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Improving the prescription of oral anticoagulants in atrial fibrillation

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Supplementary Figure 1: Sample search strategy

Database: Ovid MEDLINE

- 1 Atrial Fibrillation/ or "atrial fibrillation".mp.
- 2 Atrial Flutter/ or "atrial flutter".mp.
- 3 ("auricular fibrillation" or "heart fibrillation" or "heart atrium fibrillation").ti,ab.
- 4 1 or 2 or 3
- 5 exp Anticoagulants/tu, th [Therapeutic Use, Therapy]
- 6 ((anticoagula\$ or antithrombotic\$1) adj2 (therapy or treatment or under-treatment or uptake or underus\$ or prescri\$ or prophylaxis or management or assessment or clinic\$1)).ti,ab.
- 7 exp Factor Xa Inhibitors/tu [Therapeutic Use]
- 8 exp Antithrombins/tu [Therapeutic Use]
- 9 ("direct thrombin inhibitor\$1" or DTI\$1 or "factor Xa inhibitor\$1" or "fxa inhibitor\$1" or NOAC\$1 or "novel oral anticoagulant\$1" or "new oral anticoagulant\$1" or "non-vitamin K antagonist\$1").ti,ab.
- 10 Dabigatran/ or ("dabigatran etexilate" or dabigatran or Pradaxa).ti,ab.
- 11 Rivaroxaban/ or (rivaroxaban or Xarelto).ti,ab.
- 12 (apixaban or Eliquis).ti,ab.
- 13 (edoxaban or Lixiana or Savaysa).ti,ab.
- 14 exp Coumarins/tu, th [Therapeutic Use, Therapy]
- 15 (4-hydroxyc?umarin\$1 or "Vitamin K antagonist" or VKA\$1).mp.
- 16 Warfarin/ or (warfarin or C?umadin or Jantoven or Marevan).ti,ab.
- 17 Dicumarol/ or (dic?umarol\$ or dic?umarin or Bis-Hydroxyc?umarin or bishydroxyc?umarin or Acadyl or Acavyl or Barac?umin or Cuma or Cumid or Dic?uma\$ or Dicumol or Dikumol or Dufalone or Kumoran or Melitoxi or Temparin or Trombosan).ti,ab.
- 18 Phenprocoumon/ or (phenproc?umon\$ or fenproc?umon or phenproc?umarol or Marc?umar or Falithrom).ti,ab.
- 19 Acenocoumarol/ or (acenoc?umarol or nic?umalon\$ or Sintrom or Ascumar or Acitrom or Minisintrom or Neo-sintrom or Sinkumar or Sinthrome or Sync?umar or Synthrom or Trombostop).ti,ab.
- 20 (tioclomarol\$ or Apegmone).ti,ab.
- 21 Ethyl Biscoumacetate/ or ("ethyl bisc?umacetate" or carbethoxydic?umarol or ethyldic?umarol or dic?umacyl or Pelentan or Tromexan or Thrombolysan or Thrombarin or Neodic?umari\$).ti,ab.

- 22 ("indandione derivative\$1" or "non-c?umarin VKA\$1").ti,ab.
- 23 Phenindione/ or (phenindion\$ or fenindion\$ or Dindevan or Fenilin or Phenyline or Soluthrombine).ti,ab.
- 24 (clorindion\$ or chlorphenindone or Indaliton or Cumachlor).ti,ab.
- 25 (diphenadion\$ or difenadion\$ or diphenacin or Dipaxin or Diphac\$).ti,ab.
- 26 (fluindion\$ or Previscan).ti,ab.
- 27 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 or 14 or 15 or 16 or 17 or 18 or 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26
- 28 Health Education/ or Health Promotion/ or "Outcome and Process Assessment (Health Care)"/ or Primary Health Care/ or Health Knowledge, Attitudes, Practice/ or "Outcome Assessment (Health Care)"/ or Program Evaluation/
- 29 Education, Professional/ or Education, Professional, Retraining/
- 30 Practice Patterns, Physicians'/ or Practice Guideline/ or Practice Patterns, Nurses'/ or Practice Guidelines as Topic/ or Practice Management/ or General Practice/ or Family Practice/
- 31 Medical Audit/ or Nursing Audit/
- 32 Reminder Systems/
- 33 exp Drug Therapy, Computer-Assisted/ed, mt, nu, td, ut [Education, Methods, Nursing, Trends, Utilization]
- 34 "Marketing of Health Services"/
- 35 Guideline Adherence/
- 36 Information Dissemination/ or "dissemination tool\$1".ti,ab.
- 37 Decision Support Techniques/ or Decision Support Systems, Clinical/
- 38 exp Decision Making/de [Drug Effects]
- 39 ("decision aid\$" or "decision support" or "decision making").ti,ab.
- 40 (intervention\$ or "local consensus process" or "education\$ material" or "education\$ outreach" or "education\$ meeting\$1" or "behavio?r\$ change\$1" or "perception change\$1" or "practice change\$1" or reminder\$1 or alert\$1 or "guideline\$1 implementation" or "guideline\$1 adherence" or "practice guideline\$1" or "practice pattern\$1" or audit or feedback or "evaluation feedback" or "information dissemination" or "software enhancement" or "software tool\$1" or "medical practice management software" or "stroke prevention" or "action and monitoring").ti,ab.
- 41 ((education\$ or behavio?r\$ or prescri\$ or persuasive or informational or marketing or professional\$ or physician\$1 or clinician\$1 or doctor\$1 or practitioner\$1 or GP\$1 or pharmacist\$1 or

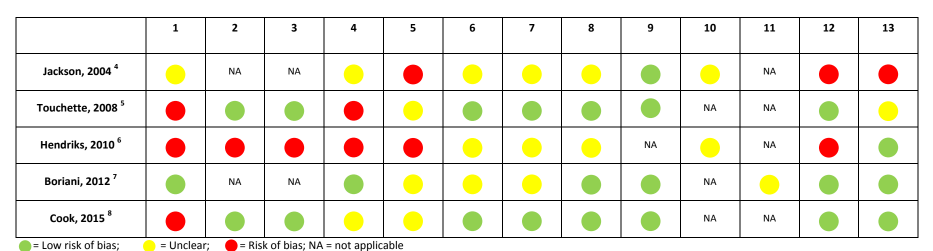
multifaceted or multidisciplinary or "patient-mediated" or "patient-driven") adj2 (intervention\$1 or strateg\$ or program\$ or initiative\$1 or incentive\$1 or improv\$)).ti,ab.

