in elderly populations (e.g. > 65 years),¹³¹ and similar effects have been reported using single-lead ECG screening in other at-risk populations.^{132, 133} Screening of elderly 538 539 540 populations (mean age 64 years) yielded a prevalence of 2.3% for chronic forms of AF in 122,571 participants 541 using either short-term ECG or pulse palpation (followed by ECG in those with an irregular pulse).¹³⁴ 542 Previously undiagnosed AF was found in 1.4% of those aged > 65 years, suggesting a number needed to screen 543 of 70. These findings encourage the further evaluation of systematic AF screening programmes in at-risk 544 populations. 545

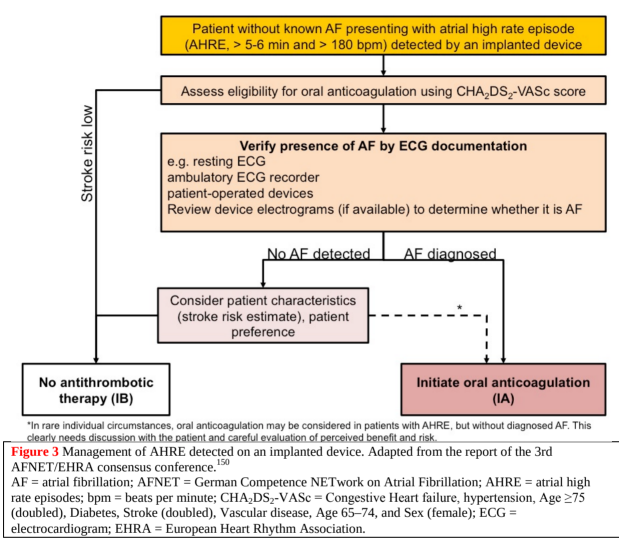
546 **5.2.2.** Prolonged monitoring for paroxysmal atrial fibrillation

Paroxysmal AF is often missed.¹²⁰ Repeated daily ECG recordings increased the detection of silent,
asymptomatic paroxysmal AF in an unselected Swedish population aged > 75 years.^{120, 135} Several patientoperated devices^{136, 137} and extended continuous ECG monitoring using skin patch recorders¹³⁸ have been
validated for detection of paroxysmal AF.¹³⁹ The detection rate of asymptomatic AF by new technologies such
as smartphone cases with ECG electrodes, smart watches, and blood pressure machines with AF detection
algorithms, has not yet been formally evaluated against an established arrhythmia detection method.¹⁴⁰

553554 5.2.3. Patients with pacemakers and implanted devices

555 Implanted pacemakers or defibrillators with an atrial lead allow continuous monitoring of atrial rhythm. Using 556 this technology, patients with atrial high rate episodes (AHRE) can be identified. Depending on the risk profile of the population studied, such AHRE are detected in 10–15% of pacemaker patients.¹⁴¹ AHRE are associated 557 558 with an increased risk of overt AF (hazard ratio [HR] 5.56; 95% confidence interval [CI] 3.78–8.17; P < 0.001) 559 and ischaemic stroke or systemic embolism (HR 2.49; 95% CI 1.28–4.85; P = 0.007). The stroke risk in AHRE patients seems lower than the stroke risk in patients with diagnosed AF, and not all AHRE represent AF.¹⁴² 560 Strokes often occur without AHRE detected within 30 days before the event.¹⁴³⁻¹⁴⁷ Consequently, it is unclear 561 whether AHRE imply the same therapeutic requirements as overt AF,¹⁴⁸ and the benefit of OAC in patients with 562 563 AHRE is being evaluation in ongoing clinical trials (e.g. ARTESiA [NCT01938248] and NOAH 564 [NCT02618577]). At present, pacemakers and implanted devices should be interrogated on a regular basis for 565 AHRE, and patients with AHRE should undergo further assessment of stroke risk factors and for overt AF,

566 including ECG monitoring (*Figure 3*).¹⁴⁹



5.2.4. Detection of atrial fibrillation in stroke survivors

Sequential stratified ECG monitoring detected AF in 24% (95% CI 17-31) of stroke survivors,¹⁵¹ and in 11.5% (95% CI 8.9%–14.3%) in another meta-analysis,¹⁷ with large variations depending on the timing, duration, and method of monitoring. AF detection is not uncommon in unselected stroke patients (6.2%, 95% CI 4.4-8.3),¹²⁸ but is more likely in patients with cryptogenic stroke implanted with loop recorders or who have had ECG monitors for several weeks.^{18, 128, 152} Cryptogenic stroke is defined as a stroke in which the cause could not be identified after extensive investigations.¹⁵³ A broader definition is embolic stroke of undetermined source.¹⁵⁴ Several studies have also found AF in patients in whom another competing cause for stroke has been identified clinically (e.g. hypertension or carotid artery stenosis).^{27, 127} Hence, prolonged ECG monitoring seems reasonable in all survivors of an ischaemic stroke without an established diagnosis of AF.

Recommendations for screening for AF

Recommendations	Class ^a	Level ^b	Ref s ^c
Opportunistic screening for AF is recommended by pulse taking or ECG rhythm strip in patients > 65 years of age	I	В	130, 134, 155
In patients with TIA or ischaemic stroke, screening for AF is recommended by short-term ECG recording followed by continuous ECG monitoring for at least 72 hours	I	В	27, 127

It is recommended to interrogate pacemakers and ICDs on a regular basis for atrial high rate episodes (AHRE). Patients with AHRE should undergo further ECG monitoring to document AF before initiating AF therapy	I	В	141, 156
In stroke patients, additional ECG monitoring by long-term non- invasive ECG monitors or implanted loop recorders should be considered to document silent AF	lla	В	18, 128
Systematic ECG screening may be considered to detect AF in patients aged > 75 years, or those at high stroke risk	llb	В	130, 135, 157

- 587 AF = atrial fibrillation; AHRE = atrial high rate episodes; ECG = electrocardiogram; ICD = implantable
- 588 cardioverter defibrillator; TIA = transient ischaemic attack.
- 589 ^aClass of recommendation.
- 590 ^bLevel of evidence.
- 591 ^cReference(s) supporting recommendations.
- 592

593 5.3. Electrocardiogram detection of atrial flutter

Right atrial isthmus-dependent flutter has a typical ECG pattern and ventricular rate.¹⁵⁸ The prevalence of atrial flutter is less than one-tenth of the prevalence of AF.¹⁵⁹ Atrial flutter often coexists with or precedes AF.¹⁶⁰ In typical, isthmus-dependent flutter, P waves will often show a 'saw tooth' morphology, especially in the inferior leads (II, III, aVF). The ventricular rate can be variable (usual ratio of atrial to ventricular contraction 4:1 to 2:1, in rare cases 1:1) and macro-reentrant tachycardias may be missed in stable 2:1 conduction. Vagal stimulation or intravenous adenosine may be helpful to unmask atrial flutter. The management of atrial flutter is discussed in Section 12.7. Left or right atrial macro-reentrant tachycardia is usually confined to patients after catheter ablation for AF, AF surgery, or after open heart surgery.¹⁵⁸

602

603 6 Classification of atrial fibrillation

604 6.1. Atrial fibrillation pattern

In many patients, AF progresses from short, infrequent episodes to longer and more frequent attacks. Over time,
 many patients will develop sustained forms of AF. In a small proportion of patients, AF will remain paroxysmal
 over several decades (2–3% of AF patients).¹⁶¹ The distribution of paroxysmal AF recurrences is not random,
 but clustered.¹⁶² AF may also regress from persistent to paroxysmal AF. Furthermore, asymptomatic recurrences
 of AF are common in patients with symptomatic AF.¹²⁰

610

611Based on presentation, duration, and spontaneous termination of AF episodes, five types of AF are612traditionally distinguished: first diagnosed, paroxysmal, persistent, long-standing persistent, and permanent AF613(*Table 5*). If patients suffer from both paroxysmal and persistent AF episodes, the more common type should be614used for classification. Clinically determined AF patterns do not correspond well to the AF burden measured by615long-term ECG monitoring. ¹⁶³ Even less is known about the response to therapy in patients with long-standing616persistent AF or long-standing paroxysmal AF. Despite these inaccuracies, the distinction between paroxysmal617and persistent AF has been used in many trials and therefore still forms the basis of some recommendations.618There is some evidence suggesting that AF burden may influence stroke risk^{44, 124, 164} and could modify

 $\begin{array}{ccc} 618 \\ 619 \\ 619 \\ 620 \end{array}$ There is some evidence suggesting that AF burden may influence stroke risk^{44, 124, 164} and could modify the response to rhythm control therapy.^{76, 165} The evidence for this is weak. Therefore, AF burden should not be a major factor in deciding on the usefulness of an intervention that is deemed suitable for other reasons.

- 621 622
- Table 5 Patterns of AF

AF pattern	Definition
First diagnosed AF	AF that has not been diagnosed before, irrespective of the duration of the arrhythmia or the presence and severity of AF-related symptoms.
Paroxysmal AF	Self-terminating, in most cases within 48 hours. Some AF paroxysms may continue for up to 7 days. ^a Most AF episodes that are cardioverted within 24-48 hours should be considered paroxysmal. ^a
Persistent AF	AF that lasts longer than 7 days, including episodes that are terminated by cardioversion, either with drugs or by direct current cardioversion, after 7 days or more.
Long-standing persistent AF	Continuous AF lasting for \geq 1 year when it is decided to adopt a rhythm control strategy.
Permanent AF	AF is accepted by the patient (and physician). Hence, rhythm control

interventions are, by definition, not pursued in patients with permanent AF. Should a rhythm control strategy be adopted, the arrhythmia would be re-classified as 'long-standing persistent AF'.

623 AF = atrial fibrillation.

^aThe distinction between paroxysmal and persistent AF is often not made correctly without access to long-term
 monitoring.¹⁶³ Hence, this classification alone is often insufficient to select specific therapies. If both persistent
 and paroxysmal episodes are present, the predominant pattern should guide the classification.

627

628 6.2. Atrial fibrillation types reflecting different causes of the arrhythmia

The risk of developing AF is increased in a variety of physiological and disease states, and the historic term 'lone AF' is probably misleading and should be avoided.¹⁶⁶ Although the pattern of AF may be the same, the mechanisms underpinning AF vary substantially between patients¹⁶⁷ (*Table 6*). This suggests that stratifying AF patients by underlying drivers of AF could inform management, for example, considering cardiac and systemic comorbidity (e.g. diabetes and obesity¹⁶⁸), lifestyle factors (e.g. activity level, smoking, alcohol intake^{169, 170}), markers of cardiac structural remodelling (e.g. fibrosis¹⁷¹⁻¹⁷³ or electrocardiographic parameters of AF complexity¹⁷⁴), or genetic background. *Table 6* provides such a taxonomy, informed by expert consensus,^{76, 120, 175} but without much evidence to underpin its clinical use.¹⁷⁶ Systematic research defining the major drivers of AF is clearly needed to better define different types of AF.¹⁷⁶

638 639

Table 6 Clinical types of AF (modified from the report on the 4 th AFNET/EHRA consensus conference ⁷⁶) ^a					
AF type	Clinical presentation	Possible pathophysiology			
AF secondary to structural heart disease	AF in patients with LV systolic or diastolic dysfunction, long-standing hypertension with LVH, and/or other structural heart diseases. The onset of AF in these patients is a common cause of hospitalization and a predictor of poor outcome	Increased atrial pressure and atrial structural remodelling, together with activation of the sympathetic and renin– angiotensin system			
Focal AF	Patients with repetitive atrial runs and frequent, short episodes of paroxysmal AF. Often highly symptomatic, younger patients with distinguishable atrial waves (coarse AF), atrial ectopy, and/or atrial tachycardia deteriorating in AF	Localized triggers, in most cases originating from the pulmonary veins, initiate AF. AF due to one or a few reentrant drivers is also considered to be part of this type of AF			
Polygenic AF	AF in carriers of common gene variants that have been associated with early onset AF	Currently under study. The presence of some gene variants may also influence treatment outcomes			
Postoperative AF	New onset of AF (usually self-terminating) after major (typically cardiac) surgery in patients who were in sinus rhythm before surgery and had no history of AF	Acute factors: inflammation, atrial oxidative stress, high sympathetic tone, electrolyte changes, and volume overload, possibly interacting with a pre-existing substrate			
AF in patients with mitral stenosis or prosthetic heart valves	AF in patients with mitral stenosis, after mitral valve surgery and in some cases other valvular disease	Left atrial pressure (stenosis) and volume (regurgitation) load are the main drivers of atrial enlargement and structural atrial remodelling in these patients			
AF in athletes	Usually paroxysmal, related to duration and intensity of training	Increased vagal tone and atrial volume			
Monogenic AF	AF in patients with inherited cardiomyopathies, including channelopathies	The arrhythmogenic mechanisms responsible for sudden death are likely to contribute to the occurrence of AF in these patients			

640 AF = atrial fibrillation; LV = left ventricular; LVH = left ventricular hypertrophy.

⁶41 ^aIt is recognized that these types of AF will overlap in clinical practice, and that their impact for management

642 needs to be evaluated systematically.

643

644 6.3. Symptom burden in atrial fibrillation

645 Patients with AF have significantly poorer quality of life than healthy controls, experiencing a variety of 646 symptoms including lethargy, palpitations, dyspnoea, chest tightness, sleeping difficulties, and psychosocial 647 distress.^{32, 177-180} Improved quality of life has been noted with both pharmacological and interventional therapies,¹⁸¹⁻¹⁸⁵ but there are limited data to compare the benefit of different treatments.^{32, 186} Assessment of 648 649 quality of life is further constrained by a lack of cross-validation of the several AF-specific quality-of-life tools.¹⁸⁷⁻¹⁹¹ With regard to symptom assessment, the European Heart Rhythm Association (EHRA) suggested the EHRA symptom scale (*Table 7*) to describe symptom severity in AF patients.¹⁹² A similar scale (the 650 651 Canadian Cardiovascular Society Severity of Atrial Fibrillation Scale) is used in Canada.¹⁹³ The EHRA scale has been used and validated.¹⁹⁴⁻¹⁹⁹ A modification was proposed in 2014, subdividing EHRA class 2 into mild 652 653 654 (2a) or moderate (2b) impact.¹⁹⁹ As symptoms in class 2b ('troubling' symptoms) identified patients with a 655 health utility benefit of rhythm control in that study, this modification may provide a threshold for potential treatment decisions, but this remains to be tested. While some AF patients had no or minimal symptoms (25-656 40%), many (15–30%) reported severe or disabling symptoms.^{194, 196} The EHRA scale should be used to guide 657 symptom-orientated treatment decisions and for longitudinal patient profiling. 658

659 660

Table 7 Modified EHRA symptom scale (modified from Wynn et al¹⁹⁹)

Modified EH	RA Symptoms	Description
score		
1	None	AF does not cause any symptoms
2a	Mild	Normal daily activity not affected by symptoms related to AF ^a
2b	Moderate	Normal daily activity not affected ^a
3	Severe	Normal daily activity affected
4	Disabling	Normal daily activity discontinued

661 AF = atrial fibrillation; EHRA = European Heart Rhythm Association.

^aEHRA class 2a and 2b can be differentiated by evaluating whether patients are functionally affected by their

AF symptoms. AF-related symptoms are most commonly fatigue/tiredness and exertional shortness of breath, or
 less frequently palpitations and chest pain.^{42, 194, 200-202}

665 666

Recommendation on use of the modified EHRA symptom scale

Recommendation	Class ^a	Level ^b	Refs ^c
Use of the modified EHRA symptom scale is recommended in clinical practice and research studies to quantify AF-related symptoms	I	С	192, 199

667 AF = atrial fibrillation; EHRA = European Heart Rhythm Association.

^aClass of recommendation.

669 ^bLevel of evidence.

670 ^cReference(s) supporting recommendations.

672 **7** Detection and management of risk factors and concomitant

673 cardiovascular diseases

Many cardiovascular diseases and concomitant conditions increase the risk of developing AF (*Table 8*),
 recurrent AF, and AF-associated complications. Identification of such conditions, their prevention and treatment
 is an important leverage to prevent AF and its disease burden. Knowledge of these factors and their management

677 is hence important for optimal management of AF patients.^{203, 204}

678 679

671

Table 8 Cardiovascular and other conditions independently associated with AF

Characteristic/comorbidity	Association with AF
Genetic predisposition (based on multiple common gene variants associated with AF) ⁶⁴	HR range 0.4–3.2

19	
Older age ¹⁹	HR:
50–59 years	1.00 (reference)
60–69 years	4.98 (95% CI 3.49–7.10)
70–79 years	7.35 (95% CI 5.28–10.2)
80–89 years	9.33 (95% CI 6.68–13.0)
Hypertension (treated) vs. none ¹⁹	HR 1.32 (95% CI 1.08–1.60)
Heart failure vs. none ¹⁹	HR 1.43 (95% CI 0.85–2.40)
Valvular heart disease vs. none ²⁰⁵	RR 2.42 (95% CI 1.62–3.60)
Myocardial infarction vs. none ¹⁹	HR 1.46 (95% CI 1.07–1.98)
Thyroid dysfunction ^{206, 207}	(reference: euthyroid)
hypothyroidism	HR 1.23 (95% CI 0.77-1.97)
subclinical hyperthyroidism	RR 1.31 (95% CI 1.19–1.44)
overt hyperthyroidism	RR 1.42 (95% CI 1.22–1.63)
Obesity ^{19, 208}	HR:
none (BMI $< 25 \text{ kg/m}^2$)	1.00 (reference)
overweight (BMI 25–30 kg/m ²)	1.13 (95% CI 0.87–1.46)
obese (BMI \ge 31 kg/m ²)	1.37 (95% CI 1.05–1.78)
Diabetes mellitus vs. none ¹⁹	HR 1.25 (95% CI 0.98–1.60)
Chronic obstructive pulmonary disease ²⁰⁹	RR:
FEV1	
$\geq 80\%$	1.00 (reference)
60-80%	1.28 (95% CI 0.79–2.06)
< 60%	2.53 (95% CI 1.45–4.42)
Obstructive sleep apnoea vs. none ²¹⁰	HR 2.18 (95% CI 1.34–3.54)
Chronic kidney disease ²¹¹	OR:
none	1.00 (reference)
stage 1 or 2	2.67 (95% CI 2.04-3.48)
stage 3	1.68 (95% CI 1.26–2.24)
stage 4 or 5	3.52 (95% CI 1.73–7.15)
Smoking ²¹²	HR:
never	1.00 (reference)
former	1.32 (95% CI 1.10–1.57)
current	2.05 (95% CI 1.71–2.47)
Alcohol consumption ²¹³	RR:
None	1.00 (reference)
1–6 drinks/week	1.01 (95% CI 0.94–1.09)
7–14 drinks/week	1.07 (95% CI 0.98–1.17)
15–21 drinks/week	1.14 (95% CI 1.01–1.28)
> 21 drinks/week	1.39 (95% CI 1.22–1.58)
Habitual vigorous exercise ²¹⁴	RR:
Non-exercisers	1.00 (reference)
<1 day/week	0.90 (95% CI 0.68-1.20)
1-2 days/week	1.09 (95% CI 0.95–1.26)
3–4 days/week	1.04 (95% CI 0.91–1.19)
5–7 days/week	1.20 (95% CI 1.02-1.41)
AF = atrial fibrillation; BMI = body mass index; CI = confid	

680

AF = atrial fibrillation; BMI = body mass index; CI = confidence interval; FEV1 = forced expiratory volume in 681 1 second; HR = hazard ratio; OR = odds ratio; RR = risk ratio

682

683 7.1. **Heart failure**

Heart failure and AF coincide in many patients.²¹⁵⁻²¹⁷ They are linked by similar risk factors and share a 684 common pathophysiology.²¹⁸ Heart failure and AF can cause and exacerbate each other through mechanisms 685 686 such as structural cardiac remodelling, activation of neurohormonal mechanisms, and rate-related impairment of left ventricular (LV) function. Patients with AF and concomitant heart failure, both with preserved ejection fraction (LV ejection fraction [LVEF] \geq 50%) and reduced ejection fraction (LVEF < 40%),^{219, 220} suffer from a worse prognosis, including increased mortality.^{16, 221} The recent ESC Guidelines on heart failure²²² have also 687 688 689 690 introduced a new category of heart failure with mid-range ejection fraction (HFmrEF; LVEF 40-49%), although 691 data on AF patients in this group are currently limited. Prevention of adverse outcomes and maintenance of a

692 good quality of life are the aims of management in all patients with AF and concomitant heart failure, regardless of LVEF.²²³ The general approach to AF management does not differ between heart failure patients and others,
 but a few considerations are worthwhile to consider. Of note, the only therapy with proven prognostic value in
 these patients is anticoagulation, and appropriate OAC should be prescribed in all patients at risk of stroke (see
 Chapter 8).

697

6987.1.1.Patients with atrial fibrillation and heart failure with reduced ejection

699 fraction

In addition to OAC, standard heart-failure therapy should be used in patients with heart failure with reduced
 ejection fraction (HFrEF), as detailed in the ESC Guidelines.²²² This includes angiotensin-converting enzyme
 (ACE) inhibitors or angiotensin receptor blockers (ARBs), mineralocorticoid antagonists, defibrillators and
 cardiac resynchronization therapy,²¹⁸ in addition to combined angiotensin receptor neprilysin inhibition (ARNI)
 in patients able to tolerate an ACE inhibitor or ARB with ongoing symptoms.²²⁴

705 Rate control of AF is discussed in detail in Chapter 9. In brief, only beta-blockers and digoxin are 706 suitable in HFrEF because of the negative inotropic potential of verapamil and diltiazem. Beta-blockers are 707 usually the first-line option in patients with clinically stable HFrEF, although a meta-analysis using individual 708 patient data from randomized controlled trials (RCTs) found no reduction in mortality from beta-blockers versus placebo in those with AF at baseline (HR 0.97, 95% CI 0.83–1.14).²³ Digoxin is commonly prescribed in 709 710 clinical practice but no head-to-head RCTs in AF patients have been performed. In a meta-analysis of 711 observational studies, digoxin had a neutral effect on mortality in patients with AF and concomitant heart failure 712 (adjusted observational studies HR 0.90, 95% CI 0.70-1.16; propensity-matched observational studies RR 1.08, 713 95% CI 0.93–1.26).²²⁵ Initial and combination rate-control therapy for AF in HFrEF should therefore take 714 account of individual patient characteristics and symptoms; beta-blocker initiation should be delayed in patients 715 with acute decompensated heart failure, and digoxin has more adverse effects in patients with renal impairment 716 (see Chapter 9).

Patients with AF and HFrEF who present with severe symptoms may require rhythm control therapy in addition to rate control therapy. For patients who develop HFrEF as a result of rapid AF (tachycardiomyopathy), a rhythm control strategy is preferred, based on several relatively small patient cohorts and trials reporting improved LV function after restoration of sinus rhythm.^{185, 226-228} The diagnosis of tachycardiomyopathy can be challenging, and at times requires restoration of sinus rhythm.²²⁹ Catheter ablation may be a useful method to restore LV function and quality of life in AF patients with HFrEF,^{185, 226-228} but further data are needed. *Figure 4* summarizes the approach to patients with AF and heart failure.

Management of patients presenting acutely with AF and heart failure

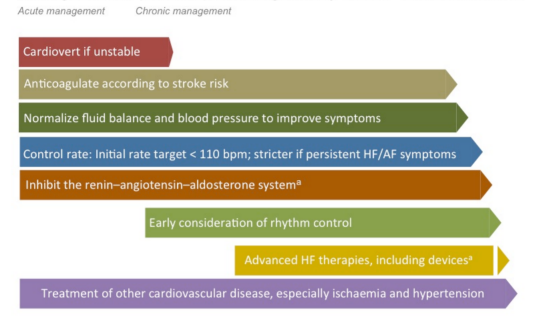


Figure 4 Initial management of newly diagnosed with AF and heart failure. Adapted from Kotecha and Piccini.²¹⁸

ACE = angiotensin-converting enzyme; AF = atrial fibrillation; ARB = angiotensin receptor blocker; ARNI = angiotensin receptor neprilysin inhibition; bpm = beats per minute; HF = heart failure.

^aIn patients with heart failure and reduced ejection fraction; also consider combined ARNI in patients able to tolerate an ACE inhibitor or ARB with ongoing symptoms.

731732 **7.1.2.** Atrial fibrillation patients with heart failure with preserved ejection fraction

733 The diagnosis of heart failure with preserved ejection fraction (HFpEF) in patients with AF is problematic 734 because of the difficulty in separating symptoms that are due to HF from those due to AF. Although diagnostic 735 differentiation can be achieved by cardioversion and clinical reassessment, this option is often not appropriate in 736 this group, particularly as a specific therapy that improves prognosis in HFpEF is currently lacking. 737 Echocardiography can support detection of HFpEF in patients with symptomatic AF by providing evidence of 738 relevant structural heart disease (e.g. LV hypertrophy [LVH]) and/or measurement of diastolic dysfunction. 739 Reduced early diastolic myocardial velocity e' by tissue Doppler reflects impaired LV relaxation, while the ratio of E/e' has demonstrated a significant correlation with invasive measurement of LV filling pressures.²³⁰⁻²³⁴ 740 741 Natriuretic peptide levels are part of the diagnostic assessment of HFpEF,²²² although natriuretic peptide levels are elevated in AF patients and the optimum diagnostic cut-off is still unknown.²³⁵ The management of patients 742 743 with AF and concomitant HFpEF should focus on control of fluid balance and concomitant conditions such as 744 hypertension and ischaemia.

745

746 **7.1.3.** Atrial fibrillation patients with heart failure with mid-range ejection

747 fraction

HFmrEF is a recently defined entity, describing patients with symptoms and signs of heart failure, LVEF 40–
49%, elevated levels of natriuretic peptides, and either LV hypertrophy, left atrial (LA) enlargement, or
evidence of diastolic dysfunction.²²² However, diagnosis is more difficult in patients with AF, as natriuretic

- 751 peptides are elevated in AF and LA dilatation is common, regardless of concomitant heart failure. LVEF is also
- variable and difficult to assess in AF patients because of AF-induced reduction in systolic LV function and

variable cardiac cycle length. Further study of this group is required before particular treatment strategies in AF
 patients with HFmrEF can be recommended.

755756 **7.1.4.** Prevention of atrial fibrillation in heart failure

Retrospective analyses from large randomized trials have reported a lower incidence of new-onset AF in patients treated with ACE inhibitors/ARBs compared with placebo.²³⁶⁻²³⁸ The reduced incidence of AF with 757 758 ACE inhibitors/ARBs is less evident in patients with HFpEF²³⁹ and is lost in patients without heart failure.²⁴⁰⁻²⁴² 759 Neprilysin inhibition does not seem to add to this effect.²²⁴ Beta-blocker therapy was associated with a 33% 760 761 reduction in the adjusted odds of incident AF in HFrEF patients pretreated with ACE inhibitors/ARBs, 762 reinforcing the importance of beta-blocker therapy in HFrEF patients in sinus rhythm.²³ Eplerenone, a 763 mineralocorticoid receptor antagonist, also reduced the risk of new-onset AF in patients with LVEF \leq 35%, 764 New York Heart Association (NYHA) Class II, and pretreatment with ACE inhibitors/ARBs and betablockers.243 765

766

767 7.2. Hypertension

768 **7.2.1.** Treatment of hypertension to prevent incident atrial fibrillation

Inhibition of the renin–angiotensin–aldosterone system can prevent structural remodelling and recurrent AF.²³⁶
 ²⁴⁴ A recent analysis of the Danish healthcare database with long-term monitoring of the effect of different antihypertensive agents on the occurrence of overt AF suggests a beneficial effect of ACE inhibitors or ARBs.²⁴⁵ Secondary analyses of ACE inhibitors or ARBs in patients with heart failure or LVH show a lower

incidence of new-onset AF.^{238, 246}

775 **7.2.2.** Blood pressure control in patients with atrial fibrillation

Hypertension is a stroke risk factor in AF, and uncontrolled high blood pressure enhances the risk of stroke and
bleeding events and may lead to recurrent AF. Good blood-pressure control should therefore form an integral
part of the management of AF patients.²⁴⁷ In patients with established AF, but without LV dysfunction or heart
failure, ARBs do not prevent recurrent AF better than placebo.^{240, 241} ACE inhibitors or ARBs may reduce
recurrent AF after cardioversion when coadministered with antiarrhythmic drug therapy compared with an
antiarrhythmic drug alone.^{248, 249} Meta-analyses driven by these studies suggested a lower risk of recurrent
AF, ^{236-238, 250} but at least one controlled trial failed to demonstrate benefit.^{240, 251}

784 7.3. Valvular heart disease

Valvular heart disease is independently associated with incident AF.²⁵² Approximately 30% of patients with AF
have some form of valvular heart disease, often detected only by echocardiography.^{201, 253-255} AF worsens
prognosis in patients with severe valvular heart disease,²⁵⁶ including those undergoing surgery or transcatheter
interventions for aortic or mitral valve disease.²⁵⁷⁻²⁶² Valvular heart disease can be associated with an increased
thromboembolic risk, which probably also adds to the stroke risk in AF patients.²⁶³ Similar to heart failure,
valvular disease and AF interact and sustain each other through volume and pressure overload,

valvular disease and AF interact and sustain each other through volume and pressure overload,
 tachycardiomyopathy, and neurohumoral factors.²⁶⁴⁻²⁷⁰ When valve dysfunction is severe, AF can be regarded as
 a marker for progressive disease, thus favouring valve repair or replacement.²⁷¹

Traditionally, patients with AF have been dichotomized into 'valvular' and 'non-valvular' AF.²⁷² Although slightly different definitions have been used, valvular AF mainly refers to AF patients that have either rheumatic valvular disease (predominantly mitral stenosis) or mechanical heart valves. In fact, while AF implies an incremental risk for thromboembolism in patients with mitral valve stenosis,^{263, 273, 274} there is no clear evidence that other valvular diseases, including mitral regurgitation or aortic valve disease, need to be considered when choosing an anticoagulant or indeed to estimate stroke risk.²⁷⁵ We have therefore decided to

- replace the historic term 'non-valvular' AF with reference to the specific underlying conditions.
- 800 801

Recommendations for patients with valvular heart disease and AF

Recommendations	Class ^a	Level ^D	Refs ^c
Early mitral valve surgery should be considered in severe mitral regurgitation, preserved LV function, and new-onset AF, even in the absence of symptoms, particularly when valve repair is feasible	lla	С	276

	Mitral valvotomy should be considered for asymptomatic patients with severe mitral stenosis and suitable valve anatomy who have new-onset AF	lla	С	
2	AF = atrial fibrillation; LV = left ventricular.			
	^a Class of recommendation.			

