# Prevalence, safety and effectiveness of oral anticoagulant use in people with and without dementia or cognitive impairment: a systematic review and meta-analysis

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Running title: Anticoagulation use and outcomes stratified by dementia

## Abstract

## Background

Differences in management and outcomes of oral anticoagulant (OAC) use may exist for people with and without dementia or cognitive impairment (CI).

# Objective

To systematically review the prevalence and safety and effectiveness outcomes of OAC use in people with and without dementia or CI.

#### Methods

MEDLINE, EMBASE and CINAHL were searched for studies reporting prevalence or safety and effectiveness outcomes of OAC use for people with and without dementia, published between 2000 to September 2017. Study selection, data extraction and quality assessment were performed by two-reviewers.

## Results

27 studies met pre-specified inclusion criteria (21 prevalence studies, six outcomes studies). People with dementia had 52% lower odds of receiving OAC compared to people without dementia. Mean OAC prevalence was 32% for people with dementia, compared to 48% without dementia. There was no difference in the composite outcome of embolic events, myocardial infarction, and all-cause death between dementia and non-dementia groups (adjusted hazard ratio (HR) 0.72, 95% CI, 0.45-1.14, p=0.155). Bleeding rate was lower for people without dementia (HR 0.56, 95% CI, 0.37-0.85). Adverse warfarin events were more common for residents of long-term care with dementia (adjusted incidence rate ratio 1.48, 95% CI, 1.20-1.82). Community-dwelling people with dementia treated with warfarin had poorer

anticoagulation control than those without dementia (mean time in the rapeutic range (TTR) %  $\pm$  SD, 38 $\pm$ 26 (dementia), 61 $\pm$ 27 (no dementia), p<0.0001).

# Conclusion

A lower proportion of people with dementia received oral anticoagulation compared with people without dementia. People with dementia had higher bleeding risk and poorer anticoagulation control when treated with warfarin.

**Key words:** anticoagulant, atrial fibrillation, dementia, cognitive impairment, prevalence, ischaemic stroke, haemorrhage, warfarin

## INTRODUCTION

Atrial fibrillation (AF), dementia and cognitive impairment (CI) are common in older adults, hence they often occur together [1]. AF is a key risk factor for stroke, and confers a nearly twofold increased probability of death [2-5]. Further, AF has been associated with an increased risk of developing dementia, with and without prior history of stroke [1, 6]. Diabetes, heart failure and hypertension are risk factors for both AF and CI [1, 6-9]. Between 26% and 51% of community and hospitalized individuals with AF have CI [10-12]. People with CI have longer durations of hospitalization, poorer post-discharge outcomes and increased risk of re-hospitalization than people without CI [13, 14].

The presence of dementia or CI affects the management of comorbid chronic disease [15, 16]. Prevention of long-term complications of chronic disease may be de-emphasized in the context of limited life expectancy and changing care goals [16]. Compared to people with AF and normal cognition people with dementia or CI and AF are less likely to receive vitamin K antagonists (VKA), even though people with dementia demonstrate similar or increased stroke risk [17-21] and increased mortality risk [22, 23]. People with dementia are at increased risk of haemorrhagic complications, such as bleeding linked to falls [24-26]. Further, due to the detrimental effects of amyloid-beta on arterial walls, people with dementia may experience increased rates of intracranial haemorrhage [27, 28]. European Society of Cardiology guidelines recommend withholding OAC in people with dementia only when medication non-adherence is suspected and cannot be assured by a caregiver [22]. American Academy of Neurology guidelines state insufficient evidence is available regarding the safety of OAC for stroke prevention in AF in moderate to severe dementia [29].

The introduction of four direct oral anticoagulants (DOACs): dabigatran, rivaroxaban, apixaban and edoxaban, has expanded the anticoagulant armamentarium for stroke

prevention in AF. Large phase III randomised controlled trials (RCTs) provide evidence of non-inferiority or superiority to warfarin for the prevention of cerebral and systemic embolic events in AF, but reduced risk of intracranial bleeding [30-34]. Well-conducted observational studies support the effectiveness and safety of DOACs compared with warfarin in more inclusive groups [35-39]. DOACs offer practical advantages over VKA therapy as DOAC dosing is based on clinical characteristics and fixed dosing regimens [40]. OAC utilization has increased considerably following DOAC introduction. There has been increasing uptake of DOACs, while the use of VKA has gradually reduced [41-45]. Increasing OAC use has been observed in women [41] and in older people, particularly octogenarians [41, 44]. However, comparative effectiveness and safety studies that include representative samples of people with dementia or CI are lacking [45]. Few people with dementia were eligible to participate in the pivotal DOAC trials [46]. The objective of this systematic review was to identify published data comparing the prevalence and safety and effectiveness outcomes of OAC use in people with AF with and without dementia or cognitive impairment, and to summarise the data using a meta-analysis.

# **METHODS**

The review was conducted and reported in accordance with the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement [47]. The review protocol was registered in the Prospero International Prospective Register of Systematic Reviews (PROSPERO Number CRD42017050663). Oral anticoagulant medications were defined as oral formulations of vitamin K antagonists, direct thrombin inhibitors and factor Xa inhibitors (Anatomical Therapeutic Chemical (ATC) codes of the World Health Organization: B01AA03 (warfarin), B01AE07 (dabigatran etexilate), B01AF01 (rivaroxaban), B01AF01 (apixaban) and B01AF03 (edoxaban) [48]. Studies of all forms of cognitive impairment and dementia were considered, including mild cognitive impairment, Alzheimer's disease, vascular dementia, mixed dementias and Lewy Body dementia.

## **Search strategy**

Studies were identified through a literature search using MEDLINE, EMBASE and Cumulative Index to Nursing and Allied Health Literature (CINAHL) databases from 1 January 2000 until 30 September 2017. This date range was selected to cover eight to 10 years before and after the introduction of the DOACs. Medical subject headings (MeSH), Emtree terms, keywords and truncated search terms related to dementia or CI (dementia, Alzheimer's disease, cognitive impairment, cognitive aging) and anticoagulants (anticoagulant, novel oral anticoagulant, NOAC, direct oral anticoagulant, DOAC, apixaban, dabigatran, rivaroxaban, edoxaban, warfarin, vitamin K antagonist, direct thrombin inhibitor and factor Xa inhibitor) were combined. Searches were limited to English-language. Reference lists of identified articles were screened for any additional studies. Full search strategies are available in Appendix 1 of Supplemental Material.

## Inclusion and exclusion criteria

Studies of all designs were eligible for inclusion. Studies were included in this review if they reported:

- original research reporting the prevalence or safety and effectiveness outcomes of oral anticoagulant use for people with and without dementia or CI;
- prevalence or safety and effectiveness outcomes data separately for people with and without dementia or CI drawn from the same study sample and presented within the study result, for example, sub-group analyses;

• prevalence data of specific oral anticoagulants or prevalence data for classes of oral anticoagulants such as vitamin K or non-vitamin K antagonists for people with and without dementia or CI;

Studies were excluded if they:

- reported the prevalence or safety and effectiveness outcomes of oral anticoagulant use in people with dementia or CI only;
- only reported aggregated results for oral and parenteral anticoagulants combined or antiplatelet and anticoagulant medications combined;
- did not present original data, or were case reports, conference proceedings, review articles, editorials or letters, or not available in English language.

## **Study selection**

One reviewer (TRA) performed the full search strategy, removed duplicates and screened article titles. Abstracts were screened independently by two reviewers (TRA, LF). Full-text copies were obtained if studies appeared to meet inclusion criteria or if it was unclear if they met inclusion criteria. Full-text articles were independently reviewed by two investigators (TRA, LF) for inclusion. Discrepancies were discussed with a third investigator (JI) until consensus was reached.

#### **Data extraction**

Data were extracted by two reviewers (TRA and LF) independently using a standardised data extraction tool. Data extracted included study details, publication year, study design, study country and setting, study sample characteristics (age, gender), sample size, data sources used, data collection period, prevalence of dementia or CI within study sample, prevalence of OAC use for the overall study sample, prevalence of OAC use among participants with dementia or CI, prevalence of OAC use among participants without dementia or CI, safety and effectiveness outcomes of OAC use for participants with dementia, OAC investigated and OAC indications(s), safety and effectiveness outcomes from OAC use for participants without dementia, dementia type and the method used to identify dementia or CI. Data were extracted separately for participants with and without dementia or CI. Prevalence results include both estimates based on individual oral anticoagulants and grouped oral anticoagulants. When prevalence of OAC use data were clearly reported for these groups, results provided by the authors were used. When data were not clearly reported, but stratification and calculations were possible using the published data, calculations were undertaken to determine prevalence of OAC use among participants with dementia or CI and those without dementia or CI. Data for safety and effectiveness outcomes from OAC use were descriptively extracted from each study and reported separately.

#### Quality assessment

Two investigators (LF, TRA) independently assessed the methodological quality of prevalence and outcomes studies using adapted versions of the Joanna Briggs Institute critical appraisal tools for analytical cross-sectional studies and cohort studies, respectively [49] (Appendix 2). Quality assessment tools were selected based on study designs of included studies. No RCTs were identified in this systematic review. For cross-sectional prevalence studies, the definition of dementia and medication use, were assessed against pre-specified quality criteria. These quality criteria were applied even when comparing the prevalence of OAC use in people with and without dementia was not the primary objective of each included study (Appendix 2). Any disagreements in assessments were resolved by a third investigator (JI).

## Mean OAC prevalence and time trends

The mean OAC prevalence for cardioembolic stroke prevention in AF for dementia/CI and non-dementia/CI groups was calculated by averaging OAC prevalence for all studies combined and stratified by community, hospital and long term care settings. Trends in OAC prevalence for cardioembolic stroke prevention in AF over the time period 2000 to 2016 were examined by plotting OAC prevalence for dementia/CI and non-dementia/CI groups by mid-year of study observation period. A linear trend line was fit to examine changes in OAC prevalence over time. Two studies did not report time of study observation period and were excluded [50, 51].

#### **Meta-analysis**

The prevalence of OAC use for people with AF both with and without dementia or CI and crude odds ratios (OR) were calculated from study data of included articles. Meta-analyses were conducted by pooling all studies, and then stratifying by healthcare settings: community, hospital and long-term care (e.g: residential aged care facilities). Meta-analyses were performed using Review Manager 5.3 [52]. Data were pooled using a random effect model as described by DerSimonian-Laird [53]. The pooled-effect of OAC use for people with and without dementia are reported as OR and 95% confidence intervals (CI). Statistical heterogeneity was assessed among studies by the I<sup>2</sup> statistic. To account for both clinical and statistical heterogeneity between studies we utilised a random-effects model. Sensitivity analyses were conducted to investigate the influence of individual studies and characteristics in the pooled ORs for OAC prevalence.

## RESULTS

Electronic database searches yielded 4081 articles, of which 27 were finally included in this review (figure 1). Of the included 27 studies, 21 studies provided results for prevalence of OAC use for cardioembolic stroke prevention in AF and six studies provided results for

safety and effectiveness outcomes from OAC use for cardioembolic stroke prevention in AF among people with and without dementia or CI.

## **Study characteristics**

Study characteristics are summarised in table 1. Studies were conducted in United States of America (n=8) [20, 23, 51, 54-58], Canada (n=3) [17, 59, 60], United Kingdom (n=4) [19, 24, 61, 62] and rest of Europe (n=11) [50, 63-72], and one study was a multicentre international study [18]. Three prevalence studies utilised data from the Stroke in Atrial Fibrillation Ensemble II (SAFE II) study (multi-site European study) [65, 66, 68].

Of the 21 studies reporting the prevalence of OAC use, 11 were conducted in a hospital setting [17, 20, 50, 55, 59, 64-68, 70], seven in a community setting [19, 24, 54, 60-62, 69] and three in long-term care [51, 57, 63]. Fifteen of the studies were cross-sectional designs, four were retrospective cohort studies, one study was a prospective cohort study and one was a series of cross-sectional studies (table1). Data from prevalence studies involved 14,734 people with dementia and 307,961 people without dementia.

Of the six studies that presented safety and effectiveness outcomes data of OAC use, four were conducted in community settings [18, 23, 56, 72], one in a hospital [71] and one in long-term care setting [58]. Four of the studies were retrospective cohort designs [23, 56, 71, 72], one study was a prospective cohort study [58] and one study undertook post-hoc analysis of a subset of data collected in a randomised controlled trial (table 1) [18].

Warfarin was the anticoagulant investigated for 20 of the 27 studies. One study included dabigatran, rivaroxaban, apixaban and warfarin [17], one study reviewed warfarin and phenprocoumaron [70], one study reviewed warfarin and acenocoumarol [71] and one study reviewed acenocoumarol alone [72]. Three studies did not specify the exact anticoagulant [50, 62, 64] but stated vitamin K antagonists were used.

The indication for OAC for 24 of the 27 studies was stroke prevention in AF alone. Further, one study included thromboembolic disease, mechanical valve replacement and stroke prevention in AF indications [58], one study included treatment of venous thromboembolism (VTE) and stroke prevention in AF indications [56] and one study did not specify the indication [50].

## Study participant characteristics

The included studies selected their patients based on the presence of AF (n=13), AF plus incident- or prior-stroke and/or TIA (n=7), AF/thromboembolic disease/mechanical valve replacement (n=2), AF plus an additional risk factor for stroke (n=1), received treatment from a cardiac provider (n=1), had sustained hip fracture secondary to high-energy fall (n=1), admitted to a geriatric unit and were receiving OAC (n=1), were aged 75 years and older with a history of cardiovascular disease (n=1) (table 1).

Age was reported as mean with standard deviation, median with range or interquartile range (IQR) and by proportions for specified age groups. Mean age ranged from  $70.9 \pm 9.5$  years to  $87.1 \pm 5.3$  years [18, 63]. Median age ranged from 73 (IQR: 64-81) to 85 years [57, 62]. Three studies stratified by age groups and included 21% of participants aged between 60-69 years [24], 9.4% aged less than 65 years [66], and 16% between 65-75 years [23]. The proportion of females ranged from 45% to 75%. The proportion of participants within each study with dementia or CI ranged from 1% to 75%.

The presence of dementia or CI was variably defined across studies. Dementia was reported for 14 studies, cognitive impairment/disorders/dysfunction was reported for 10 studies, and three studies considered both terms as distinct clinical classifications. Eleven studies identified the presence of dementia from information available in administrative data: International Classification of Diseases and Health Related Problems (ICD) codes for dementia [60, 70], Quality and Outcome Read Codes for dementia [19, 24, 61, 62], dementia diagnosis within the Minimum Data Set [57] or comorbid information/problem lists from hospital electronic medical records [54, 59], electronic nursing home database [57], or stroke registry [17]. Nine studies identified people with dementia or cognitive impairment via medical diagnoses found in medical charts and histories, where some studies specified a formal dementia or geriatric assessment and others did not [20, 23, 51, 55, 58, 63-66, 68]. Seven studies described dementia diagnosis ascertainment from validated methods such as the full or modified Mini Mental State Examination (MMSE) or Short Portable Mental Status questionnaire [18, 50, 56, 67, 69, 71, 72] (table 1).

## Methodological quality of studies

Fifteen of 21 cross-sectional prevalence studies scored the maximum on quality assessment. Comparative prevalence of OAC use in people with and without dementia was not the main outcome of interest in all 21 studies included in this review. For this reason we did not assess whether confounding factors were adequately addressed when investigating the difference in prevalence among people with and without dementia or CI. All studies for which prevalence results were obtained compared characteristics of people receiving OAC with those not receiving OAC, which was stratified by presence of dementia (sub-group analyses). Five of the 21 studies from which prevalence data were obtained did not indicate how OAC use was measured which precludes rigorous assessment of whether this was measured validly [50, 62, 64, 67, 70]. For studies that compared safety and effectiveness outcomes of OAC use between dementia and non-dementia groups, three studies scored 10 out of a maximum of 11 points [18, 58, 72] while three studies scored 7 or less points on quality assessment [23, 56, 71]. These three studies were descriptive and did not deal with confounding factors. One study did not provide adequate information to measure OAC use [23]. Full quality assessment results are available in appendix 3 of supplemental material.

#### Prevalence of oral anticoagulant use

The prevalence of OAC use for cardioembolic stroke prevention in AF was 29% (4221/14539) for people with dementia or CI and 47% (144254/306751) for people without dementia or CI when all study data were combined. Prevalence of OAC use for cardioembolic stroke prevention in AF in people with and without dementia or CI ranged from 8.3% to 64.0% and 7.0% and 75.6%, respectively (table 2). Mean prevalence of OAC use for cardioembolic stroke prevention in AF for people with dementia was 32% compared with 48% for people without dementia (figure 2). For the time period 1998 to 2014, OAC prevalence for cardioembolic stroke prevention in AF increased for both dementia and non-dementia groups across all health care settings combined (figure 3).