- 42 28 or 29 or 30 or 31 or 32 or 33 or 34 or 35 or 36 or 37 or 38 or 39 or 40 or 41
- 43 4 and 27 and 42
- 44 limit 43 to humans

Supplementary Figure 2: Risk of bias in RCTs using the Cochrane Risk of Bias Tool.

	Random sequence generation	Allocation concealment	Blinding of study participants	Blinding of investigators	Blinding of outcome assessment	Incomplete Primary outcome data	Incomplete Secondary outcome data	Handling missing data	Selective reporting
Bajorek, 2016 ¹									
Arts, 2017 ²									
Holt, 2017 ³									
= Low risk of bias	; = Unclear;	= Risk of bias	S		1			<u>'</u>	

Supplementary Figure 3: Risk of bias in controlled studies using an adjusted Cochrane Risk of Bias Tool.



Criteria used:

- 1. How were different groups selected (e.g. from the same source, at the same time).
- 2. For historical controlled studies also consider if the two sets of patients are comparable.
- 3. For historical controlled studies also consider: 1. changes in the diagnostic criteria; 2. differences in concomitant standards of care over time (e.g. new guidelines).
- 4. Were different groups similar at baseline on important characteristics that could affect outcomes (e.g., demographics, risk factors, co-morbid conditions)?
- 5. Did the authors report that the sample size was sufficiently large to be able to detect a difference in the main outcome between groups with at least 80% power?
- 6. Blinding of outcome assessment.
- 7. Was blinding of outcome assessment the same for all groups?
- 8. Incomplete primary outcome data.
- 9. Incomplete secondary outcome data.
- 10. Handling missing data (e.g. intention to treat).
- 11. Was follow-up time and method of follow-up the same in both groups?
- 12. Selective reporting (e.g. only certain outcomes, no adverse events).
- 13. Other sources of bias.

Supplementary Figure 4: Risk of bias in cross-sectional studies using the NIH National Heart, Lung and Blood Institute Quality Assessment Tool for observational cohort and cross-sectional studies.

	1	2	3	4	5	6	7	8	9	10	11	12	13	14
Falces, 2011 ⁹	✓	✓	CD	√	x	NA	√	√	√	x	√	NR	√	√

^{✓ =} YES; x = NO; CD = cannot determine; NA = not applicable; NR = not reported

Criteria used:

- 1. Was the study question or objective clearly stated?
- 2. Were eligibility/selection criteria for the study population pre-specified and clearly described?
- 3. Was the participation rate of eligible persons at least 50%?
- 4. Were all the subjects selected or recruited from the same or similar populations (including the same time period)? Were inclusion and exclusion criteria for being in the study pre-specified and applied uniformly to all participants?
- 5. Was a sample size justification, power description, or variance and effect estimates provided?
- 6. For the analyses in this paper, were the exposure(s) of interest measured prior to the outcome(s) being measured?
- 7. Was the timeframe sufficient so that one could reasonably expect to see an association between exposure and outcome if it existed?
- 8. For exposures that can vary in amount or level, did the study examine different levels of the exposure as related to the outcome (e.g., categories of exposure, or exposure measured as continuous variable)?
- 9. Were the exposure measures (independent variables) clearly defined, valid, reliable, and implemented consistently across all study participants?
- 10. Was the exposure(s) assessed more than once over time?
- 11. Were the outcome measures (dependent variables) clearly defined, valid, reliable, and implemented consistently across all study participants?
- 12. Were the outcome assessors blinded to the exposure status of participants?
- 13. Was loss to follow-up after baseline 20% or less?
- 14. Were key potential confounding variables measured and adjusted statistically for their impact on the relationship between exposure(s) and outcome(s)?

Supplementary Figure 5: Risk of bias in before-after studies using an adapted form of the NIH National Heart, Lung and Blood Institute Quality Assessment Tool for before-after studies with no control group.

	1	2	3	4	5	6	7	8	9	10	11	12	13	14
Sobreques, 2002 ¹⁰	√	х	√	CD	NR	х	CD	✓	NR	✓	CD	√	х	✓
Lowdon, 2004 ¹¹	✓	CD	√	CD	CD	✓	✓	х	CD	CD	NA	CD	х	✓
Bajorek, 2005 ¹²	✓	√	х	х	✓	✓	√	✓	CD	✓	✓	√	х	✓
Bo, 2007 ¹³	✓	√	х	✓	✓	✓	✓	✓	CD	✓	✓	✓	х	✓
Coll-Vinent, 2007 14	✓	✓	✓	CD	CD	✓	√	✓	CD	✓	✓	√	х	✓
Jackson, 2011 15	✓	✓	х	х	CD	✓	√	✓	✓	✓	CD	√	х	✓
Robson, 2014 ¹⁶	✓	✓	х	✓	CD	✓	√	✓	CD	✓	NA	√	√	✓
Oliveira, 2014 ¹⁷	✓	√	х	CD	CD	✓	CD	✓	NR	CD	х	х	х	✓
Das, 2015 ¹⁸	✓	✓	✓	х	CD	✓	√	✓	NA	✓	✓	√	х	✓
Hsieh, 2016 ¹⁹	✓	х	х	CD	CD	✓	✓	✓	CD	✓	CD	√	✓	✓
Wang, 2017 ²⁰	✓	√	х	CD	CD	✓	✓	✓	CD	✓	✓	√	х	NA

^{✓ =} YES; x = NO; CD = cannot determine; NA = not applicable; NR = not reported

Criteria used:

- 1. Was the study question or objective clearly stated?
- 2. Were eligibility/selection criteria for the study population pre-specified and clearly described?
- 3. Were the participants in the study representative of those who would be eligible for the service/intervention in the general or clinical population of interest?
- 4. Were all eligible participants that met the pre-specified entry criteria enrolled?
- 5. Was the sample size sufficiently large to provide confidence in the findings?

- 6. Was the service/intervention clearly described and delivered consistently across the study population?
- 7. Was the duration of the intervention sufficient so that one could reasonably expect to see any changes in practice and/or behaviour?
- 8. Were the outcome measures pre-specified, clearly defined, valid, reliable, and assessed consistently across all study participants?
- 9. Were the people assessing the outcomes blinded to the participants' interventions?
- 10. Was the length of follow-up sufficient so that one could reasonably expect to capture any changes in practice and/or behaviour?
- 11. Was the loss to follow-up after baseline 20% or less? Were those lost to follow-up accounted for in the analysis?
- 12. Did the statistical methods examine changes in outcome measures from before to after the intervention? Were statistical tests done that provided p values for the pre-to-post changes?
- 13. Were outcome measures of interest taken multiple times before the intervention and multiple times after the intervention (i.e., did they use an interrupted time-series design)?
- 14. If the intervention was conducted at a group level (e.g., a whole hospital, a community, etc.) did the statistical analysis take into account the use of individual-level data to determine effects at the group level?