- 803 804 ^bLevel of evidence.
- 805 ^cReference(s) supporting recommendations.
- 806
- 807

802

808 7.4. **Diabetes mellitus**

Diabetes and AF frequently coexist because of associations with other risk factors.²⁷⁷⁻²⁸³ Diabetes is a risk factor 809 for stroke and other complications in AF.²⁸⁴ In patients with AF, a longer duration of diabetes appears to confer 810 811 a higher risk of thromboembolism, albeit without greater risk of OAC-related bleeding.²⁸⁵ Unfortunately, 812 intensive glycaemic control does not affect the rate of new-onset AF,²⁸⁴ while treatment with metformin seems 813 to be associated with a decreased long-term risk of AF in diabetic patients²⁸⁶ and may even lower long-term 814 stroke risk.¹³ Diabetic retinopathy, a measure of disease severity, does not increase the risk of ocular bleeding in anticoagulated patients.28 815

816

817 7.5. **Obesity and weight loss**

818 7.5.1. Obesity as a risk factor

Obesity increases the risk for AF (risk ratio 1.5–1.8),²⁸⁸⁻²⁹¹ with a progressive increase according to body mass 819 index.^{288, 290-292} Obese patients may have more LV diastolic dysfunction, increased sympathetic activity and inflammation, and increased fatty infiltration of the atria.²⁹³⁻²⁹⁵ Obesity may also be a risk factor for ischaemic 820 821 822 stroke, thromboembolism, and death in AF patients.²⁹² 823

824 Weight reduction in obese patients with atrial fibrillation 7.5.2.

825 Intensive weight-reduction management in addition to management of other cardiovascular risk factors (in the range of 10–15 kg weight loss achieved) led to fewer AF recurrences and symptoms compared with an approach based on general advice in obese patients with AF.^{203, 204, 296} Improved cardiorespiratory fitness can further decrease AF burden in obese patients with AF.²⁹⁷ Although the findings in these studies have to be confirmed, 826 827 828 829 they underpin the positive effect of weight reduction in obese patients. 830

831 7.5.3. Catheter ablation in obese patients

Obesity may increase the rate of AF recurrence after catheter ablation,²⁹⁸⁻³⁰¹ with obstructive sleep apnoea as an 832 833 important potential confounder. Obesity has also been linked to a higher radiation dose and complication rate during AF ablation.^{302, 303} Notably, the symptomatic improvement after catheter ablation of AF in obese patients 834 seems comparable to the improvement in normal-weight patients.²⁹⁸ In view of the potential to reduce AF 835 836 episodes by weight reduction (see Section 6.5.2.), AF ablation should be offered to obese patients in conjunction 837 with lifestyle modifications that lead to weight reduction. 838

839 Recommendation for obese patients with AF

AF = atrial fibrillation

841	AF = atrial fibrillation.			
	Recommendation	Class ^a	Level ^b	Refs ^c
	In obese patients with AF, weight loss together with management of	IIa	В	204, 288, 296
	other risk factors should be considered to reduce AF burden and			
	symptoms			
017				

842 843

840

^a Class of recommendation

^b Level of evidence 844

845 ^cReference(s) supporting recommendation(s) 846

847 7.6. Chronic obstructive pulmonary disease, sleep apnoea, and other respiratory 848 diseases

AF has been associated with obstructive sleep apnoea.^{304, 305} Multiple pathophysiological mechanisms can 849 850 contribute to AF in obstructive sleep apnoea, including autonomic dysfunction, hypoxia, hypercapnia, and inflammation.^{96, 304-307} Obstructive sleep apnoea exaggerates intrathoracic pressure changes, which in itself and 851 852 via vagal activation can provoke shortening of the atrial action potential and induce AF. Risk factor reduction and continuous positive airway pressure ventilation can reduce AF recurrence.³⁰⁸⁻³¹² It seems reasonable to 853 854 consider obstructive sleep apnoea screening in AF patients with risk factors. Obstructive sleep apnoea treatment 855 should be optimized to improve AF treatment results in appropriate patients. Servo-controlled pressure support 856 therapy should not be used in HFrEF patients with predominantly central sleep apnoea (of which 25% had 857 concomitant AF).³¹³

Patients with chronic obstructive pulmonary disease often suffer from atrial tachycardias, which need
to be differentiated from AF by ECG. Agents used to relieve bronchospasm, notably theophyllines and betaadrenergic agonists, may precipitate AF and make control of the ventricular response rate difficult. Nonselective beta-blockers, sotalol, propafenone, and adenosine should be used with caution in patients with
significant bronchospasm, while they can safely be used in patients with chronic obstructive pulmonary disease.
Beta-1 selective blockers (e.g. bisoprolol, metoprolol, and nebivolol), diltiazem, and verapamil are often

tolerated and effective (see Chapter 9).

865

866 Recommendations for patients with AF and respiratory diseases

Recommendations	Class ^a	Level ^b	Ref s ^c
Correction of hypoxaemia and acidosis should be considered as initial management for patients who develop AF during an acute pulmonary illness or exacerbation of chronic pulmonary disease	IIa	С	
Interrogation for clinical signs of obstructive sleep apnoea in all AF patients should be considered	IIa	В	304, 305, 314, 315
Obstructive sleep apnoea treatment should be optimized to reduce AF recurrences and improve AF treatment results	IIa	В	307-311

- 867 AF = atrial fibrillation.
- 868 ^aClass of recommendation.
- 869 ^bLevel of evidence.
- 870 ^cReference(s) supporting recommendations.
- 871

872 7.7. Chronic kidney disease

AF is present in 15–20% of patients with CKD.³¹⁶ The definition of CKD in most AF trials is relatively strict.
Although an estimated creatinine clearance (CrCl) rate of < 60 mL/min is indicative of CKD, a number of trials
in AF patients have used CrCl < 50 mL/min to adapt NOAC dosage, usually estimated using the Cockroft–Gault
formula. CrCl in AF patients can deteriorate over time.³¹⁷ The management of OAC in patients with CKD is
discussed in Section 8.2.4.

878

879 Recommendations for patients with kidney disease and AF880

Recommendations	Class ^a	Level ^b	Refs ^c
The assessment of kidney function by serum creatinine or creatinine clearance is recommended in all AF patients to detect kidney disease and to support correct dosing of AF therapy	I	A	316, 318-321
All AF patients treated with oral anticoagulation should be considered for at least yearly renal function evaluation to detect kidney disease	lla	В	

AF = atrial fibrillation.

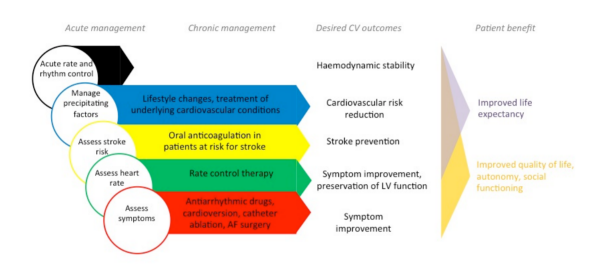
- ^aClass of recommendation.
- 883 ^bLevel of evidence.
- 884 ^cReference(s) supporting recommendations.

885

8 Integrated management of patients with atrial fibrillation 886

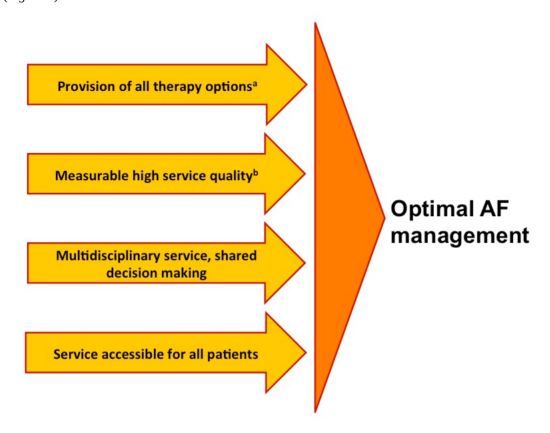
887 Most patients access the healthcare system initially through pharmacists, community health workers, or primary 888 care physicians. As AF is often asymptomatic, these healthcare professionals are important stakeholders to 889 enable adequate detection of AF and to ensure consistent management. The initial assessment should be 890 performed at the point of first contact with the healthcare system, and is feasible in most healthcare settings 891 (when an ECG is available). We propose to consider five domains in the initial assessment of patients presenting 892 with newly diagnosed AF (Figure 5). These domains are:

- 893 1. Haemodynamic instability or limiting, severe symptoms
- 894 Presence of precipitating factors (e.g. thyrotoxicosis, sepsis, or postoperative AF) and 2. 895 underlying cardiovascular conditions
- Stroke risk and need for anticoagulation 896 3.
- 897 Heart rate and need for rate control 4.
- 898 5. Symptom assessment and decision for rhythm control



899	
900	Figure 5 Acute and chronic management of AF patients, desired cardiovascular outcomes, and patient benefits.
901	Adapted from the report on the 4th AFNET/EHRA consensus conference. ⁷⁶
902	AF = atrial fibrillation; AFNET = German Competence NETwork on Atrial Fibrillation; EHRA = European
903	Heart Rhythm Association.
904	
905	An integrated, structured approach to AF care, as applied successfully to other domains of medicine, ³²²⁻³²⁴ will
906	facilitate consistent, guideline-adherent AF management for all patients ³²⁵ (Figure 6), with the potential to
907	improve outcomes. ^{42, 326, 327} Such approaches are consistent with the Innovative Care for Chronic Conditions
908	Framework proposal put forward by the World Health Organization. ³²⁸ Review by an AF service, or at least
909	referral to a cardiologist, will usually be required after the initial assessment to fully evaluate the effect of AF on
910	cardiovascular health. ³²⁹ There may also be reasons for early or urgent referral (<i>Table 9</i>). Integrated care of all
911	patients with newly diagnosed AF should help to overcome the current shortcomings of AF management, such
912	as underuse of anticoagulation, access to rate and rhythm control therapy, and inconsistent approaches to
913	cardiovascular risk reduction. Integrated AF care requires the cooperation of primary care physicians,
914	cardiologists, cardiac surgeons, AF specialists, stroke specialists, allied health practitioners and patients,

915 encompassing lifestyle interventions, treatment of underlying cardiovascular diseases and AF-specific therapy 916 (Figure 7).



917 918 Figure 6 Achieving optimal management of AF patients.

919 AF = atrial fibrillation.

920 ^aOn-site or through institutionalized cooperation.

921 ^bSafety outcomes should be collected in published and monitored central databases.

Integrated AF management

- Central role in care process
- Patient education
- Encouragement and empowerment for selfmanagement
- Advice and education on lifestyle and risk factor
- management Shared decision Making
- Informed, involved, empowered patient

- Physicians (general physicians, cardiology and stroke AF specialists, surgeons) and allied health professionals work in a collaborative practice model
- Efficient mix of communication skills. education, and experience
- Working together in a multidisciplinary chronic AF care team

Technology tools

- Information on AF Clinical decision support
- Checklist and communication tool
- Used by healthcare professionals and patients
- Monitoring of therapy adherence and

effectiveness

 Navigation system to support decision making in treatment team

- Structured support for lifestyle changes
- Anticoagulation
- Rate control
- Antiarrhythmic drugs
- Catheter and surgical interventions (ablation, LAA occluder, AF surgery, etc)
- Complex management decisions underpinned by an AF Heart Team

922 923

Figure 7 Fundamentals of integrated care in AF patients. 924 AF = atrial fibrillation; LAA = left atrial appendage. 925 926 Table 9 Clinical signs calling for urgent involvement of a specialized AF service.^a Haemodynamic instability Uncontrollable rate Symptomatic bradycardia not amenable to reduced dosing of rate control agents Severe angina or worsening left ventricular function Transient ischemic attack or stroke

927 AF = atrial fibrillation

928 ^aAnticoagulation should be initiated early in all suitable patients and will not routinely require specialist input.

929

930 8.1. **Evidence supporting integrated atrial fibrillation care**

931 Several structured approaches to AF care have been developed. Some evidence underpins their use, while more 932 research is needed into the best way of delivering integrated AF care. Integrated AF management in an RCT 933 increased the use of evidence-base care and reduced by approximately one-third the composite outcome of 934 cardiovascular hospitalization and cardiovascular death over a mean follow-up of 22 months (14.3% vs. 20.8%, 935 HR 0.65; 95% CI 0.45–0.93; P = 0.017) compared with usual care in a large tertiary care centre.³³⁰ Integrated AF management appeared cost-effective in that study.³³¹ However, an Australian RCT showed only a marginal 936 effect on unplanned admissions and death using integrated AF care limited to the initial care period, possibly 937 emphasizing the need for sustained integration of AF care.³³² Two observational studies of integrated AF care found fewer hospitalizations,^{333, 334} one study showed fewer cases of stroke,³³³ and a further non-randomized 938 939 940 study identified a trend for a lower rate of the composite outcome of death, cardiovascular hospitalization, and 941 AF-related emergency visits.³³⁵ More research is needed, and integrated AF care is likely to require different 942 designs in different healthcare settings. 943

944 **Components of integrated atrial fibrillation care** 8.2.

945 8.2.1. Patient involvement

946 Patients should have a central role in the care process. As treatment of AF requires patients to change their 947 lifestyles and adhere to chronic therapy, at times without an immediately tangible benefit, they need to 948 understand their responsibilities in the care process. Physicians and healthcare professionals are responsible for 949 providing access to evidence-based therapy, but adherence to therapy is ultimately the responsibility of informed and autonomous patients, best described as 'shared accountability'.³³⁶ Hence, information and 950 education of patients and often of their partners and relatives is indispensable to encourage a self-management 951 role and to empower patients to participate in shared decision-making, ^{326, 328} and to support their understanding 952 953 of the disease and the suggested treatments.³³⁷

955 8.2.2. Multidisciplinary atrial fibrillation teams

956 Delegation of tasks from specialists to general physicians and from physicians to allied health professionals is a 957 fundamental concept of integrated care models. A multidisciplinary AF team approach includes an efficient mix 958 of interpersonal and communication skills, education and expertise in AF management, as well as the use of 959 dedicated technology. This approach underlines the importance of redesigning daily practice in a way that 960 encourages non-specialists and allied professionals to have an important role in educating patients and 961 coordinating care, while the specialist remains medically responsible. Cultural and regional differences will 962 determine the composition of AF teams.

964 8.2.3. Role of non-specialists

AF patients often initially present to general practitioners or pharmacists. Some physicians in primary care have
 extensive expertise in stroke prevention and initial management of AF patients. Others may seek training to
 acquire such knowledge. Other components of AF management (e.g. assessment of concomitant cardiovascular
 conditions, antiarrhythmic drug therapy, or interventional treatment) often require specialist input. Integrated
 AF care structures should support treatment initiation by non-specialists where appropriate, and provide ready
 access to specialist knowledge to optimize AF care.

972 8.2.4. Technology use to support atrial fibrillation care

973 Technology, such as decision support software, has the potential to enhance the implementation of evidence974 based care and improve outcomes, when used to enhance expert advice.³³⁸ Electronic tools can also ensure
975 coherent communication within the AF team. With a view to support the wider use of such technology, this
976 Task Force is providing tools free of charge, in the form of smartphone apps, to AF healthcare professionals and

977 to AF patients. 978

954

963

979 Recommendations for an integrated approach to care

Recommendations	Class ^a	Level ^b	Ref s ^c
An integrated approach with structured organization of care and follow- up should be considered in all patients with AF, aiming to improve guideline adherence and reduce hospitalization and mortality	IIa	В	330-332
Placing patients in a central role in the decision-making should be considered in order to tailor management to patient preferences and improve adherence to chronic therapy	IIa	С	330, 332, 334

980 AF = atrial fibrillation

- 981 ^aClass of recommendation.
- 982 ^bLevel of evidence.
- 983 ^cReference(s) supporting recommendations.
- 984

985 8.3. Diagnostic workup of atrial fibrillation patients

AF is often found in patients with other, at times undiagnosed, cardiovascular conditions. Thus, all AF patients
 will benefit from a comprehensive cardiovascular assessment.³³⁹

989 **8.3.1.** Recommended evaluation in all atrial fibrillation patients

990 A complete medical history should be taken and all patients should undergo clinical evaluation that includes

- thorough assessment for concomitant conditions, establishing the AF pattern, estimation of stroke risk and AFrelated symptoms, and assessment of arrhythmia-related complications such as thromboembolism or LV
- related symptoms, and assessment of arrhythmia-related complications such as thromboembolism or LV
- 993 dysfunction. A 12-lead ECG is recommended to establish a suspected diagnosis of AF, to determine rate in AF,

994 and to screen for conduction defects, ischaemia, and signs of structural heart disease. Initial blood tests should 995 evaluate thyroid and kidney function as well as serum electrolytes and full blood count. Transthoracic

evaluate thyroid and kidney function as well as serum electrolytes and full blood count. Transthoracic
 echocardiography is recommended in all AF patients to guide treatment decisions. Transthoracic

997 echocardiography is recommended in an Air patients to guide deathent decisions. Transmonate 997 echocardiography should be used to identify structural disease (e.g. valvular disease) and assess LV size and

998 function (systolic and diastolic), atrial size, and right heart function.^{339, 340} Although biomarkers such as

999 natriuretic peptides are elevated in AF patients, there is insufficient data to suggest that blood-based parameters are independent markers for AF.³⁴¹⁻³⁴³

1001Additional investigations in selected patients with atrial fibrillation8.3.2.Additional investigations in selected patients with atrial fibrillation

Ambulatory ECG monitoring in AF patients can assess the adequacy of rate control, relate symptoms with AF recurrences, and detect focal induction of bouts of paroxysmal AF. Transoesophageal echocardiography (TOE) is useful to further assess valvular heart disease and to exclude intracardiac thrombi, especially in the LAA, to facilitate early cardioversion or catheter ablation.³⁴⁴ Patients with symptoms or signs of myocardial ischaemia should undergo coronary angiography or stress testing as appropriate. In patients with AF and signs of cerebral ischaemia or stroke, computed tomography (CT) or magnetic resonance imaging (MRI) of the brain is

1009 recommended to detect stroke and support decisions regarding acute management and long-term

anticoagulation. Delayed-enhancement MRI of the left atrium using gadolinium contrast,³⁴⁵⁻³⁴⁷ T1 mapping
 using cardiac MRI,³⁴⁷ and intracardiac echo³⁴⁸ may help to guide treatment decisions in AF, but require external
 validation in multicentre studies.

1012

1014 8.4. Structured follow-up

1015 Most AF patients need regular follow-up to ensure continued optimal management. Follow-up may be

1016 undertaken in primary care, by specially trained nurses, by cardiologists, or by AF specialists.^{325, 330} A specialist

1017 should coordinate care and follow-up. Follow-up should ensure implementation of the management plan,

1018 continued engagement of the patient, and therapy adaptation where needed.

1019

1020 **Recommendations for diagnostic workup of AF patients**

Recommendations	Class ^a	Level ^b	Refs ^c
ECG documentation is required to establish the diagnosis of AF	I	В	349
A full cardiovascular evaluation, including an accurate history, careful clinical examination, and assessment of concomitant conditions, is recommended in all AF patients	I	С	
Transthoracic echocardiography is recommended in all AF patients to guide management	I	С	339
Long-term ECG monitoring should be considered in selected patients to assess the adequacy of rate control in symptomatic patients and to relate symptoms with AF episodes	lla	С	

1021 AF = atrial fibrillation; ECG = electrocardiogram.

1022 ^aClass of recommendation.

- 1023 ^bLevel of evidence.
- 1024 ^cReference(s) supporting recommendations.
- 1025

1026 8.5. Defining goals of atrial fibrillation management

1027AF management comprises therapies with prognostic impact (anticoagulation and treatment of cardiovascular1028conditions) and therapies predominantly providing symptomatic benefit (rate control, rhythm control, *Table 10*).1029Therapies with prognostic benefit need careful explanation to patients when their benefits are not directly felt.1030Rhythm control therapy can be successful if symptoms are controlled, even when AF recurs. Explaining the1031expected benefits to each patient at the start of AF management will prevent unfounded expectations and has the1032potential to optimize quality of life.

1033

1034 **Table 10** Goal-based follow-up

Category

Intervention

Follow-up aspects

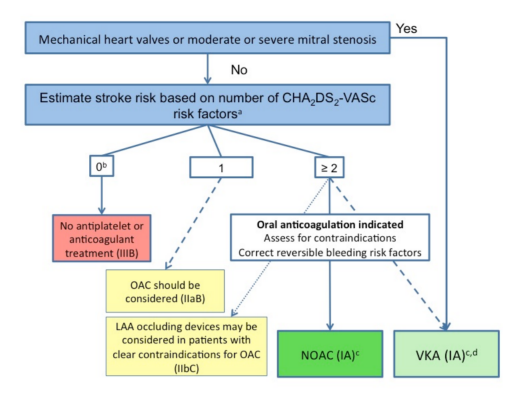
Prognostic	Comorbidity control	Obesity	Weight loss
	(relevant examples	Arterial hypertension	Blood pressure control
	given)	Heart failure	Heart failure therapy
		Coronary artery disease	Statin and antiplatelet therapy
			Revascularization
		Diabetes	Glycaemic control
		Valvular Heart Disease	Valve repair or replacement
Prognostic	Anticoagulation	Indication (risk profile; timing,	Stroke
		e.g. post-cardioversion);	Bleeding
		Adherence (NOAC or VKA)	Mortality
		and INR (if VKA);	
		NOAC dosing (co-	
		medications, age, weight, renal	
		function)	
Mainly symptomatic	Rate control	Symptoms	EHRA score
Partly prognostic		Average resting heart rate	Heart failure status
		< 110 bpm	LV function
			Exercise capacity
Symptomatic at present	Rhythm control	Symptoms vs. side-effects	Hospitalization
		Exclusion of proarrhythmia	Therapy complications
		(PR; QRS; QTc interval)	
Relevant for	Patient education and	Knowledge (about disease;	Adherence to therapy
implementation of and	self-care capabilities	about treatment; about	Directed evaluation, preferably
adherence to therapy		management goals)	based on systematic checklists
		Capabilities (what to do if)	
Relevant for chronic	Caregiver	Who? (spouse; GP; home	Directed evaluation of task
care management	involvement	nurse; pharmacist)	performance (e.g. via patient
		Clearly spelling out	card)
		participation roles	Dispensed medication
		Knowledge and capabilities	GP log of follow-up visits

1035 bpm = beats per minute; EHRA = European Heart Rhythm Association; GP = general practitioner; INR = 1036 international normalized ratio; LV = left ventricular; NOAC = non-vitamin K antagonist oral anticoagulant; 1037 VKA = vitamin K antagonist. 1038

9 Stroke prevention therapy in atrial fibrillation patients

1039 OAC therapy can prevent the majority of ischaemic strokes in AF patients and can prolong life.^{38, 39, 42, 194, 201, 329,} 1040 ³⁵⁰⁻³⁵² It is superior to no treatment or aspirin in patients with different profiles for stroke risk. ^{353, 354} The net 1041 1042 clinical benefit is almost universal, with the exception of patients at very low stroke risk, and OAC should 1043 therefore be used in most patients with AF (Figure 8). Despite this evidence, underuse or premature termination 1044 of OAC therapy is still common. Bleeding events, both severe and nuisance bleeds, a perceived 'high risk of bleeding' on anticoagulation, and the efforts required to monitor and dose-adjust VKA therapy are among the most common reasons for withholding or ending OAC.^{352, 355-359} However, the considerable stroke risk without 1045 1046 OAC often exceeds the bleeding risk on OAC, even in the elderly, in patients with cognitive dysfunction, or in patients with frequent falls or frailty.^{360, 361} The bleeding risk on aspirin is not different to the bleeding risk on VKA³⁶² or NOAC therapy,^{354, 363} while VKA and NOACs, but not aspirin, effectively prevent strokes in AF 1047 1048 1049 patients. 38, 354, 362, 363 1050

1051



1052

1004	
1053	Figure 8 Stroke prevention in AF.
1054	AF = atrial fibrillation; CHA_2DS_2 -VASc = Congestive Heart failure, hypertension, Age \geq 75 (doubled),
1055	Diabetes, Stroke (doubled), Vascular disease, Age 65–74, and Sex (female); LAA = left atrial appendage;
1056	NOAC = non-vitamin K antagonist oral anticoagulant; OAC = oral anticoagulation; VKA = vitamin K
1057	antagonist.
1058	^a Congestive heart failure, hypertension, age \geq 75 years (2 points), diabetes, prior stroke/TIA/embolus (2 points),
1059	vascular disease, age 65–74, female sex.
1060	^b Includes women without other stroke risk factors.
1061	^c IIaB for women with only one additional stroke risk factor,
1062	^d IB for patients with mechanical heart valves or mitral stenosis
1063	

1064 9.1. Prediction of stroke and bleeding risk

1065 9.1.1. Clinical risk scores for stroke and systemic embolism

Simple, clinically applicable stroke risk-stratification schemes in AF patients were developed in the late 1990s in small cohort studies and have later been refined and validated in larger populations.³⁶⁴⁻³⁶⁸ The introduction of the CHA₂DS₂-VASc score (*Table 11*) has clearly simplified the initial decision for OAC in AF patients. Since its first incorporation in the ESC guidelines in 2010,³⁶⁹ it has been widely used.³⁷⁰ We recommend estimating stroke risk in AF patients based on the CHA₂DS₂-VASc score.³⁶⁸ In general, patients without clinical stroke risk factors do not need antithrombotic therapy, while patients with stroke risk factors (i.e. CHA₂DS₂-VASc score of 1 or more for men, and 2 or more for women) are likely to benefit from OAC.

1074Table 11 Clinical risk factors for stroke, transient ischemic attack, and systemic embolism in the1075CHA2DS2-VASc score.

CHA ₂ DS ₂ -VASc risk factor	Points
Congestive heart failure	+1
Signs/symptoms of heart failure or objective evidence of reduced left-ventricular ejection fraction	

Hypertension	+1
Resting blood pressure > 140/90 mmHg on at least two occasions or current antihypertensive	
treatment	
Age 75 years or older	+2
Diabetes mellitus	+1
Fasting glucose > 125 mg/dL or treatment with oral hypoglycaemic agent and/or insulin	
Previous stroke, transient ischemic attack, or thromboembolism	+2
Vascular disease	+1
Previous myocardial infarction, peripheral artery disease, or aortic plaque	
Age 65 to 74 years	+1
Sex category (female)	+1

 CHA_2DS_2 -VASc = Congestive Heart failure, hypertension, Age >75 (doubled), Diabetes, Stroke (doubled), Vascular disease, Age 65–74, and Sex (female).

1077 1078

1076

1079 Other, less established risk factors for stroke include unstable international normalized ratio (INR) and low time 1080 in therapeutic range (TTR) in patients treated with VKAs; previous bleed or anaemia; alcohol excess and other 1081 markers for decreased therapy adherence; CKD; elevated high-sensitivity troponin T; and elevated N-terminal 1082 pro-B-type natriuretic peptide.

1083

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1084 9.1.2. Anticoagulation in patients with a CHA₂DS₂-VASc score of 1 in men and 2

in women

Controlled trials studying OAC in AF patients have been enriched for patients at high risk of stroke,^{38, 39, 42, 194, 201, 329, 351, 352} and hence there is strong evidence that patients with a CHA₂DS₂-VASc risk score of 2 or more in 1086 1087 1088 men, and 3 or more in women benefit from OAC. Fortunately, we now have a growing evidence-base regarding 1089 stroke risk in patients with one clinical risk factor (i.e. CHA2DS2-VASc score of 1 for men, and 2 for women), although this relies largely on observed stroke rates in patients not receiving OAC. In many of these patients, anticoagulation seems to provide a clinical benefit.³⁷¹⁻³⁷⁵ The rates of stroke and thromboembolism vary 1090 1091 considerably in patients with CHA_2DS_2 -VASc scores of 1 or 2 due to differences in outcomes, populations, and anticoagulation status (*Web Addenda Table 1*).^{371, 376, 377, 1041} OAC should be considered for men with a 1092 1093 1094 CHA₂DS₂-VASc score of 1 and women with a score of 2, balancing the expected stroke reduction, bleeding risk, and patient preference. Importantly, age (65 years and older) conveys a relatively high and continuously 1095 1096 increasing stroke risk that also potentiates other risk factors (such as heart failure and sex). Hence, an 1097 individualized weighing of risk, as well as patient preferences, should inform the decision to anticoagulate 1098 patients with only one CHA2DS2-VASc risk factor, apart from female sex. Female sex does not appear to increase stroke risk in the absence of other stroke risk factors (Web Addenda Table 1).^{378, 379} 1099

Measurement of cardiac troponin (high-sensitivity troponin T or I) and N-terminal pro-B-type natriuretic peptide may provide additional prognostic information in selected AF patients.³⁸⁰⁻³⁸² Biomarker-1100 1101 1102 based risk scores may in the future prove helpful to better stratify patients (e.g. those at a truly low risk of stroke).75, 382 1103

1105 9.1.3. Clinical risk scores for bleeding

1106 Several bleeding risk scores have been developed, mainly in patients on VKAs. These include HAS-BLED 1107 (hypertension, abnormal renal/liver function [1 point each], stroke, bleeding history or predisposition, labile 1108 INR, elderly [>65 years], drugs/alcohol concomitantly [1 point each]), ORBIT (Outcomes Registry for Better Informed Treatment of Atrial Fibrillation), and more recently, the ABC (age, biomarkers, clinical history) bleeding score, which also makes use of selected biomarkers.³⁸³⁻³⁸⁵ Stroke and bleeding risk factors overlap 1109 1110 (compare *Table 11* and *Table 12*). For example, older age is one of the most important predictors of both ischaemic stroke and bleeding in AF patients.^{386, 387} A high bleeding risk score should generally not result in 1111 1112 1113 withholding OAC. Rather, bleeding risk factors should be identified and treatable factors corrected (see Section 1114 8.5). Table 12 provides details of modifiable bleeding risk factors. 1115

1116 Table 12 Modifiable and non-modifiable risk factors for bleeding in anticoagulated patients based on 1117 bleeding risk scores.

Modifiable bleeding risk factors Hypertension (especially when systolic blood pressure is $> 160 \text{ mmHg})^{a,b,c}$

Medication pre	disposing to bleeding, such as antiplatelet drugs and non-steroidal anti-inflammatory drugs ^{a,}
	$(\geq 8 \text{ drinks/week})^{a,b}$
Potentially mo	difiable bleeding risk factors
Anaemia ^{b,c,d}	
Impaired renal	
Impaired liver	
	et count or function ^b
Non-modifiab	le bleeding risk factors
Age ^e (> 65 yea	$rs)^a (\geq 75 years)^{b,c,d}$
	or bleeding ^{a,b,c,d}
Previous stroke	ab
	dent CKD or renal transplant ^{a,c}
Cirrhotic liver	disease ^a
Malignancy ^b	
Genetic factors	b
Biomarker-ba	sed bleeding risk factors
High-sensitivit	y troponin T ^e
Growth differe	ntiation factor-15 ^e
Serum creatini	ne/estimated CrCL ^e

- 1118 ABC = age, biomarkers, clinical history; ATRIA = AnTicoagulation and Risk factors In Atrial fibrillation; CKD
- 1119 = chronic kidney disease; CrCl = creatinine clearance; HAS-BLED = hypertension, abnormal renal/liver
- 1120 function (1 point each), stroke, bleeding history or predisposition, labile INR, elderly (>65 years), drugs/alcohol
- 1121 concomitantly (1 point each); INR = international normalized ratio; ORBIT = Outcomes Registry for Better 1122 Informed Treatment of Atrial Fibrillation; TTR = time in therapeutic range; VKA = vitamin K antagonist.
- 1123 ^aDerived from the HAS-BLED score.³⁸⁴
- ^bDerived from the HEMORR₂HAGES score.³⁸³ 1124
- 1125 ^cDerived from the ATRIA score.^{38:}
- ^dDerived from the ORBIT score.³⁸⁸ 1126
- 1127 ^eDerived from the ABC bleeding score.³⁸⁷
- 1128

1129 Recommendations for prediction of stroke and bleeding risk

Recommendations	Class ^a	Level ^b	Refs ^c
The CHA ₂ DS ₂ -VASc score is recommended for stroke risk prediction in patients with AF	I	A	368, 371, 386
Bleeding risk scores should be considered in AF patients on oral anticoagulation to identify modifiable factors for major bleeding	lla	В	384, 386, 387, 389-392
Biomarkers such as high-sensitivity troponin and N-terminal pro- B-type natriuretic peptide may be considered to further refine stroke and bleeding risk in AF patients	llb	В	380-382, 387, 393

- 1130 AF = atrial fibrillation; CHA_2DS_2 -VASc = Congestive Heart failure, hypertension, Age \geq 75 (doubled),
- 1131 Diabetes, Stroke (doubled), Vascular disease, Age 65–74, and Sex (female); OAC = oral anticoagulation.
- 1132 ^aClass of recommendation.
- 1133 ^bLevel of evidence.
- 1134 ^cReference(s) supporting recommendations. 1135

1136 9.2. **Stroke prevention**

1137 9.2.1. Vitamin K antagonists

1138 Warfarin and other VKAs were the first anticoagulants used in AF patients. VKA therapy reduces risk of stroke 1139

- by two-thirds and mortality by one-quarter compared with control (aspirin or no therapy).³⁸ VKAs have been used in many patients throughout the world with good outcomes,³⁹⁴⁻³⁹⁶ and this is reflected in the warfarin arms 1140
- 1141 of the NOAC trials (see Section 8.2.2.). The use of VKAs is limited by the narrow therapeutic interval,
- 1142 necessitating frequent monitoring and dose adjustments, but VKAs, when delivered with adequate TTR, are

1143 effective for stroke prevention in AF patients. Clinical parameters can help to identify patients who are likely to 1144 achieve a decent TTR on VKA therapy.³⁹⁷ These have been summarized in the SAMe-TT₂R₂ score. Patients who 1145 fare well on this score, when treated with a VKA, have on average a higher TTR than patients who do not fare well on the score.^{398, 399} VKAs are currently the only treatment with established safety in AF patients with 1146 1147 rheumatic mitral valve disease and/or a mechanical heart valve prosthesis.⁴⁰⁰

1148 1149 9.2.2. Non-vitamin K antagonist oral anticoagulants

1150 NOACs, including the direct thrombin inhibitor dabigatran and the factor Xa inhibitors apixaban, edoxaban, and 1151 rivaroxaban, are suitable alternatives to VKAs for stroke prevention in AF (Table 13). Their use in clinical 1152 practice is increasing rapidly.⁴⁰¹ All NOACs have a predictable effect (onset and offset) without need for regular 1153 anticoagulation monitoring. The phase III trials have been conducted with carefully selected doses of the 1154 NOACs, including clear rules for dose reduction that should be followed in clinical practice (Table 13).