An overall meta-analysis for all healthcare settings revealed that people with dementia or CI had a significantly lower prevalence of OAC use for cardioembolic stroke prevention in AF compared to people without dementia or CI (OR 0.48, 95% CI=0.40–0.58, p<0.00001) (figure 4 (1.1.1)). Significant statistical heterogeneity between studies was found ( $I^2$ =93%). When stratified by healthcare setting, people with dementia or CI residing in the community had a significantly lower prevalence of OAC use (OR 0.40, 95% CI=0.31–0.52, p<0.00001) (figure 4 (1.1.2)), followed by the people with dementia or CI receiving care in hospital (OR 0.49, 95% CI=0.33–0.73, p<0.00001) (figure 4 (1.1.3)), then followed by residents in long-term care (OR 0.66, 95% CI=0.45–0.95, p<0.00006) (figure 4 (1.1.4)) when compared to people without dementia or CI. Sensitivity analysis revealed no significant influence of any individual studies, study characteristics or dementia classification on the prevalence of OAC in people with and without dementia (Figures 1-5 and 8-9 within Appendix 4 of Supplemental Material). Additionally, to assess increasing prevalence of OAC over time, a sensitivity analysis was conducted that included studies published during or after 2010 only which showed a similar pooled odds ratio to the overall odds ratio (Figure 2, Appendix 4 in

<u>Supplemental Material</u>). However, sensitivity analysis that included studies with  $\geq 30\%$  of the study sample with a prior history of stroke or TIA demonstrated a higher prevalence of OAC use for cardioembolic stroke prevention in AF compared to people without dementia or CI (OR 0.58, 95% CI=0.43–0.79, p<0.00001) (Figure 6 of Appendix 4 in Supplemental Material).

### Safety and effectiveness outcomes of oral anticoagulant use

Safety and effectiveness outcomes of oral anticoagulant use for cardioembolic stroke prevention in AF for people with and without dementia or CI are summarised in table 3. Differences in effectiveness and safety were reported for dementia/CI and non-dementia/CI groups. It was not possible to conduct a meta-analysis on the safety and effectiveness of OACs. Data on the safety and effectiveness of OACs from each study were reported separately.

#### Effectiveness outcomes

One study reported that the composite outcome of stroke, non-central nervous system (CNS) embolism, myocardial infarction (MI), vascular death, and all-cause death was significantly lower for people without dementia than for people with dementia (HR 0.46, 95% CI, 0.27-0.78, p=0.002). When controlled for TTR, there was no increased risk for the composite outcome in the dementia group (adjusted HR 0.72, 95% CI, 0.45-1.14, p=0.155) [18]. Results for studies of smaller samples suggested that rates of thrombosis [56], stroke, and mortality [23] were not different for dementia and non-dementia groups (table 3).

#### Safety outcomes: anticoagulation control

Four studies reported varied results regarding anticoagulation control. One study found that people with CI residing in the community had poorer anticoagulation control than people without CI. People with CI (MMSE score <24) demonstrated lower mean percentage of TTR

(mean  $\pm$  standard deviation (SD) 38 $\pm$ 26) compared to people without cognitive impairment (MMSE score >27), (mean (SD) 61 $\pm$ 27), p< 0.0001) [71]. Results of another study demonstrated that long-term warfarin users with CI monitored within a pharmacist-managed anticoagulation clinic also spent reduced TTR compared with warfarin users without CI, but the result was not statistically significant (TTR % mean (SD) 61 $\pm$ 16 (MMSE  $\leq$ 26), 65 $\pm$ 20 (MMSE >26), p=0.36 [56]. Further descriptive results in another study indicated patients monitored in an anticoagulation clinic with an MMSE score less than 23 spent 68% of TTR compared with 76% for those with an MMSE 23 and above [72]. In addition, no differences for percentage of days with subtherapeutic, therapeutic and supratherapeutic INR values were found for people with and without dementia in long-term care [58].

## Safety outcomes: adverse events

Total bleeding (minor and major) was found to be significantly lower for people without dementia than for those with dementia (HR) 0.56, 95% CI, 0.37-0.85) [18]. Although, in two studies, no significant differences were found for rates of minor and major bleeding and haemorrhage between dementia and non-dementia groups [23, 56]. Adverse warfarin events (AWEs) (injuries from warfarin) were significantly higher for residents in long-term care with dementia (adjusted incidence rate ratio (IRR) 1.48, 95% CI, 1.20-1.82). Risk of potential or preventable AWEs which constituted an INR value greater than 4.5 was also higher (adjusted IRR 1.36, 95% CI, 1.06-1.76) [58] (table 3).

Table 1. Methodological characteristics of included studies of prevalence and outcomes of oral anticoagulant use in people with and without dementia or cognitive impairment (by year of publication)									
First author (year)	Study design, country and health care setting	Population (N), description of study sample and study data source(s)	Anticoagulant reviewed and main indication(s)	Dementia type reported, data source and measurement method	Time of data collection				
	Articles relating to prevalence of oral anticoagulant use (by year of publication)								
		N=370		Condition reported:					
	Cross-sectional	Patients diagnosed with an acute stroke or TIA		Cognitive impairment					
Deplanque	Five countries: Austria, Belgium,	with known AF (paroxysmal or permanent) -	Warfarin		September 2001 - June				
(2004)[65]	France, Italy and Portugal	on admission to hospital	Stroke prevention in AF	Data source and measurement method:	2002				
	Hospital	Medical histories from a variety of sources:		Ascertainment of documentation of cognitive impairment diagnosis from					
		GP, patient interview, family, cardiologist		study data (derived from medical history taking)					
	Cross-sectional	N=117	Warfarin	Condition reported: Dementia					
Latif (2005)[51]	USA	Nursing home residents with AF	Stroke prevention in AE	Data source and measurement method:	Not specified				
	Residential Aged Care Facility	Medical charts and administrative data	Subke prevention in Ar	Dementia diagnosis within the nursing home medical charts					
Choudhry (2006)[60]	Cross-sectional Canada Community	N=116200 <sup>a</sup> Patients with an identifiable cardiac provider         Data sources:         1. Canadian Institutes of Health Information         database         2. The Ontario Drug Benefits claims database         3. Ontario Health Insurance Plan         4. Ontario Registered Persons database         5. Corporate Providers Database of the Ontario         Ministry of Health         6. Southarn Medical database		Condition reported: Dementia Data source and measurement method: Presence of dementia diagnosis coding (hospital ICD-9 codes 290.1 to 290.4, 290.8, 290.9, 294.1, 331.0, 331.1, 331.2 046.1, 046.2) in hospital administrative data	1 January, 1994 – March 31, 2002				
		N=320 (subset of Deplanque 2004[65])		Condition reported:					
	Cross-sectional	Patients with AF who have suffered ischaemic		Cognitive impairment					
Deplanque	Five countries: Austria, Belgium,	stroke and were being discharged from	Warfarin		September 2001 – June				
(2006)[66]	France, Italy and Portugal	hospital	Stroke prevention in AF	Data source and measurement method:	2002				
	Hospital	Medical histories from a variety of sources:		Ascertainment of documentation of cognitive impairment diagnosis from					
		GP, patient interview, family, cardiologist		study data (derived from medical history taking)					
U-1-1-(2007)[55]	Cross-sectional	N=405	Warfarin	Condition reported: Cognitive impairment/dementia	January 2001 -				
Hylek (2006)[55]	USA	Hospitalized patients with AF	Stroke prevention in AF	Madical discussion of domentic within the bognital madical magard	June 2003				
	Prospective cohort	N=204		Condition reported:					
Lefebyre (2006)[68]	France and Italy	Patients diagnosed with an acute stroke or TIA	Warfarin	Cognitive impairment	September 2001 – June				
2000/[00]	Hospital	with known AF (paroxysmal or permanent)	Stroke prevention in AF	Data source and measurement method:	2002				
	nospitai	Medical histories from a variety of sources:	Saoke provention in M	Ascertainment of documentation of cognitive impairment diagnosis from	2002				
1	1		1	input the decomposition of cognitive imput them diagnosis from	1				

		GP, patient interview, family, cardiologist		study data (derived from medical history taking)	
Lopponen (2006)[69]	Cross-sectional Finland Community	N=409 Patients aged 75 years and older with CVD Patient interview, laboratory and clinical examinations	Warfarin Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Two stage process: 1) MMSE, 2) Interview covering items of the Hachinski Ischaemic Scale and the Clinical Dementia Rating. Dementia was also assessed in clinical examination according to DSM-IV criteria, diagnosis of possible Alzheimer's disease according to the NINCDS-ADRDA criteria and diagnosis of possible vascular dementia according to the NINDS-AIREN criteria	1998 – 1999
Partington (2007)[59]	Cross-sectional Canada Hospital	N=196 (entire study sample) N=106 (patients eligible for anticoagulation in which dementia stratification presented) Patients with AF and acute ischaemic stroke EMR data	Warfarin Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Dementia documentation in primary diagnoses and comorbid conditions from the hospital's EMR	1999 – 2004
Doucet (2008)[67]	Cross-sectional France Hospital	N=209 Patients $\geq$ 65 years with chronic AF Medical charts	Warfarin Stroke prevention in AF	Condition reported: Dementia (mean MMSE for anticoagulant and aspirin groups provided) Data source: Medical charts	January 2004 – April 2005
De Breucker (2010)[64]	Cross-sectional Belgium Hospital	N=111 Patients admitted to an acute geriatric unit at an academic hospital Computerized medical charts	Vitamin K Antagonist (exact medication not specified) Stroke prevention in AF	Condition reported: Cognitive disorders Data source and measurement method: Documentation of cognitive disorders within comprehensive geriatric assessments	April 2006 – November 2008
Ewen (2012)[54]	Retrospective longitudinal cohort study USA Community	N=1141 Patients with AF EMR data, hospital administrative data	Warfarin Stroke prevention in AF	Condition reported: Cognitive dysfunction Data source and measurement method: EMR problem list, hospital administrative data	January 1, 1998 – June 30, 2010
Holt (2012)[62]	Longitudinal series of cross- sectional surveys United Kingdom Community	N=59804 Patients with AF QResearch database	Specific anticoagulant(s) not specified Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Read code for dementia within the QResearch database	2007-2010 (2010 data only presented in this paper)
Scowcroft (2012)[24] Mohammed	Retrospective cohort United Kingdom General Practice Cross-sectional	N=81381 Patients aged >60 years with a new diagnosis of AF United Kingdom General Practice Research Database N=50361	Warfarin Stroke prevention in AF Warfarin	Condition reported: Alzheimer's disease/dementia Data source and measurement method: Presence of dementia Read Code in the United Kingdom General Practice Research Database Condition reported: Dementia	2000 – 2009 1 May 2010
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(2013)[19]	United Kingdom	Patients with a diagnosis of AF (≥ 35 years of	Stroke prevention in AF	Data source and measurement method:	
	General Practice	age).		Dementia Read Code present within patient records of the Health	
		The Health Improvement Network (THIN)		Improvement Network (THIN) database	
		database			
				Condition reported:	
Reardon (2013)[57]	Cross-sectional	N=5211 Long-term care residents with AF	Warfarin	Dementia/cognitive impairment Data source and measurement method:	2004 and
	USA	National Nursing Home Survey and the	Stroke prevention in AF	Presence of dementia or cognitive impairment diagnosis within the minimum	1 January 2007 –
	Long-term care	AnalytiCare Long-Term Care databases		data set of the AnalytiCare Long-Term Care database or from comorbid	30 June 2009
		· ······, · · · ···· · ·····g · · · ··· · · ·		condition information in the National Nursing Home Survey database	
				Condition reported: Domentia	
	Cross sectional	N-21096		Data source and measurement method:	
Dreischulte	Scotland	Patients with AE	Warfarin	Quality and Outcomes defined Read Codes for dementia or prescription for	31 March 2007
(2014)[61]	Community	Souttish Conoral Practice date	Stroke prevention in AF	quanty and outcomes defined Read Codes for dementia of prescription for	
Community		Scottish General Fractice data		conoral practices	
		N. 1929		general practices	
		N=1828 Patients >18 years with index event of stroke		Condition reported:	
	Crear and inval	or TIA; and diagnosed AF and a minimar	Di	Dementia	
Tenisley (2014)[70]	Cross-sectional	physical impairment and direct discharge after	Phenprocountaron, wartarin	Data source and measurement method:	2004 2010
Tanislav (2014)[70]	Germany	acute treatment or referral to a renabilitation		Presence of dementia ICD-10 codes within the claims data from a nationwide	2004 - 2010
	Hospitai	racinty.	Stroke prevention in AF	statutory health insurance company (F00, F01, F02, F03, G30)	
		Registry data of the Institute of Quality			
		Assurance Hesse and Claims data from a			
		nationwide statutory health insurance company			
	Cross-sectional	N=1085		Condition reported: Cognitive impairment	
Bahri (2015)[63]	France	Nursing home residents over 75 years with a	Warfarin		March 2012
	Long-term care	documented history of AF	Stroke prevention in AF	Data source and measurement method: Documentation of dementia/cognitive	
		Medical charts		impairment with or without formal assessment from medical records	
	Cross-sectional	N=1225 Patients with hip fracture secondary to a high	Chronic anticoagulation therapy (CAT) (exact	Condition reported: Dementia Data source and measurement method:	
Formiga (2016)[50]	Spain	energy impact	medication not provided)	Short Portable Mental Status questionnaire from the comprehensive geriatric	Not provided
	Hospital	Hospital medical records	Indication not provided	assessment	
	Retrospective cohort	N=5781	Warfarin dahigatran	Condition reported: Dementia	
Shah (2016)[17]	Canada	Patients $\geq$ 65 years with AF hospitalized from	rivaroxahan and anixahan	Data source and measurement method:	1 July 2003 -
	Hoepital	ischaemic stroke or TIA	Stroke prevention in AE	Presence of dementia diagnosis within the comorbid condition information in	31 December 2011
	riospitat	Databases: Ontario Stroke Registry, Canada	Subke prevention in AF	Ontario Stroke Registry	

McGrath (2017)[20]	Retrospective cohort United States of America Hospital	Census, Ontario Drug Benefits, Canadian Institute for Health Information Discharge Abstract and the National Ambulatory Reporting System N=1405 Individuals with AF and acute ischaemic stroke surviving hospitalization Kaiser Permanente database	Warfarin Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Dementia documentation in medical records extracted from structured chart review	July 1996 – September 2003
		Articles relating to outcomes	from oral anticoagulant us	e (by year of publication)	
Van Deelen (2005)[72]	Retrospective cohort study The Netherlands Community	$\label{eq:N=152} \begin{split} N{=}152 \\ Patients \geq 70 \mbox{ years with AF treated with} \\ acenocoumarol managed by an anticoagulation} \\ service \end{split}$	Acenocoumarol (nicoumalone) Stroke prevention in AF	Condition reported: Cognitive impairment Data source and measurement method: MMSE during home visit on index date. Patients with MMSE < 23 were considered cognitively impaired.	March – May 2003
Jacobs (2009)[23]	Retrospective cohort study United State of America Community	N=106 Patients ≥ 65 years with chronic AF receiving warfarin or aspirin Medical records	Warfarin Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Documentation of dementia in medical records	2003
Flaker (2010)[18]	Post-hoc analysis of a randomized controlled trial 522 centres/31 countries Community	N=2510 Community patients with AF and an additional risk factor for stroke ACTIVE-W study data [73]	Warfarin Stroke prevention in AF	Condition reported: Cognitive impairment Data source and measurement method: Presence of cognitive impairment within clinical trial data which used a modified MMSE	June 2003 and December 2004
Khreizat (2012)[56]	Retrospective cohort study United States of America Community	N=57 Community patients aged ≥ 60 years on warfarin with target INR of 2-3. Medical charts	Warfarin Stroke prevention in AF and treatment of VTE	Condition reported: Cognitive impairment Data source and measurement method: Cognitive assessment was part of routine care using the Folstein MMSE. Cognitive impairment was defined as having a MMSE ≤ 26 . A lower cut point of MMSE ≤ 23 was also used to see if it impacted results	2006-2010
Tija (2012)[58]	Prospective cohort study (embedded within a clinical trial) United States of America Long-term care	N=435 Nursing home residents prescribed warfarin Clinical trial data (included medical charts and data abstraction by trained investigators)	Warfarin Stroke prevention in AF Thromboembolic disease Mechanical valve replacement	Condition reported: Dementia Data source and measurement method: Medical record review for dementia diagnosis	1 October 2007 to 31 December 2008
Gorzelak-Pabis (2016)[71]	Retrospective cohort study Poland Community	$\label{eq:resonance} \begin{array}{l} N{=}154\\ Persons with AF and dementia and indications\\ for OAC (CHA_2DS_2VAS_C \geq 1 and HASBLED\\ < 3) \end{array}$	Warfarin and acenocoumarol Stroke prevention in AF	Condition reported: Cognitive impairment Data source and measurement method: Cognitive skills were assessed using the Polish version of the correct MMSE.	2013-2015

		Medical charts		MMSE scores were corrected using Mungas adjustments for age and		
				education level. $MMSE < 27$ was considered cognitive impairment.		
a - study sample was larger, but this group (n-value) were people with an identifiable provider in which dementia information was available						
Abbreviations: AF = atrial fibrillation; CVD = cardiovascular disease; DOAC = direct oral anticoagulant; TIA = transient ischemic attack; MMSE = Mini-Mental State Examination; VTE = venous thromboembolism; INR = international normalised ratio; GP = general practitioner: ICD-9/ICD-10 =						

International Classification of Diseases and Health Related Problems, 9<sup>th</sup> edition; NINDS-AIREN = National Institute of Neurological and Communicative Disorders and Stroke and the Alzheimer's Disease and Related Disorders Association; NINDS-AIREN = National Institute of Neurological and Communicative Disorders and Stroke and the Association Internationale pour la Recherche et l'Enseignement en Neurosciences.