Supplementary Table 1: Excluded studies

Study	Reason for exclusion
Makowski 1994 ²¹	Full text unavailable
Steffensen, 1997 ²²	Ineligible outcome
Gaughan, 2000 ²³	Not original data
Valeti, 2000 ²⁴	Full text unavailable
Batty, 2001 ²⁵	Ineligible outcome
O'Rourke 2001 ²⁶	Ineligible study design
Elliot, 2002 ²⁷	Ineligible outcome
Jackson, 2003 ²⁸	Not original data
Batty, 2004 ²⁹	Ineligible outcome
Alberts, 2004 ³⁰	Ineligible study design
Kiechl, 2004 ³¹	Ineligible study design
Claes, 2005 ³²	Ineligible outcome
Wright, 2007 ³³	Ineligible outcome
Schwarz, 2009 ³⁴	Ineligible outcome
Albert, 2010 ³⁵	Ineligible outcome
Bishop, 2011 ³⁶	Ineligible outcome
Szabo, 2011 ³⁷	Ineligible population
Boriani, 2011 ³⁸	Not original data
Healicon, 2011 ³⁹	Full text unavailable
Hendriks, 2012 ⁴⁰	Ineligible population
Boriani, 2012 ⁴¹	Not original data
Larsen, 2012 ⁴²	Full text unavailable
Skanes, 2013 ⁴³	Ineligible intervention
Samani, 2013 ⁴⁴	Ineligible outcome
Gadzhanova, 2013 ⁴⁵	Ineligible outcome

Arts, 2013 46	Ineligible study design
Jeng, 2013 ⁴⁷	Not original data
Beadles, 2014 ⁴⁸	Ineligible intervention
Po, 2014 ⁴⁹	Ineligible outcome
Grant, 2014 ⁵⁰	Ineligible outcome
Alkhalil, 2014 ⁵¹	Full text unavailable
Skolarus, 2014 ⁵²	Ineligible outcome
Sibai, 2014 ⁵³	Ineligible study design
Das, 2014 ⁵⁴	Not original data
Fuenzalida, 2015 ⁵⁵	Ineligible outcome
Daacke, 2015 ⁵⁶	Full text unavailable
Akhavein, 2015 ⁵⁷	Full text unavailable
Garber, 2015 ⁵⁸	Ineligible study design
Zheng, 2016 ⁵⁹	Ineligible intervention
Eckman, 2016 ⁶⁰	Ineligible outcome
Eckman, 2016 ⁶¹	Ineligible outcome
Abidi, 2016 ⁶²	Ineligible outcome
Lee, 2016 ⁶³	Ineligible study design
Rao, 2016 ⁶⁴	Ineligible study design
Willis, 2016 ⁶⁵	Ineligible study design
Czernik, 2016 ⁶⁶	Full text unavailable
Cloutier, 2016 ⁶⁷	Full text unavailable
Amiri, 2017 ⁶⁸	Ineligible outcome
Barmano, 2017 ⁶⁹	Ineligible outcome
Rose, 2017 ⁷⁰	Ineligible outcome
Karlsson, 2017 ⁷¹	Ineligible study design
Virdee, 2017 ⁷²	Ineligible study design

Supplementary Table 2: Outcome measures and results

Author	Outcome measure	Baseline or pre-	Follow-up or post-	Group comparison	P-value
		intervention	intervention		
		% (n)	% (n)		
RCTs	·				
Bajorek et al. ¹	a. Use of anticoagulants	January-June 2013	12 month follow-up	a. OACs in the intervention	a. 0.02
	b. Proportion of patients	Control	Control	arm at baseline and follow-up	
Australia, 2016	recommended a different	a. (total n=187): 94.7%	a. Not reported		
	type of therapy	(177)			
	c. Proportion of GPs	Intervention	Intervention		
	agreeing with	a. (total n=206): 89.3%	a. (total n=206): 92.2%		
	recommendations	(184)	(190)		
			b. 36.4%		
			c.75.2%		
Arts et al. ²	a. Proportion of patients	01/10/2013	<u>01/09/2014</u>	a. between group difference	a. Chi sq. between group difference at
7 to ct u	treated according to the	Control	Control	at baseline: 8%	baseline:
The Netherlands,	Dutch guidelines.	a. (total n=235): 42% (99)	a. (Total n=259): 50% (130)	between group difference at	0.04
2017	b. Proportion of patients on	b. 40% (94)	b. 51% (132)	follow-up: 5%	Chi sq. between group difference at
	OACs at baseline and	Intervention	Intervention	b. between group difference	follow-up: 0.23
	follow-up	a. (total n=496): 50% (248)	a. (Total n=522): 55% (287)	at baseline: 8%	a. Cluster analysis between groups:
	·	b. 48% (238)	b. 60% (313)	between group difference at	0.21
		, ,	, ,	follow-up: 9%	b. Chi sq. between group difference at
				·	baseline:
					0.05
					Chi sq. between group difference at
					follow-up: 0.02

