

Systematic approach to outcome assessment from coded electronic healthcare records in the DaRe2THINK NHS-embedded randomized trial

Xiaoxia Wang (1,2, Alastair R. Mobley (1,2, Otilia Tica (1,1,4), Kelvin Okoth (1,2,4), Rebecca E. Ghosh⁴, Puja Myles⁴, Tim Williams⁴, Sandra Haynes⁵, Krishnarajah Nirantharakumar (1,3,4), David Shukla (1,3,6), and Dipak Kotecha (1,2,*, on behalf of the DaRe2THINK Trial Committees[†]

¹Institute of Cardiovascular Sciences, University of Birmingham, Birmingham, UK; ²Health Data Research UK Midlands, Queen Elizabeth Hospital, University Hospitals Birmingham NHS Foundation Trust, Birmingham, UK; ³Institute of Applied Health Sciences, University of Birmingham, Birmingham, UK; ⁴Clinical Practice Research Datalink, Medicines and Healthcare products Regulatory Agency, London, UK; ⁵Patient and Public Involvement team, Birmingham, UK; and ⁶Primary Care Clinical Research, NIHR Clinical Research Network West Midlands, Birmingham, UK

Received 30 May 2022; revised 15 August 2022; online publish-ahead-of-print 16 September 2022

Aims	Improving the efficiency of clinical trials is key to their continued importance in directing evidence-based patient care. Digital innovations, in particular the use of electronic healthcare records (EHRs), allow for large-scale screening and follow up of participants. However, it is critical these developments are accompanied by robust and transparent methods that can support high-quality and high clinical value research.
Methods and results	The DaRe2THINK trial includes a series of novel processes, including nationwide pseudonymized pre screening of the primary-care EHR across England, digital enrolment, remote e-consent, and 'no-visit' follow up by linking all primary- and secondary-care health data with patient-reported outcomes. DaRe2THINK is a pragmatic, healthcare-embedded rando- mized trial testing whether earlier use of direct oral anticoagulants in patients with prior or current atrial fibrillation can prevent thromboembolic events and cognitive decline (www.birmingham.ac.uk/dare2think). This study outlines the systematic approach and methodology employed to define patient information and outcome events. This includes transparency on all medical code lists and phenotypes used in the trial across a variety of national data sources, including Clinical Practice Research Datalink Aurum (primary care), Hospital Episode Statistics (secondary care), and the Office for National Statistics (mortality).
Conclusion	Co-designed by a patient and public involvement team, DaRe2THINK presents an opportunity to transform the approach to randomized trials in the setting of routine healthcare, providing high-quality evidence generation in populations representative of the community at risk.

^{*} Corresponding author. Heritage Building, Queen Elizabeth Hospital, University Hospitals Birmingham NHS Foundation Trust, Mindelsohn Way, Birmingham, B15 2GW, UK. Tel: +44 121 3718122, Email: d.kotecha@bham.ac.uk

[†]See Acknowledgements.

[©] The Author(s) 2022. Published by Oxford University Press on behalf of the European Society of Cardiology.

This is an Open Access article distributed under the terms of the Creative Commons Attribution License (https://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.

Graphical Abstract



Keywords

Randomized controlled trial • Electronic healthcare record • Coding • Atrial fibrillation • Anticoagulation • Primary care • Secondary care

Introduction

Rapid uptake of electronic healthcare record (EHR) systems across the world have led to the opportunity for large-scale, real-world, longitudinal clinical research.¹ In countries with national health systems, there is also the potential to link all EHRs for an individual patient, across both primary and secondary care, and from birth to death. Standardized coding systems provide the basis for harnessing medical information from different healthcare providers, with linkage allowing for a complete picture of each patient's history, healthcare utilization, and adverse events.² This is an attractive (and cost-efficient) prospect for clinical research, particular in countries such as the UK where the quality of EHR data is typically high due to the need for accurate assessment of healthcare utilization within the National Health Service (NHS).^{3–5}

Observational clinical research has benefited for many decades from EHR data. Evolving technology and coding systems, coupled with more complete EHR coverage, have provided the opportunity for randomized controlled trials (RCTs) to be embedded alongside EHR systems, taking advantage of existing clinical data and follow up. The coronavirus pandemic has epitomized how innovations in the design and running of RCTs are critical to address unmet clinical need. However, RCTs remain too costly, tend to recruit selective patients within high-performing centres, and frequently do not match the real population at risk in our communities. The DaRe2 approach (healthcare Data for pragmatic clinical Research in the NHS—primary 2 secondary) was designed to incorporate recent innovations in EHR and clinical research, combined with advances in mobile technology, to provide an end-to-end framework for RCTs embedded in Primary Care in England. The DaRe2THINK trial provides an exemplar of a 'remote-RCT' to address a key public health concern, requiring no physical patient visits despite being a clinical trial of an investigational medicinal product.⁶ In this study, we outline the design features of DaRe2THINK, and the systematic approach used to define patient information and outcome events that will be accrued during the trial. All medical code lists and phenotypes are presented for full transparency, and for future use by other researchers using EHR data.

The DaRe2THINK trial

Using the digital potential of the EHR, DaRe2THINK is transformational project that will underpin a new trajectory for NHS-based research for patient benefit. DaRe2THINK is led by the University of Birmingham, in collaboration with the Clinical Practice Research Datalink (CPRD; part of the Medicines and Healthcare products Regulatory Agency). The trial is supported by the University Hospitals Birmingham NHS Foundation Trust, the National Institute for Health and Care Research (NIHR) Clinical Research Network (West Midlands Primary Care), and Health Data Research UK Midlands.

DaRe2 platform features

- (1) Automated secure screening of inclusion and exclusion criteria across >13 million NHS patients, including around one in four primary-care sites in England that are part of CPRD. This provides rapid and cost-efficient screening of a diverse and representative proportion of the UK population,⁷ yet maintains patient privacy using pseudonymized records that can only be reidentified by local NHS staff.
- (2) Access to national live data on the numbers of potentially eligible patients allows for smart adaptation of trial inclusion and exclusion criteria.
- (3) Targeted enrolment of primary-care sites identified as having potentially eligible patients, streamlining recruitment by focusing resources on high-value sites.
- (4) Dynamic adaptive screening of the population at each site, with updates on a weekly basis of newly eligible patients, reducing burden on local NHS staff and simplifying recruitment.
- (5) Easy enrolment of patients without need for physical attendance, with remote e-consent using their own smartphone/tablet/computer, one-click randomization by primary-care staff, and drug prescription via usual clinical systems.
- (6) 'No-visit' follow up, utilizing NHS records linked across national primary and secondary care for capture of endpoints without the patient attending the healthcare site, and no need for NHS investigators to complete case report forms.
- (7) Regular scheduled patient-reported outcomes, with requests sent by automated text message to each participant's telephone and email, and secure data acquisition through online completion.
- (8) Automated collation of safety outcomes from the primary-care NHS record, avoiding the possibility of missed events unknown to the local investigator.
- (9) Co-design and management of all processes by a patient and public involvement (PPI) team,⁸ supporting a patient-centric approach with sustainable and valued output.

DaRe2THINK design

DaRe2THINK will be the first exemplar of this system, and is focused on the intersection of key national priorities for health and social care services—the common heart rhythm disorder called atrial fibrillation (AF) and the impact this condition has on thromboembolic events, including long-term cognitive decline and vascular dementia. Patients with AF suffer from a high rate of morbidity, with one in four stroke patients having AF as a potential cause, half of AF patients developing heart failure that responds poorly to treatment, and the rate of death doubled at all ages.^{9–11} Most patients with AF develop progressive subclinical cerebral damage over time,^{12,13} with high rates of cognitive impairment and a 40% increased risk of dementia compared with normal sinus rhythm.^{14,15} As the prevalence of AF is expected to double in the coming decades,¹⁶ this will place an unsustainable burden on healthcare and social services.