Table 2. Prevalence of oral anticoagulant use in studies of persons with and without dementia – stratified by healthcare setting (by year of publication)						
Author (year)	Age <sup>a</sup> and gender, % female	Prevalence of dementia (study sample)	Prevalence of anticoagulant use (study sample)	Prevalence of anticoagulant use in persons <u>with</u> dementia	Prevalence of anticoagulant use in persons <u>without</u> dementia	Odds ratio <sup>b</sup> (95% CI)
		Con	munity or General Practice			
Choudhry (2006)[60]	Warfarin users (n=50551) with identifiable providers = 76.2 (6.5), 48.3% Warfarin non-users (n=65649) with identifiable providers = 77.2 (7.1) 49% female	1738/116200 (2%)	50551/116200 (44%)	556/1738 (32%)	49995/114462 (43.7%)	0.61 (0.55 – 0.67)
Lopponen (2006)[69]	CVD+dem: 84.4 (5.7) CVD+no dem: 79.8 (4.4) 66% female	85/409 (21%)	Warfarin use only <sup>d</sup> 24/64 (38%)	5/20 (25%)	19/44 (43.2%)	0.44 (0.14 - 1.42)
Ewen (2012)[54]	70 (13.3) 48% female	87/1141 (8%)	764/1141 (67%)	55/87 (63%)	709/1054 (67.3%)	0.84 (0.53 – 1.32)
Holt (2012)[62]	Median age <sup>e</sup> (at AF diagnosis): 73.0 (IQR=64.0- 81.0), median age (in 2010, of 69762 registered in 2010): 80.0 years (IQR=71.0-87.0) 47% female	374/34041 (1%)	18042/34041 (53%)	108/374 (29%)	17934/33667 (53.2%)	0.36 (0.28 - 0.45)
Scowcroft (2012)[24]	60-69=17054 (21%) 70-79=30350 (37%) 80+=33977 (42%) 52% female	53825/81381 (7%)	37119/81381 (46%)	1376/5382 (26%)	35761/75999 (47.0%)	0.39 (0.36 - 0.41)
Mohammed (2013)[19]	75.6 (11.7), 44% female	2255/50361 (4%)	24064/50361 (48%)	567/2255 (25%)	23497/48106 (48.8%)	0.35 (0.32 - 0.39)
Dreischulte (2014)[61]	75.5 (no SD) 45% female	1034/21096 (5%)	8852/20443 (43% - all current anticoagulation), 11959/20443 (59% - anticoagulant ever since diagnosis)	144/1006 (14%)	8717/19437 (44.8%)	0.21 (0.17 – 0.25)
Total prevalence: community setting		Data combined: 59398/304629 (20%) Mean (%) (Std Dev): 15 (23)	Data combined: 142523/303631 (47%) Mean (%) (Std Dev): 50 (10)	Data combined: 2811/10862 (26%) Mean (%) (Std Dev): 31 (15)	Data combined: 136632/292769 (47%) Mean (%) (Std Dev): 50 (8)	0.40 (0.31 - 0.52)
		•	Hospital		·	
Deplanque (2004)[65]	Median age: 78 (range 29-101)					
Depianque (2004)[65]	58% female	82/370 (22%)	82/288 (29%)	4/41 (10%)	78/329 (24%)	0.35 (0.12 – 1.01)

Deplanque (2006)[66]	< 65: 30 (9.4%) 65-74: 85 (26.6%) ≥ 75: 205 (64.1%) 57% female 80 (no SD)	38/320 (12%)	186/320 (58%)	7/38 (18%)	179/282 (64%)	0.13 (0.06 – 0.31)
Hylek (2006)[55]	58% female	51/405 (13%)	206/405 (51%)	8/51 (16%)	198/354 (56%)	0.15 (0.07 – 0.32)
Lefebvre (2006)[68]	Median age: 78.5 years (range: 54-101), 59% female	24/204 (12%)	53/204 (26%)	2/24 (8%)	51/180 (28%)	0.23 (0.05 - 1.01)
Partington (2007) <sup>[59],e</sup>	OAC 77.7 (8.6), 47% female No OAC 82.0 (9.2), 42% female	22/106 (21%)	57/106 (29%)	12/22 (55%)	45/84 (54%)	1.04 (0.41 – 2.67)
Doucet (2008)[67]	84.7 (7) 61% female	57/209 (27%)	102/209 (49%)	23/57 (40%)	79/152 (52%)	0.63 (0.34 – 1.16)
De Breucker (2010)[64]	84 (5), 72% female	65/111 (59%)	57/111 (51%)	35/65 (54%)	22/46 (48%)	1.27 (0.60 – 2.71)
Tanislav (2014)[70]	77.61 (8.6) 58% female	241/1828 (13%)	827/1828 (45%)	67/241 (28%)	760/1587 (48%)	0.42 (0.31 – 0.56)
Formiga (2016)[50]	82.7 (6) 74% female	249/1225 (20%)	99/1225 (8%)	30/249 (12%)	69/976 (7%)	1.80 (1.14 – 2.83)
McGrath (2016)[20]	79 (9) 54% female	195/1405 (14%)	786/1405 (56%)	67/195 (34%)	719/1210 (59%)	0.36 (0.26 - 0.49)
Shah (2016)[17]	Median age (IQR) No OAC=82 (75-87), OAC=79 (73-85) Females No OAC 54.9% OAC 53%	589/5781 (10%)	4235/5781 (73%)	377/589 (64%)	3858/5102 (76%)	0.57 (0.48 – 0.69)
Total prevalence: Hospital		Data combined: 1613/11964 (13%)	Data combined: 6690/11882 (56%)	Data combined: 632/1572 (40%)	Data combined: 6058/10302 (59%)	
setting	-	Mean (%) (Std Dev): 20 (14)	Mean (%) (Std Dev): 45 (18)	Mean (%) (Std Dev): 31 (20)	Mean (%) (Std Dev): 47 (20)	0.49 (0.33 – 0.73)
			Long-Term Care			
Latif (2005)[51]	84.6 (no SD) 71% female	66/117 (56%)	54/117 (46%)	26/66 (39%)	28/51 (55%)	0.53 (0.25 – 1.12)
Reardon (2013)[57]	NNHS database - median age 85 years 70% female AnalytiCare database - median age 83 years 63% female	1457/5211 (28%)	2176/5211 (42%)	462/1457 (32%)	1714/3754 (46%)	0.55 (0.49 – 0.63)

Bahri (2015)[63]	87.1 (5.3) 73% female	777/1085 (72%)	541/1085 (50%)	357/777 (46%)	541/1085 (50%)	0.86 (0.71 - 1.03)
Total prevalence: Long- Term Care setting		Data combined: 2300/6413 (36%) Mean (%) (Std Dev): 52 (22)	Data combined: 2771/6413 (43%) Mean (%) (Std Dev): 46 (4)	Data combined: 845/2300 (37%) Mean (%) (Std Dev): 39 (7)	Data combined: 2283/4890 (47%) Mean (%) (Std Dev): 50 (5)	0.66 (0.45 - 0.95)
TOTAL FOR ALL STUDIES COMBINED		Data combined: 63311/323006 (20%) Mean (%) (Std Dev): 23 (21)	Data combined: 151984/321926 (47%) Mean (%) (Std Dev): 47 (14)	Data combined: 4288/14734 (29%) Mean (%) (Std Dev): 32 (17)	Data combined: 144793/307961 (47%) Mean (%) (Std Dev): 48 (15)	0.48 (0.40 - 0.58)

a - presented as mean (years) ± standard deviation unless otherwise indicated

b-Odds ratios are crude unless otherwise specified. Crude odds ratios were calculated with data extracted from sub-group analysis of results within research papers

c - Holt et al (2012) - age data are based on the full cohort of 99351 persons. Prevalence data include persons with a CHADS2 score >2 (n=34041) in which dementia stratification was available.

d - Includes patients using warfarin. Patients using antiplatelets excluded

e - Results provided reflect the 106 patients eligible for OAC in which dementia/no dementia stratification was available (n=196 for entire study sample)

Abbreviations: CVD = cardiovascular disease; Dem = dementia; OAC = oral anticoagulation; CI = confidence interval; Std Dev = standard deviation.

Table 3. Outcomes for oral anticoagulant use in persons with and without dementia (by year of publication)					
A		Prevalence of dementia	Outcomes reported that were stratified	Ordenne north	
Autnor (year)	Age and gender, % female	(study sample)	by dementia/non-dementia	Outcome results	
Van Deelen (2005)[72]	Age and gender stratified by % TTR INR 2-3.4 > 70% TT: 78.8 (5.3), 48.5% female INR 2-3.4 > 70% TT: 79.5 (5.3), 50% female	24/152 (15.8%)	Treatment time in therapeutic range	<u>INR with therapeutic range</u> MMSE < 23: 68% of treatment time MMSE ≥23: 76% of treatment time	
Jacobs (2009)[23]	65-75 years, n=17 (16%); 75-85, n=51 (48%); >85, n=38 (36%), 75% female	22/106 <sup>b</sup> (21%)	Mortality, haemorrhage and stroke (17 people with dementia were receiving warfarin and 73 without dementia or falls were receiving warfarin). <i>Results are</i> <i>descriptive</i> .	Mortality           Dementia: 8/17 (47.1%)           No dementia: 10/73 (13.7%) <u>Haemorrhage</u> Dementia: 1/17 (5.9%)           No dementia: 4/73 (5.5%) <u>Stroke</u> Dementia: 0/17 (0%)           No dementia: 2/73 (2.7%)	
Flaker (2010)[18]	70.9 ± 9.5, 65.5% female	365/2510 (14.5%)	Stroke, non-CNS embolism, vascular events, myocardial infarction, total bleeding (minor and major)	$\label{eq:second} \begin{array}{l} \hline Composite of stroke, vascular death, MI or non-CNS embolism\\ \\ MMSE < 26: 6.7 per 100 person-years\\ \\ MMSE \geq 26: 3.6 per 100 person-years\\ \\ Unadjusted HR (95\% CI) = 0.46 (0.27-0.78), p=0.002\\ \\ \\ Adjusted HR (95\% CI) = 0.72 (0.45-1.14), p=0.155\\ \\ \hline \\ \hline \\ \hline \\ Total bleeding (includes major and minor)\\ \\ \\ MMSE < 26: 42 per 100 person-years\\ \\ \\ MMSE \geq 26: 7 per 100 person-years\\ \\ \\ HR (95\% CI) = 0.56 (0.37-0.85), p=0.04\\ \end{array}$	
Khreizat (2012) [56]	New warfarin users MMSE score >26: 79.4 $\pm$ 9.5, 92% female MMSE score $\leq$ 26: 75.6 $\pm$ 6.3, 75% female Long-term warfarin users MMSE score >26: 81.0 $\pm$ 6.9, 68% female	30/57 (53%)	Outcomes were stratified by new warfarin users and long-term users with and without dementia/cognitive impairment Visits/days required to achieve therapeutic anticoagulation (new users); TTR/long-term anticoagulation stability; percentage of clinic visits with reported dose mishaps; frequency of in-range INRs following dose	New warfarin users (n=20; dementia=12, no dementia=8)         Visits to achieve therapeutic anticoagulation         MMSE score >26: $5.8 \pm 4.3$ MMSE score $\leq 26: 4.6 \pm 2.4$ (p=0.44).         Days to reach therapeutic anticoagulation         MMSE score >26: $35.8 \pm 30.5$ MMSE score $\leq 26: 51.6 \pm 45.7$	

	MMSE score $\le 26$ : 74.6 $\pm$ 9.3, 77%		mishaps; minor bleeding; major bleeding;	(p=0.36).
	female		thrombosis (long-term users).	
				Long term warfarin users (n=54; dementia=28, no dementia=26)
				$\underline{\text{TTR}} [\text{mean} \pm \text{SD}]$
				$MMSE \le 26: 61 \pm 16\%$
				$MMSE > 26: 65 \pm 20\%$
				(p=0.36)
				Frequency of dose mishaps
				$MMSE \le 26: 86/691$ visits
				MMSE > 26: 74/705 visits
				(p=0.18)
				In-range INRs following dose mishaps
				$MMSE \leq 26: 16\%$
				MMSE > 26: 32%
				(p=0.013)
				Minor bleeding (per patient-year)
				$MMSE \le 26: 0.20 \pm 0.42$
				$MMSE > 26: 0.28 \pm 0.54$
				(p=0.51)
				Major bleeding (per patient-year)
				$MMSE \le 26: 0.02 \pm 0.10$
				$MMSE > 26: 0.07 \pm 0.25$
				(p=0.29)
				Thrombosis (per patient-year)
				$MMSE \le 26:0$
				MMSE > 26: 0.01±0.06
				(p=N/A)
	Dementia		Number of INR tests; percentage of days	Number of INR tests, mean (SD)
	$83.6 \pm 9.3, 74\%$ female		with subtherapeutic, therapeutic and	Dementia: 24.2 (13.9)
Tija (2012)[58]	No dementia	218/435 (50%)	supratherapeutic INRs; incidence of AWEs	No dementia: 26.0 (14.5)
	$80.4 \pm 11.6, 61\%$ female		(injuries from warfarin), incidence of	(p=0.017)
			preventable and potential AWEs (INRs $>$	

			4.5), adjusted association of dementia with	INR < 2, % (SD)
			AWEs and preventable and potential AWEs	Dementia: 37.8 (23.2)
				No dementia: 37.7 (20.4)
				(p=0.95)
				<u>INR &lt; 2-3, % (SD)</u>
				Dementia: 49.5 (22.2)
				No dementia: 48.6 (19.9)
				(p=0.72)
				<u>INR &lt; 3-4.5, % (SD)</u>
				Dementia: 10.7 (9.8)
				No dementia: 11.7 (12.2)
				(p=0.34)
				<u>INR &gt;4.5, % (SD)</u>
				Dementia: 2.1 (6.7)
				No dementia: 2.0 (7.1)
				(p=0.82)
				Incidence rates of AWEs (injuries from warfarin, events per 100 resident-months)
				Dementia: 12.8
				No dementia: 9.99
				IRR (95% CI) = 1.40 (1.14-1.72) – after adjustment for resident characteristics
				IRR (95% CI) = 1.48 (1.20-1.82) – after adjustment for nursing home characteristics and case mix
				Incidence of preventable and potential AWEs (INR >4.5, events per 100 resident-months)
				Dementia:8.09
				No dementia: 6.50
				IRR (95% CI) = 1.35 (1.04-1.74) – after adjustment for resident characteristics
				IRR (95% CI) = 1.36 (1.06-1.76) – after adjustment for nursing home characteristics and case mix
	MMSE score > $27:73 \pm 0.610$			Mean TTR, % (mean ± SD):
Correlate Pabie	formula $5 \text{ COLC} \leq 27.73 \pm 7,0170$			$MMSE < 27: 38 \pm 26$
(2016)[71]	MMSE $aaara < 27, 77 \pm 11, 600/$	42/104 (40%)	Mean TTR and INR values	$MMSE \ge 27:61{\pm}27$
(2010)[/1]	formalo $< 2/1 / 1 \pm 11,09\%$			(p<0.0001)
	iciliale			

	$\underline{\text{TTR}} > 60, \text{ n (\%)}$ :				
	MMSE < 27: 12/42 (28%)				
	MMSE ≥ 27: 38/62 (61%)				
	(p<0.0001)				
	<u><i>INR</i> &lt; 2</u> , n (%):				
	MMSE < 27: 19/42 (46%)				
	MMSE ≥ 27: 37/62 (59%)				
	(p<0.05)				
	<u>INR 2-3, n (%):</u>				
	MMSE < 27: 11/42 (26%)				
	MMSE ≥ 27: 37/62 (60%)				
	(p<0.05)				
	<u><i>INR</i> &gt; 3</u> , n (%):				
	MMSE < 27: 12/42 (28%)				
	MMSE ≥ 27: 14/62 (22%)				
	(p<0.05)				
a - presented as mean (years) ± standard deviation unless otherwise indicated					
- 112 patients in study sample, but 106 undergoing antithrombotic treatment					

Abbreviations: TTR = time in therapeutic range; INR = international normalized ratio; OR = odds ratio; HR = Hazard Ratio; CI = confidence interval; N/A=not applicable; AWEs = adverse warfarin-associated events; IRR = incident rate ratio; TT = treatment time; CNS = central nervous system

## DISCUSSION

To our knowledge, this is the first systematic review to investigate the prevalence and outcomes of OAC use for cardioembolic stroke prevention in AF in people with and without dementia or CI. There are three major findings from the review. First, people with dementia had 52% lower odds of receiving OAC for embolic stroke prevention associated with AF than people without dementia. Mean OAC prevalence for people with dementia was 32% compared with 48% for people without dementia. Over the time period 1998 to 2012, OAC prevalence increased for both groups for all healthcare settings combined. Second, six studies compared safety and effectiveness outcomes of OAC use among people with and without dementia, with all studies investigating diverse outcomes. This heterogeneity precludes a meta-analysis of outcomes data to accurately determine whether people with or without dementia have different outcomes of OAC use in people with dementia. No DOAC safety or effectiveness studies identified by our search strategy have included representative samples of persons with dementia or presented sub-analyses for people with dementia.