Author	Outcome measure	Baseline or pre-	Follow-up or post-	Group comparison	P-value
		intervention	intervention		
		% (n)	% (n)		
Holt at al. ³	Proportion of patients	<u>20/02/2014</u>	a. 6 month follow-up	a. baseline to 6 months	<u>a.</u> 0.213
	eligible for OAC who were	Control	Control	adjusted for baseline	
UK, 2017	currently prescribed an	61.9% (9.89)	63.9% (<u>SD</u> 9.46)	prescribing mean difference	<u>b. 0.173</u>
	OAC (CHADS ₂ \geq 2)	Intervention	Intervention	[95% confidence interval]:	
		63.5% (8.85)	66.3% (<u>SD</u> 9.25)	1.21% [-0.72 to 3.13]	
			b. 12 month follow-up	b. baseline to 12 month	
			<u>Control: 67.8%</u>	adjusted for baseline	
			Intervention: 65.9%	prescribing mean difference	
				[95% confidence interval]:	
				1.79% [-0.82 to 4.41]	
Controlled studies					
Jackson et al. 4	a. Percentage of eligible	Pre-intervention	Post-intervention	Performed comparisons	a. On admission (pre- to post-
	patients receiving warfarin	(01/02/2001 - 31/01/2002)	(01/02/2002 - 31/01/2003)	between all different group	intervention)
Australia, 2004	upon hospital admission	Receiving warfarin:	Receiving warfarin:	combinations	total 0.05;
	b. Percentage of eligible	Intervention	Intervention		high risk 0.02;
	patients receiving warfarin	a. On admission	a. On admission		high risk without contraindications
	upon hospital discharge	total 33% (n=81/245);	total 43% (n=67/157);		0.05;
	c. All prescriptions	high risk 33% (n=64/ 192);	high risk 46% (n=58/125);		intermediate risk 0.60
	dispensed in area (DDDs	high risk without	high risk without		b. <i>On discharge</i>
	per 1000 of population)	contraindications 39%	contraindications 53%		(pre- to post-intervention)
		(n=50/127);	(n=46/87);		total <0.05;
		intermediate risk 30%	intermediate risk 36%		high risk <0.01;
		(n=11/37)	(n=9/25)		high risk without contraindications
		b. On discharge	b. On discharge		<0.05;
		total 39%	total 51%		c. Dispensed prescriptions
		high risk 40%	high risk 56%		Intervention region: pre- vs. post
		high risk without	high risk interventions 64%		<0.001
		contraindications: 49%	c. Prescriptions dispensed		Control region: pre- vs. post <0.001
		c. Prescriptions dispensed	Control region: 1149		Intervention vs control region pre-
		Control region: 1127	Intervention region: 1191		intervention: 0.34
		Intervention region: 1124			Intervention vs control region post -
					intervention: <0.001

Author	Outcome measure	Baseline or pre-	Follow-up or post-	Group comparison	P-value
		intervention	intervention		
_		% (n)	% (n)		
Touchette et al. 5	a. Percentage of patients	Not reported	Control	Between group difference	a. 0.60
	receiving warfarin in-		(01/05/2001-25/07/2001)	a. 3.7%	b. <0.01
USA, 2008	hospital at discharge		a. 41.8% (n=41/98)	b. 21.8%	c. <0.01
	b. Percentage of patients		b. 56.1% (n=55/98)	c. 21.5%; OR 2.46 [95% CI,	
	with a discharge plan for		c. 57.1% (n=56/98)	1.63-3.74]	
	warfarin use		Intervention		
	c. Percentage of patients in		<u>(20/09/2001 – 28/02/2002)</u>		
	planned or actual warfarin		a. 45.5% (n=70/154)		
	use at discharge		b. 77.9% (n=120/154)		
	(high risk according to		c. 78.6% (n=121/154)		
	Chest 2004 guidelines ⁷³)				
Hendriks et al. ⁶	The percentage of patients	Not reported	Control	Between group difference	Difference in therapy per CHADS ₂ score
	receiving VKA treatment		<u>2003-2004</u>	CHADS ₂ =0 21%	<0.001
Netherlands, 2010	(according		CHADS ₂ =0 39% (n=7/18)	CHADS ₂ =1 29%	Difference in high-risk patients
	to the ACC/AHA/ESC AF		CHADS ₂ =1 93% (n=28/30)	CHADS ₂ >1 10%	<0.05
	guidelines ⁷⁴ :- control		CHADS ₂ >1 80% (n=42/52)	In high -risk patients 18%	
	group: 2001; intervention		In high -risk patients		
	group: 2006)		appropriate treatment was		
			given to 79% (41/52)		
			Intervention		
			<u>06/2006-04/2007</u>		
			CHADS ₂ =0 18% (n=6/34)		
			CHADS ₂ =1 64% (n=25/39)		
			CHADS ₂ >1 90% (n=34/38)		
			In high -risk patients		
			appropriate treatment was		
			given to 97% (n=37/38)		

Boriani et al. ⁷ Italy, 2012	Outcome measure Percentage of patients on OAC therapy at the end of the observational period (≤ 48 months) (CHADS₂ ≥ 1)	Baseline or pre- intervention % (n) Control 46.9% (n=693/1477) Intervention 46.1% (n=904/1961)	Follow-up or post- intervention % (n) Control 56.8% (n=258/454) (Intervention during the stroke risk evaluation phase 69.4% (n=474/683) Intervention: final follow-up 72.6% (n=496/683)	Between group difference 15.8%	P-value <0.001
Cook et al. ⁸ USA, 2015	a. Prescription of warfarin in high-risk eligible patients within 30 days of AF diagnosis (CHADS₂ ≥ 2) b. Prescription of warfarin in all eligible patients within 30 days of AF diagnosis (CHADS₂ ≥ 2) c. Frequency of an appropriate medication prescription (warfarin for any warfarin-eligible patient, or aspirin for warfarin-eligible low-risk patients [CHADS2<2]).	Control: (12/2008 – 02/2009) a. High risk (CHADS2_2) 36% (n=34/94) b. Not provided c. 43% (n=85/196)	Intervention: (12/2009 – 02/2010) a. High risk (CHADS2_2) 27% (n=34/125) b. Not provided c. 45% (n=109/244)	a. Between group difference in high-risk patients OR 0.66 [95% CI, 0.37-1.17] b. Between group difference in all warfarin-eligible patients Adjusted OR 0.91 [95% CI, 0.60 -1.38] c. Between group difference in appropriate prescription OR 1.05 [95% CI, 0.72–1.54]; adjusted OR 1.12 [95% CI, 0.76–1.66]	a. Difference in high-risk patients 0.16 b. Difference in all warfarin-eligible patients 0.65 c. Between group difference in appropriate prescription 0.78 adjusted: 0.57
Cross-sectional stud	dies				
Falces et al. ⁹ Spain, 2011	Percentage prescription of anticoagulation therapy (ACC/AHA/ESC AF guidelines ⁷⁴)	Usual care: (Specialist units 01/2008 – 12/2008) 69.3% (n=201/290) (Univariate analysis)	Integrated care: Primary care 01/2008 – 12/2009; specialist units 01/2009 - 12/2009) 94.6% (n=211/223) (Univariate analysis)	Logistic regression model: adjusted OR 7.1 [95% CI, 3.8- 13.5]	<0.001 for univariate analysis and logistic regression