DaRe2THINK is a pragmatic, NHS-embedded, open-label, eventdriven, parallel-group RCT. It will test whether earlier use of anticoagulants is effective and cost-effective at preventing thromboembolic events (primary outcome) and cognitive decline (key secondary outcome) in patients with AF at low or intermediate risk of stroke. Around 3000 patients will be randomized to either starting any direct oral anticoagulant (used due to their excellent safety and efficacy profile^{17,18}), or continuing standard-of-care where anticoagulation is instituted when the patient is older with clear risk factors for stroke.¹⁹ Follow up includes yearly collection of cognitive and outcome data for 5 years. Consent also includes remote follow up at 10 years and then lifetime EHR outcomes for long-term conditions such as vascular dementia. A summary of the DaRe2THINK trial is presented in Figure 1. Due to the ability to access pseudonymized NHS records, DaRe2THINK will be able to adapt inclusion and exclusion criteria to reflect current changes in practice and extraneous factors (e.g. the coronavirus pandemic). Recruitment has England (https://www.birmingham.ac.uk/ commenced across dare2think).

Patient and public involvement has been central to the development of the DaRe2THINK trial process from conception through to delivery. DaRe2THINK follows the PPI-POSITIVE approach for implementation of genuine public engagement at every stage of the trial, including patient contact, study management, analysis, and dissemination.⁸ A plain English summary written by the PPI team is provided in *Table 1*.

Systematic coding methodology

The DaRe2 approach links and combines NHS data from primarycare, secondary-care, and other national databases in order to provide a full and complete picture of each participant's health and healthcare utilization. The UK public health system is free at the point of delivery to all citizens, with indirect reimbursement via local health authorities based on coded data for diseases and procedures performed. In this section, we detail the processes employed in the DaRe2THINK trial to systematically catalogue and incorporate all relevant codes, including for selection criteria, baseline variables, outcome endpoints. In addition, DaRe2THINK includes monthly automated searches of the primary-care record to assess for safety events. For transparency, all codes used are presented in Supplementary material online, Appendix S1 for other researchers to see, comment, update, and re-use. The current trial protocol, including details on trial processes and sample size calculations, is available in Supplementary material online, Appendix S2. The DaRe2THINK trial adheres to the CODE-EHR best practice framework for the use of structured EHRs in clinical research.² This study meets all five of the CODE-EHR minimum standards, and will in addition meet all five standards for preferred criteria once completed; further details are presented in Supplementary material online, Appendix S3. The DaRe2THINK investigators are committed to the FAIR principles (Findable, Accessible, Interoperable, and Reusable).²⁰

Data sources

Primary-care data are obtained through CPRD Aurum, a prospectively collected, population-based, pseudonymized medical record database that collects daily information directly from NHS primarycare sites that are part of the CPRD network and use the Egton





Figure 1 DaRe2THINK key innovations. Flow chart of the DaRe2 approach (healthcare data for pragmatic clinical research in the NHS—primary 2 secondary) and summary of the DaRe2THINK clinical trial. *Please see current protocol at www.birmingham.ac.uk/dare2think for full details of inclusion and exclusion criteria. CPRD, Clinical Practice Research Datalink; DOAC, direct oral anticoagulant; NHS, National Health Service.

Table 1 Plain English se	ummary of DaRe2THINK
Challenges and opportunities	The NHS is unlike any other in the world, caring for people throughout their lives both in the community and in hospitals. At the heart of DaRe2THINK is that health data collected within these services can be used for the benefit of patients. Clinical trials are an important way to understand how new treatments can be used in the NHS, but many trials struggle to find the right patients, or be relevant to their needs. The DaRe2 approach (healthcare Data for pragmatic clinical Research in the NHS—primary 2 secondary) will test a new way of running trials based at general practitioner (GP) surgeries using routine NHS information. We will include patients that do not normally take part in clinical trials and follow them up without the need to revisit their GP or attend hospital. This approach could improve the health and well-being of those treated by the NHS, while reducing the time needed from staff and patients to engage in important research.
Addressing a gap in current healthcare	As an example of this new system, DaRe2THINK will target an issue of huge importance to patients, our NHS and the social care system. AF is a common heart rhythm condition that leads to a high chance of stroke, frequent hospital admissions, and poor quality of life. Patients also have a much higher risk of cognitive decline (trouble remembering, concentrating, or making everyday decisions) and dementia. This may be due to silent 'micro-strokes' that gradually damage the brain over time. Blood thinning tablets (anticoagulants) greatly reduce the number of patients with AF that will suffer a stroke, but are usually only given to older patients or those with other health issues. This may be too late to avoid dementia. It also leaves those younger than 65 years, and some patients aged 65–75 without treatment that could prevent these devastating complications.
Aim of DaRe2THINK	A new class of blood thinning tablets are now widely used in the NHS which are more convenient for patients to take, and have a lower risk of bleeding than older treatments. These drugs could provide an effective way to prevent strokes, brain damage, and dementia in later life for a broader group of patients, but this needs to be tested in a clinical trial.
How the trial works	With the support of a PPI team and a national network of research nurses and GPs, the trial will include 3000 patients from up to 600 GP surgeries across England. Each patient will either continue their current treatment or start an additional blood thinning tablet on a random basis. Patients will be followed up automatically within the NHS to look at the difference in those who suffer from strokes, blood clots, heart attacks, other problems with the blood vessels, and dementia. Patients will self-report their memory, reaction times, and quality of life using simple questionnaires through their mobile phone or the internet, again without needing to revisit their doctor.
Results and impact	DaRe2THINK will answer important questions for a growing number of patients with AF. The combination of information from the community as well as hospitals across the NHS will allow us to see whether these blood thinning tablets should be prescribed more widely. DaRe2THINK will allow us to develop and improve this new clinical trial system so that future research in the NHS will continue to benefit those patients most in need.

Medical Information Systems software system. As of March 2021 when DaRe2THINK was initiated, CPRD Aurum included 39555 354 research acceptable patients of which 13 299 826 were actively registered (19.9% of the UK population) across 1375 active primary-care sites (15.3% of UK general practices). All baseline data for DaRe2THINK trial participants is extracted from CPRD Aurum, obviating the need for investigators to fill in case report forms. CPRD Aurum includes codes for diagnosis and non-prescription data (med-codeid), which is cross-mapped to SNOMED CT (UK edition)²¹ and Read Version 2,²² with prescriptions by product code (prodcodeid) mapped to the Dictionary of Medicines and Devices.²³

Secondary care Hospital Episode Statistics (HES) admitted patient care data, which contains information on all admissions to NHS hospitals in England, as well as NHS-funded care at independent providers, are linked at the patient level to CPRD Aurum by NHS Digital. With around 99% of hospital activity in England funded by the NHS, this provides almost total coverage of healthcare utilization for the DaRe2THINK trial, including admission and discharge date, admission type (pre planned or emergency), primary diagnosis (reason for hospitalization), secondary diagnoses, and any procedures performed during the hospital stay. HES uses the International Classification of Diseases version 10 (ICD-10) for diagnosis codes,²⁴ and the Office of Population Censuses and Surveys Classification of

Surgical Operation and Procedures (OPCS) version 4 for procedures.²⁵

Death and cause of death are obtained via linkage with the Office for National Statistics (ONS) mortality database,²⁶ which includes all deaths in England occurring both within and outside any healthcare setting. Data in ONS are coded using ICD-10.

Development of the medical code lists

To achieve a comprehensive set of contemporary codes which are suitable for defining outcome and safety events in a clinical trial, we designed and employed a four-phase systematic framework to develop code lists for data extraction from the listed data sources. The approach is summarized in *Figure 2*. Throughout the four phases, we used DExtER, an automated platform for clinical epidemiology that was developed to support clinical code list development and validation of phenotypes.²⁸

Phase 1: phenotype specification

Prior to searching for relevant medical codes, the DaRe2THINK trial selection criteria and outcome measures were first transformed into relevant phenotypes for each data source. This was an iterative process that accounted for the limitations of the data source and, where



Figure 2 DaRe2THINK coding framework. Systematic, iterative approach to defining code lists from national healthcare data sources. *Study-Within-A-Trial to be performed in a subset of patients with clinical events identified using primary and secondary healthcare records.²⁷

needed, combined phenotypes to derive a particular outcome. For example, 'major bleeding' is not recorded as such in any primary-care data set; hence, we combined a phenotype list for bleeding events with a list generated to identify concurrent hospitalization. This provided a code list for the safety outcome of 'bleeding resulting in hospitalization' based solely on CPRD Aurum data. This process was facilitated by the Data Coordination Team, a specific DaRe2THINK committee which includes clinical expertise in primary and secondary care, data scientists with knowledge and experience in using medical codes, and representatives of CPRD.