1155 1156 Apixaban

1157 In the ARISTOTLE (Apixaban for Reduction in Stroke and Other Thromboembolic Events in Atrial 1158 Fibrillation) trial,³¹⁹ apixaban reduced stroke or systemic embolism by 21% compared with warfarin, combined 1159 with a 31% reduction in major bleeding and an 11% reduction in all-cause mortality (all statistically significant). 1160 Rates of haemorrhagic stroke and intracranial haemorrhage, but not of ischaemic stroke, were lower on apixaban. Rates of gastrointestinal bleeding were similar between the two treatment arms.⁴⁰² 1161

1162 Apixaban is the only NOAC that has been compared with aspirin in AF patients: apixaban significantly 1163 reduced stroke or systemic embolism by 55% compared with aspirin, with no significant difference in rates of 1164 major bleeding or intracranial haemorrhage.^{354, 403} 1165

1166 Dabigatran

1167 In the RE-LY (Randomized Evaluation of Long-Term Anticoagulation Therapy) study,^{318, 404} dabigatran 150 mg 1168 twice daily reduced stroke and systemic embolism by 35% compared with warfarin without a significant 1169 difference in major bleeding events. Dabigatran 110 mg twice daily was non-inferior to warfarin for prevention 1170 of stroke and systemic embolism, with 20% fewer major bleeding events. Both dabigatran doses significantly 1171 reduced haemorrhagic stroke and intracranial haemorrhage. Dabigatran 150 mg twice daily significantly 1172 reduced ischaemic stroke by 24% and vascular mortality by 12%, while gastrointestinal bleeding was 1173

significantly increased by 50%. There was a non-significant numerical increase in the rate of myocardial infarction with both dabigatran doses,^{318, 404} which has not been replicated in large post-authorization

1174

1175 analyses.³⁹⁶ These data have also replicated the benefit of dabigatran over VKAs found in the RE-LY trial in 1176 patients enriched for the higher dabigatran dose (150 mg twice daily).³⁹⁶

1177 1178 Edoxaban

In the ENGAGE AF-TIMI 48 (Effective Anticoagulation with Factor Xa Next Generation in Atrial Fibrillation– Thrombolysis in Myocardial Infarction 48) trial,³²¹ edoxaban 60 mg once daily and edoxaban 30 mg once daily 1179 1180 (with dose reductions in certain patients according to *Table 13*), were compared with adjusted-dose warfarin.⁴⁰⁵ 1181 1182 Edoxaban 60 mg once daily was non-inferior to warfarin (primary outcome, HR 0.87; 97.5% CI 0.73–1.04; P = 1183 0.08). In an on-treatment analysis, edoxaban 60 mg once daily significantly reduced stroke or systemic 1184 embolism by 21% and significantly reduced major bleeding events by 20% compared with warfarin, while 1185 edoxaban 30 mg once daily was non-inferior to warfarin for prevention of stroke and systemic embolism but 1186 significantly reduced major bleeding events by 53%. Cardiovascular death was reduced in patients randomized 1187 to edoxaban 60 mg once daily or edoxaban 30 mg once daily compared with warfarin. Only the higher dose

- 1188 regimen has been approved for stroke prevention in AF.
- 1189 1190 Rivaroxaban

1191 In the ROCKET-AF (Rivaroxaban Once Daily Oral Direct Factor Xa Inhibition Compared with Vitamin K 1192 Antagonism for Prevention of Stroke and Embolism Trial in Atrial Fibrillation) trial,³²⁰ patients were 1193 randomized to rivaroxaban 20 mg once daily or VKA, with a dose adjustment to 15 mg daily for those with

1194 estimated CrCl 30-49 mL/min by the Cockroft-Gault formula. Rivaroxaban was non-inferior to warfarin for the

1195 prevention of stroke and systemic embolism in the intent-to-treat analysis, while the per-protocol on-treatment

1196 analysis achieved statistical superiority with a 21% reduction in stroke or systemic embolism compared with

1197 warfarin. Rivaroxaban did not reduce the rates of mortality, ischaemic stroke, or major bleeding events

- 1198 compared to VKA. There was an increase in gastrointestinal bleeding events, but a significant reduction in
- 1199 haemorrhagic stroke and intracranial haemorrhage with rivaroxaban compared with warfarin. Comparable event
- 1200 rates have been reported in post-authorization analyses, which are part of the post-approval risk-management
- process.406,407 1201

1202 **Table 13 NOACs compared with warfarin in controlled trials**

	Dabigatran (RE-LY)	Rivaroxaban (ROCKET-AF)	Apixaban (ARISTOTLE)	Edoxaban (ENGAGE AF-TIMI 48)
Mechanism	Oral direct thrombin inhibitor	Oral direct factor Xa inhibitor	Oral direct factor Xa inhibitor	Oral direct factor Xa inhibitor
Bioavailability, %	6	66 fasting, 80–100 with food	50	62
Time to peak levels, h	3	2-4	3	1–2
Half-life, h	12–17	5–13	9–14	10–14
Excretion	80% renal	66% liver, 33% renal	27% renal	50% renal
Dose	150 mg or 110 mg twice daily	20 mg once daily	5 mg twice daily	60 mg or 30 mg once daily
Dose reduction in selected patients		Rivaroxaban 15 mg once daily if CrCl 30–49 mL/min	Apixaban 2.5 mg twice daily if at least 2 of age \geq 80 years, body weight \leq 60 kg or serum creatinine level \geq 1.5 mg/dL (133 µmol/L)	Edoxaban 60 mg reduced to 30 mg once daily, and edoxaban 30 mg reduced to 15 mg once daily, if any of the following: CrCl 30– 50 mL/min, body weight \leq 60 kg, concomitant use of verapamil or quinidine or dronedarone
Study design	Randomized, open-label	Randomized, double-blind	Randomized, double-blind	Randomized, double-blind
Number of patients	18,113	14,264	18,201	21,105
Follow-up period, years	2	1.9	1.8	2.8
Randomized groups	Dose-adjusted warfarin vs. blinded doses of dabigatran (150 mg twice daily or 110 mg twice daily)	Dose-adjusted warfarin vs. rivaroxaban 20 mg once daily	Dose-adjusted warfarin vs. apixaban 5 mg twice daily	Dose-adjusted warfarin vs. edoxaban (60 mg once daily or 30 mg once daily)
Age, years	Mean ± SD 71.5 ± 8.7	Median 73; IQR 65–78	Median 70; IQR 63–76	Median 72; IQR 64–78
Men, %	63.6	60.3	64.5	61.9
CHADS ₂ score (mean)	2.1	3.5	2.1	2.8

1203

	Warfarin	Dabigatran 150	Dabigatran 110	Warfarin	Rivaroxaban	Warfarin	Apixaban	Warfarin	Edoxaban 60	Edoxaban 30
	<i>n</i> = 6022	<i>n</i> = 6076	<i>n</i> = 6015	<i>n</i> = 7133	<i>n</i> = 7131	<i>n</i> = 9081	<i>n</i> = 9120	<i>n</i> = 7036	<i>n</i> = 7035	<i>n</i> = 7034
	Event rate, %/year	Event rate, %/year (RR vs. warfarin)	Event rate, %/year (RR vs. warfarin)	Event rate, %/year	Event rate, %/year (HR vs. warfarin)	Event rate, %/year	Event rate, %/year (HR vs. warfarin)	Event rate, %/year	Event rate, %/year (HR vs. warfarin)	Event rate, %/year (HR vs. warfarin)
Stroke/systemic embolism	1.72	1.12 (0.65, 0.52–0.81; <i>P</i> for non- inferiority and superiority < 0.001)	1.54 (0.89, 0.73–1.09; <i>P</i> for non- inferiority < 0.001)	2.42	2.12 (0.88, 0.75–1.03; <i>P</i> for non-inferiority < 0.001, <i>P</i> for superiority = 0.12)	1.60	1.27 (0.79, 0.66–0.95; P < 0.001 for non- inferiority, P = 0.01 for superiority)	1.80	1.57 (0.87, 0.73– 1.04; <i>P</i> < 0.001 for non-inferiority, <i>P</i> = 0.08 for superiority)	2.04 (1.13, 0.96–1.34; P = 0.005 for non- inferiority, $P = 0.10$ for superiority)
Ischaemic stroke	1.22	$\begin{array}{l} 0.93 \ (0.76, \\ 0.59 - 0.97; \\ P = 0.03) \end{array}$	$\begin{array}{l} 1.34 \ (1.10, \\ 0.88 - 1.37; \\ P = 0.42) \end{array}$	1.42	$\begin{array}{l} 1.34 \ (0.94; \ 0.75-1.17; \\ P = 0.581) \end{array}$	1.05	$\begin{array}{l} 0.97 \ (0.92, \ 0.74 - 1.13; \\ P = 0.42) \end{array}$	1.25	1.25 (1.00, 0.83– 1.19; P = 0.97)	$\begin{array}{c} 1.77 \ (1.41, \\ 1.19 - 1.67; \\ P < 0.001) \end{array}$
Haemorrhagic stroke	0.38	0.10 (0.26, 0.14–0.49; <i>P</i> < 0.001)	0.12 (0.31, 0.17–0.56; <i>P</i> < 0.001)	0.44	0.26 (0.59; 0.37-0.93; P = 0.024)	0.47	$\begin{array}{c} 0.24 \ (0.51, \ 0.35 - 0.75; \\ P < 0.001) \end{array}$	0.47	0.26 (0.54, 0.38– 0.77; <i>P</i> < 0.001)	0.16 (0.33, 0.22–0.50; <i>P</i> < 0.001)
Major bleeding	3.61	3.40 (0.94, 0.82-1.08; P = 0.41)	2.92 (0.80, 0.70-0.93; P = 0.003)	3.45	3.60 (1.04; 0.90-2.30; P = 0.58)	3.09	2.13 (0.69, 0.60–0.80; <i>P</i> < 0.001)	3.43	2.75 (0.80, 0.71– 0.91; <i>P</i> < 0.001)	1.61 (0.47, 0.41–0.55; <i>P</i> < 0.001)
Intracranial bleeding	0.77	0.32 (0.42, 0.29–0.61; <i>P</i> < 0.001)	0.23 (0.29 0.19–0.45; <i>P</i> < 0.001)	0.74	0.49 (0.67; 0.47-0.93; P = 0.02)	0.80	0.33 (0.42, 0.30–0.58; <i>P</i> < 0.001)	0.85	0.39 (0.47, 0.34– 0.63; <i>P</i> < 0.001)	0.26 (0.30, 0.21–0.43; <i>P</i> < 0.001)
Gastrointestinal major bleeding	1.09	1.60 (1.48, 1.19–1.86; <i>P</i> < 0.001)	1.13 (1.04, 0.82–1.33; P = 0.74)	1.24	2.00 (1.61; 1.30-1.99; <i>P</i> < 0.001)	0.86	0.76 (0.89, 0.70–1.15; <i>P</i> = 0.37)	1.23	1.51 (1.23, 1.02– 1.50; <i>P</i> = 0.03)	0.82 (0.67, 0.53–0.83; <i>P</i> < 0.001)
Myocardial infarction	0.64	$\begin{array}{l} 0.81 \ (1.27, \\ 0.94 \text{-} 1.71; \\ P = 0.12) \end{array}$	0.82 (1.29, 0.96-1.75; P = 0.09)	1.12	0.91 (0.81; 0.63–1.06; P = 0.12)	0.61	0.53 (0.88, 0.66–1.17; P = 0.37)	0.75	0.70 (0.94, 0.74– 1.19; P = 0.60)	0.89 (1.19, 0.95–1.49; P = 0.13)
Death from any cause	4.13	3.64 (0.88, 0.77-1.00; P = 0.051)	$\begin{array}{c} 3.75 \ (0.91, \\ 0.80 - 1.03; \\ P = 0.13) \end{array}$	2.21	1.87 (0.85; 0.70–1.02; P = 0.07)	3.94	3.52 (0.89, 0.80-0.99; P = 0.047)	4.35	3.99 (0.92, 0.83– 1.01; <i>P</i> = 0.08)	3.80 (0.87, 0.79–0.96; P = 0.006)

1204 $AF = atrial fibrillation; CHADS_2 = Cardiac failure, Hypertension, Age, Diabetes, Stroke (Doubled); CrCl = creatinine clearance; HR = hazard ratio; IQR = interquartile range (25th to 1205 75th quartiles); RR = risk ratio; SD = standard deviation.$

1206 RRs and HRs compared to warfarin therapy are presented with 95% confidence intervals and *P*-values.

1207

1208 9.2.3. Non-vitamin K antagonist oral anticoagulants or vitamin K antagonists

1209 Both VKAs and NOACs are effective for the prevention of stroke in AF. A meta-analysis³⁹ based on the high-1210 dose treatment groups of the pivotal studies of warfarin versus NOACs included 42,411 patients receiving a 1211 NOAC and 29,272 receiving warfarin. NOACs in these dosages significantly reduced stroke or systemic 1212 embolic events by 19% compared with warfarin (RR 0.81; 95% CI 0.73–0.91; P < 0.0001), mainly driven by a 1213 reduction in haemorrhagic stroke (RR 0.49; 95% CI 0.38–0.64; P < 0.0001). Mortality was 10% lower in 1214 patients randomized to NOAC therapy (RR 0.90; 95% CI 0.85–0.95; P = 0.0003) and intracranial haemorrhage 1215 was halved (RR 0.48; 95% CI 0.39–0.59; P < 0.0001), while gastrointestinal bleeding events were more 1216 frequent (RR 1.25; 95% CI 1.01–1.55; P = 0.04).³⁹ The stroke reduction with NOACs was consistent in all 1217 evaluated subgroups, while there was a suggestion of greater relative reduction in bleeding with NOACs at 1218 centres with poor INR control (interaction P = 0.022). Notably, the substantial reduction in intracranial 1219 haemorrhage by NOACs compared with warfarin seems unrelated to poor or good INR control.^{408,405} 1220 1221 Oral anticoagulation in atrial fibrillation patients with chronic kidney 9.2.4.

1222 disease

CKD is associated with stroke and bleeding in large data sets.^{410, 411} Anticoagulation can be safely used in AF 1223 1224 patients with moderate or moderate-to-severe CKD (glomerular filtration rate [GFR] \geq 15 mL/min): the SPAF 1225 (Stroke Prevention in Atrial Fibrillation) III trial randomized 805/1936 participants with stage 3 CKD (estimated $GFR < 59 \text{ mL/min/1.73 m}^2$), and reported good outcomes on warfarin (INR 2–3).⁴¹² This finding is supported by 1226 1227 a large Swedish database, in which stroke risk was lower in CKD patients with AF treated with warfarin 1228 (adjusted HR 0.76; 95% CI 0.72–0.80),⁴¹³ while bleeding was also slightly increased, especially during therapy 1229 initiation.⁴¹⁴ In a meta-analysis of the major NOAC trials, patients with mild or moderate CKD suffered fewer 1230 strokes, systemic emboli, or major bleeding events on NOACs than on warfarin.⁴¹⁵ Kidney function should be 1231 regularly monitored in AF patients on OAC to allow dose adaptation for those on NOACs (Table 14) and to refine risk estimation.416 1232 1233

Table 14 Inclusion criteria, dose adjustments, and outcomes in patients with chronic kidney disease in the
 four major randomized trials comparing NOACs with warfarin in patients with AF. Adapted from Hart
 *et al.*³¹⁶

<i>ei ui</i> .				
	Dabigatran (RE-LY) ^{318, 425}	Rivaroxaban (ROCKET-AF) 320, 426	Apixaban (ARISTOTLE) ^{319, 427}	Edoxaban (ENGAGE AF- TIMI 48) ³²¹
Renal clearance	80%	35%	25%	50%
Number of patients	18,113	14,264	18,201	21,105
Dose	150 mg or 110 mg twice daily	20 mg once daily	5 mg twice daily	60 mg or 30 mg once daily
Exclusion criteria for CKD	CrCl < 30 mL/min	CrCl < 30 mL/min	Serum creatinine > 2.5 mg/dL or CrCl < 25 mL/min	CrCl < 30 mL/min
Dose adjustment with CKD	None	15 mg once daily if CrCl < 30–49 mL/min	2.5 mg twice daily if serum creatinine ≥ 1.5 mg/dL plus age ≥ 80 years or weight ≤ 60 kg	30 mg or 15 mg once daily if CrCl < 50 mL/min
Per cent of patients with CKD	20% with CrCl 30–49 mL/min	21% with CrCl 30–49 mL/min	15% with CrCl 30–50 mL/dL	19% with CrCl < 50 mL/min
Reduction of stroke and systemic embolism	No interaction with CKD status	No interaction with CKD status	No interaction with CKD status	NA
Reduction of major haemorrhages compared with warfarin	Reduction in major haemorrhage with dabigatran was greater in patients with	Major haemorrhage similar	Reduction in major haemorrhage with apixaban	NA

	estimated GFR		
	> 80 mL/min		
	with either dose		

1237 AF = atrial fibrillation; CKD = chronic kidney disease; CrCl = creatinine clearance; GFR = glomerular filtration 1238 rate; NA = not available; NOAC = non-vitamin K antagonist oral anticoagulant.

1240 **9.2.5**. Oral anticoagulation in atrial fibrillation patients on dialysis

1241 Approximately one in eight dialysis patient suffers from AF, with an incidence rate of 2.7/100 patient-years.⁴¹⁷

AF is associated with increased mortality in patients on dialysis.⁴¹⁷ There are no randomized trials assessing 1242

OAC in haemodialysis patients,⁴¹⁸ and no controlled trials of NOACs in patients with severe CKD (CrCl < 25-30 mL/min).³¹⁸⁻³²¹ Warfarin use was associated either with a neutral or increased risk of stroke in database 1243

1244 1245

analyses of patients on dialysis, ⁴¹⁹⁻⁴²¹ including a population-based analysis in Canada (adjusted HR for stroke 1.14; 95% CI 0.78–1.67, adjusted HR for bleeding 1.44; 95% CI 1.13–1.85).⁴²² In contrast, data from Denmark 1246

suggest a benefit of OAC in patients on renal replacement therapy.⁴²³ Hence, controlled studies of 1247

anticoagulants (both VKAs and NOACs) in AF patients on dialysis are needed.⁴²⁴ 1248

1249

1239

1250 **9.2.6**. Patients with atrial fibrillation requiring kidney transplantation

1251 There are no randomized trials assessing OAC in patients after kidney transplantation. The prescription of

1252 NOAC therapy should be guided by the estimated GFR of the transplanted kidney. Potential pharmacokinetic

1253 interactions of OAC with immunosuppressive agents should be considered.

1254

1255

1256 9.2.7. Antiplatelet therapy as an alternative to oral anticoagulants

The evidence supporting antiplatelet monotherapy for stroke prevention in AF is very limited.^{38, 428-430} VKA 1257 1258 therapy prevents stroke, non-central nervous system embolus, myocardial infarction, and vascular death better 1259 than single or dual antiplatelet therapy with aspirin and clopidogrel (annual risk of 5.6% for aspirin and 1260 clopidogrel vs. 3.9% with VKA therapy).⁴³¹ Even greater benefits were seen in VKA-treated patients with a high TTR.⁴³² Antiplatelet therapy increases bleeding risk, especially dual antiplatelet therapy (2.0% vs. 1.3% with 1261 antiplatelet monotherapy; P < 0.001),⁴³³ with bleeding rates that are similar to those on OAC.^{354, 362, 431, 434} Thus, 1262 antiplatelet therapy cannot be recommended for stroke prevention in AF patients. 1263 1264

1265 Recommendations for stroke prevention in patients with AF

Recommendations	Class ^a	Level ^b	Refs ^c
Oral anticoagulation therapy to prevent thromboembolism is recommended for all male AF patients with a CHA ₂ DS ₂ -VASc score of 2 or more	I	A	38, 318-321, 354, 404
Oral anticoagulation therapy to prevent thromboembolism is recommended in all female AF patients with a CHA ₂ DS ₂ -VASc score of 3 or more	I	A	38, 318-321, 354, 404
Oral anticoagulation therapy to prevent thromboembolism should be considered in male AF patients with a CHA ₂ DS ₂ - VASc score of 1, considering individual characteristics and patient preferences	lla	В	371, 375-377
Oral anticoagulation therapy to prevent thromboembolism should be considered in female AF patients with a CHA ₂ DS ₂ - VASc score of 2, considering individual characteristics and patient preferences	lla	В	371, 376, 377
Vitamin K antagonist therapy (INR 2.0–3.0 or higher) is recommended for stroke prevention in AF patients with moderate-to-severe mitral stenosis or mechanical heart valves	I	В	274, 435-440
When oral anticoagulation is initiated in a patient with AF who is eligible for a NOAC (apixaban, dabigatran, edoxaban, or rivaroxaban), a NOAC is recommended in preference to a Vitamin K antagonist	Ι	A	39, 318-321, 404
When patients are treated with a vitamin K antagonist, time	I	A	395, 432, 441-444

in therapeutic range (TTR) should be kept as high as possible and closely monitored			
AF patients already on treatment with a vitamin K antagonist may be considered for NOAC treatment if TTR is not well controlled despite good adherence, or if patient preference without contraindication (e.g. prosthetic valve)	llb	A	39, 318, 319, 404, 408
Combinations of oral anticoagulants and platelet inhibitors increase bleeding risk and should be avoided in AF patients without another indication for platelet inhibition	III (harm)	В	429, 445
In male or female AF patients without additional stroke risk factors, anticoagulant or antiplatelet therapy is not recommended for stroke prevention	III (harm)	В	368, 371, 376, 377
Antiplatelet monotherapy is not recommended for stroke prevention in AF patients, regardless of stroke risk	III (harm)	A	38, 429, 430
NOACs (apixaban, dabigatran, edoxaban, and rivaroxaban) are not recommended in patients with mechanical heart valves (Level of evidence B) or moderate-to-severe mitral stenosis (Level of evidence C)	III (harm)	B/C	318-321, 400, 404

- 1266 AF = atrial fibrillation; CHA₂DS₂-VASc = Congestive Heart failure, hypertension, Age \geq 75 (doubled),
- 1267 Diabetes, Stroke (doubled), Vascular disease, Age 65–74, and Sex (female); INR = international normalized
- 1268 ratio; NOAC = non-vitamin K antagonist oral anticoagulant; OAC = oral anticoagulation; TTR = time in
- 1269 therapeutic range; VKA = vitamin K antagonist.
- 1270 ^aClass of recommendation.
- 1271 ^bLevel of evidence.
- 1272 ^cReference(s) supporting recommendations.
- 1273

1274 **9.3**. Left atrial appendage occlusion and exclusion

1275 Left atrial appendage occlusion devices 9.3.1.

Interventional LAA occlusion, 446-449 and limited experience with percutaneous LAA ligation, has mainly been 1276 1277 reported in observational studies and registries. Only one device (Watchman®) has been compared with VKA 1278 therapy in randomized trials (PROTECT AF [Watchman Left Atrial Appendage System for Embolic Protection 1279 in Patients With AF trial], see Web Addenda Table 2; and PREVAIL [Prospective Randomized Evaluation of 1280 the Watchman LAA Closure Device In Patients with AF Versus Long Term Warfarin Therapy trial]).⁴⁴⁹⁻⁴⁵¹ In these data sets, LAA occlusion was non-inferior to VKA treatment for the prevention of stroke in AF patients with moderate stroke risk, with a possibility of lower bleeding rates in the patients who continued follow-up.⁴⁵², 1281 1282 ⁴⁵³ These data were confirmed in a patient-level meta-analysis of the two trials and their associated registries.⁴⁵³ 1283 LAA occlusion may also reduce stroke risk in patients with contraindications to OAC.^{454, 455} The implantation procedure can cause serious complications,^{446, 456-458} with high event rates reported in analyses from insurance 1284 1285 databases and systematic reviews, possibly identifying a certain degree of reporting bias.^{446, 456} A large recent 1286 European registry reported a high rate of implantation success (98%), with an acceptable procedure-related complication rate of 4% at 30 days.⁴⁵⁹ Most patients who historically would be considered unsuitable for OAC therapy seem to do relatively well on contemporarily managed OAC.^{396, 407, 460} Adequately powered controlled 1287 1288 1289 1290 trials are urgently needed to inform the best use of these devices, including LAA occluders in patients who are 1291 truly unsuitable for OAC or in patients who suffer a stroke on OAC, randomized comparisons of LAA occluders 1292 with NOACs, and assessment of the minimal antiplatelet therapy acceptable after LAA occlusion. 1293

1294 9.3.2. Surgical left atrial appendage occlusion or exclusion

1295 Surgical LAA occlusion or exclusion concomitant to cardiac surgery has been performed for many decades and 1296 with various techniques. Multiple observational studies indicate the feasibility and safety of surgical LAA 1297 occlusion/exclusion, but only limited controlled trial data are available.⁴⁶¹⁻⁴⁶⁴ Residual LAA flow or incomplete 1298 LAA exclusion can increase stroke risk.⁴⁶⁵ In most studies, LAA occlusion/exclusion was performed during other open heart surgery, and more recently in combination with surgical ablation of AF⁴⁶³ or as a stand-alone 1299 1300 thoracoscopic procedure. One randomized trial evaluating the role of concomitant AF surgery and LAA occlusion reported in 2015, without a clear benefit of LAA exclusion for stroke prevention in the subgroup 1301 undergoing AF surgery.⁴⁶⁶ A large randomized trial is currently underway.⁴⁶⁷

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- 1303

1304 Recommendations for occlusion or exclusion of the LAA

Recommendations	Class ^a	Level ^b	Refs ^c
After surgical occlusion or exclusion of the LAA, it is recommended to continue anticoagulation in at-risk patients with AF for stroke prevention	Ι	В	461, 462
LAA occlusion may be considered for stroke prevention in patients with AF and contraindications for long-term anticoagulant treatment (e.g. those with a previous life-threatening bleed without a reversible cause)	IIb	В	449, 453, 454
Surgical occlusion or exclusion of the LAA may be considered for stroke prevention in patients with AF undergoing cardiac surgery	IIb	В	463
Surgical occlusion or exclusion of the LAA may be considered for stroke prevention in patients undergoing thoracoscopic ablation surgery	IIb	В	468

- 1305 AF = atrial fibrillation; LAA = left atrial appendage.
- 1306 ^aClass of recommendation.
- 1307 ^bLevel of evidence.
- 1308 ^cReference(s) supporting recommendations.
- 1309

1310 **9.4**. **Secondary stroke prevention**

1311 The most important risk factors for stroke in patients with AF are advanced age and previous cardioembolic stroke or TIA,³⁸² emphasizing the need for OAC in these patients. The highest risk of recurrent stroke is in the

- 1312 early phase after a first stroke or TIA.469,470 1313
- 1314

1315 9.4.1. Treatment of acute ischaemic stroke

1316 Systemic thrombolysis with recombinant tissue plasminogen activator (rtPA) is an effective and approved

1317 medical treatment for acute ischaemic stroke in patients presenting within 4.5 hours of symptom onset.⁴⁷¹

Systemic thrombolysis is contraindicated in patients on therapeutic OAC.^{472, 473} Recombinant tissue 1318

plasminogen activator can be given in patients treated with a VKA if the INR is below 1.7,⁴⁷⁴ or in dabigatran-1319

treated patients with a normal activated partial thromboplastin time and last intake of drug > 48 hours previously (based on expert consensus).⁴⁷² Whether specific NOAC antidotes⁴⁷⁵ could be used followed by systemic 1320

1321

thrombolysis needs to be investigated. Thrombectomy can be performed in anticoagulated patients with distal 1322 1323 occlusion of the internal carotid artery or middle cerebral artery in a 6-hour window.⁴⁷⁶

1324

1325 9.4.2. Initiation of anticoagulation after transient ischaemic attack or ischaemic

1326 stroke

1327 Data on the optimal use of anticoagulants (heparin, low-molecular-weight heparin, heparinoid, VKA, NOAC) in

1328 the first days after a stroke are scarce. Parenteral anticoagulants seem to be associated with a non-significant

1329 reduction in recurrent ischaemic stroke when administered 7 to 14 days after the acute stroke (odds ratio [OR]

1330 0.68; 95% CI 0.44-1.06), with a significant increase in symptomatic intracranial bleeding (OR 2.89; 95% CI

1.19–7.01), and a similar rate of death or disability at final follow-up.⁴⁷⁷ It seems likely that the bleeding risk on 1331

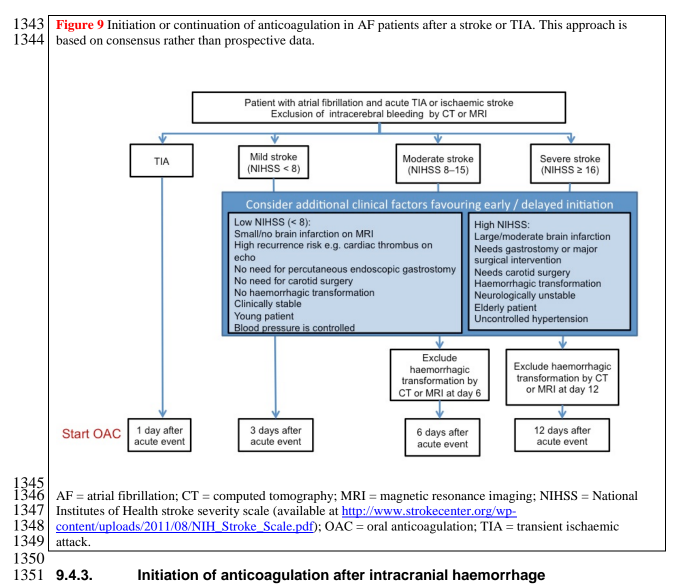
1332 parenteral anticoagulation exceeds the stroke prevention benefit in the first days after a large stroke, whereas 1333 patients with a TIA or a small stroke may benefit from early (immediate) initiation or continuation of

1334 anticoagulation. Therefore, we propose to initiate anticoagulation in AF patients between 1 and 12 days after an

- ischaemic stroke, depending on its severity (*Figure 9*).⁴⁷⁸ We suggest repeat brain imaging to determine the 1335
- optimal initiation of anticoagulation in patients with a large stroke at risk for haemorrhagic transformation. Long-term OAC with a VKA^{363, 479-481} or NOAC⁴⁸² conveys benefits in AF patients who survived a stroke. 1336
- 1337

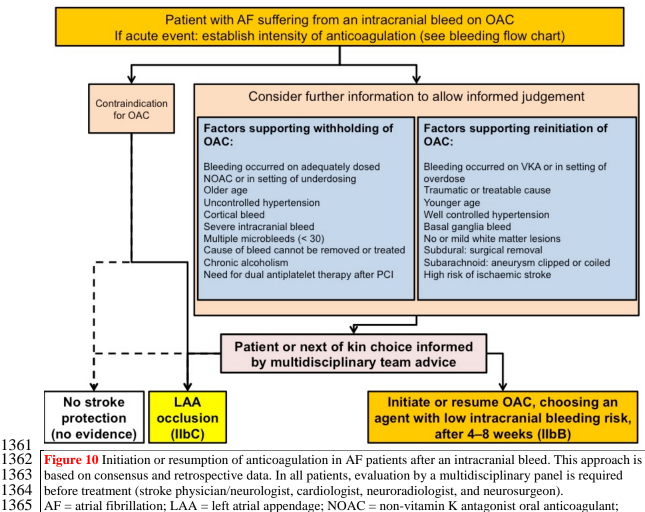
1338 NOACs seem to convey slightly better outcomes, mainly driven by fewer intracranial haemorrhages and

- haemorrhagic strokes (OR 0.44, 95% CI 0.32-0.62).⁴⁸² Detailed data for edoxaban have not yet been 1339
- 1340 published.³²¹ If a patient suffers a stroke or TIA whilst taking an anticoagulant, switching to another
- 1341 anticoagulant should be considered.
- 1342



1352 No prospective studies have investigated the benefit or risk of the initiation of OAC after intracranial haemorrhage,⁴⁸³ and patients with a history of intracranial bleeding were excluded from the randomized trials 1353 1354 comparing NOACs with VKAs. The available evidence indicates that anticoagulation in patients with AF can be 1355 reinitiated after 4-8 weeks, especially when the cause of bleeding or the relevant risk factor (e.g. uncontrolled hypertension) has been treated, and that such treatment leads to fewer recurrent (ischaemic) strokes and lower mortality.^{460, 484} If anticoagulation is resumed, it seems reasonable to consider anticoagulants with a low 1356 1357 bleeding risk.³⁹ Figure 10 depicts a consensus opinion on the initiation or resumption of OAC after an 1358 1359 intracranial haemorrhage. We recommend a multidisciplinary decision with input from stroke 1360

physicians/neurologists, cardiologists, neuroradiologists, and neurosurgeons.