People with dementia were less likely to receive OAC than people without dementia. Possible reasons for OAC underuse include: frailty, falls risk, active or prior bleeding, fear of bleeding complications, comorbidities, poor adherence, difficulties with self-monitoring, poor anticoagulation control and polypharmacy [10, 18, 51, 70, 71, 74]. Results from the European Heart Rhythm Association EP Wire survey found that 40% of respondents considered dementia as a key reason not to prescribe OAC. The only more important reason cited was prior or active bleeding or increased bleeding risk [74]. Yet it remains unclear to what extent dementia is associated with lower use of OAC independent of other factors that may contraindicate the prescription of OAC [75, 76]. Ultimately, people with dementia are more likely to experience substantial comorbidity, frailty and polypharmacy [75]. In a sample of

people with AF and dementia at high stroke risk but without increased bleeding risks or absolute contraindications to OAC, it was found that 22% of people received inadequate OAC and 39.5% received no OAC [76]. Further, at the time of dementia diagnosis, 26% of people with AF received warfarin, 37% antiplatelet therapy and 37% did not receive either antiplatelet or OAC [21]. While in people receiving warfarin therapy who were subsequently diagnosed with dementia, 16% remained on warfarin after dementia diagnosis compared with 96.7% of people who were not diagnosed with dementia [77]. Reluctance to prescribe OAC or an inclination to cease OAC in people with dementia could demonstrate that physicians perceive dementia as a limiting factor for OAC, possibly due to perceived increased bleeding risk or lack of adherence [74-76]. Moreover, high thromboembolic risk is often undervalued in ageing individuals with comorbid illness [78] and clinicians may be uncertain whether older, frail people, such as people with dementia could benefit from stroke reduction and whether this counterbalances the risk of bleeding [77, 78]. Our review demonstrates OAC under use in people with dementia and AF and possible higher bleeding risks. However, the risk-benefit of treatment for people with dementia may still provide net clinical benefit. Recent analysis of data from the Swedish Dementia Registry demonstrates lower risk of ischemic stroke and mortality, with only a small increase in any-cause haemorrhage in people with AF and dementia treated with warfarin [21]. Collectively, results may demonstrate that patients people with dementia and AF should not routinely be excluded from OAC treatment despite a slightly higher bleeding risk.

Over the time period of 1998 to 2012, increasing OAC prevalence was observed for both dementia and non-dementia groups. When stratified by healthcare setting, OAC prevalence for people with dementia in a hospital setting demonstrated the greatest increase. Medical practitioner characteristics and healthcare setting (hospital, community, long term care) have been found to influence OAC prescribing. It has been demonstrated that cardiologists have

increased guideline adherence, whereas General Practitioners (GPs) were less adherent [79]. Specialist therapeutic recommendations from neurology [70] facilitates the prescription of OAC, and follow-up by cardiologists and younger GPs were strong predictors of VKA treatment [65]. Patients treated at primary stroke centres and large academic hospitals were more likely to receive thromboprophylaxis than patients treated at smaller or general hospitals [34]. Residing in long term care is a negative predictor of being discharged from hospital with OAC [34, 66]. It is not possible to quantify the influence of practitioner characteristics and healthcare setting on our results, however future studies could confirm the effect of these factors on OAC use, particularly for people with dementia and since the introduction of the DOACs.

The results of this study reflect a low prevalence of OAC use for cardioembolic stroke prevention in AF in <u>patients-people</u> with (48%) and without dementia or CI (32%). These results suggest possible under treatment in high risk populations for stroke. These results suggest limited compliance with current stroke prevention guidelines, especially among people with dementia. Alternatively, data included in this was averaged over an extended time period (2000-2017), which could mask the possible magnitude of changing rates of anticoagulation prevalence rates. Further, only one study included in this review provided data on DOAC use in dementia and non-dementia groups. Recent Australian and Norwegian studies have suggested that the overall prevalence of OAC use has increased since the availability of DOACs, particularly for octogenarians [41, 42].

Insufficient studies were identified in this present review to provide enough comparative information or to conduct a meta-analysis for outcomes of OAC use in persons with and without dementia. Two studies demonstrated that people with dementia have poorer anticoagulation control during treatment with VKA and spend more time below therapeutic range than people without dementia [56, 71]. Results that demonstrate a relationship between

CI and low TTR should not be directly interpreted as cause and effect, as other reasons could influence low TTR, although, it is clinically intuitive. Safe administration of thromboprophylaxis is heavily reliant on self-care. Poor self-care has been identified as a major contributor to hospital readmission and poor health outcomes in patients with heart failure [80]. This could also be expected for AF. People with dementia or CI could have difficulty in acquiring knowledge of chronic disease and medications. A thorough understanding of chronic illness and intact executive function are crucial for managing chronic disease [81, 82]. Limited executive functioning influences the ability to recognise symptoms and make decisions [83], which may result in poor in-range INRs and harm for people with dementia receiving OAC.

The composite outcome of stroke, non-CNS embolism, vascular death, MI and mortality was found to be significantly higher for people with dementia than those without, but when controlled for TTR, there was no increased risk [18]. This suggests that improving TTR for people with dementia could reduce embolic events. Further, two studies found that thrombosis [56], stroke and mortality [23] were not different for dementia and non-dementia groups, however these studies were limited by small numbers. Conflicting results were found for rates of bleeding events between dementia and non-dementia groups. One study demonstrated increased risk of total bleeding in people with dementia [18] and non-significant differences were found in a further two studies [23, 56].

Poor anticoagulation control is a known deterrent for prescribing OAC [75, 77, 84]. Poor anticoagulation control is closely correlated with embolic stroke, haemorrhage and mortality [85-87]. Given potential difficulties in achieving good anticoagulation control in persons with dementia receiving VKA, this may explain why proportionally less people with cognitive impairment receive anticoagulation than do people without cognitive impairment. DOACs circumvent some limitations of warfarin, such as the need for routine monitoring, and have

more predictable pharmacokinetics [40], and are simpler to use than VKA which may improve adherence [88], hence in people with cognitive impairment DOACs could alternatively be considered [89]. Indeed, the European Society of Cardiology guidelines recommend switching those with poor INR control to DOACs [22], but as yet there is little evidence to support this recommendation. DOACs directly inhibit thrombin (dabigatran) and factor Xa (apixaban, rivaroxaban and edoxaban) [90]. DOACs have a rapid onset of action, shorter half-lives and do not affect factor VII. These mechanisms could decrease bleeding risk; particularly limiting traumatic intracranial bleeding related to falls [91] which is critical when considering OAC for people with dementia. Dementia, per se, can impair medication adherence [92], but comorbidity burden [93] and polypharmacy [94] are known to reduce medication adherence, of which there is increased occurrence in persons with AF and dementia [94]. These areas require thorough investigation to understand the risks and benefits of DOACs in people with dementia.

#### Limitations

Our study has several limitations. First, the primary data sources have limitations in that comparisons are derived from sub-group analyses of observational studies. These studies did not examine anticoagulation in relation to cognitive status as the main objective. Crude ORs were therefore calculated and no adjustments have been made for variables confounding the prevalence of OAC in dementia/CI and non-dementia/CI groups. Further, information about cognitive status may be limited. For example, dementia and CI were defined in different ways in various studies, and the severity of dementia was not consistently reported. The effect of the use of data obtained from sub-groups of large studies and the heterogeneity of dementia definitions on our findings is unknown. <u>Our meta-analyses showed substantial heterogeneity between studies demonstrated by high I<sup>2</sup> values and caution should be used when interpreting findings. Participants of the studies included in this review that were documented to have had</u>

CI may have been more likely to have marked CI for it to have been documented. Hence, the observed results may not be generalizable to all people with CI, and this could underestimate the use of OAC in persons with dementia and CI. In addition, we did not assess how the diagnosis or detection of AF occurred for each study. Variability in AF detection rates could influence prescribing of OACs, which could impact the generalizability of the findings of this review to the general population. Further, given the heterogeneity of approaches taken and various safety and effectiveness outcomes reported in the outcomes studies, it was not possible to average or meta-analyse safety and effectiveness outcomes data. The methodological quality of included studies that determined prevalence of OAC use was generally sound. Five prevalence studies did not score maximum points of quality assessment as inclusion criteria were not clearly defined, exposure and outcomes measurements were unclear, and objective, standard criteria for measurement of diagnoses and conditions were not used. Three studies evaluating outcomes of OAC use for people with and without dementia did not provide adequate information to measure exposure (OAC use) and two studies were descriptive and therefore no adjustment for confounding factors was made, which limits the quality. Further, studies were conducted in the UK, the rest of Europe and North America which may limit the generalizability of results to other countries and healthcare systems.

### CONCLUSION

People with atrial fibrillation who also have dementia are less likely to receive OAC for stroke prevention than people without dementia. There is a dearth of information regarding the outcomes of OAC use for stroke prevention in AF in people with dementia and CI. Given the increasing use of the DOACs, in particular within older age groups, the declining use of warfarin, and the limited generalizability of study findings from pivotal DOAC trials and various observational studies to people with dementia, there is an urgent need for more

information. Studies of the safety of OAC specifically in people with AF and dementia of various types, investigating the OAC type, dose, and adherence are urgently needed to guide treatment.

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## **Conflicts of interest**

The authors have no conflicts of interest to declare.

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Figure 1. Literature flow diagram of studies identified, screened and included in the metaanalysis and systematic review; OAC = oral anticoagulation.



**Figure 2.** Mean prevalence of OAC use: overall, and stratified by community, hospital and long-term care healthcare settings for dementia/CI and non-dementia/CI groups. OAC = oral *anticoagulation; CI = cognitive impairment.* 



**Figure 3.** OAC prevalence by mid-year of study observation period: overall and stratified by community, hospital and long-term care healthcare settings for dementia and non-dementia groups, by mid-year of study observation period. *Vertical-axis, prevalence of OAC (%); Horizontal-axis, publication year; Red square and trend line = non-dementia; Blue diamond and trend line = dementia/cognitive impairment; OAC = oral anticoagulation.* 



Figure 4. Forest plots of oral anticoagualtion use in people with and without dementia or cognitive impairment for 1.1.1) for all healthcare settings, and then subgroup analysis according to healthcare setting: 1.1.2) studies conducted in the community 1.1.3) studies conducted in hospitals and 1.1.4) studies conducted in long-term care

	Deme	ntia	No den	nentia		Odds Ratio		Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year	M-H, Random, 95% Cl
1.1.1 All studies for a	ill healthc	are sett	ings					
Deplanque 2004	4	41	78	329	2.2%	0.35 [0.12, 1.01]	2004	
Choudbry 2006	20 556	00 1729	28 70005	114462	3.4% 6.0%	0.53 [0.25, 1.12]	2005	
Deplanque 2006	550	38	49995	282	2,9%		2000	
Hvlek 2006	. 8	51	198	354	3.2%	0.15 [0.07, 0.32]	2006	_ <b>_</b>
Lefebvre 2006	2	24	51	180	1.3%	0.23 [0.05, 1.01]	2006	
Lopponen 2006	5	20	19	44	1.9%	0.44 [0.14, 1.42]	2006	
Partington 2007	12	22	45	84	2.6%	1.04 [0.41, 2.67]	2007	
Doucet 2008	23	57	79	152	4.0%	0.63 [0.34, 1.16]	2008	
De Breucker 2010	35 4076	5000 5000	22	4b 75000	3.3% ഭവസ	1.27 [0.60, 2.71]	2010	
Ewen 2012	1370	000Z 87	33701 709	1054	0.9% 5.0%	0.39 [0.36, 0.41]	2012	
Holt 2012	108	374	17934	33667	6.4%	0.36 [0.28, 0.45]	2012	+
Mohammed 2013	567	2255	23497	48106	6.9%	0.35 [0.32, 0.39]	2013	-
Reardon 2013	462	1457	1714	3754	6.8%	0.55 [0.49, 0.63]	2013	+
Tanislav 2014	67	241	760	1587	6.0%	0.42 [0.31, 0.56]	2014	
Dreischulte 2014	144	1006	8717	19437	6.6%	0.21 [0.17, 0.25]	2014	+
Bahri 2015	357	777	541	1085	6.6%	0.85 [0.71, 1.03]	2015	-
Shah 2016 Formigo 2016	3//	589	3858	5102	5.5% 5.0%	0.57 [0.48, 0.69]	2016	
Furriiga 2016 McGrath 2017	3U 67	249	09 710	970	5.0%	1.80 [1.14, 2.83] 0.36 [0.26, 0.49]	2010	
Subtotal (95% CI)	07	14734	715	307961	100.0%	0.48 [0.40, 0.58]	2017	♦
Total events	4288		144973					
Heterogeneity: Tau <sup>2</sup> =	0.13; Chi	²= 295.	06, df = 2	0 (P < 0.0	0001); P	= 93%		
Test for overall effect:	Z=7.55 (	P < 0.00	)001)					
4.4.2 Chudian annshi				_				
1.1.2 Studies conduc	ted in a c	ommun	ity settin	g 	47.40	0.04 10 55 0.071		
Choudhry 2006	556	1738	49995	114462	17.4%	0.61 [0.55, 0.67]	2006	
Eupponen 2006	5	20	700	1064	079.C 11.604	0.44 [0.14, 1.42]	2000	·
Scoweroft 2012	1376	5382	35761	75999	17.6%	0.39 [0.35, 1.32]	2012	
Holt 2012	108	374	17934	33667	15.7%	0.36 [0.28, 0.45]	2012	+
Mohammed 2013	567	2255	23497	48106	17.4%	0.35 [0.32, 0.39]	2013	•
Dreischulte 2014	144	1006	8717	19437	16.4%	0.21 [0.17, 0.25]	2014	* <b>_</b>
Total events	2811	10802	126622	292709	100.0%	0.40 [0.51, 0.52]		•
Heterogeneity: Tau <sup>2</sup> =	: 0.10: Chi	<sup>2</sup> =137.	100002 59. df = 6	(P < 0.00	001): I <b>P</b> =	96%		
Test for overall effect:	Z = 6.86 (	P < 0.00	0001)		,			
4.4.2 Chudian ann dua								
1.1.5 Studies conduc	ted in a n	ospital	setting	220	0.50	0.05 (0.40, 4.04)	2004	
Deplanque 2004	4	41	/8	329	0.5%	0.35 [0.12, 1.01]	2004	
Lefebyre 2006	2		51	180	4.5%		2000	
Hvlek 2006	8	51	198	354	8.4%	0.15 [0.07, 0.32]	2006	_ <b>_</b>
Partington 2007	12	22	45	84	7.3%	1.04 [0.41, 2.67]	2007	
Doucet 2008	23	57	79	152	9.7%	0.63 [0.34, 1.16]	2008	
De Breucker 2010	35	65	22	46	8.6%	1.27 [0.60, 2.71]	2010	
Tanislav 2014	67	241	760	1587	11.9%	0.42 [0.31, 0.56]	2014	
Formiga 2016	30	249	69	976	10.9%	1.80 [1.14, 2.83]	2016	
Shah 2016 MaGroth 2017	377	589	3858	5102	12.5%	0.57 [0.48, 0.69]	2016	
Subtotal (95% CI)	07	1572	/18	10302	100.0%	0.49 [0.33, 0.73]	2017	•
Total events	632		6058					•
Heterogeneity: Tau <sup>2</sup> =	: 0.31; Chi	<sup>2</sup> = 67.2	8, df = 10	(P < 0.00	001); I <sup>2</sup> =	85%		
Test for overall effect:	Z= 3.58 (	P = 0.00	)03)					
444000								
1.1.4 Studies conduc	ted in a lo	ong tern	1 care se	ating	40.40		0005	
Latif 2005 Bearden 2012	26	1457	28	51	16.1%	0.53 [0.25, 1.12]	2005	
Reardon 2013 Robri 2015	462	1457	1714	3754	43.1%	0.55 [0.49, 0.63]	2013	
Subtotal (95% CI)	357	2300	541	4890	40.8% 100.0%	0.85 [0.71, 1.03] 0.66 [0.45, 0.95]	2015	•
Total events	845	2000	2283			5155 [0140] 0100]		•
Heterogeneity: Tau <sup>2</sup> =	: 0.08; Chi	<sup>2</sup> = 14.7	2, df = 2 (	P = 0.000	6); I <sup>z</sup> = 86	i%		
Test for overall effect:	Z= 2.23 (	P = 0.03	3)					
								0.01 0.1 1 10 100
Toot for outparoup diff	oronooo:	chiz – A	57 df - 3	0.00-0.00	18-24	407		Dementia No dementia