Author	Outcome measure	Baseline or pre-	Follow-up or post-	Group comparison	P-value
		intervention	intervention		
		% (n)	% (n)		
Before-after studies	3	•		•	•
Sobreques et al. 10	Percentage of eligible	Pre-intervention	Post-intervention	Between group difference	<0.01
	patients taking	70.5% (total n=53)	88.6% (total n=53)	18.1%	
Spain, 2002	acenocumarol				
Lowdon et al. 11	Percentage of eligible	<u>Pre-intervention</u>	<u>Post-intervention</u>	Between group difference	only reported for total <0.01 and total
	patients prescribed	<u>(01/01 – 04/02)</u>	(05/02 - 12/02)	total 23.1%;	eligible <0.001
UK, 2004	anticoagulants	total 31.4% (n=38/121);	total 54.5% (n=30/55);	eligible patients only (no	
		eligible patients only (no	eligible patients only (no	contraindications) 47.2%	
		contraindications) 43.7%	contraindications) 90.9%		
		(n=38/87);	(n=30/33);		
		total high risk 39.6%	total high risk 62.8%		
		(n=36/91);	(n=27/43);		
		eligible high risk 52.2%	eligible high risk 93.1%		
		(n=36/69)	(n=27/29)		
Bajorek et al. ¹²	Percentage of patients	Pre-intervention	Post-intervention	Between group difference	<u>a.</u> 0.39
	receiving warfarin (±	On admission	<u>a.</u> At discharge	post-intervention	
Australia, 2005	aspirin)	20.7% (n=45/218)	17.4% (n=38/218)	<u>a.</u> 3.3%	
			b. At 3 month follow-up		
			<u>16.1 (30/187)</u>		
			c. At 6 month follow-up		
			<u>16.5 (30/184)</u>		

Author	Outcome measure	Baseline or pre-	Follow-up or post-	Group comparison	P-value
		intervention	intervention		
		% (n)	% (n)		
Bo et al. ¹³	Increase in appropriate	Pre-intervention	Post-intervention	Between groups absolute	no p values provided
	OAC prescription at	Year 2000	Year 2004	difference at discharge in	
Italy, 2007	discharge	On admission	On admission	OAC strongly recommended	
		OAC strongly	OAC strongly	group	
		recommended: 36.8%	recommended: 58.1%	25.3% (95%CI: 15% 35%).	
		Year 2000	Year 2004		
		At discharge	At discharge	Adjusted OR for OAC	
		OAC strongly	OAC strongly	prescription at discharge	
		recommended: 56.6% with	recommended: 81.9%	2.11 [95% CI, 1.47 3.04]	
		OAC (n=60/106)	(n=86/105)		
Coll-Vinent et al. 14	a. Percentage of patients	The pre-intervention period	The post-intervention	a. Between group difference:	a. Not reported
	receiving anticoagulation	(June 2004)	period (June 2005)	After visit between pre- and	
Spain, 2007	treatment	a. Before visit in the pre-	a. Before visit in the post-	post-intervention periods:	
	b. No anticoagulation when	intervention period: 53%	intervention period: 58%	10%	
	this is indicated	(n=154/293)	(n=155/267)	b. Between group difference:	
		a. After visit in the pre-	a. After visit in the post-	After visit between pre- and	b. Not reported
		intervention period: 52%	intervention period: 62%	post-intervention periods: -	
		(n=151/293)	(n=163/267)	14%	
		b. Before visit in the pre-	b. Before visit in the post-		
		intervention period: 25%	intervention period: 13%		
		(n=74/293)	(n=36/267)		
		b. After visit in the pre-	b. After visit in the post-		
		intervention period: 24%	intervention period: 10%		
		(69/293)	(n=25/267)		
<u> </u>					

Author	Outcome measure	Baseline or pre-	Follow-up or post-	Group comparison	P-value
		intervention	intervention		
		% (n)	% (n)		
Jackson et al. 15	Proportion of eligible	Pre-intervention	Post-intervention	Performed comparisons	Between study arms on admission: high
	patients receiving warfarin	(02/2004 - 09/2004)	(10/2004 - 02/2006)	between all different group	risk <0.01; moderate risk 0.15; low risk
Australia, 2011		On admission: high risk 31%	On admission: high risk 44%	combinations	no p-value; total 0.004
		(n=76/248); moderate risk	(n=47/107); moderate risk		Between study arms at discharge: high
		30% (n=22/73); low risk	50% (n=6/12); low risk 0%		risk <0.0001; moderate risk 0.0004; low
		16% (n=3/19); total 30%	(n=0/3); total 43%		risk no p-value; total 0.0008
		(n=101/340)	(n=53/122)		Admission vs discharge:
		At discharge: high risk 30%	At discharge: high risk 57%		Pre-intervention: no significant
		(n=76/259); moderate risk	(n=65/115); moderate risk		difference
		33% (n=26/80); low risk	80% (n=13/16); low risk 0%		Post-intervention: total 0.05; high risk
		16% (n=4/35); total 29%	(n=0/3); total 58%		0.04; moderate/low risk no p-value
		(n=106/364)	(n=78/134)		
Oliveira et al. 17	a. Percentage of patients	<u>Pre-intervention</u>	Post-intervention	Between group difference	Not reported
	prescribed appropriate	(01/05/2012 - 04/05/2012)	(03/09/2012 - 07/09/2012)	a. "This means an increase of	
Portugal, 2014	prophylactic therapy based	a: 49.5% (n=52/105)	a: 60% (n=57/95)	21.2% in the number of	
	on risk-scores (> 94% with	b: 46.4% (n=45/97)	b: 56.3% (n=49/87)	patients receiving	
	CHA ₂ DS ₂ -VASc ≥2)			appropriate therapy	
	a. antithrombotics			compared to the first	
	b. OACs			assessment"	
				b. no comparison	
Robson et al. 16	Percentage of patients with	Pre-intervention:	Post-intervention:	Between group difference:	Pre-intervention to pre-intervention:
	AF and CHA ₂ DS ₂ -VASc ≥1 on	<u>(</u> 04/2008 <u>)</u>	(04/2013)	Pre-intervention to pre-	2008 vs. 2011: 0.184
UK, 2014	anticoagulants	50.8% (n=1943/3825)	59.8% (n=2492/4168)	intervention	
		Pre-intervention: 04/2011		2008 vs. 2011: 2.2%	Pre-intervention to post-intervention:
		52.6% (n=2085/3964)		Immediately pre-intervention	2011 vs. 2013: <0.001
		(Intervention commenced		to post-intervention	
		04/2011)		2011 vs. 2013: 7.2%	
				Difference in slope of the	
				trends: 1.63 [95% CI, 1.32 -	
				1.94] p=<0.001	