Phase 2: systematic code searching

A systematic search was carried out against the agreed phenotypes using the medical code dictionaries (SNOMED CT and Read codes for CPRD Aurum, and ICD-10 codes for HES and ONS). When searching medical code dictionaries, search terms and acronyms were agreed with clinical input. The hierarchy structure of each dictionary was used to find additional relevant children or sibling codes, and these were collated into a dynamic code list. The list was supplemented by any codes identified by other researchers after screening of previously published and relevant research. To facilitate future use, or if there are updates to coding systems during the trial follow-up period, all search terms and searching scripts are saved as meta data. Medication codes required in the phenotyping algorithms were extracted from the CPRD Aurum product lists based on both British National Formulary codes and drug substance names. CPRD provided established code lists for demographics and measurements of ethnicity, height, weight, blood pressure, etc.

Phase 3: iterative and clinical review

The codes found in the systematic search went through a multi-stage review process. This included review by data and clinical scientists. Code lists were reviewed independently by two individuals, marking codes for inclusion, exclusion, or further review. Discrepancies between the reviewers and any queries were resolved by consensus agreement after adjudication by a senior investigator. All resultant code lists were then further reviewed by senior clinical experts to ensure that code lists were correct and consistent.

Phase 4: analysis of prevalence and validation

For validation purposes, the prevalence of conditions based on the final code lists was then cross-checked with prevalence estimates from relevant publications and the Quality and Outcomes Framework (a national system designed to remunerate general practices for

Table 2 Summary of code lists for selection criteria

-		
Selection criterion (CPRD Aurum)	No. of codes	Appendix table
Diagnosis of AF (previous, current, or chronic)	43	S1
Existing use of an anticoagulant		
Oral anticoagulants	54	S2
Low molecular weight heparin	80	S3
Stroke	120	S4
Transient ischaemic attack	11	S5
Arterial thromboembolism	81	S6
Stroke risk factors		
Myocardial infarction	80	S7
Peripheral arterial disease	16	S8
Aortic disease	4	S9
Diabetes mellitus	455	S10
Antidiabetic drug/insulin	433	S11
Hypertension	85	S12
Antihypertensive drug	724	S13
Heart failure	54	S14
Loop diuretic therapy	59	S15
Prior major bleeding (intracranial bleed, or	161	S16
requiring hospitalization)		
High bleeding risk		
Gastrointestinal tract ulcer	234	S17
Brain injury	568	S18
Spinal injury	1	S19
Eye injury	11	S20
Estimated glomerular filtration rate <30 mL/min	6	S21
Azole-antimycotic treatment	20	S22
Current diagnosis of dementia	194	S23

Table 3 Summary of code lists for outcome events

Outcome event	CPRD Aurum			HES		
	Number of codes reviewed	Final number of codes	Appendix table	Number of codes reviewed	Final number of codes	Appendix table
Cardiovascular mortality	-	-	-	423	353	S24
Ischaemic stroke	677	315	S25	41	10	S26
Transient ischaemic attack	62	44	S27	41	30	S28
Pulmonary and venous thromboembolism	1759	212	S29			
Arterial thromboembolism	1759	306	S30			
Thromboembolic event				144	67	S31
Myocardial infarction	215	150	S32	22	19	S33
Heart failure	430	100	S43	22	10	S44
Vascular dementia	252	58	S34	6	6	S35
Gastrointestinal bleeding	902	558	S36	42	25	S37
Other bleeding	4088	1642	S38	235	59	S39
Hospitalization event	3701	483	S40	-	-	-
Intracranial bleeding	1405	566	S41	36	31	S42

X. Wang et al.

providing good quality care to their patients). Results were discussed with the DaRe2THINK Data Coordination Team, and then suggested changes iteratively fed back to Phases 1–3 as appropriate. This process will be updated regularly through implementation so that phenotype definitions can be adjusted accordingly, for example if there is a change to coding systems. Cross-validation of endpoints will be conducted in a prospectively registered Study-Within-A-Trial.²⁷ In a subset of participants, we will compare outcome events obtained from coded primary and secondary EHR data with in-depth and granular information extracted from one of the UK's largest hospital trusts (University Hospitals Birmingham NHS Foundation Trust). This will involve extraction of text using machine learning methods from letters, clinical notes, imaging, and time-series data to determine accuracy and missingness.

Results and final code lists

Using the systematic approach discussed above, we searched a total of 1176 611 codes, including 1159 849 from primary care, and 16 762 from secondary care and national databases. This led to the creation of a total of 77 code lists across the different sections of the trial.

Participant selection criteria

The DaRe2THINK trial uses automated pre screening of the primary-care EHR to determine inclusion and exclusion according to the defined selection criteria. *Table 2* lists the main components of the selection criteria and associated CPRD Aurum code lists. A total of 3494 codes were included in these code lists using the systematic approach, including contribution from 14 published code lists.^{5,29} Individual codes for each list can be found in Supplementary material online, *Appendix S1 Tables S1–S23*.