Test for subgroup differences:  $Chi^2 = 4.57$ , df = 3 (P = 0.21), l<sup>2</sup> = 34.4%

Table 1. Methodological characteristics of included studies of prevalence and outcomes of oral anticoagulant use in people with and without dementia or cognitive impairment (by year of publication)					
First author (year)	Study design, country and health care setting	Population (N), description of study sample and study data source(s)	Anticoagulant reviewed and main indication(s)	Dementia type reported, data source and measurement method	Time of data collection
		Articles relating to prevalen	ce of oral anticoagulant use	(by year of publication)	
		N=370		Condition reported:	
	Cross-sectional	Patients diagnosed with an acute stroke or TIA		Cognitive impairment	
Deplanque	Five countries: Austria, Belgium,	with known AF (paroxysmal or permanent) -	Warfarin		September 2001 - June
(2004)[65]	France, Italy and Portugal	on admission to hospital	Stroke prevention in AF	Data source and measurement method:	2002
	Hospital	Medical histories from a variety of sources:		Ascertainment of documentation of cognitive impairment diagnosis from	
		GP, patient interview, family, cardiologist		study data (derived from medical history taking)	
	Cross-sectional	N=117	Warfarin	Condition reported: Dementia	
Latif (2005)[51]	USA	Nursing home residents with AF	Stroke prevention in AE	Data source and measurement method:	Not specified
	Residential Aged Care Facility	Residential Aged Care Facility Medical charts and administrative data		Dementia diagnosis within the nursing home medical charts	
Choudhry (2006)[60]	Cross-sectional Canada Community	N=116200 <sup>a</sup> Patients with an identifiable cardiac provider Data sources: 1. Canadian Institutes of Health Information database 2. The Ontario Drug Benefits claims database 3. Ontario Health Insurance Plan 4. Ontario Registered Persons database 5. Corporate Providers Database of the Ontario Ministry of Health 6. Southam Medical database	Warfarin Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Presence of dementia diagnosis coding (hospital ICD-9 codes 290.1 to 290.4, 290.8, 290.9, 294.1, 331.0, 331.1, 331.2 046.1, 046.2) in hospital administrative data	1 January, 1994 – March 31, 2002
		N=320 (subset of Deplanque 2004[65])		Condition reported:	
	Cross-sectional	Patients with AF who have suffered ischaemic		Cognitive impairment	
Deplanque	Five countries: Austria, Belgium,	stroke and were being discharged from	Warfarin		September 2001 – June
(2006)[66]	France, Italy and Portugal	hospital	Stroke prevention in AF	Data source and measurement method:	2002
	Hospital	Medical histories from a variety of sources:		Ascertainment of documentation of cognitive impairment diagnosis from	
		GP, patient interview, family, cardiologist		study data (derived from medical history taking)	
U-1-1- (2007)[55]	Cross-sectional	N=405	Warfarin	Condition reported: Cognitive impairment/dementia	January 2001 -
Hylek (2006)[55]	USA	Hospitalized patients with AF	Stroke prevention in AF	Madical discussion of domentic within the bognital madical magard	June 2003
	Prospective cohort	N-204		Condition reported:	
Lefebyre (2006)[68]	France and Italy	Patients diagnosed with an acute stroke or TIA	Warfarin	Cognitive impairment	September 2001 – June
2000/[00]	Hospital	with known AF (paroxysmal or permanent)	Stroke prevention in AF	Data source and measurement method:	2002
	riospitai	Medical histories from a variety of sources	Subre prevention in Al	Ascertainment of documentation of cognitive impairment diagnosis from	2002
		incurcal mistories from a variety of soulces.	1	riser annual of documentation of cognitive impairment diagnosis nom	1

		GP, patient interview, family, cardiologist		study data (derived from medical history taking)	
Lopponen (2006)[69]	Cross-sectional Finland Community	N=409 Patients aged 75 years and older with CVD Patient interview, laboratory and clinical examinations	Warfarin Stroke prevention in AF	Condition reported: dementia Data source and measurement method: Two stage process: 1) MMSE, 2) Interview covering items of the Hachinski Ischaemic Scale and the Clinical Dementia Rating. Dementia was also assessed in clinical examination according to DSM-IV criteria, diagnosis of possible Alzheimer's disease according to the NINCDS-ADRDA criteria and diagnosis of possible vascular dementia according to the NINDS-AIREN criteria	1998 – 1999
Partington (2007)[59]	Cross-sectional Canada Hospital	N=196 (entire study sample) N=106 (patients eligible for anticoagulation in which dementia stratification presented) Patients with AF and acute ischaemic stroke EMR data	Warfarin Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Dementia documentation in primary diagnoses and comorbid conditions from the hospital's EMR	1999 – 2004
Doucet (2008)[67]	Cross-sectional France Hospital	N=209 Patients $\geq$ 65 years with chronic AF Medical charts	Warfarin Stroke prevention in AF	Condition reported: dementia (mean MMSE for anticoagulant and aspirin groups provided) Data source: Medical charts	January 2004 – April 2005
De Breucker (2010)[64]	Cross-sectional Belgium Hospital	N=111 Patients admitted to an acute geriatric unit at an academic hospital Computerized medical charts	Vitamin K Antagonist (exact medication not specified) Stroke prevention in AF	Condition reported: Cognitive disorders Data source and measurement method: Documentation of cognitive disorders within comprehensive geriatric assessments	April 2006 – November 2008
Ewen (2012)[54]	Retrospective longitudinal cohort study USA Community	N=1141 Patients with AF EMR data, hospital administrative data	Warfarin Stroke prevention in AF	Condition reported: Cognitive dysfunction Data source and measurement method: EMR problem list, hospital administrative data	January 1, 1998 – June 30, 2010
Holt (2012)[62]	Longitudinal series of cross- sectional surveys United Kingdom Community	N=59804 Patients with AF QResearch database	Specific anticoagulant(s) not specified Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Read code for dementia within the QResearch database	2007-2010 (2010 data only presented in this paper)
Scowcroft (2012)[24]	Retrospective cohort United Kingdom General Practice	N=81381 Patients aged >60 years with a new diagnosis of AF United Kingdom General Practice Research Database	Warfarin Stroke prevention in AF Warfarin	Condition reported: Alzheimer's disease/dementia Data source and measurement method: Presence of dementia Read Code in the United Kingdom General Practice Research Database	2000 – 2009
wonannicu	Cross-sectional	IN-30301	vv arrarni	Condition reported. Dementia	1 Wiay 2010

(2013)[19]	United Kingdom	Patients with a diagnosis of AF (≥ 35 years of	Stroke prevention in AF	Data source and measurement method:	
	General Practice	age).		Dementia Read Code present within patient records of the Health	
		The Health Improvement Network (THIN)		Improvement Network (THIN) database	
		database			
				Condition reported:	
Reardon (2013)[57]	Cross-sectional	N=5211 Long-term care residents with AF	Warfarin	Dementia/cognitive impairment	2004 and
readon (2013)[37]	USA	National Nursing Home Survey and the	Stroke prevention in AF	Presence of dementia or cognitive impairment diagnosis within the minimum	1 January 2007 -
	Long-term care	A palvtiCare Long Term Care databases	Subke prevention in 74	data sat of the AnalytiCare Long Term Care database or from comorbid	30 June 2009
		Analyticale Long-Term Cale databases		condition information in the National Nursing Home Survey database	
		N. 21006		Condition reported: Dementia	
Dreischulte	Cross-sectional	N=21096	Warfarin	Data source and measurement method:	31 March 2007
(2014)[61]	Scotland	Patients with AF	Stroke prevention in AF	Quality and Outcomes defined Read Codes for dementia or prescription for	
	Community	Scottish General Practice data		acetylcholinesterase inhibitor) with the population database of Scottish	
				general practices	
		N=1828			
		Patients >18 years with index event of stroke		Condition reported:	
	Cross-sectional Germany Hospital	or TIA; and diagnosed AF and a minimal		Dementia	
		physical impairment and direct discharge after	Phenprocoumaron, warfarin	Data source and measurement method: Presence of dementia ICD-10 codes within the claims data from a nationwide	
Tanislav (2014)[70]		acute treatment or referral to a rehabilitation	and coumadin		2004 - 2010
		facility.	Stroke prevention in AF	statutory health insurance company (E00 E01 E02 E03 G30)	
		Registry data of the Institute of Quality		statutory nearth insurance company (100, 101, 102, 105, 050)	
		Assurance Hesse and Claims data from a			
		nationwide statutory health insurance company			
		N=1085		Condition reported: Cognitive impairment	
D 1 : (2015)(62)	Cross-sectional	Nursing home residents over 75 years with a	Warfarin		Nr. 1 2012
Bahri (2015)[63]	France	documented history of AF	Stroke prevention in AF	Data source and measurement method: Documentation of dementia/cognitive	March 2012
	Long-term care	Medical charts		impairment with or without formal assessment from medical records	
			Chronic anticoagulation		
	Cross-sectional	N=1225	therapy (CAT) (exact	Condition reported: dementia	
Formiga (2016)[50]	Spain	Patients with hip fracture secondary to a high	medication not provided)	Data source and measurement method:	Not provided
1 oninga (2010)[20]	Hospital	energy impact	Indication not provided	Short Portable Mental Status questionnaire from the comprehensive geriatric	1
	ľ	Hospital medical records	1	assessment	
		N=5781		Condition reported: Dementia	
Shah (2016)[17]	Retrospective cohort	Patients $\geq 65$ years with AF hospitalized from	Warfarin, dabigatran,	Data source and measurement method:	1 July 2003 –
	Canada	ischaemic stroke or TIA	rivaroxaban and apixaban	Presence of dementia diagnosis within the comorbid condition information in	31 December 2011
	Hospital	Databases: Ontario Stroka Pagistry, Conodo	Stroke prevention in AF	Ontario Stroka Pagietry	51 December 2011
		Databases. Offario Subke Kegisu y, Callada	1	Ontario Subre Registry	

McGrath (2017)[20]	Retrospective cohort United States of America Hospital	Census, Ontario Drug Benefits, Canadian Institute for Health Information Discharge Abstract and the National Ambulatory Reporting System N=1405 Individuals with AF and acute ischaemic stroke surviving hospitalization Kaiser Permanente database	Warfarin Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Dementia documentation in medical records extracted from structured chart review	July 1996 – September 2003
		Articles relating to outcomes	from oral anticoagulant us	e (by year of publication)	
Van Deelen (2005)[72]	Retrospective cohort study The Netherlands Community	$\begin{tabular}{l} N=152 \\ Patients \geq 70 \ years \ with \ AF \ treated \ with \\ acenocoumarol \ managed \ by \ an \ anticoagulation \\ service \end{tabular}$	Acenocoumarol (nicoumalone) Stroke prevention in AF	Condition reported: Cognitive impairment Data source and measurement method: MMSE during home visit on index date. Patients with MMSE < 23 were considered cognitively impaired.	March – May 2003
Jacobs (2009)[23]	Retrospective cohort study United State of America Community	N=106 Patients ≥ 65 years with chronic AF receiving warfarin or aspirin Medical records	Warfarin Stroke prevention in AF	Condition reported: Dementia Data source and measurement method: Documentation of dementia in medical records	2003
Flaker (2010)[18]	Post-hoc analysis of a randomized controlled trial 522 centres/31 countries Community	N=2510 Community patients with AF and an additional risk factor for stroke ACTIVE-W study data [73]	Warfarin Stroke prevention in AF	Condition reported: Cognitive impairment Data source and measurement method: Presence of cognitive impairment within clinical trial data which used a modified MMSE	June 2003 and December 2004
Khreizat (2012)[56]	Retrospective cohort study United States of America Community	N=57 Community patients aged ≥ 60 years on warfarin with target INR of 2-3. Medical charts	Warfarin Stroke prevention in AF and treatment of VTE	Condition reported: cognitive impairment Data source and measurement method: Cognitive assessment was part of routine care using the Folstein MMSE. Cognitive impairment was defined as having a MMSE ≤ 26 . A lower cut point of MMSE ≤ 23 was also used to see if it impacted results	2006-2010
Tija (2012)[58]	Prospective cohort study (embedded within a clinical trial) United States of America Long-term care	N=435 Nursing home residents prescribed warfarin Clinical trial data (included medical charts and data abstraction by trained investigators)	Warfarin Stroke prevention in AF Thromboembolic disease Mechanical valve replacement	Condition reported: Dementia Data source and measurement method: Medical record review for dementia diagnosis	1 October 2007 to 31 December 2008
Gorzelak-Pabis (2016)[71]	Retrospective cohort study Poland Community	$\label{eq:resonance} \hline $N=154$$$$$ Persons with AF and dementia and indications for OAC (CHA_2DS_2VAS_C \ge 1 and HASBLED $$<3)$$$$$$$$<$	Warfarin and acenocoumarol Stroke prevention in AF	Condition reported: Cognitive impairment Data source and measurement method: Cognitive skills were assessed using the Polish version of the correct MMSE.	2013-2015

		Medical charts		MMSE scores were corrected using Mungas adjustments for age and		
				education level. $MMSE < 27$ was considered cognitive impairment.		
a - study sample was larger, but this group (n-value) were the patients with an identifiable provider in which dementia information was available						
Abbreviations: AF = atrial fibrillation; CVD = cardiovascular disease; DOAC = direct oral anticoagulant; TIA = transient ischemic attack; MMSE = Mini-Mental State Examination; VTE = venous thromboembolism; INR = international normalised ratio; GP = general practitioner: ICD-9/ICD-10 =						

International Classification of Diseases and Health Related Problems, 9<sup>th</sup> edition or 10 edition; EMR = Electronic Medical Record; DSM-IV = Diagnostic and Statistical Manual of Mental Disorders 4<sup>th</sup> edition; NINCDS-ADRDA = National Institute of Neurological and Communicative Disorders and Stroke and the Alzheimer's Disease and Related Disorders Association; NINDS-AIREN = National Institute of Neurological and Communicative Disorders and Stroke and the Association Internationale pour la Recherche et l'Enseignement en Neurosciences.