1366 OAC = oral anticoagulation; PCI = percutaneous coronary intervention; VKA = vitamin K antagonist.

1367

1368 Recommendations for secondary stroke prevention

Recommendations	Class ^a	Level ^b	Refs ^c
Anticoagulation with heparin or low-molecular-weight heparin immediately after ischaemic stroke is not recommended in AF patients	III (harm)	A	477
In patients who suffer a transient ischemic attack or stroke while on anticoagulation, adherence to therapy should be assessed and optimized	lla	С	
In patients who suffer a moderate-to-severe ischaemic stroke while on anticoagulation, anticoagulation should be interrupted for 3–12 days based on a multidisciplinary assessment of acute stroke and bleeding risk	lla	С	
In AF patients who suffer a stroke, aspirin should be considered for prevention of secondary stroke until the initiation or resumption of oral anticoagulation.	lla	В	485
Systemic thrombolysis with a recombinant tissue plasminogen activator is not recommended if the INR is above 1.7 (or, for patients on dabigatran, if activated partial thromboplastin time is outside the normal range)	III (harm)	С	472, 474
NOACs are recommended in preference to VKAs or aspirin in AF patients with a previous stroke	I	В	363, 482
After TIA or stroke, combination therapy of OAC and an	III (harm)	В	486

antiplatelet is not recommended			
After intracranial haemorrhage, oral anticoagulation in patients with AF may be reinitiated after 4–8 weeks provided the cause of	llb	В	483, 484, 487
bleeding or the relevant risk factor has been treated or controlled			

1369

- 1370
- 1371 AF = atrial fibrillation; INR = international normalized ratio; NOAC = non-vitamin K antagonist oral
- 1372 anticoagulant; OAC = oral anticoagulation; TIA = transient ischaemic attack; VKA = vitamin K antagonist.
- 1373 ^aClass of recommendation.
- 1374 ^bLevel of evidence.
- 1375 ^cReference(s) supporting recommendations.
- 1376

1377 **9.5**. Strategies to minimize bleeding on anticoagulant therapy

1378 In a meta-analysis of 47 studies, the overall incidence of major bleeding with VKAs was 2.1 (range 0.9–3.4) per 1379 100 patient-years in controlled trials and 2.0 (range 0.2-7.6) per 100 patient-years for observational data sets.

1380 Minimizing treatable bleeding risk factors (see Table 12) seems paramount to reduce the bleeding rate on 1381 anticoagulants.

1382

1383 9.5.1. **Uncontrolled hypertension**

Uncontrolled blood pressure increases the risk of bleeding on OAC.⁵³ Hence, keeping systolic blood pressure 1384

well controlled is of particular relevance in anticoagulated patients with AF. Treatment according to current 1385 guidelines is recommended in patients with known hypertension.489

1386 1387

1388 9.5.2. **Previous bleeding event**

1389 History of bleeding events and the presence of anaemia are important parts of the assessment of all patients

1390

receiving OAC. The majority of bleeding events are gastrointestinal. Compared with warfarin, the risk of gastrointestinal bleeds was increased for dabigatran 150 mg twice daily,^{396, 490} rivaroxaban 20 mg once daily,⁴⁹¹ and edoxaban 60 mg once daily.³²¹ The risk of gastrointestinal bleeds was comparable to warfarin on dabigatran 110 mg twice daily⁴⁹⁰ and on apixaban 5 mg twice daily³¹⁹ Recent observational analyses do not replicate these findings, suggesting a smaller effect.^{396, 492, 493} In patients in whom the source of bleeding has been identified and 1391

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1393 1394

1395 corrected, OAC can be reinitiated. This also appears true for patients who have had an intracranial haemorrhage,

1396 once modifiable bleeding risk factors (e.g. uncontrolled hypertension) have been corrected.^{460, 484}

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1398 9.5.3. Labile international normalized ratio and adequate non-vitamin K

1399 antagonist oral anticoagulant dosing

1400 TTR on VKA therapy is an important predictor of major haemorrhage.^{432, 441, 494} Therefore we recommend

targeting the INR between 2.0 and 3.0 in patients on VKAs, maintaining a high TTR (e.g. $\geq 70\%^{494}$), and to 1401

1402 consider switching to a NOAC when a high TTR cannot be sustained.⁴⁴⁴ NOAC dosing should follow the dose-

1403 reduction criteria evaluated in the clinical trials, considering renal function, age, and weight. Patient information

1404 and empowerment, best delivered through integrated AF management, seem paramount to achieve this goal.

1405

1406 **9.5.4**. Alcohol abuse

Alcohol excess is a risk factor for bleeding in anticoagulated patients,³⁸⁴ mediated by poor adherence, liver 1407 1408 disease, variceal bleeding, and risk of major trauma. Severe alcohol abuse and binge drinking habits should be

1409 corrected in patients eligible for OAC.

1410 1411 **9.5.5**. Falls and dementia

1412 Falls and dementia are associated with increased mortality in AF patients,⁴⁹⁵ without evidence that these 1413 conditions markedly increase the risk of intracranial haemorrhage.^{495,496} Hence, anticoagulation should only be

1414 withheld from patients with severe uncontrolled falls (e.g. epilepsy or advanced multisystem atrophy with

1415 backwards falls), or in selected patients with dementia where compliance and adherence cannot be ensured by a 1416 caregiver.

1417

1418 **9.5.6**. **Genetic testing**

1419 In addition to food and drug interactions, multiple genetic variations affect the metabolism of VKAs.⁴⁹⁷ The 1420 systematic use of genetic information for adjustment of VKA dosage has been evaluated in several controlled

1421 clinical studies.⁴⁹⁸⁻⁵⁰⁰ Genetic testing has little effect on TTR or bleeding risk on warfarin, and is not

1422 recommended for clinical use at present.⁵⁰¹ 1423

1424 **9.5.7.** Bridging periods off oral anticoagulation

1425 Most cardiovascular interventions (e.g. percutaneous coronary intervention or pacemaker implantation) can be

1426 performed safely on continued OAC. When interruption of OAC is required, bridging does not seem to be 1427 beneficial, except in patients with mechanical heart valves. In a randomized trial of 1884 patients with AF,

1428 interruption of anticoagulation was non-inferior to heparin administration for the outcome of arterial

1429 thromboembolism (incidence of 0.4% and 0.3%, respectively) and resulted in a lower risk of major bleeding

1430 (1.3% and 3.2%, respectively).⁵⁰² A short interruption or continued OAC should be considered in patients at

- 1431 highest risk of stroke.
- 1432

1433 9.6. Management of bleeding events in anticoagulated patients with atrial fibrillation 1434 9.6.1. Management of minor, moderate, and severe bleeding

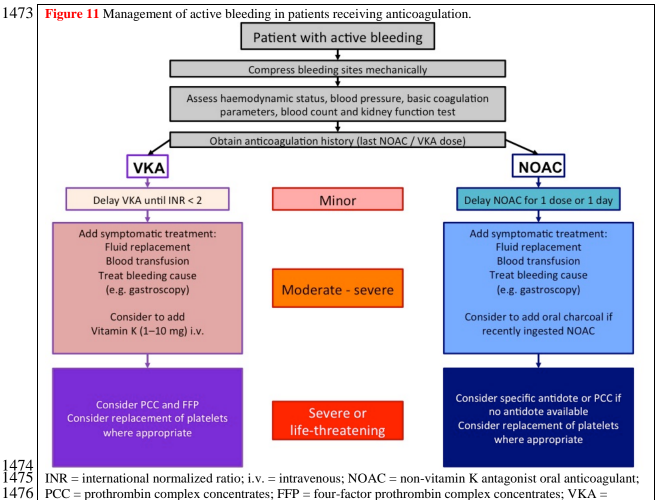
1435 General assessment of an anticoagulated patient with AF experiencing a bleeding event should include 1436 assessment of bleeding site, onset, and severity of the bleeding, the time-point of last intake of OAC and other 1437 antithrombotic drugs, and other factors influencing bleeding risk such as CKD, alcohol abuse, and concurrent 1438 medications. Laboratory tests should include haemoglobin, haematocrit, platelet count, renal function, and for 1439 VKA patients, prothrombin time, activated partial thromboplastin time, and INR. Coagulation tests do not 1440 provide much information in patients on NOACs, except for activated partial thromboplastin time in the case of 1441 dabigatran. More specific coagulation tests do exist, including diluted thrombin time (HEMOCLOT) for dabigatran and calibrated quantitative anti-factor Xa assays for factor Xa inhibitors.⁵ not always readily available and are often unnecessary for bleeding management.⁵⁰⁴ ⁰³ However, these tests are 1442 1443 1444 We propose a simple scheme to manage bleeding events in patients on OAC (Figure 11). Minor 1445 bleeding events should be treated with supportive measures such as mechanical compression or minor surgery to 1446 achieve haemostasis. In patients receiving VKAs, the next dose of VKA can be postponed. NOACs have a short 1447 plasma half-life of approximately 12 hours and improved haemostasis is expected within 12-24 hours after a 1448 delayed or omitted dose. Treatment of moderate bleeding events may require blood transfusions and fluid 1449 replacement. Specific diagnostic and treatment interventions directed against the cause of the bleeding (e.g. 1450 gastroscopy) should be performed promptly. If the intake of NOAC was recent (< 2-4 h), charcoal 1451 administration and/or gastric lavage will reduce further exposure. Dialysis clears dabigatran but is not effective 1452 for the other NOACs. 1453 Immediate reversal of the antithrombotic effect is indicated in severe or life-threatening bleeding 1454 events. An agreed, the institutional procedure for the management of life-threatening bleeds should be 1455 documented and accessible at all times to ensure adequate initial management. For VKAs, administration of 1456 fresh frozen plasma restores coagulation more rapidly than vitamin K, and prothrombin complex concentrates achieve even faster blood coagulation.⁵⁰⁵ Registry data suggest that the combination of plasma and prothrombin 1457 1458 complex concentrates is associated with the lowest case fatality following intracranial haemorrhage on VKA treatment with an INR ≥ 1.3 .⁵⁰⁶ In a multicentre randomized trial of 188 patients, four-factor prothrombin 1459 1460 complex concentrates achieved more rapid INR reversal and effective haemostasis than plasma in patients undergoing urgent surgical or invasive procedures.⁵⁰⁷ Administration of prothrombin complex concentrates may 1461 1462 also be considered for severe bleeding on NOAC treatment if specific antidotes are not available. 1463 Several antidotes to NOACs are under development. Idarucizumab (approved in 2015 by the US Food 1464 and Drug Administration and the European Medicines Agency) is a clinically available humanized antibody

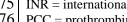
fragment that binds dabigatran and rapidly and dose-dependently reverses the effects without over-correction or thrombin generation.⁴⁷⁵ Andexanet alpha, a modified recombinant human factor Xa that lacks enzymatic activity, reverses the anticoagulant activity of apixaban and rivaroxaban in healthy probands within minutes after administration and for the duration of infusion, with a transient increase in markers of coagulation activity of uncertain clinical relevance.⁵⁰⁸ Another agent under development is ciraparantag (PER977), an antidote

1470 targeted to reverse both direct thrombin and factor Xa inhibitors as well as the indirect inhibitor enoxaparin.⁵⁰⁹

1471 The clinical usefulness of these specific antidotes needs further evaluation.

1472





vitamin K antagonist.

1477 1478

1479 **9.6.2**. Oral anticoagulation in atrial fibrillation patients at risk of or having a

1480 bleeding event

1481 While anticoagulation therapy should be paused to control active bleeding, absolute contraindications to long-1482 term OAC after a bleeding episode are rare. When nuisance bleeds are the reason to stop OAC, a change from 1483 one anticoagulant to another seems reasonable. Many causes or triggers of major bleeding events can be treated and/or eliminated, including uncontrolled hypertension, gastrointestinal ulcers, and intracranial aneurysms. Reinitiation of anticoagulation after a bleeding event is often clinically justified.^{460, 510} Difficult decisions, 1484 1485 1486 including the discontinuation and recommencement of OAC, should be taken by a multidisciplinary team, 1487 balancing estimated risk of recurrent stroke and bleeding, and considering the bleeding risk of different stroke 1488 prevention therapies. LAA exclusion or occlusion might be an alternative in selected patients.

1489

1490 **Recommendations for management of bleeding**

Recommendations	Class ^a	Level ^b	Refs ^c
Blood pressure control in anticoagulated patients with hypertension should be considered to reduce the risk of bleeding	lla	В	511
When dabigatran is used, a reduced dose of dabigatran (110 mg twice daily) may be considered in patients > 75 years to reduce the risk of bleeding	llb	В	490
In patients at high risk of gastrointestinal bleeding, a VKA or another NOAC should be preferred over dabigatran 150 mg twice daily, rivaroxaban 20 mg once daily, or edoxaban 60 mg once daily	lla	В	321, 396, 402, 405, 490, 492, 493, 512

Advice and treatment to avoid alcohol excess should be considered in all AF patients considered for OAC	lla	С	
Genetic testing before the initiation of VKA therapy is not recommended.	III (no benefit)	В	497
Reinitiation of OAC after a bleeding event should be considered in all eligible patients by a multidisciplinary AF team, considering different anticoagulants and stroke-prevention interventions, improved management of factors that contributed to bleeding, and stroke risk	lla	В	460
In AF patients with severe active bleeding events, it is recommended to interrupt OAC therapy until the underlying cause is resolved	I	С	

1491 AF = atrial fibrillation; NOAC = non-vitamin K antagonist oral anticoagulant; OAC = oral anticoagulation;

1492 VKA = vitamin K antagonist

1493 ^aClass of recommendation.

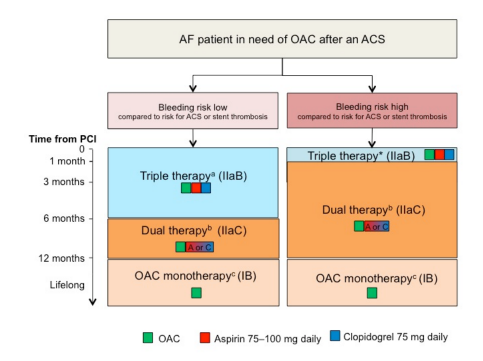
¹⁴⁹⁴ ^bLevel of evidence.

1495 ^cReference(s) supporting recommendations.

1496

1497 9.7. Combination therapy with oral anticoagulants and antiplatelets

1498 Approximately 15% of AF patients in contemporary trials⁵¹³ and registries⁵¹⁴⁻⁵¹⁶ have a history of myocardial infarction. Between 5% and 15% of AF patients will require stenting at some point in their lives. This scenario 1499 requires careful consideration of antithrombotic therapy, balancing bleeding risk, stroke risk, and risk of acute 1500 coronary syndromes (ACS).⁵¹⁶ Co-prescription of OAC with antiplatelet therapy, in particular triple therapy, increases the absolute risk of major haemorrhage.^{445, 517, 518} A recent meta-analysis involving 30,866 patients 1501 1502 with a recent ACS evaluated the effects of adding NOAC therapy to single (4135 patients) or dual (26,731 1503 patients) antiplatelet therapy.⁵¹⁹ The addition of a NOAC increased the bleeding risk by 79–134%, while 1504 1505 reducing recurrent ischaemic events only marginally in patients without AF. OAC monotherapy, and not 1506 combination therapy with antiplatelets, is recommended in AF patients with stable CAD but without an ACS 1507 and/or coronary intervention in the previous 12 months. In patients treated for ACS and in those receiving a 1508 coronary stent, short-term triple combination therapy of OAC, clopidogrel, and aspirin seems warranted (Figure 1509 12).



1510

1511 Figure 12 Antithrombotic therapy after an ACS in AF patients requiring anticoagulation. 1512 ACS = acute coronary syndrome; AF = atrial fibrillation; OAC = oral anticoagulation (using vitamin K)1513 antagonists or non-vitamin K antagonist oral anticoagulants); PCI = percutaneous coronary intervention. 1514 ^aDual therapy with OAC and aspirin or clopidogrel may be considered in selected patients, especially those not 1515 receiving a stent or patients at a longer time from the index event. 1516 ^bOAC plus single antiplatelet. 1517 ^cDual therapy with OAC and an antiplatelet agent (aspirin or clopidogrel) may be considered in patients at high 1518 risk of coronary events. 1519 1520 1521 **9.7.1**. Antithrombotic therapy after acute coronary syndromes and percutaneous 1522 coronary intervention in patients requiring oral anticoagulation 1523 The optimal combination antithrombotic therapy or duration of combination therapy for AF patients undergoing 1524 percutaneous coronary intervention is not known, but the continued bleeding risk suggests a short duration. 1525 Expert consensus,⁵²⁰ reviewed and reconsidered by this Task Force, suggests the following principles: AF 1526 patients at risk for stroke, patients with mechanical valves, and patients with recent or recurrent deep vein 1527 thrombosis or pulmonary embolism should continue OAC during and after stenting. In general, a short period of

1528 triple therapy (OAC, aspirin, clopidogrel) is recommended, followed by a period of dual therapy (OAC plus a 1529 single antiplatelet) (Figure 13). When a NOAC is used, the consensus recommendation is that the lowest dose

1530 effective for stroke prevention in AF should be considered. Dose reduction beyond the dosing regimens tested in 1531 the phase III trials is not currently recommended, and awaits assessment in ongoing controlled trials. The 1532 combination of aspirin, clopidogrel, and low-dose rivaroxaban (2.5 mg twice daily) is not recommended for

1533 stroke prevention in AF.⁵²¹

1534 The use of prasugrel or ticagrelor as part of triple therapy should be avoided unless there is a clear need for these agents (e.g. stent thrombosis on aspirin plus clopidogrel), given the lack of evidence and the greater risk of major bleeding compared with clopidogrel.^{522, 523} Ongoing trials will inform about such combination 1535 1536 1537 therapies in the future.

1538 The omission of aspirin while maintaining clopidogrel and OAC has been evaluated in the WOEST 1539 (What is the Optimal antiplatElet and anticoagulant therapy in patients with oral anticoagulation and coronary 1540 StenTing) trial, in which 573 anticoagulated patients undergoing percutaneous coronary intervention (70% with

1541 AF) were randomized to either dual therapy with OAC and clopidogrel (75 mg once daily) or to triple therapy

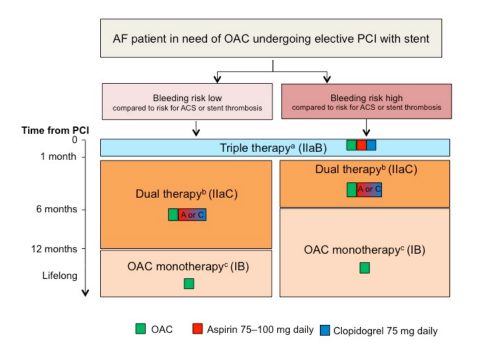
1542 with OAC, clopidogrel, and aspirin.⁵²⁴ Bleeding was lower in the dual versus triple therapy arm, driven by fewer

1543 minor bleeding events. The rates of myocardial infarction, stroke, target vessel revascularization, and stent

thrombosis did not differ (albeit with low event numbers), but all-cause mortality was lower in the dual therapy group at 1 year (2.5% vs. triple 6.4%). Although the trial was too small to assess ischaemic outcomes, dual

1546 therapy with OAC and clopidogrel may emerge in the future as an alternative to triple therapy in patients with

1547 AF and ACS and/or coronary intervention.⁵²⁵



1548

15-0	
1549	Figure 13 Antithrombotic therapy after percutaneous intervention in AF patients requiring anticoagulation.
1550	ACS = acute coronary syndrome; AF = atrial fibrillation; OAC = oral anticoagulation (using vitamin K
1551	antagonists or non-vitamin K antagonist oral anticoagulants); PCI = percutaneous coronary intervention.
1552	^a Dual therapy with OAC and aspirin or clopidogrel may be considered in selected patients.
1553	^b OAC plus single antiplatelet.
1554	^c Dual therapy with OAC and an antiplatelet agent (aspirin or clopidogrel) may be considered in patients at high
1555	risk of coronary events.
1556	
1557	Recommendations for combination therapy with oral anticoagulants and antiplatelets

1558

Recommendations	Class ^a	Level ^b	Refs ^c
After elective coronary stenting for stable coronary artery disease in AF patients at risk of stroke, combination triple therapy with aspirin, clopidogrel and an oral anticoagulant should be considered for 1 month to prevent recurrent coronary and cerebral ischaemic events	Па	В	522, 524
After an ACS with stent implantation in AF patients at risk of stroke, combination triple therapy with aspirin, clopidogrel, and an oral anticoagulant should be considered for 1–6 months to prevent recurrent coronary and cerebral ischaemic events	Па	С	520

After an ACS without stent implantation in AF patients at risk of stroke, dual therapy with an oral anticoagulant and aspirin or clopidogrel should be considered for up to 12 months to prevent recurrent coronary and cerebral ischaemic events	IIa	С	
The duration of combination antithrombotic therapy, especially triple therapy, should be kept to a limited period, balancing the estimated risk of recurrent coronary events and bleeding	IIa	В	520
Dual therapy with any oral anticoagulant plus clopidogrel 75 mg/day may be considered as an alternative to initial triple therapy with aspirin in selected patients.	IIb	С	524, 525

1559 ACS = acute coronary syndromes; AF = atrial fibrillation

- 1560 ^aClass of recommendation.
- 1561 ^bLevel of evidence.

1562 ^cReference(s) supporting recommendations.

1563

10 Rate control therapy in AF 1564

1565 Rate control is an integral part of the management of AF patients, and is often sufficient to improve AF-related

1566 symptoms. Compared with stroke prevention and rhythm control, very little robust evidence exists to inform the

1567 best type and intensity of rate control treatment, with the majority of data derived from short-term crossover trials and observational studies.^{41, 526-528} Pharmacological rate control can be achieved for acute or long-term rate

- 1568 1569
- control with beta-blockers, digoxin, the calcium channel blockers diltiazem and verapamil, or combination 1570
- therapy (Table 15). A number of antiarrhythmic drugs also have rate-limiting properties (amiodarone,
- 1571 dronedarone, sotalol, and to some extent propafenone), but they should only be used in patients needing rhythm 1572 control therapy (see Chapter 10).
- 1573

1574 10.1. Acute rate control

1575 In the setting of acute new-onset AF, patients are often in need of heart rate control. Physicians should evaluate

1576 underlying causes of elevated heart rate, such as infection, endocrine imbalance, anaemia, and pulmonary

1577 embolism. For acute rate control, beta-blockers and diltiazem/verapamil are preferred over digoxin because of

their rapid onset of action and effectiveness at high sympathetic tone.⁵²⁸⁻⁵³² The choice of drug (*Table 15*) and 1578 1579

target heart rate will depend on patient characteristics, symptoms, LVEF and haemodynamics, but a lenient 1580

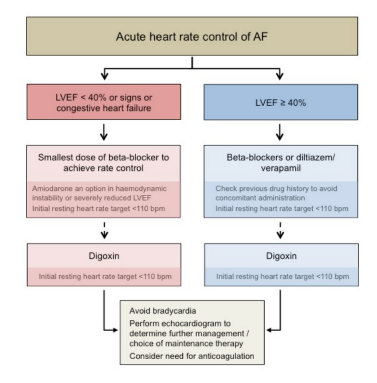
initial approach to heart rate seems acceptable. Combination therapy may be required (Figure 14). In patients with evidence of HFrEF, beta-blockers, digitalis (digoxin or digitoxin), or their combination should be used,²¹⁸,

1581 ⁵³³ as diltiazem and verapamil can have negative inotropic effects in patients with LVEF < 40%.^{222, 534, 535} In 1582

1583 critically ill patients and those with severely impaired LV systolic function, intravenous amiodarone can be used

where excess heart rate is leading to haemodynamic instability.⁵³⁶⁻⁵³⁸ Urgent cardioversion should be considered 1584

1585 in unstable patients (see Chapter 10.2).



1586

1590

1587 Figure 14 Acute heart rate control of AF.

1588 See *Table 15* for medication dosage. Digitoxin is a suitable alternative to digoxin, where available.

1589 AF = atrial fibrillation; bpm = beats per minute; LVEF = left ventricular ejection fraction.

1591 **10.2**. Long-term pharmacological rate control

1592 10.2.1. **Beta-blockers**

1593 Beta-adrenoreceptor blocker monotherapy is often the first-line rate-controlling agent, ⁵³⁹ largely based on 1594 observations of better acute heart rate control than digoxin. Interestingly, the prognostic benefit of beta-blockers 1595 seen in HFrEF patients with sinus rhythm is lost in those with AF. In an individual patient-level meta-analysis 1596 of RCTs, beta-blockers did not reduce all-cause mortality compared to placebo in those with AF at baseline (HR 1597 0.97; 95% CI 0.83–1.14; P = 0.73), whereas there was a clear benefit in patients with sinus rhythm (HR 0.73; 1598 95% CI 0.67–0.80; P < 0.001).²³ The study, which included 3066 participants with HFrEF and AF, showed 1599 consistency across all subgroups and outcomes, with no heterogeneity between the 10 RCTs included ($I^2 = 0\%$). 1600 Despite this lack of prognostic benefit in HFrEF, this Task Force still considers beta-blockers as a useful first-1601 line rate control agent across all AF patients, based on the potential for symptomatic and cardiac function improvement as a result of rate control, the lack of harm from published studies, and the good tolerability profile 1602 across all ages in sinus rhythm and in AF.^{23, 540} 1603

1604

Non-dihydropyridine calcium channel blockers 1605 10.2.2.

Verapamil or diltiazem provides reasonable rate control in AF patients.⁵⁴¹ They should be avoided in patients with HFrEF because of their negative inotropic effects.^{222, 534, 535} Verapamil or diltiazem can improve 1606 1607 arrhythmia-related symptoms, ⁵²⁶ in comparison with beta-blockers, which reduced exercise capacity and 1608 1609 increased B-type natriuretic peptide in one small trial of low-risk patients with preserved LVEF.⁵⁴²

1610 1611 10.2.3. Digitalis

Cardiac glycosides such as digoxin and digitoxin have been in use for over two centuries, although prescriptions have been declining steadily over the past 15 years.⁵⁴³ In the randomized Digitalis Investigation Group (DIG) 1612

- 1613
- 1614 trial, digoxin had no effect on mortality compared to placebo in HFrEF patients in sinus rhythm (RR 0.99; 95%

- 1615 CI 0.91–1.07), but reduced hospital admissions (RR 0.72; 95% CI 0.66–0.79).^{544, 545} There have been no head-
- to-head RCTs of digoxin in AF patients.⁵⁴⁶ Observational studies have associated digoxin use with excess mortality in AF patients,⁵⁴⁷⁻⁵⁴⁹ but this association is likely due to selection and prescription biases rather than harm caused by digoxin,⁵⁵⁰⁻⁵⁵³ particularly as digoxin is commonly prescribed to sicker patients.²²⁵ In a 1616 1617
- 1618
- 1619 crossover mechanistic trial of 47 patients with HFrEF and AF, there were no differences in heart rate, blood
- 1620 pressure, walking distance, or LVEF between carvedilol and digoxin, although beta-blockers did result in higher
- 1621 B-type natriuretic peptide levels, combination carvedilol/digoxin improved LVEF, and digoxin withdrawal
- 1622 reduced LVEF.⁵⁵⁴ Comparisons with other rate control therapies are based on small, short-duration studies that
- 1623 identify no or marginal differences in exercise capacity, quality of life, or LVEF compared to digoxin. 526, 554-550
- 1624 Lower doses of digoxin ($\leq 250 \mu g$ once daily), corresponding to serum digoxin levels of 0.5–0.9 ng/mL, may be
- 1625 associated with better prognosis.
- 1626

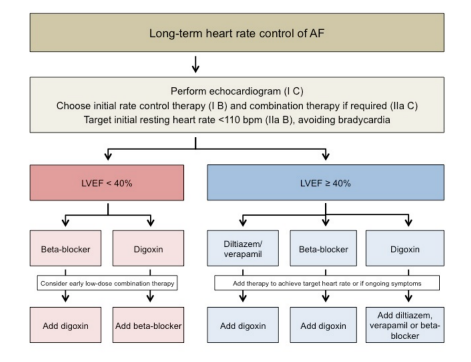
1627 10.2.4. Amiodarone

1628 Amiodarone can be useful for rate control as a last resort. The wide array of extracardiac adverse effects

1629 associated with amiodarone renders it a reserve agent in patients whose heart rate cannot be controlled with

1630 combination therapy (e.g. beta-blocker or verapamil/diltiazem combined with digoxin).