Table 4 Summary of code lists for baseline characteristics

Ethnicity 327 S45 Smoking status 104 S46 Measurements (height, weight, body mass index, blood pressure, and heart rate) 58 S477 Blood tests (creatinine, eGFR, haemoglobin, HbA1c, and lipids) 117 S48 AF 43 S1 Prior AF ablation 42 S49 Heart failure 100 S43 Hypertension 85 S12 Myocardial infarction 80 S32 Stroke 120 S4 Venous thromboembolism 212 S29 Arterial thromboembolism 306 S30 Peripheral artery disease and aortic 20 S8 and S9 plaque/atherosclerosis Diabetes mellitus 455 S10 Hypertyroidism 73 S50 Hypothyroidism 90 S51 Chronic obstructive pulmonary disease 191 S52 Eye-related diseases and procedure 293 S53-S60 Gastrointestinal bleeding 566 S41 Other bleeding 1642 S38	Characteristic (CPRD Aurum)	Number of codes	Appendix table
Ethnicity 327 343 Smoking status 104 S46 Measurements (height, weight, body mass index, blood pressure, and heart rate) 58 S47 Blood tests (creatinine, eGFR, haemoglobin, HbA1c, and lipids) 117 S48 AF 43 S1 Prior AF ablation 42 S49 Heart failure 100 S43 Hypertension 85 S12 Myocardial infarction 80 S32 Stroke 120 S4 Venous thromboembolism 212 S29 Arterial thromboembolism 306 S30 Peripheral artery disease and aortic 20 S8 and S9 plaque/atherosclerosis Diabetes mellitus 455 S10 Hyperthyroidism 73 S50 S51 Chronic obstructive pulmonary disease 191 S52 Eye-related diseases and procedure 293 S53-S60 Gastrointestinal bleeding 566 S41 Other bleeding 1642 S38 H	Fall of the s		C4F
Smoking status 104 546 Measurements (height, weight, body mass index, blood pressure, and heart rate) 58 547 Blood tests (creatinine, eGFR, haemoglobin, HbA1c, and lipids) 117 548 AF 43 51 Prior AF ablation 42 549 Heart failure 100 543 Hypertension 85 512 Myocardial infarction 80 532 Stroke 120 54 Venous thromboembolism 212 529 Arterial thromboembolism 306 530 Peripheral artery disease and aortic 20 58 and 59 plaque/atherosclerosis 510 44 Hyperthyroidism 73 550 Hypothyroidism 90 551 Chronic obstructive pulmonary disease 191 552 Eye-related diseases and procedure 293 553–560 Gastrointestinal bleeding 566 541 Other bleeding 1642 538 Hosptalization 483 4		327	545
Preasurements (neight, weight, body mass58547index, blood pressure, and heart rate)Blood tests (creatinine, eGFR, haemoglobin, HbA1c, and lipids)117S48AF43S1Prior AF ablation42S49Heart failure100S43Hypertension85S12Myocardial infarction80S32Stroke120S4Venous thromboembolism212S29Arterial thromboembolism306S30Peripheral artery disease and aortic20S8 and S9plaque/atherosclerosisDiabetes mellitus455S10Hyperthyroidism73S50Hypothyroidism90S51Chronic obstructive pulmonary disease191S52Eye-related diseases and procedure293S53-S60Gastrointestinal bleeding566S41Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62-S64S67Angiotensin II receptor antagonists116Alosterone receptor antagonists116S68Beta-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitors54S76<	Smoking status	104	546
Index, blood pressure, and neart rate)Blood tests (creatinine, eGFR, haemoglobin, HbA1c, and lipids)117S48AF43S1Prior AF ablation42S49Heart failure100S43Hypertension85S112Myocardial infarction80S32Stroke120S4Venous thromboembolism212S29Arterial thromboembolism306S30Peripheral artery disease and aortic plaque/atherosclerosisS10Diabetes mellitus455S10Hyperthyroidism73S50Hypothyroidism90S51Chronic obstructive pulmonary disease191S52Eye-related diseases and procedure293S53-S60Gastrointestinal bleeding566S41Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers155S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitors34S76inhibitors14S75Sodium-glucose tra	Measurements (height, weight, body mass	58	547
Biood tests (creatinine, eCFR, 117 548 haemoglobin, HbA1c, and lipids) AF 43 51 Prior AF ablation 42 549 Heart failure 100 543 Hypertension 85 512 Myocardial infarction 80 532 Stroke 120 54 Venous thromboembolism 212 529 Arterial thromboembolism 306 530 Peripheral artery disease and aortic 20 58 and 59 plaque/atherosclerosis Diabetes mellitus 455 510 Hyperthyroidism 73 550 Hypothyroidism 90 551 Chronic obstructive pulmonary disease 191 552 Eye-related diseases and procedure 293 553–560 Gastrointestinal bleeding 558 536 Intracranial bleeding 558 536 Intracranial bleeding 558 536 Intracranial bleeding 558 536 Antiplatelet therapy 79 561 Diuretics 243 515 and 562– 564 Calcium channel blockers 54 567 Angiotensin II receptor antagonists 116 568 Beta-blockers 58 570 Aldosterone receptor antagonist 32 571 Other antihypertensive drugs 76 572 Antiarrhythmic drugs (Classes 1 and 3) 94 573 and 574 Digoxin 14 575 Sodium–glucose transport protein 2 34 576 inhibitors	index, blood pressure, and heart rate)	447	C 40
AF 43 S1 Prior AF ablation 42 S49 Heart failure 100 S43 Hypertension 85 S12 Myocardial infarction 80 S32 Stroke 120 S4 Venous thromboembolism 212 S29 Arterial thromboembolism 306 S30 Peripheral artery disease and aortic 20 S8 and S9 plaque/atherosclerosis Diabetes mellitus 455 S10 Hyperthyroidism 73 S50 Hypothyroidism 90 S51 Chronic obstructive pulmonary disease 191 S52 Eye-related diseases and procedure 293 S53-S60 Gastrointestinal bleeding 558 S36 Intracranial bleeding 566 S41 Other bleeding 1642 S38 Hospitalization 483 S40 Antiplatelet therapy 79 S61 Diuretics 267 S65 and S66 ACE inhibitors 54 S67 Angiotensin II receptor antagonists <	Blood tests (creatinine, eGFR,	117	548
AF4351Prior AF ablation42\$49Heart failure100\$43Hypertension85\$12Myocardial infarction80\$32Stroke120\$4Venous thromboembolism212\$29Arterial thromboembolism20\$8 and \$9plaque/atherosclerosis20\$8 and \$9Diabetes mellitus455\$10Hyperthyroidism73\$50Hypothyroidism90\$51Chronic obstructive pulmonary disease191\$52Eye-related diseases and procedure293\$53-\$60Gastrointestinal bleeding566\$41Other bleeding1642\$38Hospitalization483\$400Antiplatelet therapy79\$61Diuretics243\$15 and \$62-Calcium channel blockers267\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers155\$70Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors34\$76inhibitors34\$76inhibitors34\$76inhibitors34\$76Non-steroidal anti-inflammatory drugs86\$77<	naemogiobin, HDATC, and lipids)	40	64
Prior AF ablation42549Heart failure100S43Hypertension85S12Myocardial infarction80S32Stroke120S4Venous thromboembolism212S29Arterial thromboembolism306S30Peripheral artery disease and aortic20S8 and S9plaque/atherosclerosis90S51Diabetes mellitus455S10Hyperthyroidism73S50Hypothyroidism90S51Chronic obstructive pulmonary disease191S52Eye-related diseases and procedure293S53-S60Gastrointestinal bleeding566S41Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62-KS67S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers155S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitors34S76inhibitors34S76inhibitors34S76inhibitors34S76inhibitors34S76Anta		43	51
Heart tailure100\$43Hypertension85\$12Myocardial infarction80\$322Stroke120\$4Venous thromboembolism212\$29Arterial thromboembolism306\$30Peripheral artery disease and aortic20\$8 and \$9plaque/atherosclerosis20\$8 and \$9Diabetes mellitus455\$10Hyperthyroidism73\$50Hypethyroidism90\$51Chronic obstructive pulmonary disease191\$52Eye-related diseases and procedure293\$53-\$60Gastrointestinal bleeding566\$41Other bleeding1642\$38Hospitalization483\$40Antiplatelet therapy79\$61Diuretics243\$15 and \$62-Sé4\$67\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers115\$69Alpha-blockers58\$70Aldosterone receptor antagonist32\$71Other anthypertensive drugs76\$72Antairrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors34\$76inhibitors34\$76inhibitors34\$76	Prior AF ablation	42	549
Hypertension85512Myocardial infarction80\$32Stroke120\$4Venous thromboembolism212\$29Arterial thromboembolism306\$30Peripheral artery disease and aortic20\$8 and \$9plaque/atherosclerosis20\$8 and \$9Diabetes mellitus455\$10Hyperthyroidism73\$50Hypethyroidism90\$51Chronic obstructive pulmonary disease191\$52Eye-related diseases and procedure293\$53-\$60Gastrointestinal bleeding558\$36Intracranial bleeding566\$41Other bleeding1642\$38Hospitalization483\$400Antiplatelet therapy79\$61Diuretics267\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers15\$69Alpha-blockers58\$70Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors34\$76inhibitors34\$76inhibitors34\$76	Heart failure	100	543
Myocardial infarction80\$32Stroke120\$4Venous thromboembolism212\$29Arterial thromboembolism306\$30Peripheral artery disease and aortic20\$8 and \$9plaque/atherosclerosis20\$8 and \$9Diabetes mellitus455\$10Hyperthyroidism73\$50Hypothyroidism90\$51Chronic obstructive pulmonary disease191\$52Eye-related diseases and procedure293\$53-\$60Gastrointestinal bleeding566\$41Other bleeding1642\$38Hospitalization483\$40Antiplatelet therapy79\$61Diuretics243\$15 and \$62- \$64Calcium channel blockers267\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers15\$69Alpha-blockers58\$70Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors54\$76Non-steroidal anti-inflammatory drugs86\$77	Hypertension	85	\$12
Stroke12054Venous thromboembolism212S29Arterial thromboembolism306S30Peripheral artery disease and aortic20S8 and S9plaque/atherosclerosis20S8 and S9Diabetes mellitus455S10Hyperthyroidism73S50Hypothyroidism90S51Chronic obstructive pulmonary disease191S52Eye-related diseases and procedure293S53-S60Gastrointestinal bleeding566S41Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62-Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers15S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitors34S76inhibitors34S76	Myocardial infarction	80	\$32
Venous thromboembolism212\$29Arterial thromboembolism306\$30Peripheral artery disease and aortic20\$8 and \$9plaque/atherosclerosis21\$5 \$10Diabetes mellitus455\$10Hyperthyroidism73\$50Hypothyroidism90\$51Chronic obstructive pulmonary disease191\$52Eye-related diseases and procedure293\$53-\$60Gastrointestinal bleeding558\$36Intracranial bleeding566\$41Other bleeding1642\$38Hospitalization483\$40Antiplatelet therapy79\$61Diuretics243\$15 and \$62-Calcium channel blockers267\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers115\$69Alpha-blockers58\$70Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75\$0dium-glucose transport protein 234S76inhibitors34\$76inhibitors34\$76\$72Non-steroidal anti-inflammatory drugs86\$77	Stroke	120	S4
Arterial thromboembolism306\$30Peripheral artery disease and aortic plaque/atherosclerosis20\$8 and \$9Diabetes mellitus455\$10Hyperthyroidism73\$50Hypothyroidism90\$51Chronic obstructive pulmonary disease191\$52Eye-related diseases and procedure293\$53-\$60Gastrointestinal bleeding558\$36Intracranial bleeding566\$41Other bleeding1642\$38Hospitalization483\$40Antiplatelet therapy79\$61Diuretics243\$15 and \$62-\$64Calcium channel blockers267\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers58\$70Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors34\$76Non-steroidal anti-inflammatory drugs86\$77	Venous thromboembolism	212	\$29
Peripheral artery disease and aortic20S8 and S9plaque/atherosclerosisDiabetes mellitus455S10Hyperthyroidism73S50Hypothyroidism90S51Chronic obstructive pulmonary disease191S52Eye-related diseases and procedure293S53-S60Gastrointestinal bleeding558S36Intracranial bleeding566S41Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62- S64Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitors38S70	Arterial thromboembolism	306	\$30
Diabetes mellitus455S10Hyperthyroidism73S50Hypothyroidism90S51Chronic obstructive pulmonary disease191S52Eye-related diseases and procedure293S53–S60Gastrointestinal bleeding558S36Intracranial bleeding566S41Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62–Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium–glucose transport protein 234Non-steroidal anti-inflammatory drugs86S77	Peripheral artery disease and aortic	20	S8 and S9
Diabetes memory155510Hyperthyroidism73S50Hypothyroidism90S51Chronic obstructive pulmonary disease191S52Eye-related diseases and procedure293S53–S60Gastrointestinal bleeding558S36Intracranial bleeding566S41Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62–Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium–glucose transport protein 234S76inhibitors36S77	Disbetes mellitus	455	\$10
Hypertrijvertijvertijvertijvertijvertijvertijvertijvertijvertij75550Hypothyroidism90\$51Chronic obstructive pulmonary disease191\$52Eye-related diseases and procedure293\$53-\$60Gastrointestinal bleeding558\$36Intracranial bleeding566\$41Other bleeding1642\$38Hospitalization483\$40Antiplatelet therapy79\$61Diuretics243\$15 and \$62-Calcium channel blockers267\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers58\$70Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors\$6\$77	Hyperthyroidism	73	\$50
Typerin outsin7.65.51Chronic obstructive pulmonary disease191S52Eye-related diseases and procedure293S53–S60Gastrointestinal bleeding558S36Intracranial bleeding566S41Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62– S64Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium–glucose transport protein 234S76inhibitors36S77	Hypothyroidism	90	S50 S51
Chilomic obstructive pullificities131332Eye-related diseases and procedure293\$53-\$60Gastrointestinal bleeding558\$36Intracranial bleeding566\$41Other bleeding1642\$38Hospitalization483\$40Antiplatelet therapy79\$61Diuretics243\$15 and \$62- \$64Calcium channel blockers267\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers58\$70Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors\$86\$77	Chronic obstructivo pulmonany disease	101	557
Lycenetated diseases and procedure273353-500Gastrointestinal bleeding558S36Intracranial bleeding566S41Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62-S64S65 and S66S67Angiotensin II receptor antagonists116S68Beta-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitors36S77	Evo related diseases and procedure	202	553 540
Clash Onlecting5.565.56Intracranial bleeding566541Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62 S64Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitorsS77	Castrointestinal blooding	550	536
Intract aniar bleeding300341Other bleeding1642S38Hospitalization483S40Antiplatelet therapy79S61Diuretics243S15 and S62 S64Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitors36S77		566	530 541
Other bleeding1042330Hospitalization483\$40Antiplatelet therapy79\$61Diuretics243\$15 and \$62 \$64Calcium channel blockers267\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers115\$69Alpha-blockers58\$70Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors\$6\$77	Other blooding	1642	511
Antiplatelet therapy79S61Diuretics243S15 and S62- S64Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitors	Hospitalization	493	530 540
Antipiatelet therapy77501Diuretics243\$15 and \$62- \$64Calcium channel blockers267\$65 and \$66ACE inhibitors54\$67Angiotensin II receptor antagonists116\$68Beta-blockers115\$69Alpha-blockers58\$70Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors\$77		705	S41
Did edds243313 and 302- S64Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitorsS77	Diurotics	243	S15 and S62
Calcium channel blockers267S65 and S66ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitorsS77	Dial etics	273	515 and 502-
ACE inhibitors54S67Angiotensin II receptor antagonists116S68Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitorsNon-steroidal anti-inflammatory drugs86S77	Calcium channel blockers	267	S65 and S66
Angiotensin II receptor antagonists116S68Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitorsNon-steroidal anti-inflammatory drugs86S77		54	567
Augustain in receptor antagonists110500Beta-blockers115S69Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitorsNon-steroidal anti-inflammatory drugs86S77	Angiotensin II receptor antagonists	116	568
Alpha-blockers58S70Aldosterone receptor antagonist32S71Other antihypertensive drugs76S72Antiarrhythmic drugs (Classes 1 and 3)94S73 and S74Digoxin14S75Sodium-glucose transport protein 234S76inhibitors	Beta-blockers	115	569
Aldosterone receptor antagonist32570Aldosterone receptor antagonist32\$71Other antihypertensive drugs76\$72Antiarrhythmic drugs (Classes 1 and 3)94\$73 and \$74Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors	Alpha-blockers	58	570
Other antihypertensive drugs 76 S72 Antiarrhythmic drugs (Classes 1 and 3) 94 S73 and S74 Digoxin 14 S75 Sodium-glucose transport protein 2 34 S76 inhibitors Non-steroidal anti-inflammatory drugs 86 S77	Aldosterone receptor antagonist	30	570 571
Antiarrhythmic drugs (Classes 1 and 3) 94 \$73 and \$74 Digoxin 14 \$75 Sodium-glucose transport protein 2 34 \$76 inhibitors Non-steroidal anti-inflammatory drugs 86 \$77	Other antibypertensive drugs	76	\$72
Digoxin14\$75Sodium-glucose transport protein 234\$76inhibitors\$77	Antiarrhythmic drugs (Classes 1 and 3)	94	\$73 and \$74
Sodium-glucose transport protein 2 34 S76 inhibitors Non-steroidal anti-inflammatory drugs 86 S77		14	\$75 and \$71
inhibitors Non-steroidal anti-inflammatory drugs 86 \$77	Sodium_glucose transport protein ?	24	575
Non-steroidal anti-inflammatory drugs 86 S77	inhibitors	т	570
	Non-steroidal anti-inflammatory drugs	86	S77