Author	Outcome measure	Baseline or pre-	Follow-up or post-	Group comparison	P-value
		intervention	intervention		
		% (n)	% (n)		
Das et al. ¹⁸	Overall proportion of	Pre-intervention	Post-intervention	Between group difference	<0.0001
	eligible patients receiving	77% (n=4187/5471)	95% (n=5207/5471)	18%	
UK, 2015	anticoagulation (CHADS ₂ or				
	CHA ₂ DS ₂ -VASc ≥1)				
Hsieh et al. ¹⁹	Percentage of discharge	<u>Pre-intervention</u>	<u>During- intervention</u>	Between group difference	<0.001
	prescription of oral	(05/2006 - 07/2008)	(08/2010 - 07/2011)	32%	
Taiwan, 2016	anticoagulants for eligible	32.1% (total n=9612)	64.1% (total n=7492)		
	AF				
Wang et al. ²⁰	a. Proportion of	Pre-intervention	Post-intervention	Change in patient use of	a. Change in Warfarin use p<0.001
_	participants receiving OACs	a. Total n=251	a. Total n=251	OACs (in eligible patients	a. Change in NOAC use p<0.001
Australia, 2017	(Warfarin and NOACs)	OAC 50.5% (126):	OAC 70.0% (176):	according to risk assessment	b. Agreement between HCPs and tool
	b. Level of HCP agreement	Warfarin 30.3% (76)	Warfarin 40.0% (76)	tool)	re use of anticoagulants vs other
	with tools	NOAC 20.0% (50)	NOAC 30.0% (54)		therapy p<0.001
	recommendations		(Interpreted from a graph)		
			b. Agreed whether eligible		
			for OACs:199 (79.3%)		
			b. Agreed with		
			recommended therapy 132		
			(52.6%)		
I					

Abbreviations: AF, atrial fibrillation; CHADS₂ and CHA₂DS₂-VASc, scoring schemes for stroke risk assessment; CI, confidence interval; OAC, oral anticoagulants; OR, odds ratio ; SD, standard deviation; VKA, Vitamin K antagonists; HCP, healthcare professional.

References

- 1. Bajorek BV, Magin PJ, Hilmer SN, Krass I. Optimizing stroke prevention in patients with atrial fibrillation: A cluster-randomized controlled trial of a computerized antithrombotic risk assessment tool in australian general practice, 2012-2013. Prev Chronic Dis. 2016;13:E90
- 2. Arts DL, Abu-Hanna A, Medlock SK, van Weert HC. Effectiveness and usage of a decision support system to improve stroke prevention in general practice: A cluster randomized controlled trial. PLoS One. 2017;12(2):e0170974
- 3. Holt TA, Dalton A, Marshall T, et al. Automated software system to promote anticoagulation and reduce stroke risk: Cluster-randomized controlled trial. Stroke. 2017;48(3):787-790
- 4. Jackson SL, Peterson GM, Vial JH. A community-based educational intervention to improve antithrombotic drug use in atrial fibrillation. Ann Pharmacother. 2004;38(11):1794-1799
- 5. Touchette DR, McGuinness ME, Stoner S, Shute D, Edwards JM, Ketchum K. Improving outpatient warfarin use for hospitalized patients with atrial fibrillation. Pharm Pract. 2008;6(1):43-50
- 6. Hendriks JLM, Nieuwlaat R, Vrijhoef HJM, de Wit R, Crijns HJGM, Tieleman RG. Improving guideline adherence in the treatment of atrial fibrillation by implementing an integrated chronic care program. Neth Heart J. 2010;18(10):471-477
- 7. Boriani G, Santini M, Lunati M, et al. Improving thromboprophylaxis using atrial fibrillation diagnostic capabilities in implantable cardioverter-defibrillators the multicentre italian ANGELS of AF project. Circ Cardiovasc Qual Outcomes. 2012;5(2):182-188
- 8. Cook DA, Enders F, Caraballo PJ, Nishimura RA, Lloyd FJ. An automated clinical alert system for newly-diagnosed atrial fibrillation. PloS one. 2015;10(4):e0122153
- 9. Falces C, Andrea R, Heras M, et al. [Integration between cardiology and primary care: impact on clinical practice]. Rev Esp Cardiol. 2011;64(7):564-571
- 10. Sobreques J, Espinasa J, Cebria J. Effectiveness of an intervention programme to improve oral anti-coagulation treatment for patients with chronic auricular fibrillation in a health district. Aten Primaria. 2002;30(9):588-589
- 11. Lowdon DW, Harper JR, Gillespie ND. Improving thromboprophylaxis in elderly patients with non-valvular atrial fibrillation. Scott Med J. 2004;49(4):148-150
- 12. Bajorek BV, Krass I, Ogle SJ, Duguid MJ, Shenfield GM. Optimizing the use of antithrombotic therapy for atrial fibrillation in older people: a pharmacist-led multidisciplinary intervention. J Am Geriatr Soc. 2005;53(11):1912-1920
- 13. Bo S, Valpreda S, Scaglione L, et al. Implementing hospital guidelines improves warfarin use in non-valvular atrial fibrillation: a before-after study. BMC Public Health. 2007;7:203
- 14. Coll-Vinent B, Pacheco G, Junyent M, et al. [Impact of implementing common guidelines at different care levels in a healthcare area on the improvement of atrial fibrillation treatment]. Rev Esp Cardiol. 2007;60(4):392-403
- 15. Jackson SL, Peterson GM. Stroke risk assessment for atrial fibrillation: Hospital-based stroke risk assessment and intervention program. J Clin Pharm Ther. 2011;36(1):71-79