Table 2. Outcomes for oral anticoagulant use in persons with and without dementia (by year of publication)					
		Prevalence of dementia	Outcomes reported that were stratified		
Author (year)	Age" and gender, % female	(study sample)	by dementia/non-dementia	Outcome results	
	Age and gender stratified by %TTR				
Van Deelen	INR 2-3.4 > 70% TT:			INR with therapeutic range	
(2005)[72]	78.8 (5.3), 48.5% female	24/152 (15.8%)	Treatment time in therapeutic range	MMSE < 23: 68% of treatment time	
(2003)[72]	INR 2-3.4 > 70% TT:			MMSE $\geq$ 23: 76% of treatment time	
	79.5 (5.3), 50% female				
				Mortality	
				Dementia: 8/17 (47.1%)	
				No dementia: 10/73 (13.7%)	
	65.75 years $p=17.(16\%)$ :		Mortality, haemorrhage and stroke		
	$75 \ 95 \ p=51 \ (49\%)$		(17 people with dementia were receiving	Haemorrhage	
Jacobs (2009)[23]	(3-63, 11-31, (4670),	22/106 <sup>b</sup> (21%)	warfarin and 73 without dementia or falls	Dementia: 1/17 (5.9%)	
	75% formale		were receiving warfarin). Results are	No dementia: 4/73 (5.5%)	
	7.570 ICIIIAIC		descriptive.		
				Stroke	
				Dementia: 0/17 (0%)	
				No dementia: 2/73 (2.7%)	
				Composite of stroke, vascular death, MI or non-CNS embolism	
		365/2510 (14 5%)		MMSE < 26: 6.7 per 100 person-years	
				$MMSE \ge 26: 3.6 \text{ per } 100 \text{ person-years}$	
			Stroke non-CNS embolism vascular	Unadjusted HR (95% CI) = 0.46 (0.27-0.78), p=0.002	
Flaker (2010)[18]	70.9 + 9.5, 65.5% female		events myocardial infarction total bleeding	Adjusted HR (95% CI) = 0.72 (0.45-1.14), p=0.155	
		000/2010 (110/0)	(minor and major)		
			(minor and major)	Total bleeding (includes major and minor)	
				MMSE < 26: 42 per 100 person-years	
				$MMSE \ge 26:7 per 100 person-years$	
				HR (95% CI) = 0.56 (0.37-0.85), p=0.04	
	New warfarin users		Outcomes were stratified by new warfarin	New warfarin users (n=20; dementia=12, no dementia=8)	
	MMSE score >26: 79.4 ± 9.5, 92%		users and long-term users with and without	Visits to achieve therapeutic anticoagulation	
	female		dementia/cognitive impairment	MMSE score >26: $5.8 \pm 4.3$	
	MMSE score $\leq 26$ : 75.6 ± 6.3, 75%			MMSE score $\leq 26$ : 4.6 $\pm$ 2.4	
Khreizat (2012) [56]	female	30/57 (53%)	Visits/days required to achieve therapeutic	(p=0.44).	
			anticoagulation (new users); TTR/long-term		
	Long-term warfarin users		anticoagulation stability; percentage of	Days to reach therapeutic anticoagulation	
	MMSE score >26: 81.0 $\pm$ 6.9, 68%		clinic visits with reported dose mishaps;	MMSE score >26: 35.8 ± 30.5	
	female		frequency of in-range INRs following dose	MMSE score $\le 26: 51.6 \pm 45.7$	

	MMSE score $\leq 26$ : 74.6 $\pm$ 9.3, 77%		mishaps; minor bleeding; major bleeding;	(p=0.36).
	female		thrombosis (long-term users).	
				Long term warfarin users (n=54; dementia=28, no dementia=26)
				$\underline{\text{TTR}} [\text{mean} \pm \text{SD}]$
				$MMSE \le 26: 61 \pm 16\%$
				$MMSE > 26: 65 \pm 20\%$
				(p=0.36)
				Frequency of dose mishaps
				MMSE ≤ 26: 86/691 visits
				MMSE > 26: 74/705 visits
				(p=0.18)
				In-range INRs following dose mishaps
				MMSE ≤ 26: 16%
				MMSE > 26: 32%
				(p=0.013)
				Minor bleeding (per patient-year)
				$MMSE \le 26: 0.20\pm0.42$
				$MMSE > 26: 0.28 \pm 0.54$
				(p=0.51)
				Major bleeding (per patient-year)
				$MMSE \le 26: 0.02\pm0.10$
				$MMSE > 26: 0.07 \pm 0.25$
				(p=0.29)
				Thrombosis (per patient-year)
				$MMSE \le 26:0$
				MMSE > 26: 0.01±0.06
				(p=N/A)
	Dementia		Number of INR tests; percentage of days	Number of INR tests, mean (SD)
	$83.6 \pm 9.3, 74\%$ female		with subtherapeutic, therapeutic and	Dementia: 24.2 (13.9)
Tija (2012)[58]	No dementia	218/435 (50%)	supratherapeutic INRs; incidence of AWEs	No dementia: 26.0 (14.5)
	$80.4 \pm 11.6, 61\%$ female		(injuries from warfarin), incidence of	(p=0.017)
			preventable and potential AWEs (INRs >	

			4.5), adjusted association of dementia with	<u>INR &lt; 2, % (SD)</u>
			AWEs and preventable and potential AWEs	Dementia: 37.8 (23.2)
				No dementia: 37.7 (20.4)
				(p=0.95)
				INR < 2-3, % (SD)
				Dementia: 49.5 (22.2)
				No dementia: 48.6 (19.9)
				(p=0.72)
				INR < 3-4.5, % (SD)
				Dementia: 10.7 (9.8)
				No dementia: 11.7 (12.2)
				(p=0.34)
				<u>INR &gt;4.5, % (SD)</u>
				Dementia: 2.1 (6.7)
				No dementia: 2.0 (7.1)
				(p=0.82)
				Incidence rates of AWEs (injuries from warfarin, events per 100 resident-months)
				Dementia: 12.8
				No dementia: 9.99
				IRR (95% CI) = 1.40 (1.14-1.72) – after adjustment for resident characteristics
				IRR (95% CI) = 1.48 (1.20-1.82) – after adjustment for nursing home characteristics and case mix
				Incidence of preventable and potential AWEs (INR >4.5, events per 100 resident-months)
				Dementia:8.09
				No dementia: 6.50
				IRR (95% CI) = 1.35 (1.04-1.74) – after adjustment for resident characteristics
				IRR (95% CI) = 1.36 (1.06-1.76) – after adjustment for nursing home characteristics and case mix
	NO.407			Mean TTR, % (mean ± SD):
	MMSE score $\ge 27$ : $73 \pm 9$ , 61%			MMSE < 27: 38±26
Gorzelak-Pabis	temale	42/104 (40%)	Mean TTR and INR values	MMSE ≥ 27: 61±27
(2016)[71]	MMSE score < 27: 77 $\pm$ 11, 69%			(p<0.0001)
	temale			

	$\frac{11R > 60, n(\%)}{2}$			
	MMSE < 27: 12/42 (28%)			
	MMSE ≥ 27: 38/62 (61%)			
	(p<0.0001)			
	<u><i>INR</i> &lt; 2, n (%):</u>			
	MMSE < 27: 19/42 (46%)			
	MMSE ≥ 27: 37/62 (59%)			
	(p<0.05)			
	<u>INR 2-3, n (%):</u>			
	MMSE < 27: 11/42 (26%)			
	MMSE ≥ 27: 37/62 (60%)			
	(p<0.05)			
	<u>INR &gt; 3, n (%):</u>			
	MMSE < 27: 12/42 (28%)			
	MMSE ≥ 27: 14/62 (22%)			
	(p<0.05)			
a - presented as mean (years) ± standard deviation unless otherwise indicated	·			
112 patients in study sample, but 106 undergoing antithrombotic treatment				

Abbreviations: TTR = time in therapeutic range; INR = international normalized ratio; OR = odds ratio; HR = Hazard Ratio; CI = confidence interval; N/A=not applicable; AWEs = adverse warfarin-associated events; IRR = incident rate ratio; TT = treatment time; CNS = central nervous system

Table 3. Outcomes for oral anticoagulant use in persons with and without dementia (by year of publication)					
A		Prevalence of dementia	Outcomes reported that were stratified	Ordenne north	
Autnor (year)	Age and gender, % female	(study sample)	by dementia/non-dementia	Outcome results	
Van Deelen (2005)[72]	Age and gender stratified by % TTR INR 2-3.4 > 70% TT: 78.8 (5.3), 48.5% female INR 2-3.4 > 70% TT: 79.5 (5.3), 50% female	24/152 (15.8%)	Treatment time in therapeutic range	<u>INR with therapeutic range</u> MMSE < 23: 68% of treatment time MMSE ≥23: 76% of treatment time	
Jacobs (2009)[23]	65-75 years, n=17 (16%); 75-85, n=51 (48%); >85, n=38 (36%), 75% female	22/106 <sup>b</sup> (21%)	Mortality, haemorrhage and stroke (17 people with dementia were receiving warfarin and 73 without dementia or falls were receiving warfarin). <i>Results are</i> <i>descriptive</i> .	Mortality           Dementia: 8/17 (47.1%)           No dementia: 10/73 (13.7%) <u>Haemorrhage</u> Dementia: 1/17 (5.9%)           No dementia: 4/73 (5.5%) <u>Stroke</u> Dementia: 0/17 (0%)           No dementia: 2/73 (2.7%)	
Flaker (2010)[18]	70.9 ± 9.5, 65.5% female	365/2510 (14.5%)	Stroke, non-CNS embolism, vascular events, myocardial infarction, total bleeding (minor and major)	$\label{eq:composite of stroke, vascular death, MI or non-CNS embolism} \\ MMSE < 26: 6.7 per 100 person-years \\ MMSE \ge 26: 3.6 per 100 person-years \\ Unadjusted HR (95% CI) = 0.46 (0.27-0.78), p=0.002 \\ Adjusted HR (95% CI) = 0.72 (0.45-1.14), p=0.155 \\ \hline Total bleeding (includes major and minor) \\ MMSE < 26: 42 per 100 person-years \\ MMSE \ge 26: 7 per 100 person-years \\ HR (95% CI) = 0.56 (0.37-0.85), p=0.04 \\ \hline$	
Khreizat (2012) [56]	New warfarin users MMSE score >26: 79.4 $\pm$ 9.5, 92% female MMSE score $\leq$ 26: 75.6 $\pm$ 6.3, 75% female Long-term warfarin users MMSE score >26: 81.0 $\pm$ 6.9, 68% female	30/57 (53%)	Outcomes were stratified by new warfarin users and long-term users with and without dementia/cognitive impairment Visits/days required to achieve therapeutic anticoagulation (new users); TTR/long-term anticoagulation stability; percentage of clinic visits with reported dose mishaps; frequency of in-range INRs following dose	New warfarin users (n=20; dementia=12, no dementia=8)         Visits to achieve therapeutic anticoagulation         MMSE score >26: $5.8 \pm 4.3$ MMSE score $\leq 26: 4.6 \pm 2.4$ (p=0.44).         Days to reach therapeutic anticoagulation         MMSE score >26: $35.8 \pm 30.5$ MMSE score $\leq 26: 51.6 \pm 45.7$	

	MMSE score $\le 26$ : 74.6 $\pm$ 9.3, 77%		mishaps; minor bleeding; major bleeding;	(p=0.36).
	female		thrombosis (long-term users).	
				Long term warfarin users (n=54; dementia=28, no dementia=26)
				$\underline{\text{TTR}} [\text{mean} \pm \text{SD}]$
				$MMSE \le 26: 61 \pm 16\%$
				$MMSE > 26: 65 \pm 20\%$
				(p=0.36)
				Frequency of dose mishaps
				$MMSE \le 26: 86/691$ visits
				MMSE > 26: 74/705 visits
				(p=0.18)
				In-range INRs following dose mishaps
				$MMSE \leq 26: 16\%$
				MMSE > 26: 32%
				(p=0.013)
				Minor bleeding (per patient-year)
				$MMSE \le 26: 0.20 \pm 0.42$
				$MMSE > 26: 0.28 \pm 0.54$
				(p=0.51)
				Major bleeding (per patient-year)
				$MMSE \le 26: 0.02 \pm 0.10$
				$MMSE > 26: 0.07 \pm 0.25$
				(p=0.29)
				Thrombosis (per patient-year)
				$MMSE \le 26:0$
				MMSE > 26: 0.01±0.06
				(p=N/A)
	Dementia		Number of INR tests; percentage of days	Number of INR tests, mean (SD)
	$83.6 \pm 9.3, 74\%$ female		with subtherapeutic, therapeutic and	Dementia: 24.2 (13.9)
Tija (2012)[58]	No dementia	218/435 (50%)	supratherapeutic INRs; incidence of AWEs	No dementia: 26.0 (14.5)
	$80.4 \pm 11.6, 61\%$ female		(injuries from warfarin), incidence of	(p=0.017)
			preventable and potential AWEs (INRs >	

			4.5), adjusted association of dementia with	<u>INR &lt; 2, % (SD)</u>
			AWEs and preventable and potential AWEs	Dementia: 37.8 (23.2)
				No dementia: 37.7 (20.4)
				(p=0.95)
				<u>INR &lt; 2-3, % (SD)</u>
				Dementia: 49.5 (22.2)
				No dementia: 48.6 (19.9)
				(p=0.72)
				<u>INR &lt; 3-4.5, % (SD)</u>
				Dementia: 10.7 (9.8)
				No dementia: 11.7 (12.2)
				(p=0.34)
				<u>INR &gt;4.5, % (SD)</u>
				Dementia: 2.1 (6.7)
				No dementia: 2.0 (7.1)
				(p=0.82)
				Incidence rates of AWEs (injuries from warfarin, events per 100 resident-months)
				Dementia: 12.8
				No dementia: 9.99
				IRR (95% CI) = 1.40 (1.14-1.72) – after adjustment for resident characteristics
				IRR (95% CI) = 1.48 (1.20-1.82) – after adjustment for nursing home characteristics and case mix
				Incidence of preventable and potential AWEs (INR >4.5, events per 100 resident-months)
				Dementia:8.09
				No dementia: 6.50
				IRR (95% CI) = 1.35 (1.04-1.74) – after adjustment for resident characteristics
				IRR (95% CI) = 1.36 (1.06-1.76) – after adjustment for nursing home characteristics and case mix
	MMSE score > $27:73 \pm 9.61\%$			Mean TTR, % (mean ± SD):
Correlate Pabie	formula $5 \text{ COLC} \le 27.73 \pm 7,0170$			MMSE < 27: 38±26
(2016)[71]	$MMSE_{000ro} < 27, 77 \pm 11, 600/$	42/104 (40%)	Mean TTR and INR values	$MMSE \ge 27: 61 \pm 27$
(2010)[/1]	formula $< 2/1 / 1 \pm 11,09\%$			(p<0.0001)
	ICIIIAIC			

		<u>TTR &gt; 60, n (%):</u>
		MMSE < 27: 12/42 (28%)
		$MMSE \ge 27: 38/62 (61\%)$
		(p<0.0001)
		<u><i>INR</i> &lt; 2, n (%):</u>
		MMSE < 27: 19/42 (46%)
		MMSE ≥ 27: 37/62 (59%)
		(p<0.05)
		<u>INR 2-3, n (%):</u>
		MMSE < 27: 11/42 (26%)
		$MMSE \ge 27: 37/62 (60\%)$
		(p<0.05)
		<u><i>INR</i> &gt; 3, n (%):</u>
		MMSE < 27: 12/42 (28%)
		$MMSE \ge 27: 14/62 (22\%)$
		(p<0.05)
a - presented as mean (year	rs) ± standard deviation unless otherwise indicated	
b – 112 patients in study s	umple, but 106 undergoing antithrombotic treatment	
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Abbreviations: TTR = time in therapeutic range; INR = international normalized ratio; OR = odds ratio; HR = Hazard Ratio; CI = confidence interval; N/A=not applicable; AWEs = adverse warfarin-associated events; IRR = incident rate ratio; TT = treatment time; CNS = central nervous system

## SUPPLEMENTAL MATERIAL

Appendix 1. Database search strategies (EMBASE, Medline, and CINAHL)

Appendix 2a. Methodological quality of studies checklist for prevalence studies

Appendix 2b. Methodological quality of studies checklist for outcomes studies

Appendix 3a. Results of quality assessment for prevalence studies

Appendix 3b. Results of quality assessment for outcomes studies

**Appendix 4.** Sensitivity analyses investigating sources of heterogeneity in meta-analyses of oral anticoagulant use in people with and without dementia or mild cognitive impairment for all healthcare settings

## Appendix 1. Database search strategies (EMBASE, Medline and CINAHL)

#### EMBASE

Dementia/ 2. dementia.mp. 3. Alzheimer Disease/ 4. alzheimer\*.mp. 5. Cognition Disorders/ 6. cognition disorder\*.mp.
 7. Cognitive Aging/ 8. cognitive aging.mp. 9. Memory Disorders/ 10. memory disorder\*.mp. 11. Mild Cognitive Impairment/ 12. mild cognitive impairment.mp.