ACE, angiotensin-converting enzyme.

Outcome events and safety search

Outcome events in the DaRe2THINK trial are collated from primary-care, secondary-care, and national mortality data. Automated safety searches are also performed on a monthly basis in the primary-care EHR. *Table 3* provides a summary of the code lists used for outcome events, including 4434 codes from CPRD Aurum and 610 codes from HES and ONS. For safety events, 3081 CPRD

Aurum codes were used to determine ischaemic stroke, intracranial bleeding, gastrointestinal bleeding, and other bleeding, with 483 codes to define associated hospitalization. Code lists were supplemented after review of 27 code lists from existing published literature,^{30–43} with the final list of individual codes detailed in Supplementary material online, *Appendix S1, Tables S24–S44*.

Baseline characteristics

To remove the need for case report forms, baseline characteristics in the DaRe2THINK trial are extracted from the primary-care EHR. *Table 4* provides a summary of these code lists, with a total of 6802 codes from CPRD Aurum. Measurements and blood results are extracted as the latest value in the past 12 months, medications during the last 12 months, and health conditions throughout the participant's primary-care EHR. For heart failure, hypertension, and diabetes, diagnosis is classified in two ways: (i) use of one or more of the relevant codes identified in the EHR and (ii) a 'secure' diagnosis based on the code identified *in addition* to an associated medication in the last 12 months (loop diuretic therapy, antihypertensive medications, or oral antidiabetic/insulin, respectively). Codes are provided in Supplementary material online, *Appendix S1, Tables S1–S77*.

Discussion

The DaRe2 framework was designed to link together a series of innovations for a digital clinical trial, creating a patient-centred approach with high quality output, yet low burden on healthcare staff. The ability to pre screen from over 13 million NHS patient records allows for targeted and efficient recruitment, and simplifies enrolment of patients that are representative of the real-world population. Combining all health data from primary and secondary care sources nationally enables a 'remote' RCT, with no requirement for baseline or follow-up visits. These processes are supported by advances in digital technology, including remote e-consent and patient reported outcomes. The DaRe2THINK trial is using this sequence of innovations to test a hypothesis of critical importance to public health, with the aim of preventing the long-term consequences of AF, including cognitive decline and vascular dementia.