- 1631
- 1632 In summary, there is equipoise for the use of different rate control agents in AF. The choice of beta-blocker.
- 1633 diltiazem/verapamil, digoxin, or combination therapy should be made on an individual basis, after consideration
- 1634 of patient characteristics and patient preference. All available therapies have the potential for adverse effects and 1635
- patients should initially be treated with a low dose and uptitrated to achieve symptom improvement. In practice, 1636 achieving a heart rate < 110 bpm will often require combination therapy (*Figure 15*). The benefit of different
- rate control strategies on symptoms, quality of life, and other intermediate outcomes is under investigation.⁵⁵⁹ 1637



1638

- 1639 Figure 15 Long-term heart rate control of AF.
- 1640 See *Table 15* for medication dosage. Digitoxin is a suitable alternative to digoxin, where available.
- AF = atrial fibrillation; bpm = beats per minute; LVEF = left ventricular ejection fraction. 1641
- 1642

Heart rate targets in atrial fibrillation 1643 **10.3**.

1644 The optimal heart rate target in AF patients is unclear. The RACE (Rate Control Efficacy in Permanent Atrial

- 1645 Fibrillation) II study randomized 614 patients with permanent AF to either a target heart rate < 80 bpm at rest 1646 and < 110 bpm during moderate exercise, or to a lenient heart rate target of < 110 bpm. There was no difference
- in a composite of clinical events (14.9% in the strict rate control group, 12.9% in the lenient group),⁵⁶⁰ NYHA 1647
- class, or hospitalizations.^{560, 561} Similar results were found in a pooled analysis of the AFFIRM (Atrial 1648
- Fibrillation Follow-up Investigation of Rhythm Management) and RACE trials (1091 participants), albeit with 1649
- 1650 smaller heart rate differences and without randomization.⁵⁶² It is worthwhile to note that many 'adequately rate-
- 1651 controlled' patients (resting heart rate 60-100 bpm) are severely symptomatic, calling for additional
- 1652 management.¹⁹⁴ Nonetheless, lenient rate control is an acceptable initial approach, regardless of heart failure
- 1653 status, unless symptoms call for stricter rate control.
- 1654

Atrioventricular node ablation and pacing 1655 10.4.

1656 Ablation of the atrioventricular node/His bundle and implantation of a VVI pacemaker can control ventricular rate when medications fail to control rate and symptoms. It is a relatively simple procedure with a low complication rate and low long-term mortality risk,^{563,564} especially when the pacemaker is implanted a few weeks before the AV nodal ablation and the initial pacing rate after ablation is set at 70–90 bpm.^{565,566} The procedure does not worsen LV function⁵⁶⁷ and may even improve LVEF in selected patients.⁵⁶⁸⁻⁵⁷⁰ In some 1657 1658 1659 1660 1661 patients in heart failure treated with biventricular pacing (cardiac resynchronization therapy), AF can

1662 terminate,⁵⁷¹ although such a 'rhythm control' effect of cardiac resynchronization therapy is likely to be small

1663 and clearly needs confirmation.⁵⁷² AV nodal ablation renders patients pacemaker-dependent for the rest of their

- 1664 lives, limiting AV nodal ablation and pacing to patients whose symptoms cannot be managed by rate controlling
- 1665 medication or by reasonable rhythm control interventions. The choice of pacing therapy (right ventricular or
- 1666 biventricular pacing with or without an implantable defibrillator) will depend on individual patient characteristics, including LVEF.573,574
- 1667 1668
- 1669 Recommendations for rate control

Recommendations	Class ^a	Level ^b	Refs ^c
Beta-blocker, digoxin, diltiazem, or verapamil are recommended to control heart rate in AF patients with LVEF \ge 40%	I	В	225, 526, 528, 531, 532, 541, 555, 575
Beta-blocker and/or digoxin are recommended to control heart rate in AF patients with LVEF < 40%	I	В	23, 225, 526, 533, 554, 575, 576
Combination therapy comprising different rate controlling agents should be considered if a single agent does not achieve the necessary heart rate target	lla	С	23, 554, 577
In cases of haemodynamic instability or severe depression in LVEF, amiodarone may be considered for acute control of heart rate	llb	В	536-538
In patients with permanent AF (i.e. where no attempt to restore sinus rhythm is planned), antiarrhythmic drugs should not routinely be used for rate control	III (harm)	A	41, 578, 579
A resting heart rate of < 110 bpm (i.e. lenient rate control) should be considered as the initial heart rate target for rate control therapy	lla	В	560
Rhythm rather than rate control strategies should be considered as the preferred management in pre-excited AF and AF during pregnancy	lla	С	
Atrioventricular node ablation should be considered to control heart rate in patients unresponsive or intolerant to intensive rate and rhythm control therapy, accepting that these patients will become pacemaker dependent	lla	В	184, 564, 569

- 1670 AF = atrial fibrillation; bpm = beats per minute; LVEF = left ventricular ejection fraction.
- 1671 Digitoxin is a suitable alternative to digoxin, where available. In patients with heart failure with reduced ejection
- 1672 fraction (LVEF < 40%), recommended beta-blockers are bisoprolol, carvedilol, long-acting metoprolol, and
- 1673 nebivolol.
- 1674 ^aClass of recommendation.

^bLevel of evidence.
^c Reference(s) supporting recommendations.
Table 15 Rate control therapy in AF

		ontrol therapy in A		Side offect profile	Commonto
Thera		Acute intravenous rate control	Long-term oral rate control	Side-effect profile	Comments
Beta-	blockers	а			
Bisop	orolol	Not available	1.25–20 mg once daily or split	Most common reported adverse	Bronchospasm is rare – in cases of
Carve		Not available	3.125–50 mg twice daily	symptoms are lethargy, headache,	asthma, recommend beta-1 selective
	prolol	2.5–10 mg intravenous bolus (repeated as required)	100–200 mg total daily dose (according to preparation)	peripheral oedema, upper respiratory tract symptoms, gastrointestinal upset,	agents (avoid carvedilol). Contraindicated in acute cardiac failure
Nebiv		N/A	2.5–10 mg once daily or split	and dizziness. Adverse effects	and a history of severe
Esmo	lol	0.5 mg intravenous bolus over 1 min; then 0.05– 0.25 mcg/kg/min		include bradycardia, atrioventricular block, and hypotension	bronchospasm
Calciu	um-chan	nel blockers			
Diltia	zem	15–25 mg intravenous bolus (repeated as required)	60 mg three times daily up to 360 mg total daily dose (120–360 mg once daily modified release)	Most common reported adverse symptoms are dizziness, malaise, lethargy, headache, hot flushes,	Use with caution in combination with beta-blockers. Reduce dose with hepatic impairment and start with smaller
Veraț	pamil	2.5–10 mg intravenous bolus (repeated as required)	40–120 mg three times daily (120– 480 mg once daily modified release)	gastrointestinal upset, and oedema. Adverse effects include bradycardia, atrioventricular block, and hypotension (prolonged hypotension possible with verapamil)	dose in renal impairment. Contraindicated in LV failure with pulmonary congestion or LVEF < 40%
Cardi	ac glyco	sides			
Digo>		0.5 mg intravenous bolus (0.75–1.5 mg over 24 h in divided doses)	0.0625–0.25 mg daily dose	Most common reported adverse symptoms are gastrointestinal upset, dizziness, blurred	High plasma levels associated with increased risk of death. Check renal function before
Digito	oxin ific indica	0.4–0.6 mg intravenous bolus	0.05–0.3 mg daily dose	vision, headache, and rash. In toxic states (serum levels > 2 ng/mL), digoxin is proarrhythmic and can aggravate heart failure, particularly with coexistent hypokalaemia	starting and adapt dose in patients with CKD. Contraindicated in accessory conducting pathways, ventricular tachycardia, and hypertrophic cardiomyopathy with outflow tract obstruction
	darone	300 mg	200 mg daily	Hypotension,	Suggested as

intravenously diluted in 250 mL 5% dextrose over 30–60 min (preferably via central venous cannula) ^b	bradycardia, nausea, QT prolongation, pulmonary toxicity, skin discolouration, thyroid dysfunction, corneal deposits, and cutaneous reaction with extravasation	adjunctive therapy in patients where heart rate control cannot be achieved using combination therapy
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- 1679 AF = atrial fibrillation; CKD = chronic kidney disease; LV = left ventricular; LVEF = left ventricular ejection 1680 fraction.
- 1681 ^aA number of other beta-blockers are also available, but are not recommended as specific rate control therapy in
- 1682 AF. These include atenolol (25–100 mg once daily with a short biological half-life), propranolol (non-selective,
- 1683 1 mg over 1 min and repeat up to 3 mg at 2-min intervals [acute] or 10–40 mg three times daily [long-term]), or
- 1684 labetalol (non-selective, 1–2 mg/min [acute]).
- 1685 ^bIf ongoing requirement for amiodarone, follow with 900 mg intravenous over 24 hours diluted in 500–1000 mL 1686 via a central venous cannula.
- 1687

1688 **11 Rhythm control therapy in atrial fibrillation**

- 1689
- 1690
- 1691
- Restoring and maintaining sinus rhythm is an integral part of AF management. Antiarrhythmic drugs approximately double the rate of sinus rhythm compared with placebo.⁵⁸⁰⁻⁵⁸⁴ Catheter ablation or combination therapy is often effective when antiarrhythmic drugs fail.^{226, 585-587} Although many clinicians believe that maintaining sinus rhythm can improve outcomes in AF patients,⁵⁸⁸ all trials that have compared rhythm control 1692
- 1693 to rate control (with appropriate anticoagulation) therapy have resulted in neutral outcomes.^{41,5}
- 1694 Whether modern rhythm control management involving catheter ablation, combination therapy, and early
- 1695 therapy leads to a reduction in major cardiovascular events (e.g. stroke and cardiovascular death) is currently
- 1696 under investigation (e.g. in the EAST [Early treatment of Atrial fibrillation for Stroke prevention Trial] -AFNET 4⁴⁰ and CABANA [Catheter Ablation vs Anti-arrhythmic Drug Therapy for Atrial Fibrillation Trial]⁵⁹⁴ 1697
- 1698 trials). For now, rhythm control therapy is indicated to improve symptoms in AF patients who remain
- 1699 symptomatic on adequate rate control therapy.
- 1700

1701 **11.1**. Acute restoration of sinus rhythm

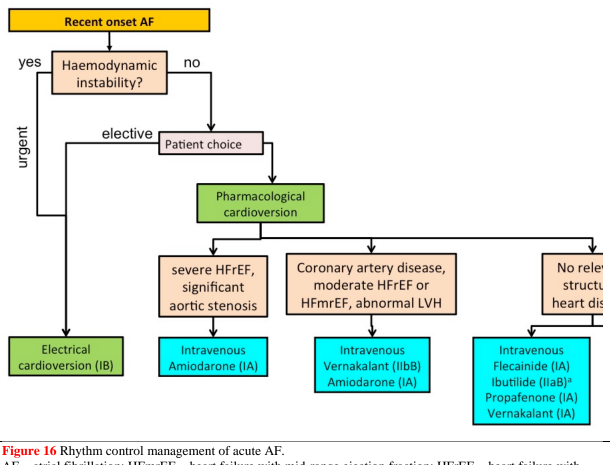
1702 11.1.1. Antiarrhythmic drugs for acute restoration of sinus rhythm

('pharmacological cardioversion') 1703

1704 Antiarrhythmic drug can restore sinus rhythm in patients with AF (pharmacological cardioversion) as

shown in small controlled trials, meta-analyses,^{41, 584, 595, 596} and in a few larger controlled trials.⁵⁹⁷⁻⁶⁰ 1705

- Outside of Europe, dofetilide is available and can convert recent-onset AF.⁶⁰⁶ Pharmacological cardioversion 1706
- restores sinus rhythm in approximately 50% of patients with recent-onset AF (Table 16).607-609 In the short term, 1707
- electrical cardioversion restores sinus rhythm quicker and more effectively than pharmacological cardioversion and is associated with shorter hospitalization.⁶⁰⁹⁻⁶¹³ Pharmacological cardioversion, conversely, does not require 1708 1709
- 1710 sedation or fasting (Figure 16).
- Flecainide and propafenone are effective for pharmacological cardioversion,^{595, 602-605, 614, 615} but their 1711 use is restricted largely to patients without structural heart disease. Ibutilide is an alternative where available, but carries a risk of torsades de pointes.⁶¹⁵ Vernakalant⁶⁰²⁻⁶⁰⁵ can be given to patients with mild heart failure 1712 1713
- 1714
- (NYHA Class I or II), including those with ischaemic heart disease, provided they do not present with hypotension or severe aortic stenosis.⁶¹⁶⁻⁶¹⁸ Amiodarone can be used in patients with heart failure and in patients 1715
- 1716 with ischaemic heart disease (although patients with severe heart failure were excluded in most of the AF
- cardioversion trials).⁵⁹⁶ Amiodarone also slows heart rate by 10-12 bpm after 8-12 hours when given 1717
- intravenously.⁵⁹⁶ Both amiodarone and flecainide appear more effective than sotalol in restoring sinus rhythm.^{600, 601, 619} 1718
- 1719



AF = atrial fibrillation; HFmrEF = heart failure with mid-range ejection fraction; HFrEF = heart failure with reduced ejection fraction.

1724 ^aIbutilide should not be used in patients with long QT interval.

1725

 $\begin{array}{c} 1720\\1721 \end{array}$

1726 **11.1.2.** 'Pill in the pocket' cardioversion performed by patients

1727 In selected patients with infrequent symptomatic episodes of paroxysmal AF, a single bolus of oral flecainide 1728 (200–300 mg) or propafenone (450–600 mg) can be self-administered by the patient at home ('pill in the pocket'

therapy) to restore sinus rhythm, after safety has been established in the hospital setting.⁶²⁰ This approach seems

marginally less effective than hospital-based cardioversion,⁶²¹ but is practical and provides control and

1731 reassurance to selected patients.

1732

1733 **Table 16** Antiarrhythmic drugs for pharmacological cardioversion

Drug	Route	First dose	Follow-up dose	Risks	Referenc es
Flecainide	Oral	200–300 mg	N/A	Avoid in patients with IHD and/or significant structural heart disease. Hypotension, atrial flutter with 1:1	595, 598
	IV	1.5–2 mg/kg over 10 min		conduction, QT prolongation	
Amiodarone	IV ^a	5–7 mg/kg over 1–2 h	50 mg/h to a maximum of 1.0 g over 24 h	Phlebitis, hypotension, bradycardia/AV block. Will slow ventricular rate. Delayed conversion to sinus rhythm (8–12 h)	596-601
Propafenone	IV	1.5–2 mg/kg over 10 min		Avoid in patients with IHD and/or significant structural heart disease. Hypotension, atrial flutter with 1:1	622-625

	Oral	450–600 mg		conduction, QRS prolongation (mild)	
Ibutilide ^b	IV	1 mg over 10 min	1 mg over 10 min after waiting for 10 min	Avoid in patients with QT prolongation, hypokalemia, severe LVH, or low ejection fraction. QT prolongation, polymorphic ventricular tachycardia/torsades de pointes (3–4% of patients). Will slow ventricular rate	614, 615
Vernakalant	IV	3 mg/kg over 10 min	2 mg/kg over 10 min after waiting for 15 min	Avoid in patients with systolic blood pressure < 100 mmHg, recent (< 30 days) ACS, NYHA Class III and IV heart failure, QT interval prolongation (uncorrected QT > 440 ms), and severe aortic stenosis. Hypotension, non-sustained ventricular arrhythmias, QT and QRS prolongation	602-605, 61

1734 ACS = acute coronary syndromes; IHD = ischaemic heart disease; IV = intravenous; LVH = left ventricular1735

hypertrophy; NYHA = New York Heart Association.

1736 ^aUse a large peripheral vessel and change to oral amiodarone within 24 h of IV (central line) administration. ^bIbutilide is only available in selected European countries.

1737 1738

1739 **Electrical cardioversion** 11.1.3.

1740 Synchronized direct current electrical cardioversion quickly and effectively converts AF to sinus rhythm and is

1741 the method of choice in severely haemodynamically compromised patients with new-onset AF (Figure 16).⁶²⁶⁻

1742 ⁶²⁸ Electrical cardioversion can be performed safely in sedated patients treated with intravenous midazolam

and/or propofol. Continuous monitoring of blood pressure and oximetry during the procedure is important.⁶²⁹ 1743 1744

Skin burns may occasionally be observed. Intravenous atropine or isoproterenol or temporary transcutaneous 1745

pacing should be available to mitigate post-cardioversion bradycardia. Biphasic defibrillators are more effective than monophasic waveforms, and have become industry standard.^{626, 628} Anterior–posterior electrode positions 1746

1747 generate a stronger shock field in the left atrium than anterolaterally positioned electrodes, and restore sinus

rhythm more effectively. 626, 627, 630 1748

Pretreatment with amiodarone (requiring a few weeks of therapy),^{631, 632} sotalol,⁶³¹ ibutilide,⁶³³ or 1749

vernakalant⁶³⁴ can improve efficacy of electrical cardioversion, and similar effects are likely for flecainide⁵⁸⁴ and propafenone.⁶³⁵ Beta-blockers,⁶³⁶ verapamil, diltiazem,⁶³⁷⁻⁶³⁹ and digoxin^{640, 641} do not reliably terminate AF or facilitate electrical cardioversion. When antiarrhythmic drug therapy is planned to maintain sinus rhythm 1750 1751

1752

1753 after cardioversion, it seems prudent to start therapy 1–3 days before cardioversion (amiodarone: a few weeks)

1754 to promote pharmacological conversion and to achieve effective drug levels.^{584, 601} 1755

1756 11.1.4. Anticoagulation in patients undergoing cardioversion

Cardioversion carries an inherent risk of stroke in non-anticoagulated patients,⁶⁴² which is reduced substantially 1757 by the administration of anticoagulation.⁶⁴³ Immediate initiation of anticoagulation is important in all patients 1758 scheduled for cardioversion.⁶⁴⁴⁻⁶⁴⁶ Patients who have been in AF for longer than 48 hours should start OAC at 1759

1760 least 3 weeks before cardioversion and continue it for 4 weeks afterwards (in patients without a need for long-

1761 term anticoagulation), and continue it indefinitely in patients at risk of stroke. This practice has never been

evaluated in controlled trials, but seemed safe in a large observational data set from Finland.⁶⁴⁷ When early 1762

1763 cardioversion is desired, TOE can exclude the majority of left atrial thrombi, allowing immediate

cardioversion.^{648, 649} Ongoing studies will inform about the safety and efficacy of newly initiated anticoagulation 1764 1765 using NOACs in patients scheduled for electrical cardioversion.

1766

Long-term antiarrhythmic drug therapy 1767 11.2.

The aim of antiarrhythmic drug therapy is improvement in AF-related symptoms.^{41, 580} Hence, the decision to 1768 1769 initiate long-term antiarrhythmic drug therapy needs to balance symptom burden, possible adverse drug

1770 reactions, and patient preferences. The principles of antiarrhythmic drug therapy outlined in the 2010 ESC AF guidelines³⁶⁹ are still relevant and should be observed: 1771

- 1772 Treatment is aimed at reducing AF-related symptoms; 1.
- 1773 Efficacy of antiarrhythmic drugs to maintain sinus rhythm is modest; 2.

1774 3. Clinically successful antiarrhythmic drug therapy may reduce rather than eliminate the recurrence of 1775 AF:

- 1776 4. If one antiarrhythmic drug 'fails', a clinically acceptable response may be achieved with another agent; 1777
 - Drug-induced proarrhythmia or extra-cardiac side-effects are frequent; 5.
 - Safety rather than efficacy considerations should primarily guide the choice of antiarrhythmic drug. 6.
- 1778 1779

1780 Antiarrhythmic drug therapy approximately doubles sinus rhythm maintenance compared with no therapy.⁵⁸⁰ There is no appreciable effect on mortality or cardiovascular complications, but rhythm control therapy can slightly increase the risk of hospitalizations (often for AF).^{41, 578, 579, 582, 589-593} To reduce the risk of side-1781 1782 1783 effects,^{201, 580} a shorter duration of antiarrhythmic drug therapy seems desirable. As an example, short-term 1784 treatment (4 weeks) with flecainide for 4 weeks after cardioversion of AF was well-tolerated and prevented most (80%) AF recurrences when compared with long-term treatment.⁵⁸⁴ Short-term antiarrhythmic drug 1785 therapy is also used to avoid early AF recurrences after catheter ablation⁶⁵⁰ and may be reasonable in patients 1786 1787 deemed at increased risk of antiarrhythmic drug side-effects or in those with a low perceived risk of recurrent 1788 AF. 1789 In addition to antiarrhythmic drug therapy and catheter ablation (see Section 10.3), management of

1790 concomitant cardiovascular conditions can reduce symptom burden in AF and facilitate maintenance of sinus rhvthm.^{203, 204, 296, 312} This includes weight reduction, blood pressure control, heart failure treatment, increasing 1791 1792 cardiorespiratory fitness, and other measures (see Chapter $\hat{6}$). 1793

1794 11.2.1. Selection of antiarrhythmic drugs for long-term therapy: Safety first!

1795 Usually, the safety of antiarrhythmic drug therapy determines the initial choice of antiarrhythmic drugs (Figure 1796 17). The following major antiarrhythmic drugs are available to prevent AF:

1797

1798 Amiodarone is an effective multichannel blocker, reduces ventricular rate, and is safe in patients with heart 1799 failure.^{582, 651} Torsades de pointes proarrhythmia can occur, and QT interval and TU waves should be monitored on therapy (see Table 17).⁶⁵² Amiodarone often causes extracardiac side-effects, especially on long-term 1800 therapy, ^{653, 654} rendering it a second-line treatment in patients who are suitable for other antiarrhythmic drugs. 1801 1802 Amiodarone appears less suitable to episodic short-term therapy (unless after catheter ablation),⁶⁵⁵ probably 1803 because of its long biological half-life.

1804

1805 Dronedarone maintains sinus rhythm, reduces ventricular rate, and prevents cardiovascular hospitalizations (mostly due to AF) and cardiovascular death in patients with paroxysmal or persistent AF or flutter who had at least one relevant cardiovascular comorbidity.^{583, 588, 656} Dronedarone increases mortality in patients with 1806 1807 recently decompensated heart failure (with or without AF)⁶⁵⁷ and in patients with permanent AF in whom sinus 1808 rhythm is not restored.⁶⁵⁸ Dronedarone moderately increases serum creatinine, reflecting a reduction in 1809 creatinine excretion rather than a decline in kidney function.⁶⁵⁹ 1810

1811

Flecainide and propafenone are effective in preventing recurrent AF.^{581, 584, 620} They should only be used in 1812 patients without significant ischaemic heart disease or heart failure to avoid the risk of life-threatening 1813 ventricular arrhythmias.⁶⁶⁰ High ventricular rates resulting from the conversion of AF into atrial flutter with 1:1 1814

1815 conduction by flecainide or propafenone can be prevented by preadministering a beta-blocker, verapamil, or diltiazem. 1816

1817

1818 Quinidine and disopyramide have been associated with an increase in all-cause mortality (OR 2.39; 95% CI 1.03–5.59; number needed to harm 109; 95% CI 34–4985) at 1-year follow-up,^{580, 661} likely due to ventricular arrhythmias (torsades de pointes).^{580, 661} Although this proarrhythmic effect is more common at higher doses, 1819 1820 1821 they are less commonly used for rhythm control in AF. Disopyramide may be useful in 'vagally mediated' AF 1822 (e.g. AF occurring in athletes and/or during sleep⁷⁶), and has been shown to reduce LV outflow gradient and improve symptoms in patients with hypertrophic cardiomyopathy. 662-664 1823 1824

1825 Sotalol has a relevant risk of torsades de pointes (1% in the Prevention of Atrial Fibrillation After Cardioversion 1826 [PAFAC] trial¹¹⁸). Its d-enantiomer is associated with an increased mortality compared to placebo in patients with LV dysfunction post-myocardial infarction,⁶⁶⁵ probably due to ventricular arrhythmias (OR 2.47; 95% CI 1.2–5.05; number needed to harm 166; 95% CI 61–1159).^{580, 665} On the other hand, d,l sotalol has been used in AF patients without safety signals in two controlled trials.^{581, 601} 1827 1828 1829

1830

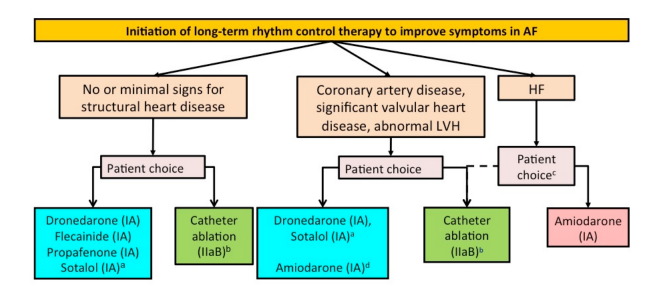
1831 **Dofetilide** is another potassium channel blocker that is mainly available outside of Europe. Dofetilide restores

and maintains sinus rhythm in heart failure patients⁶⁶⁶ and occasionally in patients refractory to other 1832

antiarrhythmic drugs.66 1833

1834

- 1835 Overall, it seems prudent to limit the use of quinidine, disopyramide, dofetilide, and sotalol to specific
- 1836 situations. Similarly, combinations of QT-prolonging antiarrhythmic drugs should generally be avoided (Table 1837 17).



1838				
1839	Figure 17 Initiation of rhythm control therapy in symptomatic patients.			
1840	AF = atrial fibrillation; HF = heart failure; LVH = left ventricular hypertrophy;			
1841	^a Sotalol requires careful evaluation of proarrhythmic risk.			
1842	^b Catheter ablation should isolate pulmonary veins and can be performed using radiofrequency or cryoballoon			
1843	catheters.			
1844	^c Catheter ablation as a first-line therapy is usually reserved for heart failure patients with tachycardiomyopathy.			
1845	^d Amiodarone is a second-choice therapy in many patients because of its extracardiac side-effects.			
1846				
1847	11.2.2. Twelve-lead electrocardiogram as a tool to identify patients at risk of			
1848	proarrhythmia			
1849	Identifying patients at risk of proarrhythmia can help to mitigate the proarrhythmic risk of antiarrhythmic			
1850	drugs. ⁶⁶⁸ In addition to the clinical characteristics mentioned above, monitoring PR, QT, and QRS durations			
1851	during initiation of antiarrhythmic drug therapy can identify patients at higher risk of drug-induced			

1852 1853

during initiation of antiarrhythmic drug therapy can identify patients at higher risk of drug-induced proarrhythmia on longer-term treatment.⁶⁶⁹⁻⁶⁷¹ In addition, the presence of 'abnormal TU waves' is a sign of imminent torsades de pointes.⁶⁵² Periodic ECG analysis for proarrhythmia signs has been used successfully in recent antiarrhythmic drug trials.^{118, 584, 672} Specifically, ECG monitoring was used systematically on days 1–3 in 1854

patients receiving flecainide, propafenone, or sotalol to identify those at risk of proarrhythmia.^{118, 584, 601} Based 1855

1856 on this evaluated practice, we suggest to record an ECG in all patients before initiation of antiarrhythmic drugs.

1857 Scheduled ECGs during the initiation period seem reasonable (Table 17).

1858

1859 Table 17 Oral antiarrhythmic drugs used for maintaining sinus rhythm after cardioversion.

ESC AF Guidelines

Drug	Dose	Main contraindications and precautions	Warning signs warranting discontinuation	Atrioventricu lar nodal slowing	Suggested ECG monitoring during initiation
Amiodarone	600 mg in divided doses for 4 weeks, 400 mg for 4 weeks, then 200 mg once daily	Caution when using concomitant therapy with QT-prolonging drugs and in patients with sinoatrial node or atrioventricular node and conduction disease. The dose of VKAs and of digitalis should be reduced. Increased risk of myopathy with statins. Caution in patients with pre- existing liver disease	QT prolongation > 500 ms	10–12 bpm in AF	Baseline, 1 week, 4 weeks
Dronedarone	400 mg twice daily	Contraindicated in NYHA class III or IV or unstable heart failure, during concomitant therapy with QT-prolonging drugs, or powerful CYP3A4 inhibitors (e.g. verapamil, diltiazem, azole antifungal agents), and when CrCl < 30 mg/mL. The dose of digitalis, beta-blockers, and of some statins should be reduced. Elevations in serum creatinine of 0.1–0.2 mg/dL are common and do not reflect a decline in renal function. Caution in patients with pre- existing liver disease	QT prolongation > 500 ms	10–12 bpm in AF	Baseline, 1 week, 4 weeks
Flecainide Flecainide slow release	100–150 mg twice daily 200 mg once daily	Contraindicated if CrCl < 50 mg/mL, liver disease, IHD, or reduced LVEF. Caution in the presence of sinoatrial node or atrioventricular node or conduction system disease. CYP2D6 inhibitors (e.g. fluoxetine, tricyclic) increase plasma concentration	QRS duration increases > 25% above baseline	None	Baseline, day 1, day 2–3
Propafenone	150–300 mg three times daily	Contraindicated in IHD or reduced LV ejection fraction. Caution in the presence of sinoatrial node or atrioventricular node and conduction system disease, renal or liver impairment, and asthma. Increases concentration of digitalis and warfarin	QRS duration increase > 25% above baseline	Slight	Baseline, day 1, day 2–3
Propafenone SR	225–425 mg twice daily				
d,l sotalol	80–160 mg twice daily	Contraindicated in the presence of significant LV hypertrophy, systolic heart failure, asthma, pre-existing QT prolongation, hypokalaemia, CrCl < 50 mg/mL. Moderate renal dysfunction requires careful adaptation of dose	QT interval > 500 ms, QT prolongation by > 60 ms upon therapy initiation	Similar to high-dose blockers	Baseline, day 1, day 2–3

1860AF = atrial fibrillation; bpm = beats per minute; CrCl = creatinine clearance; ECG = electrocardiogram; IHD =1861ischaemic heart disease; LV = left ventricular; LVEF = left ventricular ejection fraction; NYHA = New York1862Heart Association; VKA = vitamin K antagonist.

1863

1864 **11.2.3.** New antiarrhythmic drugs

Several compounds that inhibit the ultrarapid potassium current (I_{Kur}) and other inhibitors of atypical ion channels are in clinical development.⁶⁷³⁻⁶⁷⁵ They are not available for clinical use at present. The antianginal 1865 1866 compound ranolazine inhibits potassium and sodium currents and increases glucose metabolism at the expense of free fatty acid metabolism, thereby enhancing efficient use of oxygen.^{676, 677} Ranolazine was safe in patients 1867 1868 1869 with non-ST-segment elevation myocardial infarction and unstable angina evaluated in the MERLIN 1870 (Metabolic Efficiency With Ranolazine for Less Ischemia in Non ST-Elevation Acute Coronary Syndrome) trial.⁶⁷⁸ In a post-hoc analysis of continuous ECG recordings obtained during the first 7 days after 1871 1872 randomization, patients assigned to ranolazine had a trend towards fewer episodes of AF than those on placebo (75 [2.4%] vs. 55 [1.7%] patients; P = 0.08).⁶⁷⁹ In the HARMONY (A Study to Evaluate the Effect of 1873 1874 Ranolazine and Dronedarone When Given Alone and in Combination in Patients With Paroxysmal Atrial 1875 Fibrillation) trial, the highest tested dose of a combination of ranolazine (750 mg twice daily) and dronedarone 1876 (225 mg twice daily) slightly reduced AF burden in 134 subjects with paroxysmal AF and dual-chamber pacemakers.⁶⁸⁰ Small, open-label studies suggest that ranolazine might enhance the antiarrhythmic effect of 1877 amiodarone for cardioversion,⁶⁸¹⁻⁶⁸³ whereas the results from a controlled trial of ranolazine and the ranolazine– 1878 dronedarone combination to prevent AHRE in pacemaker patients were ambiguous.⁶⁸⁴ At present, there is 1879 1880 insufficient evidence to recommend ranolazine as an antiarrhythmic drug, alone or in combination with other 1881 antiarrhythmic drugs. Of note, the 'funny channel blocker' ivabradine, which is used for angina and heart 1882 failure, increases the risk of AF.685 1883 1884 Antiarrhythmic effects of non-antiarrhythmic drugs 11.2.4.