- 16. Robson J, Dostal I, Mathur R, et al. Improving anticoagulation in atrial fibrillation: Observational study in three primary care trusts. Br J Gen Pract. 2014;64(622):e275-281
- 17. Oliveira R, Grilo S, Moreira C, et al. A quality study to improve prophylactic antithrombotic therapy prescribed to patients with atrial fibrillation. Rev Port Cardiol. 2014;33(2):89-94
- 18. Das M, Panter L, Wynn GJ, et al. Primary Care Atrial Fibrillation Service: Outcomes from consultant-led anticoagulation assessment clinics in the primary care setting in the UK. BMJ open. 2015;5(12):e009267
- 19. Hsieh F-I, Jeng J-S, Chern C-M, et al. Quality improvement in acute ischemic stroke care in taiwan: The breakthrough collaborative in stroke. PLoS One. 2016;11(8):e0160426
- 20. Wang Y, Bajorek B. Pilot of a Computerised Antithrombotic Risk Assessment Tool Version 2 (CARATV2.0) for stroke prevention in atrial fibrillation. Cardiol J. 2017;24(2):176-187
- 21. Makowski TM, Jacobson AK, Hildebrand SW, Ferry DR, Heywood JT. Impact of computerized tracking on increasing the proportion of atrial-fibrillation patients receiving warfarin anticoagulation for stroke prevention. Circulation. 1994;90(4):473
- 22. Steffensen FH, Sorensen HT, Olesen F. Impact of local evidence-based clinical guidelines-a Danish intervention study. Fam Pract. 1997;14(3):209-215
- 23. Gaughan GL. The use of practitioner education and a warfarin monitoring service to improve anticoagulation practices. Arch Intern Med. 2000;160(15):2401-2402
- 24. Valeti V, Horton L, Schwartz J, Lopez-Candales A. A simple intervention to improve physician practices regarding antithrombotic therapy in patients with atrial fibrillation. J Am Coll Cardiol. 2000;35(2):102A
- 25. Batty G, Oborne CA, Hooper R, Jackson S. Investigation of intervention strategies to increase the appropriate use of antithrombotics in elderly hospital inpatients with atrial fibrillation. Journal of Clinical Governance. 2001;9(3):115-122
- 26. O'Rourke A, Fox N, Field R. A survey of education and training for clinical governance leads and Caldicott Guardians in primary care in Trent Region. Journal of Clinical Governance. 2001;9(3):137-142 136p
- 27. Elliott RA, Woodward MC, Oborne CA. Antithrombotic prescribing in atrial fibrillation: application of a prescribing indicator and multidisciplinary feedback to improve prescribing. Age Ageing. 2002;31(5):391-396
- 28. Jackson SL, Vial JH, Peterson GM. Community-based educational intervention improves Antithrombotic drug use in atrial fibrillation. Circulation. 2003;108(17):786
- 29. Batty GM, Grant RL, Aggarwal R, et al. National clinical sentinel audit of evidence-based prescribing for older people. J Eval Clin Pract. 2004;10(2):273-279
- 30. Alberts MJ, Easton JD. Stroke best practices: A team approach to evidence-based care. J Natl Med Assoc. 2004;96(4 SUPPL.):5S-20S
- 31. Kiechl S. Prevention of stroke in atrial fibrillation: Implementation of study results in practice. Wien Klin Wochenschr. 2004;116(24):817-819
- 32. Claes N, Buntinx F, Vijgen J, et al. The Belgian improvement study on oral anticoagulation therapy: A randomized clinical trial. Eur Heart J. 2005;26(20):2159-2165
- Wright J, Bibby J, Eastham J, et al. Multifaceted implementation of stroke prevention guidelines in primary care: cluster-randomised evaluation of clinical and cost effectiveness. Qual Saf Health Care. 2007;16(1):51-59 59p

- 34. Schwarz K, Morike K, Meisner C, Fux R, Gleiter CH. Improving guideline adherence of antithrombotic therapy in patients with chronic atrial fibrillation: A prospective interventional cohort study. Basic Clin Pharmacol. 2009;104(6):491-492
- 35. Albert NM, Fonarow GC, Yancy CW, et al. Outpatient cardiology practices with advanced practice nurses and physician assistants provide similar delivery of recommended therapies (findings from IMPROVE HF). Am J Cardiol. 2010;105(12):1773-1779
- 36. Bishop TF, Federman AD, Ross JS. Physician incentives to improve quality and delivery of highqualityambulatorymedical care. J Gen Intern Med. 2011;26:S68
- 37. Szabo I, Mihaly GY, Erdelyi ZS. Antithrombotic therapy in chronic atrial fibrillation: Can enlightenment and education for stroke prevention in a busy emergency department be effective? Eur Heart J. 2011;32:462
- 38. Boriani G, Lunati M, Gasparini M, et al. A medical care program to improve oral anticoagulation use by leveraging atrial fibrillation diagnostic capabilities in implantable cardioverter defibrillators: the ANGELS of AF Project. Eur Heart J. 2011;32:562
- 39. Healicon R, Barrett J, Fay M, et al. The use of oral anti-coagulants in the management of atrial fibrillation. National data from guidance on risk assessment and stroke prevention in atrial fibrillation. Europace. 2011;13:iv3
- 40. Hendriks JML, De Wit R, Crijns HJGM, et al. Nurse-led care vs. usual care for patients with atrial fibrillation: Results of a randomized trial of integrated chronic care vs. routine clinical care in ambulatory patients with atrial fibrillation. Eur Heart J. 2012;33(21):2692-2699
- 41. Boriani G, Santini M, Lunati M, et al. Improving thromboprophylaxis using atrial fibrillation diagnostic capabilities in implantable cardioverter defibrillators. The multicentre Italian ANGELS of AF project. Eur Heart J. 2012;33:1060-1061
- 42. Larsen SL, Sonderstrup D, Sonderstrup J, Toft JC. [Clinical guidelines for antithrombotic therapy of patients with atrial fibrillation can be implemented in general practice by means of data capture]. Ugeskr Laeger. 2012;174(38):2227-2229
- 43. Skanes A, Bell A. Knowledge transfer at point-of-care: Investigating new strategies for disseminating guideline recommendations. Can J Cardiol. 2013;29(10 SUPPL. 1):S259
- 44. Samani A, Das S, Khan S. Improved awareness and management of thromboprophylaxis in atrial fibrillation in an elderly care population over a seven year period. Age Ageing. 2013;42:ii6
- 45. Gadzhanova SV, Roughead EE, Bartlett MJ. Improving cardiovascular disease management in Australia: NPS MedicineWise. Med J Aust. 2013;199(3):192-195
- 46. Arts DL, Abu-Hanna A, Buller HR, Peters RJG, Eslami S, van Weert HCPM. Improving stroke prevention in patients with atrial fibrillation. Trials. 2013;14:193
- 47. Jeng JS, Lien LM, Lee TH, et al. Quality improvement in acute ischemic stroke care in taiwan: The breakthrough collaborative in stroke. Stroke. 2013;44(2 MeetingAbstract):no pagination
- 48. Beadles CA, Hassmiller Lich K, Viera AJ, Greene SB, Brookhart MA, Weinberger M. A non-experimental study of oral anticoagulation therapy initiation before and after national patient safety goals. BMJ open. 2014;4(2):e003960
- 49. Po HL, Yu HF, Lin HC, et al. Improvement in prescription frequency of oral anticoagulant for secondary stroke prevention in the breakthrough series-stroke. Cerebrovasc Dis. 2014;38:97