This study outlines our approach to defining phenotypes that support all aspects of the trial, from understanding the characteristics of recruited participants, through to determination of clinical endpoints. Unlike conventional trials with case report forms, DaRe2THINK extracts all relevant information from structured and coded healthcare data sources. Hence, it is critical for dissemination and transparency that all coding schema for this trial are pre published and available for evaluation (plus re-use) by other researchers and clinicians. In a published review of 450 EHR-based studies, only 19 (5.1%) were accompanied by a full set of clinical codes,⁴⁴ severely limiting the value of those studies, and their implementation to routine care. The emergence of EHR systems across the world provides an important opportunity for healthcare-embedded clinical research,¹ but only when this is accompanied by a clear pathway from data collection to interpretation.²

The broad scope of coded healthcare data has been a limiting factor in prior attempts at EHR-embedded clinical trials. As evidenced by the large number of codes screened for this trial (over a million), a robust and systematic approach is required to ensure that patient events are not missed. The UK NHS is ideally suited to operationalize this sort of innovative trial, with a publicly funded health system that removes opportunity for personal financial gain from over or undercoding, while having systems in place to incentivize and monitor accurate coding. Structured healthcare data are used for quality and reimbursement purposes⁴ and there is an ability to link across different healthcare sources.⁴⁵ Linkage is critical to understanding the full healthcare utilization of each participant,³ in this case, combining historical and future primary-care, secondary-care, and death data. EHR-based trials may have an advantage in this regard compared with traditional trials, where investigators are relied on to document safety and outcome events. In the modern era of multiple heath providers for a multitude of different health conditions, sourcing safety and outcome events from national data have the potential to avoid missing endpoints, and potentially eliminate ascertainment bias in unblinded research. Although the value of trials based on routine EHR data is likely to vary depending on the disease being researched,⁴⁶ it is clear that the direction of travel is to enhance existing clinical trial infrastructure with digital innovations. In this regard, DaRe2THINK is the vanguard for cost-efficient RCTs embedded in clinical care. Previous trials have shown that although endpoints based on structured healthcare data may not replicate the traditional adjudication committee output, the estimated intervention effect is almost identical for adjudicated compared with EHR follow up.^{47,48}

There remain substantial limitations to effective use of EHR data for clinical trials, including ongoing changes to coding systems. For example, ICD-10 includes 68 000 codes (a number of countries have their own editions, with ICD-11 launched this year), and SNOMED CT has over 300 000 clinical concepts. New codes can also be created within primary healthcare databases, highlighting the importance of iterative review and updates as exemplified by our coding framework. Universal limitations, such as missing data, variation in data quality, reliability of coding, and imprecision in the choice of code used, are common across real-world data sources. These and specific areas of bias such as lack of representativeness and loss to follow up are substantially minimized in DaRe2THINK due to the NHS being the de facto provider of almost all healthcare in the UK, and CPRD providing broad and representative coverage. In this setting, concordance between primary and secondary care health data is high. For example, in a random sample of 50 000 patients in CPRD Aurum, 94% of the 1260 patients with a code for myocardial infarction had corroborating evidence in the secondary care HES records.³ Similar to our transparent approach to coding, the process of integrating primary and secondary healthcare sources in DaRe2THINK will be documented in an open-access, prospectively published Statistical Analysis Plan.

The DaRe2 approach was only possible with the support and guidance of our PPI team. We are indebted to the group for their constructive criticism throughout the design process, and their ongoing contribution to trial management to produce sustainable systems focused on the needs and wants of patients. Achieving a social licence for data-related clinical research is critical, and only achievable with high-level involvement by the public. In this case, we have used the PPI-POSITIVE approach,⁸ within the context of the CODE-EHR best practice framework for using healthcare data in clinical research.²

Conclusions

The DaRe2THINK trial is using a series of digital innovations to reshape the deployment of an adaptive randomized clinical trial embedded in routine healthcare, with minimal burden for staff and patients. Integrating national healthcare data from primary and secondary sources, the system will provide evidence for a key public health concern, the prevention of cognitive decline and dementia in the rapidly growing number of patients with AF. Systematic and transparent methodology in the use of structured healthcare data, together with a patient and public mandate, are critical for evolution of these novel approaches in order to improve healthcare in our communities.

Author contributions

The coding framework was designed and implemented by X.W., O.T., K.O., K.N., D.S., and D.K. The manuscript was drafted by X.W., A.R.M. and D.K., with all other authors editing the manuscript for intellectual content. D.K. provided supervision and was responsible for the decision to submit the manuscript. This study is based in part on data from the Clinical Practice Research Datalink obtained under licence from the UK Medicines and Healthcare products Regulatory Agency. The data are provided by patients and collected by the NHS as part of their care and support. The interpretation and conclusions contained in this study are those of the author/s alone.

Ethical approval and registration

The trial has received ethical/Health Research Authority approval (REC number: 21/NE/0021; IRAS project ID: 290420), regulatory approval (MHRA CTA 21761/0364) and is registered at Clinicaltrials.gov (NCT04700826), ISRCTN (21157803), and EudraCT (2020-005774-10).

Supplementary material

Supplementary material is available at European Heart Journal – Digital Health online.

Acknowledgements

In memory of Jaqueline Jones, member of the Patient and Public Involvement team. The authors acknowledge all members of the DaRe2THINK team and trial committees:

Trial Management Group: Susan Beatty (Clinical Practice Research Datalink, MHRA), Samir Mehta (Birmingham Clinical Trials Unit), Sophie Breeze (Clinical Practice Research Datalink, MHRA), Karen Lancaster (Clinical Practice Research Datalink, MHRA), and Stuart Fordyce (Clinical Practice Research Datalink, MHRA).

Data Coordination Team: Naomi Allen (University of Oxford, UK Biobank), Melanie Calvert (Centre for Patient Reported Outcomes Research and Birmingham Health Partners Centre for Regulatory Science and Innovation), Alastair Denniston (University Hospitals Birmingham NHS Foundation Trust and University of Birmingham), and George Gkoutos (University of Birmingham and University Hospitals Birmingham NHS Foundation Trust).

Health Economics Team: Sahan Jayawardana (London School of Economics and Political Science) and Elias Mossialos (London School of Economics and Political Science).

Expert Advisory Group: Simon Ball (University Hospitals Birmingham NHS Foundation Trust and Health Data Research UK Midlands), Colin Baigent (University of Oxford), Peter Brocklehurst (Birmingham Clinical Trials Unit), Will Lester (University Hospitals Birmingham NHS Foundation Trust), Richard McManus (University of Oxford), Stefano Seri (Aston University), and Janet Valentine (Innovate UK, UK Research and Innovation).

Independent Trial Steering Committee: A. John Camm (St. George's University of London), Sandra Haynes MBE (Patient and Public Involvement team lead), Dame Julie Moore (University of Warwick), Amy Rogers (University of Dundee), and Mary Stanbury (Patient and Public Involvement).

Independent Data Monitoring Committee: Marcus Flather (University of East Anglia), Suzy Walker (General Practitioner, Fortrose Medical Practice, Highland), and Duolao Wang (Liverpool School of Tropical Medicine).

Funding

The DaRe2THINK trial is funded by the National Institute for Health and Care Research (NIHR130280). Medications in the trial are funded by the UK Department of Health and Social Care. The opinions expressed in this study are those of the authors and do not represent the NIHR or the UK Department of Health and Social Care.