ACE inhibitors or ARBs appear to prevent new-onset AF in patients with LV dysfunction and in hypertensive patients with LV hypertrophy.^{219, 236, 237, 239, 246, 250, 686} Neprilysin inhibition needs to be studied further, but does 1885 1886 not seem to enhance this effect.²²⁴ A Danish cohort study also suggested that initial treatment of uncomplicated 1887 hypertension with ACE inhibitors or ARBs reduces incident AF compared with other hypertensive agents.²⁴⁵ 1888 ARB therapy did not reduce the AF burden in patients with AF without structural heart disease.²⁴¹ Thus, ACE 1889 inhibitors or ARBs are unlikely to have a relevant direct antiarrhythmic effect. However, it might be justified to 1890 1891 consider adding ACE inhibitors or ARB therapy to antiarrhythmic drugs to reduce AF recurrences after cardioversion.^{248, 249, 687} 1892 Compared with placebo, beta-blockers are associated with a reduced risk of new-onset AF in patients 1893

1893 Compared with placebo, beta-blockers are associated with a reduced risk of new-onset AF in patients 1894 with reduced ejection fraction and sinus rhythm.²³ Beta-blockers have also been reported to reduce symptomatic 1895 AF recurrences,^{580, 636, 688} but this finding may be driven by the beneficial effect of rate control, which will

1896 render AF more often asymptomatic.

Perioperative statin therapy appeared to reduce the risk of postoperative AF in a number of small
 RCTs^{689, 690}; however, an adequately powered placebo-controlled trial has shown no effect of perioperative
 rosuvastatin therapy on postoperative AF.⁶⁹¹ Statin treatment does not prevent AF in other settings.^{692, 693}
 Similarly, polyunsaturated fatty acids failed to show convincing benefit.^{241, 694-698} The role of aldosterone

antagonists in the management of AF has not been extensively investigated in humans; although preliminary

1902 evidence from trials of eplerenone is encouraging for primary prevention,²⁴³ at present there is no robust

evidence to make any recommendation for the use of aldosterone antagonists for secondary prevention of AF.⁶⁹⁹⁻ 701

- 1905
- 1906 **Recommendations for rhythm control therapy**

Recommendations	Class ^a	Level ^b	Refs ^d
General recommendations			
Management of cardiovascular risk factors and avoidance of AF triggers should be pursued in patients on rhythm control therapy to facilitate maintenance of sinus rhythm	lla	В	203, 204, 296, 312
Rhythm control therapy is indicated for symptom improvement in patients with AF	I	В	120, 586, 601
With the exception of AF associated with haemodynamic instability, the choice between electrical and pharmacological cardioversion	lla	С	

should be guided by patient and physician preferences			
Cardioversion of AF	1		
Electrical cardioversion of AF is recommended in patients with acute naemodynamic instability to acutely restore cardiac output	1	В	612, 702- 704
Cardioversion of AF (either electrical or pharmacological) is ecommended in symptomatic patients with persistent or long- standing persistent AF as part of rhythm control therapy	I	В	584, 601, 627, 628, 648, 705
Pretreatment with amiodarone, flecainide, ibutilide, or propafenone should be considered to enhance success of electrical cardioversion and prevent recurrent AF	lla	В	248, 584, 633
n patients with no history of ischaemic or structural heart disease, lecainide, propafenone, or vernakalant are recommended for pharmacological cardioversion of new-onset AF	I	A	602-605, 614, 618, 622, 706, 707
n patients with no history of ischaemic or structural heart disease, butilide should be considered for pharmacological conversion of AF	lla	В	
n selected patients with recent-onset AF and no significant structural or ischaemic heart disease, a single oral dose of flecainide or propafenone (the 'pill in the pocket' approach) should be considered or patient-led cardioversion, following safety assessment	lla	В	620, 621
n patients with ischaemic and/or structural heart disease, amiodarone is recommended for cardioversion of AF	I	A	597-601
/ernakalant may be considered as an alternative to amiodarone for oharmacological conversion of AF in patients without hypotension, severe heart failure, or severe structural heart disease (especially aortic stenosis)	llb	В	602-605, 616, 618
Stroke prevention in patients designated for cardioversion of AF			
Anticoagulation with heparin or a NOAC should be initiated as soon as possible before every cardioversion of AF or atrial flutter	lla	В	708, 709
For cardioversion of AF/atrial flutter, effective anticoagulation is ecommended for a minimum of 3 weeks before cardioversion	I	В	648, 708
Fransoesophageal echocardiography (TOE) is recommended to exclude cardiac thrombus, as an alternative to preprocedural anticoagulation when early cardioversion is planned	I	В	648, 708
Early cardioversion can be performed without TOE in patients with a definite duration of $AF < 48$ hours	lla	В	648
n patients at risk for stroke (e.g. presence of CHA ₂ DS ₂ -VASc actors), anticoagulant therapy should be continued long-term after cardioversion according to the long-term anticoagulation ecommendations, irrespective of the method of cardioversion or the apparent maintenance of sinus rhythm. In patients without stroke risk actors, anticoagulation is recommended for 4 weeks after cardioversion	I	В	353, 710
n patients where thrombus is identified on TOE, effective anticoagulation is recommended for at least 3 weeks	I	С	
A repeat TOE to ensure thrombus resolution should be considered	lla	С	

The choice of antiarrhythmic drug needs to be carefully evaluated, taking into account the presence of comorbidities, cardiovascular risk and potential for serious proarrhythmia, extracardiac toxic effects, patient preferences, and symptom burden	1	A	41, 580
Dronedarone, flecainide, propafenone, or sotalol are recommended for prevention of recurrent symptomatic AF in patients with normal left ventricular function and without pathological left ventricular hypertrophy.	1	A	581, 583, 584, 588, 601
Dronedarone is recommended for prevention of recurrent symptomatic AF in patients with stable coronary artery disease, and without heart failure	I	A	583, 588
Amiodarone is recommended for prevention of recurrent symptomatic AF in patients with heart failure	1	В	596-598
Amiodarone is more effective in preventing AF recurrences than other antiarrhythmic drugs but extracardiac toxic effects are common and increase with time. For this reason, other antiarrhythmic drugs should be considered first	lla	С	596-598
Patients on antiarrhythmic drug therapy should be periodically evaluated to confirm their eligibility for treatment	lla	С	583, 588, 657, 658, 660
ECG recording during the initiation of antiarrhythmic drug therapy should be considered to monitor heart rate, detect QRS and QT interval prolongation, and the occurrence of atrioventricular block	lla	В	584 582, 583, 588, 601
Antiarrhythmic drug therapy is not recommended in patients with prolonged QT interval (> 0.5 s) or with significant sinoatrial node disease or atrioventricular node dysfunction who do not have a functioning permanent pacemaker	III (harm)	С	
Adding atrial-based bradycardia pacing to drug treatment that induces or exacerbates sinus node dysfunction should be considered to allow continuation of antiarrhythmic drug therapy in patients in whom AF ablation is declined or not indicated	lla	В	/11, /12
Continuation of antiarrhythmic drug therapy beyond the blanking period after AF ablation should be considered to maintain sinus rhythm when recurrences seem likely	lla	В	713
Antiarrhythmic effects of non-antiarrhythmic drugs			
ACE inhibitors, ARBs, and beta-blockers should be considered for prevention of new-onset AF in patients with heart failure and reduced ejection fraction	lla	A	23, 219, 236 237, 239, 250, 714
ACE inhibitors and ARBs should be considered for prevention of new- onset AF in patients with hypertension, particularly with LV hypertrophy	lla	В	238, 246, 686, 714
Pretreatment with ACE inhibitors or ARBs may be considered in patients with recurrent AF undergoing electrical cardioversion and receiving antiarrhythmic drug therapy	llb	В	236, 237, 248, 249
ARBs or ACE inhibitors are not recommended for the secondary prevention of paroxysmal AF in patients with little or no underlying heart disease.	III (no benefit)	В	241, 697
ACE = angiotensin-converting enzyme: $AE =$ atrial fibrillation: $ABB =$ angiotensi		1.1	CILL DC

1907 ACE = angiotensin-converting enzyme; AF = atrial fibrillation; ARB = angiotensin receptor blocker; CHA_2DS_2 -

1908 VASc = Congestive Heart failure, hypertension, Age \geq 75 (doubled), Diabetes, Stroke (doubled), Vascular

1909 disease, Age 65–74, and Sex (female); ECG = electrocardiogram; NOAC = non-vitamin K antagonist oral

1910 anticoagulant; TOE = transoesophageal echocardiography.

1911 ^aClass of recommendation.

1912 ^bLevel of evidence.

1913 ^cReference(s) supporting recommendations.

1914

1915 11.3. Catheter ablation

1916 Since the initial description of triggers in the pulmonary veins that initiate paroxysmal AF,¹⁰⁸ catheter ablation 1917 of AF has developed from a specialized, experimental procedure into a common treatment to prevent recurrent 1918 AF.^{587, 715} This is primarily achieved through isolation of the pulmonary veins, probably requiring complete 1919 isolation for full effectiveness,⁷¹⁶ and additional ablation in the posterior left atrial wall. AF ablation, when 1920 performed in experienced centres by adequately trained teams, is more effective than antiarrhythmic drug 1921 therapy in maintaining sinus rhythm, and the complication rate, though not negligible, is similar to the

1922 complication rate for antiarrhythmic drugs.^{585, 717,} 1923

1924 **11.3.1**. Indications

1925 Catheter ablation of AF is effective in restoring and maintaining sinus rhythm in patients with symptomatic 1926 paroxysmal, persistent, and probably long-standing persistent AF - in general as second-line treatment after 1927 failure of or intolerance to antiarrhythmic drug therapy. In such patients, catheter ablation is more effective than 1928 antiarrhythmic drug therapy.^{185, 586, 713, 717-720} As first-line treatment for paroxysmal AF, randomized trials 1929 showed only modestly improved rhythm outcome with catheter ablation compared to antiarrhythmic drug 1930 therapy.^{585, 721-723} Complication rates were similar, but ablation was performed in expert centres, justifying 1931 catheter ablation as first-line therapy in selected patients with paroxysmal AF who ask for interventional 1932 therapy. Fewer data are available reporting the effectiveness and safety of catheter ablation in patients with 1933 persistent or long-standing persistent AF, but all point to lower recurrence rates after catheter ablation compared to antiarrhythmic drug therapy with or without cardioversion.^{185, 717, 723-726, 1039} In patients who experience 1934 symptomatic recurrences of AF despite antiarrhythmic drug therapy, all RCTs showed better sinus rhythm maintenance with catheter ablation than on antiarrhythmic drugs.^{586, 713, 727, 728} There is no current indication for 1935 1936 catheter ablation to prevent cardiovascular outcomes (or desired withdrawal of anticoagulation), or to reduce 1937 hospitalization.40, 594 1938 1939

1940 11.3.2. Techniques and technologies

1941 Complete pulmonary vein isolation (PVI) on an atrial level is the best documented target for catheter ablation,^{716, 729-731} achievable by point-by-point radiofrequency ablation, linear lesions encircling the pulmonary veins, or cryoballoon ablation, with similar outcomes.⁷³²⁻⁷³⁴ Complete isolation of the pulmonary veins has better rhythm outcomes than incomplete isolation.⁷¹⁶ PVI was initially tested in patients with paroxysmal AF, 1942 1943 1944 1945 but appears to be non-inferior to more extensive ablation in persistent AF as well.^{729, 735} More extensive 1946 ablations have been used in patients with persistent AF, but there are insufficient data to guide the use of these at present.^{117, 718, 719, 735-737} Extended ablation procedures (beyond PVI) consistently require longer procedures and 1947 more ionizing radiation, potentially creating risk for patients. Left atrial macro-reentrant tachycardia is relatively 1948 uncommon after PVI (\approx 5%). It also seems even less common after cryoballoon ablation,⁷³⁴ but may occur in up 1949 to 25% of patients after left atrial substrate modification ablation, often due to incomplete ablation lines. Thus, 1950 1951 for patients with persistent AF, ablation of complex fractionated electrograms, ablation of rotors, or routine deployment of linear lesions or other additional ablations does not seem justified in the first procedure.^{735, 738, 739} 1952 1953 However, additional ablation on top of complete PVI⁷¹⁶ may be considered in patients with recurrent AF after the initial ablation procedure.^{719, 740, 741} In patients with documented right atrial isthmus-dependent flutter 1954 1955 undergoing AF ablation, right atrial isthmus ablation is recommended. Adenosine testing to identify patients in need of additional ablation remains controversial after evaluation in several reports.^{739, 742-744} Ablation of so-1956 1957 called 'rotors' guided by body surface mapping or endocardial mapping is under evaluation and cannot be 1958 recommended for routine clinical use at present.

1959

1960 **11.3.3.** Outcome and complications

The rhythm outcome after catheter ablation of AF is difficult to predict in individual patients.^{173, 227, 713, 728} Most 1961 patients require more than one procedure to achieve symptom control.^{713, 726, 728} In general, better rhythm 1962 1963 outcome and lower procedure-related complications can be expected in younger patients with a short history of AF and frequent, short AF episodes in the absence of significant structural heart disease.⁷⁴⁵ Catheter ablation is 1964 1965 more effective than antiarrhythmic drug therapy in maintaining sinus rhythm (Web Addenda Figure 2).⁷⁴⁶ 1966 Sinus rhythm without severely symptomatic recurrences of AF is found in up to 70% of patients with paroxysmal AF, and around 50% in persistent AF.^{713, 728, 735, 1042} Very late recurrence of AF after years of sinus 1967 1968 rhythm is not uncommon and may reflect disease progression, with important implications for continuation of 1969 AF therapies.⁷²⁸ Multiple variables have been identified as risk factors for recurrence after catheter ablation of 1970 AF, but their predictive power is weak. The decision for catheter ablation thus should be based on a shared 1971 decision-making process⁷⁴⁷ (see Chapter 7), following thorough explanation of the potential benefits and risks, 1972 and of the alternatives such as antiarrhythmic drug or acceptance of current symptoms without rhythm control 1973 therapy.¹⁷⁵

1974 Complications of catheter ablation for AF

1975 There is a clear need to systematically capture complications in clinical practice to improve the quality of AF 1976 ablation procedures.¹⁷⁵ The median length of hospital stay in AF patients undergoing their first ablation as part

1977 of the EURObservational Research Programme (EORP) was 3 days (interquartile range 2-4 days), based on

1978 data from 1391 patients from hospitals performing at least 50 ablations per year. Five to seven per cent of

1979

patients will suffer severe complications after catheter ablation of AF, and 2–3% will experience life-threatening but usually manageable complications.^{727, 748-750} Intraprocedural death has been reported, but is rare (< 0.2%).⁷⁵¹ 1980

1981 The most important severe complications are stroke/TIA (<1%), cardiac tamponade (1-2%), pulmonary vein

1982 stenosis, and severe oesophageal injury leading to atrio-oesophageal fistula weeks after ablation (Table 18).

1983 'Silent strokes' (i.e. white matter lesions detectable by brain MRI), have been observed in around 10% of 1984 patients treated with radiofrequency and cryoballoon ablation.⁷⁵² The clinical relevance of this observation is

unclear.⁷⁴⁹ Post-procedure complications include stroke, with the highest risk within the first week,⁷⁵³ late pericardial tamponade several days after catheter ablation,⁷⁵¹ and oesophageal fistulas, which usually become 1985

1986

1987 apparent 7-30 days after ablation. Timely detection of atrio-oesophageal fistulas can be life-saving and should 1988 be based on the typical triad of infection without a clear focus, retrosternal pain, and stroke or TIA.⁷⁴

1989

1990	Table 18 Complications related to catheter ablation of AF
------	---

Complication severity	Complication type	Rate ^{727, 748, 750, 754-759}
	Periprocedural death	< 0.2%
	Oesophageal injury (perforation/fistula) ^a	< 0.5%
Life-threatening complications	Periprocedural stroke (including TIA/air embolism)	< 1%
	Cardiac tamponade	1–2%
	Pulmonary vein stenosis	< 1%
Course courseling time	Persistent phrenic nerve palsy	1–2%
Severe complications	Vascular complications	2–4%
	Other severe complications	≈ 1%
Other moderate or minor complications		1–2%
Unknown significance	Asymptomatic cerebral embolism (silent stroke) ^b	5–20%
	Radiation exposure	

1991 AF = atrial fibrillation; TIA = transient ischaemic attack.

1992 ^aOesophageal fistula should be suspected in patients presenting with the triad of unspecific signs of infection,

1993 chest pain, and stroke or TIA in the first weeks after an ablation procedure. It requires immediate therapy.

1994 b < 10% for cryoablation or radiofrequency ablation, > 20% for phased radiofrequency ablation

1995

1996 Anticoagulation – before, during, and after ablation 11.3.4.

Patients anticoagulated with VKAs should continue therapy during ablation (with an INR of 2–3).⁷⁶⁰ Anticoagulation with NOACs is an alternative to warfarin.^{478, 761-765} There is no safety signal from observational 1997 1998 1999 cohorts treated with uninterrupted NOAC therapy undergoing catheter ablation in experienced centres.^{761, 763, 766,} ⁷⁶⁷ The first controlled trial, enrolling around 200 patients, has recently been published, ⁷⁶⁸ as well as several 2000 observational data sets.^{761, 769, 770} Ongoing studies compare uninterrupted VKA with NOAC therapy in AF 2001 2002 patients undergoing ablation (e.g. AXAFA - AFNET 5 [Apixaban During Atrial Fibrillation Catheter Ablation: 2003 Comparison to Vitamin K Antagonist Therapy – Anticoagulation using the direct factor Xa inhibitor apixaban 2004 during Atrial Fibrillation catheter Ablation: Comparison to vitamin K antagonist therapy; NCT02227550] and 2005 RE-CIRCUIT [Randomized Evaluation of dabigatran etexilate Compared to warfarin in pulmonaRy vein

ablation: assessment of different peri-proCedUral anticoagulation sTrategies; NCT02348723]). During ablation,
heparin should be given to maintain an activated clotting time > 300 seconds. Anticoagulation should be
maintained for at least 8 weeks after ablation for all patients. The true incidence of thromboembolic events after
catheter ablation has never been systematically studied and the expected stroke risk has been adopted from nonablation AF cohorts. Although observational studies suggest a relatively low stroke rate in the first few years

2011 after catheter ablation of AF, $^{737, 771-776}$ the long-term risk of recurrent AF and the safety profile of

2012 anticoagulation in ablated patients need to be considered. In the absence of controlled trial data, OAC after

2013 catheter ablation should follow general anticoagulation recommendations, regardless of the presumed rhythm 2014 outcome.

2015

2016 **11.3.5.** Ablation of atrial fibrillation in heart failure patients

2017 Catheter ablation, compared with amiodarone therapy, significantly reduces recurrent AF in AF patients with 2018 HFrEF.⁷⁷⁷ Selected patients with HFrEF and AF can achieve recovery of LV systolic function after catheter 2019 ablation (probably reflecting tachycardiomyopathy). Several smaller trials suggest improved LV function after 2020 catheter ablation in HFrEF patients^{185, 226-228, 778, 779} and reduced hospitalizations,^{720, 777} especially in patients 2021 without a previous myocardial infarction.⁷⁸⁰ Larger trials are warranted to confirm these findings. Catheter 2022 ablation can be demanding in these patients. Thus, indications for catheter ablation in HFrEF patients should be 2023 carefully balanced, and the procedures performed in experienced centres.

2025 **11.3.6.** Follow-up after catheter ablation

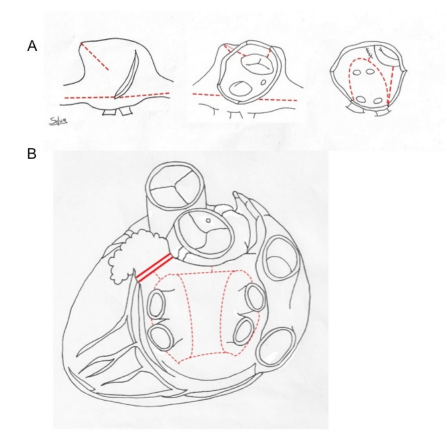
Patients and physicians involved in the follow-up after catheter ablation should know the signs and symptoms of late complications to allow swift referral for treatment. Patient should also be aware that symptomatic and asymptomatic AF recurrences are frequent after catheter ablation.^{119, 781, 782} In line with the primary goal of rhythm control therapy, asymptomatic episodes should generally not trigger further rhythm control therapy. Patients should be seen at least once by a rhythm specialist in the first 12 months after ablation. Further rhythm control options should be considered in patients with symptomatic recurrences, including discussion in a Heart Team (*Figure 17*).

2032 Team (2033

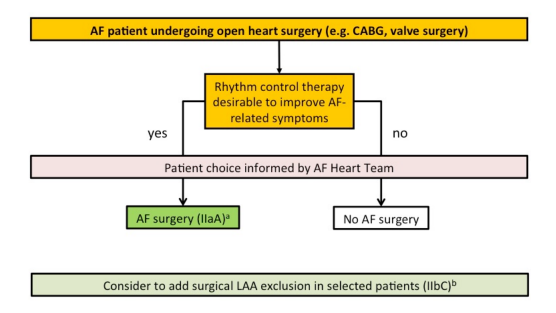
2034 11.4. Atrial fibrillation surgery

2035 **11.4.1.** Concomitant atrial fibrillation surgery

2036 The Cox maze procedure was first performed 30 years ago as a 'cut-and-sew' technique, including isolation of 2037 the posterior left atrium, a connection to the posterior mitral annulus, a cavotricuspid connection, a cavocaval connection, and exclusion of the LAA (Figure 18).⁷⁸³ Thereby, the Cox maze procedure creates an electrical 2038 2039 labyrinth (maze) of passages through which the sinoatrial node impulse finds a route to the atrioventricular node 2040 while preventing fibrillatory conduction. The Cox maze procedure and other, often simpler, forms of AF surgery have mainly been used in patients undergoing other open heart surgical procedures.^{461, 466, 784-798} In a systematic 2041 2042 review commissioned for these guidelines, concomitant AF surgery resulted in greater freedom from AF, atrial 2043 flutter, and atrial tachycardia (RR 1.94, 95% CI 1.51–2.49; n = 554 from seven RCTs) (Web Addenda Figure 2044 3).¹⁰⁴⁰ Patients undergoing the Cox maze procedure required pacemaker implantation more often (RR 1.69, 95% 2045 CI 1.12–2.54; n = 1631 from 17 RCTs), without a detectable difference in other outcomes or complications. 2046 These findings are underpinned by an analysis of Society of Thoracic Surgeons database comprising 67,389 2047 patients in AF: mortality or major morbidity was not affected by concomitant AF surgery (adjusted OR 1.00; 95% CI 0.83–1.20), but pacemaker implantation was more frequent (adjusted OR 1.26; 95% CI 1.07–1.49).⁷⁹⁹ 2048 2049 Predictors of AF recurrence after surgery include left atrial dilatation, older age, > 10-year history of AF, and non-paroxysmal AF.⁸⁰⁰⁻⁸⁰⁴ Regarding AF type, surgical PVI seems effective in paroxysmal AF.⁸⁰⁵ Biatrial lesion patterns may be more effective in persistent and long-standing persistent AF.^{797, 803, 806} The suggested 2050 2051 2052 management of patients with AF-related symptoms undergoing cardiac surgery is displayed in Figure 19, with 2053 an important contribution of the AF Heart Team to advise and inform patient choice.



- 2054
 2055 Figure 18 A. Surgical lesion sets for
 2056 Right panel: left atrial lesions.
 2057 B: Left atrial lesions in a thoracosco
 2058 appendage exclusion (double line). Figure 18 A. Surgical lesion sets for the biatrial Cox maze procedure. Left and middle panel: right atrial lesions.
- B: Left atrial lesions in a thoracoscopic minimally invasive surgical procedure (dashed lines), including left



2059

2039	
2060	Figure 19 Surgical rhythm control in patients undergoing cardiac surgery.
2061	AF = atrial fibrillation; CABG = coronary artery bypass graft; LAA = left atrial appendage; PVI = pulmonary
2062	vein isolation.
2063	^a AF surgery may be PVI in paroxysmal AF and biatrial maze in persistent or long-standing persistent AF.
2064	^b Oral anticoagulation should be continued in patients at risk of stroke irrespective of AF surgery or LAA
2065	exclusion.
2066	
2067	

2068 **11.4.2.** Stand-alone rhythm control surgery

Current technology (e.g. bipolar radiofrequency or cryothermy) renders the procedure easier and more reproducible and feasible via a mini-thoracotomy.^{786, 807, 808} Thoracoscopic PVI with bipolar radiofrequency 2069 2070 prevents recurrence of paroxysmal AF (69–91% freedom from arrhythmias at 1 year, see Figure 18B for lesion 2071 set),^{468, 809, 810} and seems effective in patients refractory to catheter ablation.⁸¹¹ The average length of hospital stay for thoracoscopic ablation varies from 3.6 to 6.0 days.^{468, 812, 813} The FAST (Atrial Fibrillation Catheter 2072 2073 Ablation vs Surgical Ablation Treatment) trial,⁴⁶⁸ and another smaller trial,⁸¹⁴ suggested that thoracoscopic AF 2074 surgery could be more effective than catheter ablation for the maintenance of sinus rhythm,^{468, 814} while also causing more complications (*Table 19*).⁸¹⁵ To improve results,^{468, 816-818} more extensive lesion sets have been 2075 2076 performed, connecting lines between the PVI encircling and towards the mitral annulus.^{812, 819-822} To improve the 2077 generation of transmural lesions,⁷¹⁶ endo-epicardial ablation strategies have recently been proposed.^{812, 823-825} 2078 2079 Although preliminary experience with hybrid simultaneous ablation shows promise, procedural time and rates of 2080 bleeding complications are higher.^{812, 823} 2081

2082 Table 19 Complications of thoracoscopic AF surgery
Complication

Complication	Rate ^{468, 815, 822, 826}
Conversion to sternotomy	0–1.6%
Pacemaker implantation	0–3.3%
Drainage for pneumothorax	0–3.3%
Pericardial tamponade	0–6.0%
Transient ischaemic attack ^a	0–3.0%

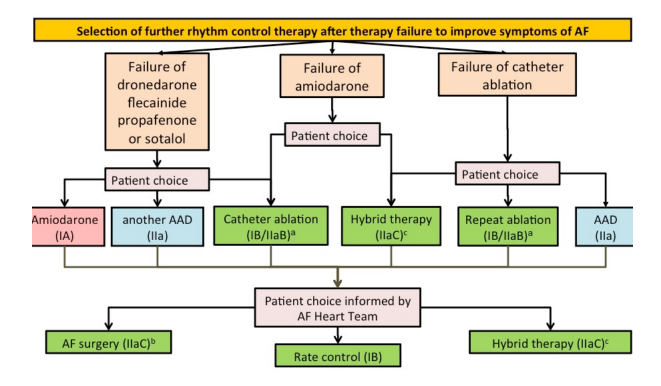
2083 AF = atrial fibrillation.

2084 ^aThe rate of asymptomatic cerebral embolism is unknown

2085

2086 11.5. Choice of rhythm control following treatment failure

2087 There is insufficient evidence on which to base clear recommendations on how to treat patients with recurrent 2088 AF after catheter ablation. Early recurrences of AF or atrial tachycardias after ablation (occurring within 8 2089 weeks) may be treated with cardioversion. Many of the published series of patients undergoing AF ablation 2090 included those who failed antiarrhythmic drug therapy. Thus, considering ablation therapy in patients who have 2091 symptomatic recurrences on antiarrhythmic drug therapy is often reasonable. Alternatively, trialling another 2092 antiarrhythmic drug can be considered. Combining antiarrhythmic drug with ablation ('hybrid therapy', see 2093 Section 11) should be considered based on the different and possibly synergistic effects of these drugs with AF 2094 ablation, possibly benefitting patients in whom either treatment alone was previously ineffective. Rate control 2095 without rhythm control, surgical ablation, or repeat catheter ablation should be considered as well as third-line 2096 options (Figure 20). Patient preferences and local access to therapy are important considerations to inform the 2097 therapy choice in patients who are in need of further rhythm control therapy after an initial therapy failure.



2098 2099 Figure 20 Choice of rhythm control approaches following treatment failure. 2100 AAD = antiarrhythmic drug; AF = atrial fibrillation; PVI = pulmonary vein isolation. 2101 ^a catheter ablation should target PVI. Class I level B for paroxysmal AF and Class IIa level B for persistent AF. ^b AF surgery may be PVI (e.g. in paroxysmal AF) or maze surgery (e.g. in therapy-refractory or long-standing 2102 2103 persistent AF). 2104 ^c Hybrid therapy involves combination of antiarrhythmic drugs, catheter ablation, and/or AF surgery. 2105 2106 11.6. The atrial fibrillation Heart Team 2107 In view of the complexity of the different treatment options in patients with failed rhythm control therapy but 2108 who still require or demand further rhythm control therapy, this Task Force proposes that decisions involving

AF surgery or extensive AF ablation should be based on advice from an AF Heart Team. This will also apply to

- 2110 reversal to a rate control strategy in patients with severe (EHRA III or IV) AF symptoms. An AF Heart Team 2111 should consist of a cardiologist with expertise in antiarrhythmic drug therapy, an interventional
- 2111 should consist of a cardiologist with expertise in antiarrhythmic drug therapy, an interventional 2112 electrophysiologist, and a cardiac surgeon with expertise in appropriate patient selection, techniques, and

- technologies for interventional or surgical AF ablation. Such AF Heart Teams and a collaborative
 - infrastructure supporting a continued interaction between physicians delivering continued care, AF
- cardiologists, interventional electrophysiologists, and AF surgeons should be established to provide optimal
- 2113 2114 2115 2116 2117 2118 2110 advice and ultimately to improve rhythm outcomes for patients in need of advanced and complex rhythm control
- interventions.