**13.** 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12

14. Anticoagulants/ 15. anticoag\*.mp. 16. NOAC.mp. 17. DOAC.mp. 18. Antithrombins/ 19. direct thrombin inhibitor.mp. 20. Warfarin/ 21. warfarin.mp. 22. Dabigatran/ 23. dabigatran.mp. 24. apixaban.mp. 25. Rivaroxaban/ 26. rivaroxaban.mp. 27. edoxaban.mp. 28. VKA.mp. 29. vitamin k antagonist.mp. 30. novel oral anticoagulant.mp. 31. direct oral anticoagulant.mp. 32. Factor Xa Inhibitors/ 33. factor Xa inhibitor\*.mp. 34. 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 or 27 or 28 or 29 or 30 or 31 or 32 or 33 35. 13 and 34

#### MEDLINE

1. dementia/ 2. dementia.mp. 3. Alzheimer disease/ 4. alzheimer.mp. 5. alzheimer\*.mp. 6. cognitive defect/ 7. cognitive defect.mp. 8. memory disorder/9. memory disorder.mp. 10. cognitive aging/11. cognitive aging.mp. 12. mild cognitive impairment/ 13. mild cognitive impairment.mp. 14. 1 or 2 or 3 or 4 or 5 or 6 or 7 or 8 or 9 or 10 or 11 or 12 or 13 16. anticoagulant.mp. 17. anticoag\*.mp. **18.** NOAC.mp. **19.** DOAC.mp. 15. anticoagulant agent/ 20. direct thrombin inhibitor.mp. 21. thrombin inhibitor/ 22. warfarin/ 23. warfarin.mp. 24. dabigatran/ 25. dabigatran etexilate/ 26. dabigatran.mp. 27. apixaban/ 28. apixaban.mp. 29. rivaroxaban/ 30. rivaroxaban.mp. 31. edoxaban/ 32. edoxaban.mp. 33. VKA.mp. 34. antivitamin K/ 35. vitamin k antagonist.mp. **36.** novel oral anticoagulant.mp. 37. direct oral anticoagulant.mp. 38. blood clotting factor 10a inhibitor/ 39. factor Xa inhibitor.mp. **40.** 15 or 16 or 17 or 18 or 19 or 20 or 21 or 22 or 23 or 24 or 25 or 26 or 27 or 28 or 29 or 30 or 31 or 32 or 33 or 34 or 35 or 36 or 37 or 38 or 39 **41.** 14 and 40

#### CINAHL

(MH "Dementia+") OR "dementia" 2. (MH "Alzheimer's Disease") OR "alzheimer" 3. alzheimer\* 4. (MH "Cognition Disorders") 5. (MH "Memory Disorders") 6. "memory disorder" 7. "cognition disorder\*" 8. "cognitive aging" 9. "cognitive impairment"
 S1 OR S2 OR S3 OR S4 OR S5 OR S6 OR S7 OR S8 OR S9
 (MH "Anticoagulants") 12. "anticoag\*" 13. "NOAC" 14. "DOAC" 15. "direct thrombin inhibitor" 16. "antithrombin" 17. (MH "Warfarin") 18. "warfarin" 19. (MH "Dabigatran Etexilate") 20. "dabigatran" 21. "apixaban" 22. (MH "Rivaroxaban") 23. "rivaroxaban" 24. "edoxaban" 25. "VKA" 26. "vitamin k antagonist" 27. "antivitamin k" 28. "novel oral anticoagulant" 29. "direct oral anticoagulant" 30. "factor Xa inhibitor" 31. "blood clotting factor 10a inhibitor" (SmartText Searching)
 33. S11 OR S12 OR S13 OR S14 OR S15 OR S16 OR S17 OR S18 OR S19 OR S20 OR S21 OR S22 OR S23 OR S24 OR S25 OR S26 OR S27 OR S28 OR S29 OR S30 OR S31 OR S32

# **Appendix 2. Risk of bias assessment tools**

# <u>Appendix 2a.</u> Adapted Joanna Briggs Institute Critical Appraisal Checklist for Analytical Cross Sectional Studies

This tool was used to assess the quality of **prevalence studies**. One point was awarded if the criterion was satisfied. A maximum of 5 points could be awarded for each study that provided oral anticoagulation prevalence estimates as sub-group analyses in results - as criteria 5, 6 and 8 are not applicable to sub-group results. A maximum of 8 points could be awarded for each study that assessed oral anticoagulation prevalence as the primary research question (ie: criteria 5, 6 and 8 become applicable). This checklist has been adapted from the original version and provides a description for how each criterion were applied and assessed.

1. Were the criteria for inclusion in the sample clearly defined?	$\Box$ Yes (1 point)
To score a 'yes,' authors should have provided clear and comprehensive	$\Box$ No (0 points)
inclusion and exclusion criteria for study sample selection and which were	$\Box$ Unclear (0 points)
developed prior to recruitment of the study participants.	Not Applicable
2. Were the study subjects and the setting described in detail?	$\Box$ Yes (1 point)
To score a 'yes,' authors should have described the study sample in sufficient	$\Box$ No (0 points)
detail including a clear description of the population from which the study	$\Box$ Unclear (0 points)
participants were selected or recruited, including demographics, location and	Not Applicable
healthcare setting, and time period.	
3. Was the exposure measured in a valid and reliable way?	$\Box$ Yes (1 point)
Authors should have clearly described the method of measurement of exposure.	$\Box$ No (0 points)
Note: for prevalence studies - exposure is dementia or mild cognitive	$\Box$ Unclear (0 points)
impairment. To score a 'yes,' a standard criterion for identifying the presence	Not Applicable
of dementia should have been reported.	
Standard criteria include:	
• Dementia codes available in administrative data (e.g: International	
Classification of Diseases and Health Related Problems codes (ICD))	
• Validated diagnostic criteria (e.g: Diagnostic and Statistical Manual	
of Mental Disorders, Mini-Mental State Exam)	
Medical diagnosis	
Medical record review or structured interview	
Standard criteria do not include:	
• self-report / patient-report / family or carer-report	
• <i>no description of a standard criteria</i>	
4. Were objective, standard criteria used for measurement of the	$\Box$ Yes (1 point)
condition?	$\Box$ No (0 points)
<i>This criterion is useful to determine if patients were included in the study based</i>	□ Unclear (0 points)
on either a specified diagnosis or definition. This is more likely to decrease the	□ Not Applicable
risk of bias. To score a 'yes,' the authors should have provided the method or	
criteria for which specific inclusion and exclusion criteria relating to	
disease/conditions were measured.	
5. Were confounding factors identified?	□ Yes (1 point)
To score a "yes," confounding factors for oral anticoagulant use or	□ No (0 points)
contraindications to oral anticoagulant use should be identified and provided	□ Unclear (0 points)
by the authors.	□ Not Applicable
Answer "not applicable" if oral anticoagulant use estimates were derived from	
sub-group analyses of results.	
6. Were strategies to deal with confounding factors stated?	□ Yes (1 point)
To score a "ves." confounding factors should be controlled for by multivariate	$\square$ No (0 points)

analysis including logistic regression stratification restricting or matching	$\Box$ Unclear (0 points)
mathods	$\Box$ Not Applicable
Answer "not applicable" if oral anticoggulant use estimates were derived from	
sub-aroun analyses of results	
7 Ware the autoemag management in a walid and valiable way?	$\Box$ Vac (1 point)
7. were the outcomes measured in a valid and reliable way?	$\Box$ res (r point)
Note: outcome measure is oral anticoaguiant use.	
To score a yes, a standard criterion for identifying oral anticoagulant use	$\Box$ Unclear (0 points)
should have been reported. In addition, oral anticoagulant use should have	$\Box$ Not Applicable
been measured in the same way for dementia and non-dementia groups.	
Standard criteria include:	
Medication charts (paper or electronic)	
• Linkage of medication records (prescribing or dispensing data)	
Structured interview	
Standard criteria do not include:	
• self-report / patient-report / family or carer-report	
• no description of a standard criteria	
8. Was appropriate statistical analysis used?	$\Box$ Yes (1 point)
To score a "yes," the methods section should have been detailed enough to	$\Box$ No (0 points)
identify analytical techniques used, for example logistic regression or	$\Box$ Unclear (0 points)
stratification and how specific confounders were identified, measured and	□ Not Applicable
controlled for. In studies using logistic regression, explanation of how	
variables were included in the logistic regression model and their relation to	
the outcome should have been provided.	
Answer "not applicable" if oral anticoagulant use estimates were derived from	
sub-group analyses of results.	

## Appendix 2b. Adapted Joanna Briggs Institute Critical Appraisal Checklist for Cohort Studies

This tool was used to assess the quality of **outcomes studies**. One point was awarded if the criterion was satisfied. A maximum of 11 points could be awarded for each study. This checklist has been adapted from the original version and provides a description for how each criterion were applied and assessed.

1. Were the two groups similar and recruited from the same population?	$\Box$ Yes (1 point)
To score a 'yes,' the dementia and non-dementia groups should have been	$\Box$ No (0 points)
selected from the same study population and be as similar as possible in	$\Box$ Unclear (0 points)
all characteristics except for the presence of dementia. The authors should	□ Not Applicable
have provided clear inclusion and exclusion criteria that were developed	
prior to recruitment of the study participants.	
2. Were the exposures measured similarly to assign people to both	$\Box$ Yes (1 point)
exposed and unexposed groups?	□ No (0 points)
Note: exposure is oral anticoagulant use. Description of how the exposure	□ Unclear (0 points)
was measured should have been described in sufficient detail.	Not Applicable
To score a 'yes' – both a standard criteria should have been used and oral	
anticoagulation use should have been measured in the same way for dementia	
and non-dementia groups.	
Standard criteria include:	
Medication charts (paper or electronic)	
• Linkage of medication records (prescribing or dispensing data)	
Structured interview	
Standard criteria do not include:	
• self-report / patient-report / family or carer-report	
• no description of a standard criteria	
3. Was the exposure measured in a valid and reliable way?	$\Box$ Yes (1 point)

	Note: exposure is oral anticoagulant use. To score a 'yes,' the study should have clearly described the method of measurement of exposure (above) and in addition provided evidence of the	<ul> <li>No (0 points)</li> <li>Unclear (0 points)</li> <li>Not Applicable</li> </ul>
	validity and reliability of the measurement method.	
	Validity refers to the percentage of cases in which the exposure is true	
	(correctly identified) when verified with an independent, 'gold standard' data	
	source (reference standard).	
	Reliability refers to the processes included in an epidemiological study to	
	check repeatability of measurements of the exposures.	
	Evidence of variary could include:	
	<ul> <li>Validation studies</li> <li>Systematic reviews of validation studies</li> </ul>	
	• Systematic reviews of validation studies Evidence of reliability could include (relevant for medication chart and	
	structured interviews only):	
	Intra-observer reliability	
	• Inter-observer reliability	
	4. Were confounding factors identified?	□ Yes (1 point)
	Confounding occurs when the estimated intervention exposure effect is biased	$\Box$ No (0 points)
	by the presence of some difference between the comparison groups (apart	□ Unclear (0 points)
	from the exposure investigated/of interest). Typical confounders include	□ Not Applicable
	baseline characteristics, prognostic factors, or concomitant exposures (e.g.	
	smoking).	
	To score a "yes," confounding factors should have been identified and	
÷	reported by the authors.	$\Box$ Vas (1 point)
	To score a "ves" confounding factors should have been controlled for by	$\square$ No (0 points)
	statistical analysis using validated methods, including: logistic regression.	$\Box$ Unclear (0 points)
	stratification, restricting or matching methods. Sufficient description of	$\Box$ Not Applicable
	statistical methods employed should have been provided by the authors.	11
	6. Were the groups/participants free of the outcome at the start of the	□ Yes (1 point)
	study (or at the moment of exposure)?	$\Box$ No (0 points)
	To score a 'yes,' authors should report whether the participants were free of	$\Box$ Unclear (0 points)
	the outcomes of interest at the start of the study. The methods section of	□ Not Applicable
	research papers should include: descriptions of participant/sample	
	recruitment, definitions of variables, and inclusion/exclusion criteria.	$\Box$ Vac (1 noint)
	Note: outcomes are events from oral anticoagulant use (embolic events	$\square$ Yes (1 point) $\square$ No (0 points)
	bleeding events or anticoagulation control)	$\Box$ Unclear (0 points)
	To score a 'ves,' a standard criterion for identifying outcomes should be	$\Box$ Not Applicable
	specified by the authors and some evidence of validity and reliability of the	II III
	measurement method reported (half a point for each).	
	Standard criteria include:	
	• Disease codes available in administrative data (e.g: International	
	Classification of Diseases and Health Related Problems codes (ICD))	
	Validated diagnostic criteria or algorithms	
	Medical diagnosis	
	Medical record review or structured interview	
	Standard criteria do not include:	
	• self-report / patient-report / family or carer-report	
	• no description of a standard criteria Evidence of validity and valiability for all included standard evidence of validity	
	Evidence of valially and reliability for all included standard criteria should also be described	
	Evidence of validity could include:	
	Validation studies	
	<ul> <li>Systematic reviews of validation studies</li> </ul>	
	- Jan 1.	

Evidence of reliability could include (relevant for medical record review or	
structured interview):	
• Intra-observer reliability	
• Inter-observer reliability	
• Evidence of specific training of those involved in collecting data	
• Evidence of more than one data collector	
8. Was the follow up time reported and sufficient to be long enough for	□ Yes (1 point)
outcomes to occur?	□ No (0 points)
To score a 'yes,' follow up time should be reported and $\geq 1$ month for all	$\Box$ Unclear (0 points)
outcomes.	□ Not Applicable
9. Was follow up complete, and if not, were the reasons to loss to follow	□ Yes (1 point)
up described and explored?	$\Box$ No (0 points)
To score a 'yes,' the proportion of patients followed up should be reported	$\Box$ Unclear (0 points)
and be greater than 80%. If follow up was less than 80% but the follow-up	Not Applicable
period was long (greater than 2 years) and sufficient details regarding efforts	
for follow up are described, then a score of 'yes' can also be awarded.	
10. Were strategies to address incomplete follow up utilized?	$\Box$ Yes (1 point)
To score a 'yes,' appropriate strategies to deal with incomplete follow-up	□ No (0 points)
should have been described and employed by the authors. For example, rates	$\Box$ Unclear (0 points)
calculated as person-years at risk and intention to treat analysis.	□ Not Applicable
11. Was appropriate statistical analysis used?	$\Box$ Yes (1 point)
To score a "yes," the methods section should have been detailed enough to	□ No (0 points)
identify analytical techniques used, for example logistic regression or	$\Box$ Unclear (0 points)
stratification and how specific confounders were identified, measured and	Not Applicable
controlled for. In studies using logistic regression, explanation of how	
variables were included in the logistic regression model and their relation to	
the outcome should have been provided.	

Appendix 3a. Results	of quality asses	ssment for pr	evalence studies	s (n=21)					
Author (Year)	Clearly defined inclusion criteria	Study subjects and setting well described	Exposure measured in a valid and reliable way	Objective, standard criteria used for condition measurement	Confounding factors identified	Strategies used to deal with confounding factors	Outcomes measured in a valid and reliable way	Appropriate statistical analysis used	TOTAL SCORE*
Bahri (2015)[63]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Choudhry (2006)[60]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
De Breucker (2010)[64]	Yes	Yes	Yes	Yes	N/A	N/A	No	N/A	4/5
Deplanque (2004)[65]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Deplanque (2006)[66]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Doucet (2008)[67]	Yes	Yes	Yes	Yes	N/A	N/A	No	N/A	4/5
Dreischulte (2014)[61]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Ewen (2012)[54]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Formiga (2016)[50]	Unclear	Yes	Yes	Unclear	N/A	N/A	Unclear	N/A	2/5
Holt (2012)[62]	Yes	Yes	Unclear	Yes	N/A	N/A	No	N/A	3/5
Hylek (2006)[55]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Latif (2005)[51]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Lefebvre (2006)[68]	Yes	Yes	Yes	Yes	Yes	Yes	Yes	N/A	5/5
Lopponen (2006)[69]	Yes	Yes	Yes	Yes	Yes	Yes	Yes	N/A	5/5
McGrath (2016)[20]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Mohammed (2013)[19]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Partington (2007)[59]	Yes	No (3/4 criteria met)	Yes	Yes	N/A	N/A	Yes	N/A	4/5
Reardon (2013)[57]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Scowcroft (2012)[24]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Shah (2016)[17]	Yes	Yes	Yes	Yes	N/A	N/A	Yes	N/A	5/5
Tanislav (2014)[70]	Yes	No	Yes	No	N/A	N/A	No	N/A	3/5

Appendix	Appendix 3b. Results of quality assessment for outcomes studies (n=6)														
Author (Year)	Similar study groups recruited from same population	Exposures measured similarly in assignment of exposed and unexposed groups	Exposure measured in a valid and reliable way	Confounding factors identified	Strategies to deal with confounding factors used	Groups/participants free of the outcome at the start of the study	Outcomes measured in a valid and reliable way	Follow-up time reported and sufficient to measure outcomes	Complete follow up. If not, reasons for incomplete follow up discussed	Strategies to address incomplete follow up used	Statistical analysis appropriate	TOTAL SCORE			
Flaker (2010)[18]	Yes	Yes <sup>a</sup>	Yes <sup>a</sup>	Yes	Yes	Unclear	Yes	Yes	Yes	Yes	Yes	10/11			
Gorzelak- Pabis (2016)[71]	Yes	No	No	No	No	Unclear	Yes	Yes	Yes	Yes	No	5/11			
Jacobs (2009)[23]	Yes	Unclear	Unclear	No	No	Unclear	Yes	Yes	Yes	Yes	No	4/11			
Khreizat (2012)[56]	Yes	Yes	Yes	No	No	Unclear	Yes	Yes	Yes	Yes	No	7/11			
Tija (2012)[58]	Yes	Yes <sup>b</sup>	Yes	Yes	Yes	Unclear	Yes	Yes	Yes	Yes	Yes	10/11			
Van Deelen (2005)[72]	Yes	Yes	Yes	Yes	Yes	Unclear	Yes	Yes	Yes	Yes	Yes	10/11			
a – study proto	ocol details publi	shed elsewhere['	73]; b – study pr	otocol details pub	lished elsewhere	[95]									

**Appendix 4.** Sensitivity analyses investigating sources of heterogeneity in meta-analyses of oral anticoagulant use in people with and without dementia or mild cognitive impairment for <u>all healthcare</u> <u>settings</u>. Types of studies included or excluded are indicated above each forest plot.