Conflict of interest: All authors have completed the ICMJE uniform disclosure form (www.icmje.org/coi_disclosure.pdf) and declare:

X.W. and A.R.M. posts are funded by the National Institute for Health and Care Research (NIHR) (130280). O.T. post was funded by the EU/EFPIA Innovative Medicines Initiative (BigData@Heart 116074) and a grant from Amomed Pharma, outside the submitted work. K.O. has nothing to disclose. R.E.G. reports grants from NIHR, during the conduct of the study. P.M. has nothing to disclose. T.W. has nothing to disclose. S.H. reports a grants from NIHR (130280), during the conduct of the study. K.N. reports grants from NIHR, during the conduct of the study; and research grants from UKRI/MRC, Kennedy Trust for Rheumatology Research, Health Data Research UK, Wellcome Trust, European Regional Development Fund, Institute for Global Innovation, Boehringer Ingelheim, Action Against Macular Degeneration Charity, Midlands Neuroscience Teaching and Development Funds, South Asian Health Foundation, Vifor Pharma, College of Police, and CSL Behring, outside the submitted work; and consulting fees from Boehringer Ingelheim, Sanofi, CEGEDIM, MSD and holds a leadership/fiduciary role with NICST, a charity and OpenClinical, a Social Enterprise; outside the submitted work. D.S. reports grants from NIHR (130280), during the conduct of the study; and grants from Pfizer outside the submitted work. D.K. reports grants from NIHR (NIHR CDF-2015-08-074 RATE-AF; NIHR130280 DaRe2THINK; NIHR132974 D2T-NeuroVascular), the British Heart Foundation (PG/17/55/33087, AA/18/2/34218 and FS/CDRF/21/ 21032), the EU/EFPIA Innovative Medicines Initiative (BigData@Heart 116074), and the European Society of Cardiology supported by educational grants from Boehringer Ingelheim/BMS-Pfizer Alliance/Bayer/ Daiichi Sankyo/Boston Scientific, the NIHR/University of Oxford Biomedical Research Centre and British Heart Foundation/University of Birmingham Accelerator Award (STEEER-AF NCT04396418). In addition, he has received a research grant to his institution from Pfizer; and research grants and advisory board fees from Bayer, Amomed and Protherics Medicines Development; all outside the submitted work.

Data availability

No additional data are available at this time.

References

- Studer R, Sartini C, Suzart-Woischnik K, Agrawal R, Natani H, Gill SK, Wirta SB, Asselbergs FW, Dobson R, Denaxas S, Kotecha D. Identification and mapping realworld data sources for heart failure, acute coronary syndrome, and atrial fibrillation. *Cardiology* 2022;**147**:98–106.
- Kotecha D, Asselbergs FW, European Society of Cardiology, BigData@Heart consortium, CODE-EHR international consensus group. CODE-EHR best practice framework for the use of structured electronic healthcare records in clinical research. BMJ 2022;378:e069048.
- Persson R, Sponholtz T, Vasilakis-Scaramozza C, Hagberg KW, Williams T, Kotecha D, Myles P, Jick SS. Quality and completeness of myocardial infarction recording in Clinical Practice Research Datalink Aurum. *Clin Epidemiol* 2021;**13**:745–753.
- Burns EM, Rigby E, Mamidanna R, Bottle A, Aylin P, Ziprin P, Faiz OD. Systematic review of discharge coding accuracy. J Public Health (Oxf) 2012;34:138–148.
- Persson R, Vasilakis-Scaramozza C, Hagberg KW, Sponholtz T, Williams T, Myles P, Jick SS. CPRD Aurum database: assessment of data quality and completeness of three important comorbidities. *Pharmacoepidemiol Drug Saf* 2020;29:1456–1464.
- 6. Kotecha D, Shukla D, Clinical Practice Research Datalink. Preventing stroke, premature death and cognitive decline in a broader community of patients with atrial fibrillation using healthcare data for pragmatic research: a randomised controlled trial (DaRe2THINK). https://www.birmingham.ac.uk/dare2think (30 March 2022).
- Wolf A, Dedman D, Campbell J, Booth H, Lunn D, Chapman J, Myles P. Data resource profile: Clinical Practice Research Datalink (CPRD) Aurum. Int J Epidemiol 2019;48:1740–1740g.
- Bunting KV, Stanbury M, Tica O, Kotecha D. Transforming clinical research by involving and empowering patients - the RATE-AF randomized trial. *Eur Heart J* 2021;42: 2411–2414.
- Kirchhof P, Benussi S, Kotecha D, Ahlsson A, Atar D, Casadei B, Castella M, Diener HC, Heidbuchel H, Hendriks J, Hindricks G, Manolis AS, Oldgren J, Popescu BA, Schotten U, Van Putte B, Vardas P. 2016 ESC guidelines for the management of atrial fibrillation developed in collaboration with EACTS. *Eur Heart J* 2016;**37**:2893–2962.
- Kotecha D, Holmes J, Krum H, Altman DG, Manzano L, Cleland JG, Lip GY, Coats AJ, Andersson B, Kirchhof P, von Lueder TG, Wedel H, Rosano G, Shibata MC, Rigby A, Flather MD, Beta-Blockers in Heart Failure Collaborative Group. Efficacy of beta blockers in patients with heart failure plus atrial fibrillation: an individual-patient data meta-analysis. *Lancet* 2014;**384**:2235–2243.
- Tica O, Khamboo W, Kotecha D. Breaking the cycle of HFpEF and AF. Card Fail Rev 2022. doi:10.15420/cfr.2022.03.
- 12. Kotecha D, Breithardt G, Camm AJ, Lip GYH, Schotten U, Ahlsson A, Arnar D, Atar D, Auricchio A, Bax J, Benussi S, Blomstrom-Lundqvist C, Borggrefe M, Boriani G, Brandes A, Calkins H, Casadei B, Castella M, Chua W, Crijns H, Dobrev D, Fabritz L, Feuring M, Freedman B, Gerth A, Goette A, Guasch E, Haase D, Hatem S, Haeusler KG, Heidbuchel H, Hendriks J, Hunter C, Kaab S, Kespohl S, Landmesser U, Lane DA, Lewalter T, Mont L, Nabauer M, Nielsen JC, Oeff M, Oldgren J, Oto A, Pison L, Potpara T, Ravens U, Richard-Lordereau I, Rienstra M, Savelieva I, Schnabel R, Sinner MF, Sommer P, Themistoclakis S, Van Gelder IC, Vardas PE, Verma A, Wakili R, Weber E, Werring D, Willems S, Ziegler A, Hindricks G, Kirchhof P. Integrating new approaches to atrial fibrillation management: the 6th AFNET/EHRA Consensus Conference. *Europace* 2018;20:395–407.
- Conen D, Rodondi N, Muller A, Beer JH, Ammann P, Moschovitis G, Auricchio A, Hayoz D, Kobza R, Shah D, Novak J, Schlapfer J, Di Valentino M, Aeschbacher S, Blum S, Meyre P, Sticherling C, Bonati LH, Ehret G, Moutzouri E, Fischer U, Monsch AU, Stippich C, Wuerfel J, Sinnecker T, Coslovsky M, Schwenkglenks M, Kuhne M, Osswald S. Relationships of overt and silent brain lesions with cognitive function in patients with atrial fibrillation. J Am Coll Cardiol 2019;**73**:989–999.
- Islam MM, Poly TN, Walther BA, Yang HC, Wu CC, Lin MC, Chien SC, Li YC. Association between atrial fibrillation and dementia: a meta-analysis. *Front Aging Neurosci* 2019;11:305.