2119 Recommendations for catheter ablation of AF and AF surgery

Recommendations	Class ^a	Level ^b	Refs ^c
Catheter ablation of symptomatic paroxysmal AF is recommended to improve AF symptoms in patients who have symptomatic recurrences of AF on antiarrhythmic drug therapy (amiodarone, dronedarone, flecainide, propafenone, sotalol) and who prefer further rhythm control therapy, when performed by an electrophysiologist who has received appropriate training and is performing the procedure in an experienced centre	I	A	585-587, 713, 727
Ablation of common atrial flutter should be considered to prevent recurrent flutter as part of an AF-ablation procedure if previously documented or occurring during the AF ablation	lla	В	827
Catheter ablation of AF should be considered as first-line therapy to prevent recurrent AF and to improve symptoms in selected patients with symptomatic paroxysmal AF as an alternative to antiarrhythmic drug therapy, considering patient choice, benefit, and risk	lla	В	585
All patients should receive oral anticoagulation for stroke prevention for at least 8 weeks after catheter (IIaB) or surgical (IIaC) ablation.	lla	B/C	/2/
Anticoagulation for stroke prevention should be continued indefinitely after apparently successful catheter or surgical ablation of AF in patients at high risk of stroke	lla	С	
When catheter ablation of AF is planned, continuation of oral anticoagulation with VKA (IIaB) or NOAC (IIaC) should be considered during the procedure, maintaining effective anticoagulation	lla	B/C	760, 768
Catheter ablation should target complete isolation of the pulmonary veins using radiofrequency ablation or cryothermy balloon catheters	lla	В	585, 715, 716, 734, 735
AF ablation should be considered in symptomatic patients with AF and heart failure with reduced ejection fraction to improve symptoms and cardiac function when tachycardiomyopathy is suspected	lla	С	185, 226-228, 720, 777-779, 828
AF ablation should be considered as a strategy to avoid pacemaker implantation in patients with AF-related bradycardia	lla	С	829, 830
Catheter or surgical ablation should be considered in patients with symptomatic persistent or long-standing persistent AF refractory to antiarrhythmic drug therapy to improve symptoms, considering patient choice, benefit and risk, supported by an AF Heart Team	lla	С	468, 735, 777, 831, 832, <mark>1040</mark>
Minimally invasive surgery with epicardial pulmonary vein isolation should be considered in patients with symptomatic AF when catheter ablation has failed. Decisions on such patients should be supported by an AF Heart Team	lla	В	468 812, 819, 823
Maze surgery, possibly via a minimally invasive approach, performed by an adequately trained operator in an experienced centre, should be considered by an AF Heart Team as a treatment option for patients with symptomatic refractory persistent AF or post-ablation AF to improve symptoms	lla	С	808, 832

Maze surgery, preferably biatrial, should be considered in patients undergoing cardiac surgery to improve symptoms attributable to AF, balancing the added risk of the procedure and the benefit of rhythm control therapy	lla	A	461, 466, 790, 791, 796, 797
Concomitant biatrial maze or pulmonary vein isolation surgery may be considered in asymptomatic AF patients undergoing cardiac surgery	llb	С	796, 797, 833

- 2120 AF = atrial fibrillation; NOAC = non-vitamin K antagonist oral anticoagulant; VKA = vitamin K antagonist.
- 2121 ^aClass of recommendation.
- 2122 ^bLevel of evidence.
- 2123 ^cReference(s) supporting recommendations.
- 2124

2125 12 Hybrid rhythm control therapy

- 2126 AF has many different drivers, which are only partially targeted by antiarrhythmic drug or catheter ablation.⁹⁶
- Hence, combination or 'hybrid' rhythm control therapy seems reasonable, although there is little evidence supporting its use.
- 2128 supporting 2129

2130 12.1. Combining antiarrhythmic drugs and catheter ablation

2131 Antiarrhythmic drug therapy is commonly given for 8-12 weeks after ablation to reduce early recurrences of AF 2132 after catheter ablation, supported by a recent controlled trial where amiodarone halved early AF recurrences compared with placebo.⁶⁵⁰ Prospective studies have not been done, but a meta-analysis of the available (weak) 2133 2134 evidence suggests slightly better prevention of recurrent AF in patients treated with antiarrhythmic drugs after catheter ablation.⁷¹³ Many patients are treated with antiarrhythmic drug therapy after catheter ablation (most 2135 often amiodarone or flecainide),⁵⁸⁷ and this seems a reasonable option in patients with recurrent AF after 2136 2137 ablation. It seems common sense to consider antiarrhythmic drug therapy in patients who are in need of further 2138 rhythm control therapy after catheter ablation, but controlled trials to confirm this are desirable. 2139 Combining cavotricuspid isthmus ablation and antiarrhythmic drugs may lead to improved rhythm

2139 Combining cavotricuspid isthmus ablation and antiarrhythmic drugs may lead to improved rhythm 2140 control without the need for left atrial ablation in patients who develop 'drug-induced atrial flutter' on therapy 2141 with flecainide, propafenone, or amiodarone,⁸³⁴⁻⁸³⁶ although recurrent AF is a concern in the long term.^{837, 838} 2142

2143 12.2. Combining antiarrhythmic drugs and pacemakers

In selected patients with sick sinus syndrome and fast ventricular response during AF paroxysms requiring rate control therapy, the addition of a pacemaker not only optimizes rate control but may also help to control rhythm.^{711, 712} Moreover, when antiarrhythmic drug treatment leads to sinus node dysfunction and bradycardia, pacing may permit uptitration of the antiarrhythmic drug dose. Such strategies have never been prospectively investigated and the existing populations studied are highly selected.^{839, 840} Some patients with AF-induced bradycardia may benefit from catheter ablation of AF, obviating the need for antiarrhythmic drugs and pacemaker implantation.^{829, 830}

2151

2152 13 Specific situations

2153 13.1. Frail and 'elderly' patients

2154 Many AF patients present at an older age (e.g. > 75 or > 80 years). There are no studies suggesting that cardiovascular risk reduction is less effective in these 'elderly' AF patients than in younger patients. Rather, age 2155 is one of the strongest predictors/risk factors for ischaemic stroke in AF (Table 11).³⁸² Good data are available to 2156 2157 support the use of anticoagulants in older patients from BAFTA (Birmingham Atrial Fibrillation Treatment of 2158 the Aged Study),³⁶² the NOAC trials,³⁹ and from analyses in elderly Americans (Medicare).³⁹⁶ Elderly AF patients are at higher risk of stroke and thus are more likely to benefit from OAC than younger patients,⁸⁴¹ and yet OAC is still underutilized in the elderly.^{220, 842} Although the evidence base is smaller for other treatment 2159 2160 2161 options in AF, the available data support the use of available rate and rhythm control interventions, including 2162 pacemakers and catheter ablation, without justification to discriminate by age group. Individual patients at older 2163 age may present with multiple comorbidities including dementia, a tendency to falls, CKD, anaemia, 2164 hypertension, diabetes, and cognitive dysfunction. Such conditions may limit quality of life more than AF-2165 related symptoms. Impairment of renal and hepatic function and multiple simultaneous medications make drug 2166 interactions and adverse drug reactions more likely. Integrated AF management and careful adaptation of drug dosing seem reasonable to reduce complications of AF therapy in such patients.⁸⁴³ 2167

2168

2169 13.2. Inherited cardiomyopathies, channelopathies, and accessory pathways

2170 Several inherited cardiac conditions are associated with early-onset AF (Table 20). Treatment of the underlying 2171 cardiac condition is an important contribution to AF management in these young patients (see also ESC 2172 guidelines on the sudden cardiac death⁸⁴⁴ and hypertrophic cardiomyopathy⁸⁴⁵).

2173 2174

Table 20 Inherited cardiomyopathies, channelopathies, and pathways associated with AF

2175

Syndrome Gene **Functional alteration** AF References prevalence 846-850 IKs 🗌 Long QT syndrome KCN01 5-10% IKr 🗌 KCNH2 SCN5A INa 🗌 INa,K 🗌 ANK2 Various effects others 851-855 INa 🗌 Brugada syndrome 10-20% SCN5A GPDIL INa 🗌 SCN1B INa 🗌 CACNA1C ICa 🗌 CACNB2b ICa 🗌 others others 853, 856-858 Short QT syndrome IKr 🗌 Up to 70% KCNH2 IKs 🗌 KCN01 IK1 🗍 KCNJ2 CACNA1C ICa \square CACNB2b ICa 859-861 Abnormal Ca²⁺ release from Catecholaminergic RYR2 Variable but ventricular tachycardia CASQ2 sarcoplasmic reticulum significant 862-864 Hypertrophic 5-15% Sarcomeric cardiomyopathy genes 865 Wolff-Parkinson-White PRKAG Variable syndrome 866 Holt-Oram syndrome TBX5 Variable 867, 868 Arrhythmogenic right Several >40% in ventricular desmosomal patients with cardiomyopathy genes. VTs unknown gene loci

2176 AF = atrial fibrillation.

2177

2178 13.2.1. Wolff–Parkinson–White syndrome

2179 Patients with pre-excitation and AF are at risk of rapid conduction across the accessory pathway, resulting in a 2180 fast ventricular rate, possibly ventricular fibrillation, and sudden death. In AF patients with evidence of an antegrade accessory pathway, catheter ablation of the pathway is recommended.^{869, 870} This procedure is safe and 2181 effective and may be considered as a prophylactic treatment strategy.^{871, 872} In AF patients surviving a sudden death event with evidence of an accessory pathway, urgent catheter ablation of the pathway is recommended.⁸⁶⁹ 2182 2183 2184 A documented short pre-excited RR interval (< 250 ms) during spontaneous or induced AF is one of the risk 2185 markers for sudden death in Wolff-Parkinson-White syndrome (WPW) syndrome, in addition to a history of 2186 symptomatic tachycardia, the presence of multiple accessory pathways, and Ebstein's anomaly. Intravenous procainamide, propafenone, or ajmaline can be used to acutely slow ventricular rate,^{873, 874} whereas digoxin, verapamil, and diltiazem are contraindicated.⁸⁷⁵ Intravenous amiodarone should be used with caution, as there 2187 2188 2189 are case reports of accelerated ventricular rhythms and ventricular fibrillation in patients with pre-excited AF 2190 receiving intravenous amiodarone infusion.8 2191

2192 13.2.2. Hypertrophic cardiomyopathy

2193 AF is the most common arrhythmia in patients with hypertrophic cardiomyopathy, affecting approximately onequarter of this population.⁸⁷⁷ Observational data highlight a high stroke risk in hypertrophic cardiomyopathy 2194

patients with AF, confirming the need for OAC.⁸⁷⁸ While there is more experience with VKAs, there are no data 2195 to suggest that NOACs cannot be used in these patients.⁸⁴⁵ Studies of rate or rhythm control medications in 2196 2197 patients with hypertrophic cardiomyopathy are relatively scarce. Beta-blockers and diltiazem or verapamil seem 2198 reasonable treatment options for rate control in these patients. In the absence of significant LV outflow tract obstruction, digoxin can be used alone or in combination with beta-blockers.⁸⁴⁵ Amiodarone seems a safe 2199 antiarrhythmic drug in AF patients with hypertrophic cardiomyopathy,⁸⁷⁹ and expert opinion suggests that 2200 disopyramide may be beneficial in those with outflow tract obstruction. AF ablation is effective to suppress symptomatic AF recurrences.⁸⁸⁰⁻⁸⁸⁴ Surgical treatment of AF may be appropriate in patients with hypertrophic 2201 2202 2203 cardiomyopathy undergoing surgery (e.g. for LV outflow tract obstruction or mitral valve surgery), but 2204 experience is limited.

2204 exj

2206 13.2.3. Channelopathies and arrhythmogenic right ventricular cardiomyopathy

2207 Many channelopathies and inherited cardiomyopathies are associated with AF. AF prevalence ranges from 5% 2208 to 20% in patients with long QT syndrome or Brugada syndrome, and is up to 70% in short QT syndrome (Table 20).^{853, 856-858} Penetrance of disease phenotype including AF is variable.^{61, 852, 885, 886} Both shortening as 2209 2210 well as prolongation of the atrial action potential have been demonstrated as likely mechanisms underlying AF 2211 in these diseases. It seems reasonable to consider antiarrhythmic drugs that reverse the suspected channel defect in AF patients with inherited cardiomyopathies (e.g. a sodium channel blocker in LQT3⁸⁵² and quinidine in 2212 2213 Brugada syndrome⁸⁸⁷). More importantly, new-onset AF in young, otherwise healthy individuals should trigger 2214 a careful search for such inherited conditions, including clinical history, family history, ECG phenotype, and 2215 echocardiography and/or other cardiac imaging.

2216 Monogenic defects only account for 3–5% of all patients with AF, even in younger populations.^{846, 848,} 2217 ⁸⁸⁸⁻⁸⁹⁰ Furthermore, there is no clear link between detected mutations and specific outcomes or therapeutic needs. 2218 For these reasons, genetic testing is not recommended in the general AF population.⁷⁷ Other guidelines have

2219 described the indications for genetic testing in patients with inherited arrhythmogenic diseases.^{844, 85}

2220

2221 **Recommendations for inherited cardiomyopathies** 2222

Recommendations	Class ^a	Level ^b	Refs ^c
WPW syndrome			
Catheter ablation of the accessory pathway in WPW patients with AF and rapid conduction over the accessory pathway is recommended to prevent sudden cardiac death	I	В	892-894
Catheter ablation of the accessory pathway is recommended without delay in WPW patients who survive sudden cardiac death	Ι	С	869
Asymptomatic patients with overt pre-excitation and AF should be considered for accessory pathway ablation after careful counselling	IIa	В	872, 895
Hypertrophic cardiomyopathy			
Lifelong oral anticoagulation to prevent stroke is recommended in hypertrophic cardiomyopathy patients who develop AF	Ι	В	878
Restoration of sinus rhythm by electrical or pharmacological cardioversion to improve symptoms is recommended in hypertrophic cardiomyopathy patients with symptomatic new-onset AF	I	В	845
In haemodynamically stable hypertrophic cardiomyopathy patients with AF, ventricular rate control using beta-blockers and diltiazem/verapamil is recommended	I	С	845
Treatment of LV outflow tract obstruction should be considered in AF patients with hypertrophic cardiomyopathy to improve symptoms	IIa	В	896
Amiodarone should be considered to achieve rhythm control and maintain sinus rhythm in hypertrophic cardiomyopathy patients	IIa	С	845, 897
Inherited cardiomyopathies and channelopathies			
Targeted genetic testing should be considered in patients with AF and a suspicion of inherited cardiomyopathies or channelopathies based on clinical history, family history, or electrocardiographic phenotype	IIa	А	852

AF = atrial fibrillation; LV = left ventricular; WPW = Wolff–Parkinson–White syndrome.

2224 ^aClass of recommendation.

2225 ^bLevel of evidence.

2226 ^cReference(s) supporting recommendations.

2227

2228 13.3. Sports and atrial fibrillation

2229 Physical activity improves cardiovascular health, which translates into a lower risk of AF.⁸⁹⁸ Therefore, physical activity is a cornerstone of preventing AF. Intensive sports practice, especially endurance sports (> 1500 h of endurance sports practice), ⁸⁹⁹ increases the risk of AF later in life, ⁹⁰⁰⁻⁹⁰² probably mediated by altered autonomic tone, volume load during exercise, atrial hypertrophy, and dilatation. ^{903, 904} This results in a U-shaped relationship of physical activity and incident AF. ^{214, 898, 902, 905, 906} Detraining can reduce AF in models⁹⁰⁴ and 2230 2231 2232 2233 reduces ventricular arrhythmias in athletes,⁹⁰⁷ but the role of detraining for AF in human athletes is unknown. 2234 2235 The management of athletes with AF is similar to general AF management, but requires a few special 2236 considerations. Clinical risk factors will determine the need for anticoagulation. Sports with direct bodily 2237 contact or prone to trauma should be avoided in patients on OAC. Beta-blockers are not well tolerated and at 2238 times prohibited, and digoxin, verapamil, and diltiazem are often not potent enough to slow heart rate during 2239 exertional AF. Catheter ablation for AF probably has similar outcomes in athletes as in non-athletes, ^{908, 909} but 2240 further data are needed. Pill-in-the-pocket therapy has been used as well.⁶²⁰ After ingestion of flecainide or 2241 propafenone as pill-in-the-pocket, patients should refrain from sports as long as AF persists and until two half-2242 lives of the antiarrhythmic drug have elapsed. Prophylactic ablation of the flutter circuit may be considered in 2243 athletes treated with sodium channel blockers.⁹

2244

2245 **Recommendations for physical activity in patients with AF**

Recommendations	Class ^a	Level ^b	Refs ^c
Moderate regular physical activity is recommended to prevent AF, while athletes should be counselled that long-lasting, more intense sports participation can promote AF	I	А	214, 898, 900- 902, 905, 906
AF ablation should be considered to prevent recurrent AF in athletes	IIa	В	908, 909
The ventricular rate while exercising with AF should be evaluated in every athlete (by symptoms and/or by monitoring), and titrated rate control should be instituted	IIa	С	
After ingestion of pill-in-the-pocket Class I antiarrhythmic drugs, patients should refrain from sports as long as AF persists and until two half-lives of the antiarrhythmic drug have elapsed	IIa	С	620

2246 AF = atrial fibrillation.

- 2247 ^aClass of recommendation.
- 2248 ^bLevel of evidence.
- 2249 ^cReference(s) supporting recommendations.
- 2250

2251 **13.4. Pregnancy**

AF in pregnant women is rare and is usually associated with pre-existing heart disease. AF is associated with increased complications for the mother and foetus.^{911, 912} Better treatment of congenital heart diseases will probably increase the incidence of AF during pregnancy in the future.⁹¹³ Pregnant women with AF should be managed as high-risk pregnancies in close collaboration with cardiologists, obstetricians, and neonatologists.

2256

2257 **13.4.1.** Rate control

2258 Owing to a lack of specific data, beta-blockers, verapamil, diltiazem, and digoxin all carry a US Food and Drug 2259 Administration pregnancy safety category of C (benefits may outweigh risk), except for atenolol (category D: 2260 positive evidence of risk). Their use should be at the lowest dose and for the shortest time required. None of the agents are teratogenic, but they readily cross the placenta.⁹¹⁴ Beta-blockers are commonly used in clinical 2261 2262 practice (e.g. for management of gestational hypertension and pre-eclampsia), but may be associated with intrauterine growth retardation,⁹¹⁵ and hence growth scans after 20 weeks gestation are recommended. Digoxin 2263 2264 is considered safe for maternal and foetal arrhythmias.⁹¹⁶ There are insufficient data to comment on verapamil or 2265 diltiazem, hence rate control using beta-blockers and/or digoxin is recommended.917 With regards to

breastfeeding, all rate control agents are present in breast milk, although levels of beta-blockers, digoxin, and

- 2267 verapamil are too low to be considered harmful. Diltiazem will be present at high levels and should be
- 2268 considered second-line treatment.⁹¹⁸
- 2269

2270 13.4.2. Rhythm control

Rhythm control therapy in pregnant patients with AF has only been reported in case studies. Amiodarone is associated with severe adverse foetal side-effects and should only be considered for emergency situations.⁹¹⁹ 2271

2272 Flecainide and sotalol can both be used for conversion of foetal arrhythmias without major adverse effects,⁹²⁰ 2273

2274 and thus are likely to be safe to treat maternal symptomatic AF. Electrical cardioversion can be effective for

2275 restoration of sinus rhythm when tachyarrhythmia is causing haemodynamic instability, with low rates of

2276 adverse outcomes for both mother and foetus.⁹²¹ However, in view of the risk of foetal distress, electrical 2277

cardioversion should only be carried out where facilities are available for foetal monitoring and emergency 2278 caesarean section. As with other emergencies during pregnancy, patients should receive 100% oxygen,

2279 intravenous access should be established early, and the mother should be positioned in the left lateral position to

2280 improve venous return.922 2281

2282 Anticoagulation 13.4.3.

2283 VKAs should be avoided in the first trimester because of teratogenic effects, and in the 2-4 weeks preceding 2284 delivery to avoid foetal bleeding. Low-molecular-weight heparins are a safe substitute, as they do not cross the 2285 placenta.⁹²³ In the third trimester, frequent laboratory checks for adequate anticoagulation (e.g. every 10-14 2286 days) and corresponding dose adjustments are advised, given that in some women high doses of both VKA and 2287 heparin may be needed to maintain adequate anticoagulation. Pregnant patients with AF and mechanical 2288 prosthetic valves who elect to stop VKA treatment in consultation with their specialist team between 6 and 12

2289 weeks of gestation, should receive continuous, dose-adjusted unfractionated heparin or dose-adjusted

2290 subcutaneous low-molecular-weight heparin. As only limited data are available about teratogenesis for NOACs, 2291 these drugs should be avoided during pregnancy.

2292

2293 Recommendations during pregnancy

Recommendations	Class ^a	Level ^b	Refs ^c
Electrical cardioversion can be performed safely at all stages of pregnancy, and is recommended in patients who are haemodynamically unstable due to AF, and whenever the risk of ongoing AF is considered high, for the mother or the foetus	I	С	
Anticoagulation is recommended in pregnant patients with AF at risk of stroke. To minimize teratogenic risk and intrauterine bleeding, dose-adjusted heparins are recommended during the first trimester of pregnancy and in the 2–4 weeks before delivery. Vitamin K antagonists or heparin can be used in the remaining parts of the pregnancy	I	В	923
NOACs should be avoided in pregnancy and in women planning a pregnancy	III (harm)	С	

- 2294 NOAC = non-vitamin K antagonist oral anticoagulants
- 2295 ^aClass of recommendation.
- 2296 ^bLevel of evidence.
- 2297 ^cReference(s) supporting recommendations.
- 2298

2299 **Postoperative atrial fibrillation** 13.5.

AF is common after cardiac surgery (occurring in 15–45% of patients),⁹²⁴⁻⁹²⁶ and is associated with increased length of hospital stay and higher rates of complications and mortality.⁹²⁷ Postoperative AF is also not 2300 2301

2302 uncommon after other major surgery, especially in elderly patients. The treatment of postoperative AF is mainly

2303 based on studies of patients undergoing cardiac surgery, with much less evidence in the non-cardiac surgery

- 2304 setting.
- 2305

Prevention of postoperative atrial fibrillation 2306 13.5.1.

2307 Beta-blockers reduce postoperative AF and supraventricular tachycardias, albeit with high heterogeneity and

2308 moderate risk of bias in a systematic review of published studies (the most commonly studied drug was

propranolol, with AF in 16.3% of the treatment group vs. 31.7% in the control group).⁹²⁵ In the majority of these 2309

2310 studies, beta-blockers were administered postoperatively, a regimen supported in a recent meta-analysis.⁹²⁸

2311 Amiodarone reduced the incidence of postoperative AF compared to a beta-blocker regimen in several meta-

analyses, also reducing hospital stay.^{925, 929-931} 2312

- Despite initial reports from meta-analyses,^{689, 932, 933} preoperative treatment with statins did not prevent postoperative AF in a prospective controlled trial.⁹³⁴ Other therapies have also been studied in small, hypothesisgenerating trials, but have not demonstrated clear beneficial effects. These include magnesium,^{925, 935, 936} n-3 polyunsaturated fatty acids,^{937,945} colchicine,⁹⁴⁶ corticosteroids,^{947, 948} and posterior pericardectomy.⁹⁴⁹ Postoperative overdrive biatrial pacing has not gained widespread use despite some suggestion of prophylactic
- 2318 effects.^{925, 950}
- 2319

2320 13.5.2. Anticoagulation

Postoperative AF is associated with an increased early stroke risk, increased morbidity, and 30-day mortality.⁹²⁷ 2321 2322 ^{951, 952} In the long term, patients with an episode of postoperative AF have a twofold increase in cardiovascular 2323 mortality and a substantially increased risk of future AF and ischaemic stroke compared with patients that remain in sinus rhythm after surgery.⁹⁵²⁻⁹⁵⁸ OAC at discharge has been associated with a reduced long-term 2324 mortality in patients with postoperative AF,⁹⁵⁹ without evidence from controlled trials. Good quality data are 2325 needed to determine whether long-term anticoagulation can prevent strokes in patients with postoperative AF at 2326 high stroke risk, $^{368, 386}$ and to assess whether short episodes of postoperative AF (e.g. < 48 h) carry a similar risk 2327 as longer episodes.⁹⁶⁰ The indication and timing of OAC in postoperative AF patients should take into 2328 2329 consideration the risk of postoperative bleeding. 2330

2331 **13.5.3.** Rhythm control therapy in postoperative atrial fibrillation

2332 In haemodynamically unstable patients, cardioversion and consideration of antiarrhythmic drugs is

2333 recommended. Amiodarone or vernakalant have been efficient in converting postoperative AF to sinus

2334 rhythm.^{603, 950, 961} A recent medium-sized trial randomizing patients with postoperative AF to either rhythm

2335 control therapy with amiodarone or to rate control did not find a difference in hospital admissions during a 60-

2336 day follow-up,⁹⁶² underpinning that the aim of rhythm control therapy should be to improve AF-related

symptoms in postoperative AF. In asymptomatic patients and in those with acceptable symptoms, rate control or

2338 deferred cardioversion preceded by anticoagulation is a reasonable approach.

2339

2340 Recommendations for preventing postoperative AF

Recommendations	Class ^a	Level ^b	Refs ^c
Perioperative oral beta-blocker therapy is recommended for the prevention of postoperative AF after cardiac surgery	I	В	925, 928
Restoration of sinus rhythm by electrical cardioversion or antiarrhythmic drugs is recommended in postoperative AF with haemodynamic instability	I	С	
Long-term anticoagulation should be considered in patients with AF after cardiac surgery at risk for stroke, considering individual stroke and bleeding risk	lla	В	368, 386
Antiarrhythmic drugs should be considered for recurrent or symptomatic postoperative AF after cardiac surgery in an attempt to restore sinus rhythm	lla	С	
Perioperative amiodarone should be considered for prophylactic therapy to prevent AF after cardiac surgery	lla	А	925
Intravenous vernakalant may be considered for cardioversion of postoperative AF in patients without severe heart failure, hypotension, or severe structural heart disease (especially aortic stenosis)	llb	В	603
Asymptomatic postoperative AF should initially be managed with rate control and anticoagulation	lla	В	962

2341 AF = atrial fibrillation.

- 2342 ^aClass of recommendation.
- 2343 ^bLevel of evidence.
- 2344 ^cReference(s) supporting recommendations.
- 2345

2346 13.6. Atrial arrhythmias in grown-up patients with congenital heart disease

Atrial arrhythmias (AF, atrial flutter, atrial tachycardias) often occur late after surgical repair of congenital heart defects, occurring in 15–40% of grown-up patients with congenital heart disease (GUCH). They are associated with heart failure, syncope, thromboembolic events, and sudden death.⁹⁶³⁻⁹⁶⁷ The pathophysiological substrate is complex, associated with hypertrophy, fibrosis, hypoxaemia, chronic haemodynamic overload, and surgical scars and patches. Additionally, related primary anomalies in the conduction pathways can lead to reentrant atrial and ventricular tachycardia, heart block, and sinus node dysfunction.⁹⁶³ Macro-reentrant atrial tachycardia or atypical atrial flutter may be seen after nearly any surgical procedure involving atriotomy or atrial patches.

2355 **13.6.1.** General management of atrial arrhythmias in grown-up patients with

2356 congenital heart disease

2357 The conventional stroke risk factors should be used to inform decisions on long-term anticoagulation in GUCH

patients with AF. In addition, anticoagulation should be considered in GUCH patients with atrial arrhythmias when they present with intracardial repair, cyanosis, Fontan palliation, or systemic right ventricle, in addition to those with conventional stroke risk factors.⁹⁶⁸ Beta-blockers, verapamil, diltiazem, and digitalis can be used. Care should be taken to avoid bradycardia and hypotension.

2362 Sodium channel blockers suppress approximately half of atrial arrhythmias in Fontan patients.⁹⁶⁹ 2363 Amiodarone is more effective, but long-term treatment with an antiarrhythmic drugs carries a high risk of 2364 extracardiac side-effects in this relatively young population. Intracardiac thrombi are common in GUCH 2365 patients undergoing cardioversion for AF, but also in patients with atrial tachycardias or atrial flutter.⁹⁷⁰ 2366 Therefore, both a TOE and anticoagulation for a few weeks before the planned cardioversion should be 2367 considered.⁹⁶⁴ Radiofrequency ablation may be a good option for symptomatic GUCH patients with atrial 2368 arrhythmias, especially in those with atrial flutter and other macro-reentrant tachycardias. Interventions should 2369 be performed in adequately qualified centres by specialized teams.

2370

2371 13.6.2. Atrial tachyarrhythmias and atrial septal defects

Atrial flutter and fibrillation occur in 14–22% of adults with unoperated atrial septal defects, especially in older patients,⁹⁷¹ and can lead to heart failure.⁹⁷² Early repair can reduce but not eliminate the risk of AF.⁹⁷³ Biatrial volume overload,⁹⁷⁴ pulmonary hypertension,⁹⁷⁵ and possibly the arrhythmogenic effect of atrial patches can contribute to these arrhythmias.⁹⁷⁶ Anticoagulation should be decided based on stroke risk factors. In patients with a history of paroxysmal or persistent AF, AF surgery could be considered at the time of surgical closure, or catheter ablation in patients undergoing interventional atrial septal defect closure. Catheter ablation of late atrial arrhythmias has shown to be effective in 46 consecutive patients after surgical atrial septal defect.⁹⁷⁷

2380 **13.6.3.** Atrial tachyarrhythmias after Fontan operation

2381 Atrial arrhythmias occur in up to 40% of patients with a Fontan circulation, and can manifest as atrial flutter, 2382 primary atrial tachycardia, AF, and accelerated junctional rhythm or junctional tachycardia⁹⁷⁸ with or without sinoatrial node dysfunction.⁹⁷⁹ Patients with atriopulmonary anastomoses (possibly due to higher atrial volume 2383 2384 and pressure load) and those with early postoperative atrial arrhythmias are more likely to develop long-term 2385 atrial arrhythmias.⁹⁸⁰ Atrial arrhythmias can also be the first manifestation of obstruction of the atriopulmonary 2386 anastomosis, a complication that must be identified. Right atrial thrombus formation is common in Fontan patients with atrial arrhythmias and requires oral anticoagulation.⁹⁸¹ Operative conversion to total 2387 cavopulmonary artery connection with concomitant arrhythmia surgery can in some patients improve heart failure symptoms and reduce recurrent arrhythmias,^{969, 982} with low recurrence rates of clinically apparent atrial 2388 2389 arrhythmias in the first few years after repeat surgery.⁹⁸³⁻⁹⁸⁵ Catheter ablation of atrial arrhythmia in Fontan 2390 2391 patients has been successful in selected patients.⁹ 2392

2393 13.6.4. Atrial tachyarrhythmias after tetralogy of Fallot correction

Approximately one-third of patients after repair of tetralogy of Fallot develop atrial arrhythmias, including intraatrial reentrant tachycardia, focal atrial tachycardia, and AF.⁹⁸⁷ Circuits involving the cavotricuspid isthmus and areas of presumed surgical right atrial scaring have been described as responsible for atrial arrhythmias.