Figure 1. Studies with less than 100 people with dementia were excluded

	Deme	ntia	No der	nentia		Odds Ratio			Odds	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year		M-H, Rando	m, 95% Cl	
Choudhry 2006	556	1738	49995	114462	9.1%	0.61 [0.55, 0.67]	2006		+		
Ewen 2012	55	87	709	1054	6.6%	0.84 [0.53, 1.32]	2012		-+	-	
Holt 2012	108	374	17934	33667	8.5%	0.36 [0.28, 0.45]	2012		-		
Scowcroft 2012	1376	5382	35761	75999	9.3%	0.39 [0.36, 0.41]	2012		•		
Mohammed 2013	567	2255	23497	48106	9.2%	0.35 [0.32, 0.39]	2013		•		
Reardon 2013	462	1457	1714	3754	9.0%	0.55 [0.49, 0.63]	2013		+		
Tanislav 2014	67	241	760	1587	7.9%	0.42 [0.31, 0.56]	2014				
Dreischulte 2014	144	1006	8717	19437	8.8%	0.21 [0.17, 0.25]	2014		-		
Bahri 2015	357	777	541	1085	8.7%	0.85 [0.71, 1.03]	2015		-		
Formiga 2016	30	249	69	976	6.6%	1.80 [1.14, 2.83]	2016				
Shah 2016	377	589	3858	5102	8.7%	0.57 [0.48, 0.69]	2016		-		
McGrath 2017	67	195	719	1210	7.7%	0.36 [0.26, 0.49]	2017				
Total (95% CI)		14350		306439	100.0%	0.50 [0.40, 0.62]			•		
Total events	4166		144274								
Heterogeneity: Tau <sup>2</sup> =	0.13; Chi	i <sup>≈</sup> = 266.3	32, df = 1	1 (P < 0.0	0001); I <sup>z</sup> :	= 96%		L			- 400
Test for overall effect:	Z= 6.35 (	(P < 0.00	001)					0.01	Dementia	No dementia	100

Figure 2. Studies published before 2010 were excluded

	Deme	ntia	No den	nentia		Odds Ratio		Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year	M-H, Random, 95% Cl
Holt 2012	108	374	17934	33667	9.3%	0.36 [0.28, 0.45]	2012	+
Scowcroft 2012	1376	5382	35761	75999	10.2%	0.39 [0.36, 0.41]	2012	•
Ewen 2012	55	87	709	1054	7.3%	0.84 [0.53, 1.32]	2012	
Mohammed 2013	567	2255	23497	48106	10.1%	0.35 [0.32, 0.39]	2013	•
Reardon 2013	462	1457	1714	3754	9.9%	0.55 [0.49, 0.63]	2013	+
Tanislav 2014	67	241	760	1587	8.7%	0.42 [0.31, 0.56]	2014	-
Dreischulte 2014	144	1006	8717	19437	9.6%	0.21 [0.17, 0.25]	2014	+
Bahri 2015	357	777	541	1085	9.6%	0.85 [0.71, 1.03]	2015	-
Shah 2016	377	589	3858	5102	9.6%	0.57 [0.48, 0.69]	2016	+
Formiga 2016	30	249	69	976	7.3%	1.80 [1.14, 2.83]	2016	
McGrath 2017	67	195	719	1210	8.5%	0.36 [0.26, 0.49]	2017	-
Total (95% CI)		12612		191977	100.0%	0.49 [0.39, 0.62]		•
Total events	3610		94279					
Heterogeneity: Tau <sup>2</sup> =	0.13; Chi	<sup>2</sup> = 219.1	19, df = 1	0 (P < 0.0	0001); I <sup>z</sup> :	= 95%		
Test for overall effect:	Z=6.15 (	P ≺ 0.00	1001)					U.UI U.I I IU IUU Dementia No dementia
								Dementia No dementia

Figure 3. Studies reporting less than 50% of the study sample as female were excluded

	Demei	ntia	No dem	entia		Odds Ratio		Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year	M-H, Random, 95% Cl
Deplanque 2004	4	41	78	329	3.4%	0.35 [0.12, 1.01]	2004	
Latif 2005	26	66	28	51	5.3%	0.53 [0.25, 1.12]	2005	
Deplanque 2006	7	38	179	282	4.5%	0.13 [0.06, 0.31]	2006	<b>_</b>
Lopponen 2006	5	20	19	44	3.0%	0.44 [0.14, 1.42]	2006	
Lefebvre 2006	2	24	51	180	2.1%	0.23 [0.05, 1.01]	2006	
Hylek 2006	8	51	198	354	5.0%	0.15 [0.07, 0.32]	2006	
Doucet 2008	23	57	79	152	6.2%	0.63 [0.34, 1.16]	2008	
De Breucker 2010	35	65	22	46	5.1%	1.27 [0.60, 2.71]	2010	<del></del>
Scowcroft 2012	1376	5382	35761	75999	10.3%	0.39 [0.36, 0.41]	2012	•
Reardon 2013	462	1457	1714	3754	10.1%	0.55 [0.49, 0.63]	2013	•
Tanislav 2014	67	241	760	1587	9.0%	0.42 [0.31, 0.56]	2014	-
Bahri 2015	357	777	541	1085	9.8%	0.85 [0.71, 1.03]	2015	-
Formiga 2016	30	249	69	976	7.6%	1.80 [1.14, 2.83]	2016	
Shah 2016	377	589	3858	5102	9.8%	0.57 [0.48, 0.69]	2016	+
McGrath 2017	67	195	719	1210	8.8%	0.36 [0.26, 0.49]	2017	
Total (95% CI)		9252		91151	100.0%	0.51 [0.40, 0.64]		◆
Total events	2846		44076					
Heterogeneity: Tau <sup>2</sup> =	0.15; Ch	i <sup>2</sup> = 149	9.81, df=	14 (P < 0	0.00001);	I <sup>2</sup> = 91%		
Test for overall effect:	Z = 5.53 (	(P < 0.0	00001)					Domontia No domontia
								Demenua No demenua

	Deme	ntia	No der	nentia		Odds Ratio		Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year	M-H, Random, 95% Cl
Latif 2005	26	66	28	51	3.7%	0.53 [0.25, 1.12]	2005	
Deplanque 2006	7	38	179	282	3.1%	0.13 [0.06, 0.31]	2006	
Choudhry 2006	556	1738	49995	114462	8.0%	0.61 [0.55, 0.67]	2006	•
Hylek 2006	8	51	198	354	3.5%	0.15 [0.07, 0.32]	2006	<b>_</b>
Doucet 2008	23	57	79	152	4.4%	0.63 [0.34, 1.16]	2008	
De Breucker 2010	35	65	22	46	3.6%	1.27 [0.60, 2.71]	2010	<b>-</b>
Ewen 2012	55	87	709	1054	5.6%	0.84 [0.53, 1.32]	2012	
Scowcroft 2012	1376	5382	35761	75999	8.1%	0.39 [0.36, 0.41]	2012	•
Holt 2012	108	374	17934	33667	7.4%	0.36 [0.28, 0.45]	2012	-
Reardon 2013	462	1457	1714	3754	7.9%	0.55 [0.49, 0.63]	2013	-
Mohammed 2013	567	2255	23497	48106	8.0%	0.35 [0.32, 0.39]	2013	•
Dreischulte 2014	144	1006	8717	19437	7.7%	0.21 [0.17, 0.25]	2014	-
Tanislav 2014	67	241	760	1587	6.9%	0.42 [0.31, 0.56]	2014	
Bahri 2015	357	777	541	1085	7.6%	0.85 [0.71, 1.03]	2015	+
Shah 2016	377	589	3858	5102	7.7%	0.57 [0.48, 0.69]	2016	+
McGrath 2017	67	195	719	1210	6.7%	0.36 [0.26, 0.49]	2017	-
Total (95% CI)		14378		306348	100.0%	0.45 [0.37, 0.54]		•
Total events	4235		144711					
Heterogeneity: Tau <sup>2</sup> =	0.12; Ch	i <sup>z</sup> = 252.						
Test for overall effect:	Z= 8.19 (	(P < 0.00	001)					Dementia No dementia

**Figure 4.** Studies reporting less than 40% prevalence of oral anticoagulation use overall (dementia and non-dementia groups combined) <u>were excluded</u>

Figure 5. Studies reporting less than 20% prevalence of dementia were excluded

	Demer	ntia	No dem	entia		Odds Ratio		Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year	M-H, Random, 95% CI
Deplanque 2004	4	41	78	329	5.6%	0.35 [0.12, 1.01]	2004	
Latif 2005	26	66	28	51	8.4%	0.53 [0.25, 1.12]	2005	
Lopponen 2006	5	20	19	44	4.9%	0.44 [0.14, 1.42]	2006	
Partington 2007	12	22	45	84	6.5%	1.04 [0.41, 2.67]	2007	
Doucet 2008	23	57	79	152	9.8%	0.63 [0.34, 1.16]	2008	
De Breucker 2010	35	65	22	46	8.3%	1.27 [0.60, 2.71]	2010	_ <b>-</b> _
Reardon 2013	462	1457	1714	3754	15.6%	0.55 [0.49, 0.63]	2013	-
Bahri 2015	357	777	541	1085	15.1%	0.85 [0.71, 1.03]	2015	-
Formiga 2016	30	249	69	976	12.0%	1.80 [1.14, 2.83]	2016	
McGrath 2017	67	195	719	1210	13.7%	0.36 [0.26, 0.49]	2017	-
Total (95% CI)		2949		7731	100.0%	0.70 [0.51, 0.95]		•
Total events	1021		3314					
Heterogeneity: Tau <sup>2</sup> =	: 0.16; Chi	i <sup>2</sup> = 53.9						
Test for overall effect:	Z= 2.28 (	(P = 0.0	12)					0.01 0.1 1 10 100 Dementia No dementia

**Figure 6.** Studies reporting  $\geq$  30% of study participants with a prior history of stroke or transient ischaemic attack <u>were included</u>

	Dementia		No dementia		Odds Ratio			Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year	M-H, Random, 95% Cl
Latif 2005	26	66	28	51	7.3%	0.53 [0.25, 1.12]	2005	
Lopponen 2006	5	20	19	44	4.4%	0.44 [0.14, 1.42]	2006	
Lefebvre 2006	2	24	51	180	3.2%	0.23 [0.05, 1.01]	2006	
Partington 2007	12	22	45	84	5.8%	1.04 [0.41, 2.67]	2007	
Doucet 2008	23	57	79	152	8.5%	0.63 [0.34, 1.16]	2008	
Scowcroft 2012	1376	5382	35761	75999	13.0%	0.39 [0.36, 0.41]	2012	•
Tanislav 2014	67	241	760	1587	11.6%	0.42 [0.31, 0.56]	2014	
Bahri 2015	357	777	541	1085	12.4%	0.85 [0.71, 1.03]	2015	-
Formiga 2016	30	249	69	976	10.1%	1.80 [1.14, 2.83]	2016	
Shah 2016	377	589	3858	5102	12.5%	0.57 [0.48, 0.69]	2016	+
McGrath 2017	67	195	719	1210	11.4%	0.36 [0.26, 0.49]	2017	-
Total (95% CI)		7622		86470	100.0%	0.58 [0.43, 0.79]		•
Total events	2342		41930					
Heterogeneity: Tau <sup>2</sup> =	0.18; Ch	i <sup>≥</sup> = 118	3.90, df =	10 (P < I	).00001);	I <sup>2</sup> = 92%		
Test for overall effect:	Z = 3.48	(P = 0.0	1005)					Dementia No dementia

**Figure 7.** Studies reporting < 30% of study participants with a prior history of stroke or transient ischaemic attack *were included* 

	Demei	ntia	No der	nentia		Odds Ratio		Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year	M-H, Random, 95% Cl
Deplanque 2004	4	41	78	329	5.3%	0.35 [0.12, 1.01]	2004	
Hylek 2006	8	51	198	354	7.5%	0.15 [0.07, 0.32]	2006	_ <b>-</b>
Deplanque 2006	7	38	179	282	6.9%	0.13 [0.06, 0.31]	2006	
Choudhry 2006	556	1738	49995	114462	15.4%	0.61 [0.55, 0.67]	2006	•
De Breucker 2010	35	65	22	46	7.8%	1.27 [0.60, 2.71]	2010	<b>-</b>
Scowcroft 2012	1376	5382	35761	75999	15.6%	0.39 [0.36, 0.41]	2012	•
Ewen 2012	55	87	709	1054	11.5%	0.84 [0.53, 1.32]	2012	
Reardon 2013	462	1457	1714	3754	15.2%	0.55 [0.49, 0.63]	2013	•
Dreischulte 2014	144	1006	8717	19437	14.8%	0.21 [0.17, 0.25]	2014	+
Total (95% CI)		9865		215717	100.0%	0.41 [0.30, 0.55]		◆
Total events	2647		97373					
Heterogeneity: Tau <sup>2</sup> =	0.15; Ch	i <sup>z</sup> = 167	.60, df=	8 (P < 0.0	0001); I <sup>z</sup> :	= 95%		
Test for overall effect:	Z= 5.86	(P < 0.0	0001)					Dementia No dementia

**Figure 8.** Studies reporting dementia <u>were included</u> (studies reporting cognitive impairment <u>were</u> <u>excluded</u>)

	Dementia		No dementia		Odds Ratio			Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year	r M-H, Random, 95% Cl
Latif 2005	26	66	28	51	4.6%	0.53 [0.25, 1.12]	2005	5
Choudhry 2006	556	1738	49995	114462	10.2%	0.61 [0.55, 0.67]	2006	6 •
Lopponen 2006	5	20	19	44	2.5%	0.44 [0.14, 1.42]	2006	5 <del>  </del>
Partington 2007	12	22	45	84	3.4%	1.04 [0.41, 2.67]	2007	7
Doucet 2008	23	57	79	152	5.5%	0.63 [0.34, 1.16]	2008	3
Holt 2012	108	374	17934	33667	9.4%	0.36 [0.28, 0.45]	2012	2 -
Scowcroft 2012	1376	5382	35761	75999	10.4%	0.39 [0.36, 0.41]	2012	2 •
Mohammed 2013	567	2255	23497	48106	10.3%	0.35 [0.32, 0.39]	2013	3 •
Dreischulte 2014	144	1006	8717	19437	9.8%	0.21 [0.17, 0.25]	2014	4 🛨
Tanislav 2014	67	241	760	1587	8.7%	0.42 [0.31, 0.56]	2014	4
Formiga 2016	30	249	69	976	7.1%	1.80 [1.14, 2.83]	2016	5
Shah 2016	377	589	3858	5102	9.7%	0.57 [0.48, 0.69]	2016	3 +
McGrath 2017	67	195	719	1210	8.5%	0.36 [0.26, 0.49]	2017	7
Total (95% CI)		12194		300877	100.0%	0.47 [0.38, 0.58]		•
Total events	3358		141481					
Heterogeneity: Tau <sup>2</sup> =	0.11; Chi	<sup>2</sup> = 190.9						
Test for overall effect:	Z=7.01 (	P ≺ 0.00	0001)					Dementia No dementia

**Figure 9.** Studies reporting cognitive impairment were included (studies reporting dementia <u>were</u> <u>excluded</u>)

	Dementia		Dementia No dementia			Odds Ratio		Odds Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	M-H, Random, 95% Cl	Year	M-H, Random, 95% Cl
Deplanque 2004	4	41	78	329	8.0%	0.35 [0.12, 1.01]	2004	<b>-</b>
Hylek 2006	8	51	198	354	11.0%	0.15 [0.07, 0.32]	2006	_ <b>-</b>
Lefebvre 2006	2	24	51	180	5.1%	0.23 [0.05, 1.01]	2006	
Deplanque 2006	7	38	179	282	10.1%	0.13 [0.06, 0.31]	2006	<b>_</b>
De Breucker 2010	35	65	22	46	11.3%	1.27 [0.60, 2.71]	2010	
Ewen 2012	55	87	709	1054	15.7%	0.84 [0.53, 1.32]	2012	
Reardon 2013	462	1457	1714	3754	19.6%	0.55 [0.49, 0.63]	2013	•
Bahri 2015	357	777	541	1085	19.2%	0.85 [0.71, 1.03]	2015	-
Total (95% CI)		2540		7084	100.0%	0.48 [0.33, 0.71]		•
Total events	930		3492					
Heterogeneity: Tau <sup>2</sup> =	0.19; Ch	i² = 48.	74, df = 7	(P < 0.0	0001); P	= 86%		
Test for overall effect:	Z = 3.67	(P = 0.0	)002)					Dementia No dementia