- 15. Kirchhof P, Haeusler KG, Blank B, De Bono J, Callans D, Elvan A, Fetsch T, Van Gelder IC, Gentlesk P, Grimaldi M, Hansen J, Hindricks G, Al-Khalidi HR, Massaro T, Mont L, Nielsen JC, Nolker G, Piccini JP, De Potter T, Scherr D, Schotten U, Themistoclakis S, Todd D, Vijgen J, Di Biase L. Apixaban in patients at risk of stroke undergoing atrial fibrillation ablation. *Eur Heart J* 2018;**39**:2942–2955.
- Lane DA, Skjoth F, Lip GYH, Larsen TB, Kotecha D. Temporal trends in incidence, prevalence, and mortality of atrial fibrillation in primary care. J Am Heart Assoc 2017; 6:e005155.
- European Medicines Agency Committee for Medicinal Products for Human Use. Assessment report: Direct oral anticoagulants (DOACs). EMA/194375/2020; Procedure number: EMEA/H/A-5(3)/1487. 2020.
- Kotecha D, Pollack CV Jr, De Caterina R, Renda G, Kirchhof P. Direct oral anticoagulants halve thromboembolic events after cardioversion of AF compared with warfarin. J Am Coll Cardiol 2018;72:1984–1986
- National Institute for Health and Care Excellence. Atrial fibrillation: diagnosis and management. NICE clinical guideline 196. 2021. www.nice.org.uk/guidance/ng196.
- Wise J, de Barron AG, Splendiani A, Balali-Mood B, Vasant D, Little E, Mellino G, Harrow I, Smith I, Taubert J, van Bochove K, Romacker M, Walgemoed P, Jimenez RC, Winnenburg R, Plasterer T, Gupta V, Hedley V. Implementation and relevance of FAIR data principles in biopharmaceutical R&D. *Drug Discov Today* 2019;**24**: 933–938.
- NHS Digital. SNOMED CT. 2021. https://digital.nhs.uk/services/terminology-andclassifications/snomed-ct (11 May 2022).
- NHS Digital. Read codes. 2020. https://digital.nhs.uk/services/terminology-andclassifications/read-codes (11 May 2022).
- NHS Business Services Authority. Dictionary of medicines and devices (dm + d). 2021. https://www.nhsbsa.nhs.uk/pharmacies-gp-practices-and-appliance-contractors/ dictionary-medicines-and-devices-dmd (11 May 2022).
- 24. World Health Organization. International classification of disease version 10. 2019. https://icd.who.int/browse10/2019/en#/XXI (11 May 2022).
- NHS. OPCS classification of interventions and procedures. 2022. https://www. datadictionary.nhs.uk/data_elements/opcs-4_code.html (11 May 2022).
- Office for National Statistics. User guide to mortality statistics. 2021. https://www. ons.gov.uk/peoplepopulationandcommunity/birthsdeathsandmarriages/deaths/ methodologies/userguidetomortalitystatisticsjuly2017 (11 May 2022).
- 27. Wang X, Gkoutos GV, Kotecha D. SWAT 158: validation of clinical events in the DaRe2THINK data-enabled clinical trial. MRC Northern Ireland Hub for Trials Methodology Research: Study-Within-A-Trial (SWAT) Repository Store. 2019. https:// www.qub.ac.uk/sites/TheNorthernIrelandNetworkforTrialsMethodologyResearch/ FileStore/Filetoupload,1218963,en.pdf (May 2022).
- Gokhale KM, Chandan JS, Toulis K, Gkoutos G, Tino P, Nirantharakumar K. Data extraction for epidemiological research (DExtER): a novel tool for automated clinical epidemiology studies. *Eur J Epidemiol* 2021;**36**:165–178.
- Jick SS, Hagberg KW, Persson R, Vasilakis-Scaramozza C, Williams T, Crellin E, Myles P. Quality and completeness of diagnoses recorded in the new CPRD Aurum database: evaluation of pulmonary embolism. *Pharmacoepidemiol Drug Saf* 2020;29: 1134–1140.
- Wu J, Mamas MA, Mohamed MO, Kwok CS, Roebuck C, Humberstone B, Denwood T, Luescher T, de Belder MA, Deanfield JE, Gale CP. Place and causes of acute cardiovascular mortality during the COVID-19 pandemic. *Heart* 2021;**107**:113–119.
- McCormick N, Lacaille D, Bhole V, Avina-Zubieta JA. Validity of heart failure diagnoses in administrative databases: a systematic review and meta-analysis. *PLoS One* 2014;9:e104519.
- McCormick N, Bhole V, Lacaille D, Avina-Zubieta JA. Validity of diagnostic codes for acute stroke in administrative databases: a systematic review. *PLoS One* 2015;10: e0135834.

- 33. Lix LM, Sobhan S, St-Jean A, Daigle JM, Fisher A, Yu OHY, Dell'Aniello S, Hu N, Bugden SC, Shah BR, Ronksley PE, Alessi-Severini S, Douros A, Ernst P, Filion KB. Validity of an algorithm to identify cardiovascular deaths from administrative health records: a multi-database population-based cohort study. *BMC Health Serv Res* 2021; 21:758.
- Nedkoff L, Lopez D, Hung J, Knuiman M, Briffa TG, Murray K, Davis E, Aria S, Robinson K, Beilby J, Hobbs MST, Sanfilippo FM. Validation of ICD-10-AM coding for myocardial infarction subtype in hospitalisation data. *Heart Lung Circ* 2022;**31**: 849–858.
- Ando T, Ooba N, Mochizuki M, Koide D, Kimura K, Lee SL, Setoguchi S, Kubota K. Positive predictive value of ICD-10 codes for acute myocardial infarction in Japan: a validation study at a single center. *BMC Health Serv Res* 2018;**18**:895.
- Nagamine T, Gillette B, Pakhomov A, Kahoun J, Mayer H, Burghaus R, Lippert J, Saxena M. Multiscale classification of heart failure phenotypes by unsupervised clustering of unstructured electronic medical record data. *Sci Rep* 2020;**10**:21340.
- Bosco E, Hsueh L, McConeghy KW, Gravenstein S, Saade E. Major adverse cardiovascular event definitions used in observational analysis of administrative databases: a systematic review. BMC Med Res Methodol 2021;21:241.
- Miao B, Hernandez AV, Alberts MJ, Mangiafico N, Roman YM, Coleman CI. Incidence and predictors of major adverse cardiovascular events in patients with established atherosclerotic disease or multiple risk factors. J Am Heart Assoc 2020;9:e014402.
- Groenendyk JW, Rivera AS, Sinha A, Lloyd-Jones DM, Feinstein MJ. Changes in proportionate cardiovascular mortality in patients with chronic infectious and inflammatory conditions in the United States, 1999–2018. *Sci Rep* 2021;**11**:23985.
- Carrero JJ, de Jager DJ, Verduijn M, Ravani P, De Meester J, Heaf JG, Finne P, Hoitsma AJ, Pascual J, Jarraya F, Reisaeter AV, Collart F, Dekker FW, Jager KJ. Cardiovascular and noncardiovascular mortality among men and women starting dialysis. *Clin J Am Soc Nephrol* 2011;6:1722–1730.
- McDonald L, Sammon CJ, Samnaliev M, Ramagopalan S. Under-recording of hospital bleeding events in UK primary care: a linked Clinical Practice Research Datalink and Hospital Episode Statistics study. *Clin Epidemiol* 2018;**10**:1155–1168.
- Mehta NK, Abrams LR, Myrskyla M. US life expectancy stalls due to cardiovascular disease, not drug deaths. Proc Natl Acad Sci U S A 2020;117:6998–7000.
- Miniño A, Klein R. Health Mortality from Major Cardiovascular Diseases. Health E-Stats (2007). Hyattsville, Maryland: National Center for Health Statistics; 2010.
- Springate DA, Kontopantelis E, Ashcroft DM, Olier I, Parisi R, Chamapiwa E, Reeves D. Clinicalcodes: an online clinical codes repository to improve the validity and reproducibility of research using electronic medical records. *PLoS One* 2014;9:e99825.
- 45. Cowie MR, Blomster JI, Curtis LH, Duclaux S, Ford I, Fritz F, Goldman S, Janmohamed S, Kreuzer J, Leenay M, Michel A, Ong S, Pell JP, Southworth MR, Stough WG, Thoenes M, Zannad F, Zalewski A. Electronic health records to facilitate clinical research. *Clin Res Cardiol* 2017;**106**:1–9.
- 46. McCord KA, Ewald H, Agarwal A, Glinz D, Aghlmandi S, Ioannidis JPA, Hemkens LG. Treatment effects in randomised trials using routinely collected data for outcome assessment versus traditional trials: meta-research study. *BMJ* 2021;**372**:n450.
- Kjoller E, Hilden J, Winkel P, Galatius S, Frandsen NJ, Jensen GB, Fischer Hansen J, Kastrup J, Jespersen CM, Hildebrandt P, Kolmos HJ, Gluud C, Group CT. Agreement between public register and adjudication committee outcome in a cardiovascular randomized clinical trial. *Am Heart J* 2014;**168**:197–204. e191–194.
- Bowman L, Mafham M, Wallendszus K, Stevens W, Buck G, Barton J, Murphy K, Aung T, Haynes R, Cox J, Murawska A, Young A, Lay M, Chen F, Sammons E, Waters E, Adler A, Bodansky J, Farmer A, McPherson R, Neil A, Simpson D, Peto R, Baigent C, Collins R, Parish S, Armitage J. Effects of aspirin for primary prevention in persons with diabetes mellitus. N Engl J Med 2018;**379**:1529–1539.