2397

2398 **Recommendations in patients with GUCH**

Recommendations	Class ^a	Level ^b	Refs ^c
Atrial septal defect closure should be considered before the fourth decade of life to diminish the chance of atrial flutter and fibrillation	lla	С	971, 972, 974

In patients who need surgical closure of an atrial septal defect and who have a history of symptomatic atrial arrhythmia, atrial ablation should be considered at the time of surgical closure	lla	С	204, 988, 989
Cox maze surgery should be considered in patients with symptomatic AF and an indication for corrective repair of congenital heart defects. All such surgery should be done in experienced centres	lla	С	988, 990
Oral anticoagualtion should be considered in all adult patients with intracardiac repair, cyanosis, Fontan palliation, or systemic right ventricle and a history of AF, atrial flutter, or intra-atrial reentrant tachycardia. In all other congenital heart disease patients with AF, anticoagulation should be considered if the CHA ₂ DS ₂ - VAS _C score is ≥ 1	lla	С	968
Catheter ablation of atrial arrhythmias associated with congenital heart defects may be considered when performed in experienced centres	llb	С	991
In patients with congenital heart disease, transoesophageal echocardiography may be considered together with 3-week anticoagulation therapy before cardioversion	llb	С	964, 970, 988, 990

- 2399 AF = atrial fibrillation; CHA₂DS₂-VASc = Congestive Heart failure, hypertension, Age \geq 75 (doubled),
- 2400 Diabetes, Stroke (doubled), Vascular disease, Age 65–74, and Sex (female); GUCH = grown-up patients with 2401 congenital heart disease; OAC = oral anticoagulation; TOE = transoesophageal echocardiography.
- 2402 ^aClass of recommendation.
- 2403 ^bLevel of evidence.
- 2404 ^cReference(s) supporting recommendations.
- 2405

2406 13.7. **Management of atrial flutter**

The goals for the management of atrial flutter are similar to those for AF.⁹⁹² Based on the available evidence, the 2407 stroke risk in patients with atrial flutter is not much different from that in AF.⁸²⁷ Furthermore, many patients diagnosed with atrial flutter develop AF.⁹⁹³⁻⁹⁹⁵ Thus, anticoagulation should be used in patients with atrial flutter 2408 2409 2410 similar to that in patients with AF. Rate control in atrial flutter is achieved with the same medications as in AF, 2411 but is often more difficult to achieve. Flecainide, propafenone, dofetilide, and intravenous ibutilide are useful for 2412 cardioversion of atrial flutter. They should be combined with a rate-controlling agent to avoid 1:1 conduction of slowing flutter waves to the ventricles. Ibutilide is more effective for conversion of atrial flutter than AF, whereas vernakalant is less effective in converting typical atrial flutter.^{996, 997} Electrical cardioversion of atrial flutter can be performed using lower energies (50–100 J) than for AF.^{998, 999} Atrial overdrive pacing through pacemaker leads or endocardial or transesophageal catheters can convert atrial flutter to sinus rhythm.^{1000, 1001} 2413 2414 2415

- 2416
- 2417 Anticoagulation and transoesophageal echocardiography around cardioversion or overdrive pacing should be 2418 used similar to that in AF.
- 2419 Ablation of the cavotricuspid isthmus for isthmus-dependent right atrial flutter (either the common 2420 counter-clockwise atrial flutter or the less-common clockwise atrial flutter) restores and maintains sinus rhythm with a success rate of 90–95%.¹⁰⁰² It may also reduce AF recurrences in selected patients,^{1003, 1004} and help to prevent hospitalizations.^{1004, 1005} Isthmus ablation is comparably safe and more effective than antiarrhythmic drug therapy, and is recommended for recurrent atrial flutter.^{585-587, 713} Catheter ablation of left atrial macro-2421 2422 2423 reentrant tachycardia is more complex, with lower success rates and higher recurrence rates.^{1006, 100} 2424
- 2425

2426 Recommendations for management of atrial flutter

Recommendations	Class ^a	Level ^b	Ref s ^c
For patients with atrial flutter, antithrombotic therapy is recommended according to the same risk profile used for AF	Ι	В	827
Overdrive atrial pacing of atrial flutter should be considered as an alternative to electrical cardioversion, depending on local availability and experience	IIa	В	1000, 1001

Management of typical atrial flutter with ablation of the cavotricuspid isthmus is recommended for patients failing antiarrhythmic drug therapy or as first-line treatment considering patient preference	Ι	В	158
If atrial flutter has been documented before AF ablation, ablation of the cavotricuspid isthmus should be considered as part of the AF ablation procedure	IIa	С	

2427 AF = atrial fibrillation.

- 2428 ^aClass of recommendation.
- 2429 ^bLevel of evidence.
- 2430 ^cReference(s) supporting recommendations.
- 2431

2432 14 Patient involvement, education and self-management

- 2433 A fundamental aspect of a structured AF management programme is the focus on patient-centred care.
- 2434

2435 **14.1**. **Patient-centred care**

2436 Autonomous, informed patients are better placed to adhere to long-term therapy, and it is very likely that long-2437 term management of chronic conditions such as AF will benefit from informed patients involved in the disease

management who are aware of their own responsibilities.³²⁸ Shared decision-making⁷⁴⁷ and patient-centred 2438

organization of care can help to ensure adherence to management and empower patients, and respect individual patient preferences, needs, and values (see Chapter 7.2).^{326, 1008, 1009} Patients in an active role tend to have better 2439 2440

2441 health outcomes and care experiences, and engagement itself can be considered as an intermediate outcome,

- 2442 particularly where related to improved clinical outcomes.¹⁰¹⁰
- 2443

2444 14.2. **Integrated patient education**

2445 Education is a prerequisite for informed, involved patients and patient-centred care. However, lack of AFrelated knowledge in patients is common, even in those who have received verbal and written information,^{32, 1011,} 2446

¹⁰¹² indicating the need to further develop structured patient education. Several patient-information tools have been developed, largely focusing on oral anticoagulation.¹⁰¹³⁻¹⁰¹⁶ Understanding patients' perceptions and 2447

2448

attitudes towards AF and its management can improve AF management and related outcomes.¹⁰¹⁷ This includes 2449 tailored patient education focusing on the disease, symptom recognition, therapy, modifiable risk factors for AF, and self-management activities.^{1018, 1019}

- 2450 2451
- 2452

2453 14.3. Self-management and shared decision-making

2454 Self-management is primarily focused on tasks to manage the condition, such as adhering to a therapeutic

regimen or modifying behaviour (e.g. resulting in smoking cessation or weight loss).¹⁰²⁰ It requires 2455

understanding of the treatment modalities and goals.³⁵⁰ Within a multidisciplinary team, allied health 2456

2457 professionals can guide this interactive process in which communication, trust, and reciprocal respect foster

patient engagement.¹⁰²¹ Shared decision-making should be considered as a routine part of the decision-making process,⁷⁴⁷ supported by decision aids where applicable.¹⁰²² Models of care that integrate education, engagement, and shared decision making are now available,¹⁰²³ and may be of particular value in the 2458

2459

2460 2461 management of AF.

2462

2463 Recommendations for patient involvement, education, and self-management

Recommendations	Class ^a	Level ^b	Refs ^c
Tailored patient education is recommended in all phases of AF management to support patients' perception of AF and to improve management	I	С	1014, 1017
Patient involvement in the care process should be considered to encourage self-management and responsibility for lifestyle changes	lla	С	328, 1010
Shared decision-making should be considered to ensure that care is based on the best available evidence and fits the needs, values, and preferences of the patient	lla	С	747

2464 AF = atrial fibrillation.

2465 ^aClass of recommendation.

2466 ^bLevel of evidence.

- 2467 ^cReference(s) supporting recommendations.
- 2468

2469 15 Gaps in evidence

2470 There are some areas of AF management that are supported by excellent evidence from multiple, adequately

2471 powered randomized trials (e.g. oral anticoagulation. Other areas, such as rhythm control therapy, integrated AF

2472 management, and lifestyle modifications are clearly developing the required evidence, while areas such as rate control are in dire need of better studies to underpin future guidelines. Here we identify areas in need of further

- 2473 2474 research.
- 2475

2476 Major health modifiers causing atrial fibrillation 15.1.

- Atrial fibrillation has different causes in different patients. More research is needed into the major causes (and electrophysiological mechanisms) of AF in different patient groups.^{176, 1024} Such research should consider the 2477
- 2478
- 2479 major comorbidities associated with AF, and characterize the response to AF therapy in patients with different, pathophysiologically distinct types of AF.
- 2480 2481

2482 15.2. How much atrial fibrillation constitutes a mandate for therapy?

2483 Technological advances allow screening for an irregular pulse using patient-operated ECG devices,

2484 smartphones, and a variety of other technologies. These may be very useful to detect silent, undiagnosed AF.¹⁵⁷

- 2485 Adequately powered studies evaluating the diagnostic accuracy of such technologies, the diagnostic yield in
- 2486 different populations, the shortest duration of atrial arrhythmias conveying a stroke risk, and ideally the effect of
- 2487 ECG screening on outcomes are needed.
- 2488

2489 15.3. Atrial high-rate episodes and need for anticoagulation

2490 All of the information on the benefit of OAC has been in patients with AF diagnosed by ECG. Technological 2491 advances allow ready detection of AHRE in patients with implanted devices and an atrial lead. Such patients are 2492 at increased stroke risk, but it is unclear whether they benefit from OAC. Controlled trials evaluating OAC in 2493 AHRE patients are ongoing and will provide evidence on the best antithrombotic therapy in these patients.

2494

2495 15.4. Stroke risk in specific populations

2496 Several specific AF groups should be studied to better characterize their risk for AF, stroke, and other AF-2497 related complications (e.g. patients with one stroke risk factor, and non-Caucasian patients). Confounding 2498 factors (e.g. different therapy of concomitant cardiovascular diseases) may help to explain the variability in the 2499 reported rates of incident AF, prevalent AF, and AF complications. This also applies to the effect of gender in 2500 AF patients.47 2501

2502 15.5. Anticoagulation in patients with severe chronic kidney disease

2503 The use of NOACs has not been tested in patients with creatinine clearance < 30 mL/min, and there is very little 2504 evidence on the effects of OAC in patients on haemodialysis or on other forms of renal-replacement therapy. 2505 Studies evaluating OAC in patients with severe chronic kidney disease are needed to inform the best

- 2506 management in this patient group at high risk for stroke and bleeding.
- 2507

2508 15.6. Left atrial appendage occlusion for stroke prevention

2509 The most common justification for LAA occlusion devices in clinical practice is a perceived high bleeding risk

and, less often, contraindications for OAC.⁴⁵⁹ Unfortunately, LAA occluders have not been tested in such 2510

2511 populations. Furthermore, LAA occluders have not been compared with NOAC therapy in patients at risk for

- 2512 bleeding, or with thoracoscopic LAA clipping. There is a clear need to conduct adequately designed and 2513 powered trials to define the clinical role of LAA occluders compared with NOAC therapy in patients with
- 2514 relative or absolute contraindications for anticoagulation, and/or in those suffering from an ischaemic stroke on
- 2515 anticoagulant therapy.
- 2516

2517 Anticoagulation in atrial fibrillation patients after a bleeding or stroke event 15.7.

2518 At least 2% of anticoagulated patients with AF will experience a serious bleeding event per year. Observational

- data suggest that OAC can be reinitiated even after an intracerebral bleeding event. 460, 484 Controlled studies 2519 2520 evaluating different anticoagulation and stroke prevention interventions are urgently needed to provide evidence
- 2521 on the best management of patients who have suffered a bleeding event that would usually lead to withholding

2522 OAC. Some studies (e.g. APACHE II¹⁰²⁵) are ongoing, but adequately powered trials are needed. Similarly,

- prospectively collected data are needed on the efficacy and bleeding risk following (re-)initiation of OAC after stroke or intracranial bleeding.
- 2525

2526 15.8. Anticoagulation and optimal timing of non-acute cardioversion

Based on retrospective data, previous recommendations on the safe time-window in which a cardioversion can be performed in new-onset AF used \leq 48 hours as the 'gold standard' for non-protected cardioversion. However, new evidence has emerged that initiating precardioversion anticoagulation in patients with AF episodes of < 24 hours or even < 12 hours would provide even better safety.^{642, 647, 1026-1028} Further research is needed to establish a clear safety margin in this clinical situation.

2532

2533 15.9. Competing causes of stroke or transient ischaemic attack in atrial fibrillationpatients

Prospective RCTs have demonstrated the superiority of carotid endarterectomy compared to stenting in patients with symptomatic high-degree stenosis of the internal carotid artery.¹⁰²⁹ As endartectomy minimizes the need for combination therapy with OAC and antiplatelets,¹⁰³⁰ this approach has appeal in patients with AF to reduce bleeding risk. However, few of these studies included patients with AF. In a large observational study, the composite of in-hospital mortality, post-procedural stroke, and cardiac complications was higher in AF patients undergoing carotid stenting (457/7668; 6.0%) compared with endarterectomy (4438/51320; 8.6%; *P* <

2541 0.0001).¹⁰³¹ Despite adjustment for baseline risk, this may just reflect the type of patients referred for each

procedure, and further randomized studies are needed to confirm the optimal treatment strategy in AF patients with carotid disease.

2543

254515.10. Anticoagulation in patients with biological heart valves (including transcatheter2546aortic valve implantation) and non-rheumatic valve disease

2547 The optimal antithrombotic therapy in the first months after biological valve replacement (including after 2548 catheter-based valve replacement) is not known. VKAs remain the mainstay during the initial postoperative 2549 period; NOACs probably deliver the same protection. In patients without AF, many centres use platelet 2550 inhibitors only. NOACs appear to be equally effective as VKAs in patients with moderate aortic stenosis, based 2551 on a subanalysis from the ROCKET-AF (Rivaroxaban Once Daily Oral Direct Factor Xa Inhibition Compared with Vitamin K Antagonism for Prevention of Stroke and Embolism Trial in Atrial Fibrillation) trial¹⁰³² as well as the Loire Valley AF project.¹⁰³³ Further data would be helpful to confirm these observations.¹⁰³⁴ The safety 2552 2553 2554 and efficacy of NOACs in patients with rheumatic mitral valve disease has not been evaluated and should be 2555 studied.

2556

2557 15.11. Anticoagulation after 'successful' catheter ablation

2558 In view of the long-term recurrence rates of AF, this Task Force recommends to continue OAC in AF patients 2559 after 'successful' catheter ablation. Nonetheless, observational data suggest that the stroke risk may be lower

2560 after catheter ablation of AF compared with other AF patients. The ongoing EAST (Early treatment of Atrial

2561 fibrillation for Stroke prevention Trial) trial will inform in a more general way whether rhythm control therapy

2562 can reduce stroke rates in anticoagulated AF patients. If confirmed, there may be a place for a controlled trial

- 2563 evaluating the termination of OAC therapy at an interval after 'successful' catheter ablation.
- 2564

2565 15.12. Comparison of rate control agents

Although the use of rate control therapy is very common in AF patients, robust data comparing rate control therapies are scant, with the majority of studies being small uncontrolled trials over short periods of follow up

therapies are scant, with the majority of studies being small uncontrolled trials over short periods of follow-up. Some studies are funded (e.g. RATE-AF [Rate Control Therapy Evaluation in Permanent Atrial Fibrillation]⁵⁵⁹ and will investigate the potential benefits of different rate controlling agents, characteristics, or biomarkers that can help to personalize the use of rate control, and the adverse-event profile of specific drugs in defined groups

- 2571 of patients (e.g. AF with HFrEF).
- 2572

2573 15.13. Catheter ablation in persistent and long-standing persistent AF

2574 While a few recent randomized studies support the use of catheter or surgical ablation in patients with persistent

2575 AF and long-standing persistent AF, there is a clear need for more data evaluating this intervention in

2576 adequately powered randomized trials.

2577

2578 15.14. Optimal technique for repeat catheter ablation

PVI emerges as the most important target for catheter ablation of AF. Although a plethora of different additional ablation techniques have been published, their added value is questionable in patients undergoing a first catheter ablation, including those with persistent AF.⁷³⁵ Many patients are in need of multiple catheter-ablation procedures, and such interventions often follow local or operator-specific protocols without clear evidence to support the choice of ablation target or intervention. There is a clear clinical need to define the best approach in

- 2584 patients who are in need of a second ablation procedure.
- 2585

2586 15.15. Combination therapy for maintenance of sinus rhythm

2587 In the follow-up after initially successful catheter ablation, even when done in experienced centres, many

- 2588 patients will experience symptomatic recurrences of AF. These patients are often managed with antiarrhythmic
- 2589 drugs. There is a surprising paucity of data evaluating different rhythm control interventions in patients with
- 2590 recurrent AF after catheter ablation. Such studies seem reasonable and feasible.
- 2591

2592 15.16. Can rhythm control therapy convey a prognostic benefit in atrial fibrillationpatients?

2594 The progress in rhythm control therapy (catheter ablation, new antiarrhythmic drugs) and observational long-

2595 term analyses suggest that rhythm control therapy may have a prognostic benefit. Ongoing trials such as

2596 CABANA and EAST - AFNET 4 will provide initial answers to this important question, but more data are

- 2597 needed, in addition to trials of surgical ablation techniques.
- 2598

2599 15.17. Thoracoscopic 'stand-alone' atrial fibrillation surgery

2600 Minimally invasive epicardial ablation surgery for the treatment of stand-alone AF was reported a decade 2601 ago.¹⁰³⁵ The procedure has since evolved towards a totally thoracoscopic procedure,¹⁰³⁶ and lesion sets were

2602 extended to a complete left atrial maze.⁸²² With such rapid development and the coexistence of different

2603 techniques and lesion sets, scientific evidence on long-term results is still limited. Randomized trials using a

- 2604 standardized procedure are urgently needed to clearly define the benefits and risks of thoracoscopic AF ablation,
- 2605 and to further support decisions of the AF Heart Team.
- 2606

2607 15.18. Surgical exclusion of the left atrial appendage

2608 Exclusion of the LAA has been performed by cardiothoracic surgeons for decades, but prospective randomized

2609 studies comparing the rate of ischaemic stroke with or without left appendage exclusion are presently lacking.

2610 The LAAOS (Left Atrial Appendage Occlusion Study) III is currently randomizing cardiac surgery patients with

2611 AF to undergo concomitant occlusion or no occlusion of the appendage.⁴⁶⁷ More data are also needed to confirm

the safety and efficacy of thoracoscopic exclusion, following early positive observational data.¹⁰³⁷

2613

2614 15.19. Concomitant atrial fibrillation surgery

Adequately powered randomized trials are needed, employing systematic follow-up, uniform lesion sets and energy sources to evaluate the benefits and risks of concomitant AF surgery in symptomatic AF patients. An

2010 energy sources to evaluate the benefits and fisks of concomitant AF surgery in symptomatic AF patients. An 2617 RCT on non-uniform lesion sets with long-term follow-up is due to publish shortly.¹⁰³⁸ These will assist the AF

2617 Ker on non-uniform resion sets with long-term ronow-up is due to publish shortly. These will assist the AP 2618 Heart Team to decide on optimal therapy for individual patients, including the full repertoire of medical and

- 2619 surgical options for the treatment of AF.
- 2620

2621 16 To do and not to do messages from the Guidelines

Recommendations for diagnosis and screening of AF	Class	Level
ECG documentation is required to establish the diagnosis of AF	Ι	В
Opportunistic screening for AF is recommended by pulse taking or ECG rhythm strip in patients > 65 years of age	Ι	В
In patients with TIA or ischaemic stroke, screening for AF is recommended by short-term ECG recording followed by continuous ECG monitoring for at least 72 hours	I	В
It is recommended to interrogate pacemakers and ICDs on a regular basis for atrial high rate episodes (AHRE). Patients with AHRE should undergo further ECG monitoring to document AF before initiating AF therapy	Ι	В
Recommendations for general management of AF	Class	Level
Tailored patient education is recommended in all phases of AF management to support patients' perception of AF and to improve management	Ι	С
A full cardiovascular evaluation, including an accurate history, careful clinical examination, and assessment of concomitant conditions, is recommended in all AF patients	Ι	С
Use of the modified EHRA symptom scale is recommended in clinical practice and research studies to quantify AF-related symptoms	Ι	С
Transthoracic echocardiography is recommended in all AF patients to guide management	Ι	C
The assessment of kidney function by serum creatinine or creatinine clearance is recommended in all AF patients to detect kidney disease and to support correct dosing of AF therapy	Ι	А
Recommendations for stroke prevention in AF	Class	Level
		Lever
The CHA ₂ DS ₂ -VASc score is recommended for stroke risk prediction in patients with AF	I	A
with AF Oral anticoagulation therapy to prevent thromboembolism is recommended for	I	
		A
with AF Oral anticoagulation therapy to prevent thromboembolism is recommended for all male AF patients with a CHA ₂ DS ₂ -VASc score of 2 or more Oral anticoagulation therapy to prevent thromboembolism is recommended in all	I	A
with AF Oral anticoagulation therapy to prevent thromboembolism is recommended for all male AF patients with a CHA ₂ DS ₂ -VASc score of 2 or more Oral anticoagulation therapy to prevent thromboembolism is recommended in all female AF patients with a CHA ₂ DS ₂ -VASc score of 3 or more When oral anticoagulation is initiated in a patient with AF who is eligible for a non vitamin-K-antagonist oral anticoagulant (NOAC, apixaban, dabigatran, edoxaban, or rivaroxaban), a NOAC is recommended in preference to a Vitamin K antagonist Vitamin K antagonist therapy (INR 2.0–3.0 or higher) is recommended for stroke prevention in AF patients with moderate-to-severe mitral stenosis or mechanical	I	A A A
with AF Oral anticoagulation therapy to prevent thromboembolism is recommended for all male AF patients with a CHA ₂ DS ₂ -VASc score of 2 or more Oral anticoagulation therapy to prevent thromboembolism is recommended in all female AF patients with a CHA ₂ DS ₂ -VASc score of 3 or more When oral anticoagulation is initiated in a patient with AF who is eligible for a non vitamin-K-antagonist oral anticoagulant (NOAC, apixaban, dabigatran, edoxaban, or rivaroxaban), a NOAC is recommended in preference to a Vitamin K antagonist Vitamin K antagonist therapy (INR 2.0–3.0 or higher) is recommended for stroke prevention in AF patients with moderate-to-severe mitral stenosis or mechanical heart valves NOACs (apixaban, dabigatran, edoxaban, and rivaroxaban) are not recommended in patients with mechanical heart valves (Level of evidence B) or moderate-to-	I I I	A A A A
with AF Oral anticoagulation therapy to prevent thromboembolism is recommended for all male AF patients with a CHA ₂ DS ₂ -VASc score of 2 or more Oral anticoagulation therapy to prevent thromboembolism is recommended in all female AF patients with a CHA ₂ DS ₂ -VASc score of 3 or more When oral anticoagulation is initiated in a patient with AF who is eligible for a non vitamin-K-antagonist oral anticoagulant (NOAC, apixaban, dabigatran, edoxaban, or rivaroxaban), a NOAC is recommended in preference to a Vitamin	I I I I	A A A A B

inhibition		
In male or female AF patients without additional stroke risk factors, anticoagulant or antiplatelet therapy is not recommended for stroke prevention	III (harm)	В
Antiplatelet monotherapy is not recommended for stroke prevention in AF patients, regardless of stroke risk	III (harm)	А
After surgical occlusion or exclusion of the left atrial appendage, it is recommended to continue anticoagulation in at-risk patients with AF for stroke prevention	Ι	В
Genetic testing before the initiation of vitamin K antagonist therapy is not recommended.	III (no benefit)	В
In AF patients with severe active bleeding events, it is recommended to interrupt oral anticoagulation therapy until the underlying cause is resolved	Ι	С
NOACs should be avoided in pregnancy and in women planning a pregnancy	III (harm)	С
For patients with atrial flutter, antithrombotic therapy is recommended according to the same risk profile used for AF	Ι	В
Management of typical atrial flutter with ablation of the cavotricuspid isthmus is recommended for patients failing antiarrhythmic drug therapy or as first-line treatment considering patient preference	I	В
Lifelong oral anticoagulation to prevent stroke is recommended in hypertrophic cardiomyopathy patients who develop AF	Ι	В
Anticoagulation with heparin or low-molecular-weight heparin immediately after ischaemic stroke is not recommended in AF patients	III (harm)	А
Systemic thrombolysis with a recombinant tissue plasminogen activator is not recommended if the INR is above 1.7 (or, for patients on dabigatran, if activated partial thromboplastin time is outside the normal range)	III (harm)	С
After TIA or stroke, combination therapy of OAC and an antiplatelet is not recommended	III (harm)	В
Recommendations for rate control of AF	Class	Level
Beta-blocker, digoxin, diltiazem, or verapamil is recommended to control heart rate in AF patients with LVEF $\geq 40\%$	Ι	В
Beta-blocker and/or digoxin is recommended to control heart rate in AF patients with LVEF $<40\%$	Ι	В
In patients with permanent AF (i.e. where no attempt to restore sinus rhythm is planned), antiarrhythmic drugs should not routinely be used for rate control	III (harm)	А
Recommendations for rhythm control of AF	Class	Level
Rhythm control therapy is indicated for symptom improvement in patients with AF	Ι	В
Cardioversion of AF (either electrical or pharmacological) is recommended in symptomatic patients with persistent or long-standing persistent AF as part of rhythm control therapy	I	В
In patients with no history of ischaemic or structural heart disease, flecainide,		А

Ι	А
Ι	В
Ι	В
I	А
I	А
Ι	А
I	В
III (harm)	С
Ι	A
III (no benefit)	В
I	А
	I I I I I I III (harm) I III (no benefit)

2623 2624 2625

2626 2627	17 A	short summary of the management of AF patients
2628 2629 2630		e provide 17 simple rules to guide diagnosis and management of AF patients according to the 2016 CTS/ESO Guidelines for the management of atrial fibrillation
2630 2631 2632	1.	Use ECG screening in at risk populations for atrial fibrillation, especially stroke survivors and the Elderly.
2633	2.	Document AF by ECG before starting treatment.
2634 2635	3.	Evaluate all AF patients by clinical evaluation, ECG, and echocardiogram for underlying cardiovascular conditions such as hypertension, heart failure, valvular heart disease, and others.
2636 2637	4.	Provide tailored information and education to AF patients to empower them to support AF management.
2638	5.	Propose life style changes to all suitable AF patients to make their management more effective.
2639 2640 2641	6.	Treat underlying cardiovascular conditions adequately, e.g. valve repair or replacement in AF patients with significant valvular heart disease, treatment of heart failure, or management of hypertension, among others.
2642 2643	7.	Use oral anticoagulation in all AF patients unless they are at low risk for stroke based on the CHA_2DS_2VASc score or have true contraindications for anticoagulant therapy.
2644 2645	8.	Anticoagulate patients with atrial flutter similar to atrial fibrillation. Offer isthmus ablation to symptomatic flutter patients.
2646 2647 2648 2649	9.	Reduce all modifiable bleeding risk factors in all AF patients on oral anticoagulation, e.g. by treating hypertension, minimising the duration and intensity of concomitant antiplatelet and NSAID therapy, treating anaemia and eliminating causes for blood loss, maintaining stable INR values in patients on vitamin K antagonists, and moderating alcohol intake
2650 2651	10.	Check ventricular rate in all AF patients and use rate control medications to achieve lenient rate control.
2652 2653 2654	11.	Evaluate AF-related symptoms in all AF patients using the modified EHRA score. Whenever patients have AF-related symptoms, aim to improve symptoms by adjustment of rate control therapy and by offering antiarrhythmic drugs, cardioversion, or catheter or surgical ablation.
2655 2656	12.	Select antiarrhythmic drugs based on their safety profile and consider catheter or surgical ablation when antiarrhythmic drugs fail.
2657 2658	13.	Do not offer routine genetic testing in AF patients unless there is a suspicion for an inherited cardiac condition.
2659	14.	Do not use antiplatelet therapy for stroke prevention in AF.
2660 2661	15.	Do not permanently discontinue oral anticoagulation in AF patients at increased risk of stroke unless such a decision is taken by a multidisciplinary team.
2662 2663	16.	Do neither use rhythm control therapy in asymptomatic AF patients, nor in patients with permanent AF.
2664 2665	17.	Do not perform cardioversion or catheter ablation without anticoagulation unless an atrial thrombus has been ruled out by transesophageal echocardiogram.
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2669 18 Web Addenda

All Web figures and Web tables are available in the Web addenda, available at European Heart Journal online
and also via the ESC Website (www.escardio.org/guidelines).

2673 **19 Appendix**

2674 ESC Committee for Practice Guidelines (CPG): Jose Luis Zamorano (Chairperson) (Spain), Victor Aboyans 2675 (France), Stephan Achenbach (Germany), Stefan Agewall (Norway), Lina Badimon (Spain), Gonzalo Barón-2676 Esquivias (Spain), Helmut Baumgartner (Germany), Jeroen J. Bax (The Netherlands), Héctor Bueno (Spain), 2677 Scipione Carerj (Italy), Veronica Dean (France), Çetin Erol (Turkey), Donna Fitzsimons (UK), Oliver 2678 Gaemperli (Switzerland), Paulus Kirchhof (UK/Germany), Philippe Kolh (Belgium), Patrizio Lancellotti 2679 (Belgium), Gregory Y. H. Lip (UK), Petros Nihoyannopoulos (UK), Massimo F. Piepoli (Italy), Piotr 2680 Ponikowski (Poland), Marco Roffi (Switzerland), Adam Torbicki (Poland), António Vaz Carneiro (Portugal), 2681 Stephan Windecker (Switzerland). 26822683 ESC National Cardiac Societies actively involved in the review process of the 2016 ESC Guidelines for the 2684 management of atrial fibrillation developed in collaboration with EACTS 2685 2686 Armenia: Armenian Cardiologists Association, Hamlet G. Hayrapetyan; Austria: Austrian Society of 2687 Cardiology, Franz Xaver Roithinger; Azerbaijan: Azerbaijan Society of Cardiology, Farid Aliyev; Belarus: 2688 Belorussian Scientific Society of Cardiologists, Alexandr Chasnoits; Belgium: Belgian Society of Cardiology, 2689 Georges H. Mairesse; Bosnia and Herzegovina: Association of Cardiologists of Bosnia and Herzegovina, 2690 Daniela Loncar Matičević; Bulgaria: Bulgarian Society of Cardiology, Tchavdar Shalganov; Croatia: Croatian 2691 Cardiac Society, Boško Skorić; Cyprus: Cyprus Society of Cardiology, Loizos Antoniades; Czech Republic: 2692 Czech Society of Cardiology, Milos Taborsky; Denmark: Danish Society of Cardiology, Steen Pehrson; 2693 Egypt: Egyptian Society of Cardiology, Said Khaled; Estonia: Estonian Society of Cardiology, Priit Kampus; 2694 Finland: Finnish Cardiac Society, Antti Hedman; The Former Yugoslav Republic of Macedonia: 2695 Macedonian FYR Society of Cardiology, Lidija Poposka; France: French Society of Cardiology, Jean-Yves Le 2696 Heuzey; Georgia: Georgian Society of Cardiology, Kakhaber Estadashvili; Germany: German Cardiac 2697 Society, Dietmar Bänsch; Hungary: Hungarian Society of Cardiology, Zoltán Csanádi; Icelandi: Icelandic 2698 Society of Cardiology, David O. Arnar; Ireland: Irish Cardiac Society, David Keane; Israel: Israel Heart 2699 Society, Roy Beinart; Italy: Italian Federation of Cardiology, Francesco Romeo; Kazakhstan: Association of 2700 Cardiologists of Kazakhstan, Kulzida Koshumbayeva; Kosovo: Kosovo Society of Cardiology, Gani Bajraktari; 2701 **Kyrgyzstan:** Kyrgyz Society of Cardiology, Aibek Mirrakhimov, **Latvia:** Latvian Society of Cardiology, 2702 Oskars Kalejs; Lebanon: Lebanese Society of Cardiology, Samer Nasr; Lithuania: Lithuanian Society of 2703 Cardiology, Germanas Marinskis; Luxembourg: Luxembourg Society of Cardiology, Carlo Dimmer; Malta: 2704 Maltese Cardiac Society, Mark Sammut; Moldova: Moldavian Society of Cardiology, Aurel Grosu; Morocco: 2705 Moroccan Society of Cardiology, Salima Abdelali; The Netherlands: Netherlands Society of Cardiology, 2706 Martin E. W. Hemels; Norway: Norwegian Society of Cardiology, Ole-Gunnar Anfinsen; Poland: Polish 2707 Cardiac Society, Beata Sredniawa; Portugal: Portuguese Society of Cardiology, Pedro Adragao; Romania: 2708 Romanian Society of Cardiology, Gheorghe-Andrei Dan; Russian Federation: Russian Society of Cardiology, 2709 Evgeny N. Mikhaylov; San Marino: San Marino Society of Cardiology, Marco Zavatta; Serbia: Cardiology 2710 Society of Serbia, Tatjana Potpara; Slovakia: Slovak Society of Cardiology, Peter Hlivak Slovenia: Slovenian 2711 Society of Cardiology, Igor Zupan; Spain: Spanish Society of Cardiology, Angel Arenal; Sweden: Swedish 2712 Society of Cardiology, Frieder Braunschweig; Switzerland: Swiss Society of Cardiology, Dipen Shah; Tunisia: 2713 Tunisian Society of Cardiology and Cardio-Vascular Surgery, Ag Sana Ouali; Turkey: Turkish Society of 2714 Cardiology, Mesut Demir; Ukraine: Ukrainian Association of Cardiology, Oleg Sychov; United Kingdom: 2715 British Cardiovascular Society, Ed Duncan. 2716

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