- 50. Grant AM, Guthrie B, Dreischulte T. Developing a complex intervention to improve prescribing safety in primary care: Mixed methods feasibility and optimisation pilot study. BMJ Open. 2014;4(1):no pagination
- 51. Alkhalil M, Cromie N. Education in atrial fibrillation. Ir J Med Sci. 2014;183(8 SUPPL. 1):S419
- 52. Skolarus LE, Morgenstern LB, Scott PA, et al. An emergency department intervention to increase warfarin use for atrial fibrillation. J Stroke Cerebrovasc Dis. 2014;23(2):199-203
- 53. Sibai M-S, Bellarbre F, Ghazali N, et al. [Decisional algorithm to prescribe vitamin K antagonist in geriatric patients with atrial fibrillation]. Geriatr Psychol Neuropsychiatr Vieil. 2014;12(1):20-24
- Das M, Panter L, Connor N, Mills J, Gupta D. The primary care atrial fibrillation (PCAF) service: Consultant-led anticoagulation assessment clinics in the primary care setting increase the uptake of anticoagulation therapy in AF patients at high-risk of stroke. Europace. 2014;16:iii16
- 55. Fuenzalida C, Coll-Vinent B, Navarro M, et al. [Temporal evolution of treatment of patients with atrial fibrillation in a urban health care area]. Med Clin (Barc). 2015;144(11):483-486
- 56. Daacke I, Hau N, Williams J, Natarajan I. Atrial fibrillation and anti-coagulation service run by a clinical nurse specialist. Value Health. 2015;18(7):A400
- 57. Akhavein RM, Sklenar J, Minnier J, Heitner S. Efficacy of clinical notifications to health care providers in management of heart failure patients. Circulation. 2015;132:no pagination
- 58. Garber JL, Willenborg KL, Rose AE. Analysis of anticoagulant prescribing in non-valvular atrial fibrillation and development of a clinical tool for guiding anticoagulant selection. J Thromb Thrombolysis. 2015;40(2):248-254
- 59. Zheng Q, Aneke-Nash C, Halperin JL, Vorchheimer DA. Novel oral anticoagulant availability increases appropriate use of anticoagulants for nonvalvular atrial fibrillation in clinical practice. J Am Coll Cardiol. 2016;67(13 SUPPL. 1):787
- 60. Eckman MH, Lip GY, Wise RE, et al. Impact of an atrial fibrillation decision support tool (AFDST) on thromboprophylaxis for atrial fibrillation. J Gen Intern Med. 2016;31(2 SUPPL. 1):S258-S259
- 61. Eckman MH, Lip GYH, Wise RE, et al. Impact of an Atrial Fibrillation Decision Support Tool on thromboprophylaxis for atrial fibrillation. Am Heart J. 2016;176:17-27
- 62. Abidi S, Cox J, Abusharekh A, Hashemian N, Abidi SS. A digital health system to assist family physicians to safely prescribe NOAC medications. Eur J Epidemiol. 2016;31:S9
- 63. Lee TM, Ivers NM, Bhatia S, et al. Improving stroke prevention therapy for patients with atrial fibrillation in primary care: protocol for a pragmatic, cluster randomized trial. Implement Sci. 2016;11:13
- Rao MP, Ciobanu AO, Lopes RD, et al. A clustered randomized trial to IMProve treatment with AntiCoagulanTs in patients with Atrial Fibrillation (IMPACT-AF): design and rationale. Am Heart J. 2016;176:107-113
- 65. Willis TA, Hartley S, Glidewell L, et al. Action to Support Practices Implement Research Evidence (ASPIRE): protocol for a cluster-randomised evaluation of adaptable implementation packages targeting 'high impact' clinical practice recommendations in general practice. Implement Sci. 2016;11:25

- 66. Czernik Z, Cheung D, Cumbler EU. Accelerating the pace of change: Standardizing anticoagulation practice for the elderly in the era of direct oral anticoagulants. J Gen Intern Med. 2016;31(2 SUPPL. 1):S862-S863
- 67. Cloutier J, Khoo C, Hiebert B, Wassef A, Seifer C. Physician decision making in anticoagulating atrial fibrillation: A prospective survey evaluating a physician alert system for atrial fibrillation detected on cardiac implantable electronic devices. J Am Coll Cardiol. 2016;67(13 SUPPL. 1):837
- 68. Amiri M, Kargar M, Borhanihaghighi A, Soltani F, Zare N. The effect of nurse-led care on stability time in therapeutic range of INR in ischemic stroke patients receiving warfarin. Appl Nurs Res. 2017;33:96-101
- 69. Barmano N, Walfridsson U, Walfridsson H, Karlsson JE. Structured care of patients with atrial fibrillation improves guideline adherence. J Atr Fibrillation. 2017;9(4)
- 70. Rose AJ, Park A, Gillespie C, et al. Results of a regional effort to improve warfarin management. Ann Pharmacother. 2017;51(5):373-379
- 71. Karlsson LO, Nilsson S, Charitakis E, et al. Clinical decision support for stroke prevention in atrial fibrillation (CDS-AF): Rationale and design of a cluster randomized trial in the primary care setting. Am Heart J. 2017;187:45-52
- 72. Virdee MS, Stewart D. Optimizing the use of oral anticoagulant therapy for atrial fibrilation in primary care: a pharmacist-led intervention. Int J Clin Pharm. 2017;39(1):173-180
- 73. Singer DE, Albers GW, Dalen JE, Go AS, Halperin JL, Manning WJ. Antithrombotic therapy in atrial fibrillation: The seventh ACCP conference on antithrombotic and thrombolytic therapy. Chest. 2004;126(3 Suppl):429s-456s
- 74. Fuster V, Ryden LE, Cannom DS, et al. ACC/AHA/ESC 2006 guidelines for the management of patients with atrial fibrillation: full text: a report of the American College of Cardiology/American Heart Association Task Force on practice guidelines and the European Society of Cardiology Committee for Practice Guidelines (Writing Committee to Revise the 2001 guidelines for the management of patients with atrial fibrillation) developed in collaboration with the European Heart Rhythm Association and the Heart Rhythm Society. Europace. 2006;8(9):651-745