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Title:
Identification and mapping real-world data sources for heart failure, acute coronary syndrome, and atrial fibrillation

Running title: Identification and mapping real-world data sources

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Abstract

Background: Transparent and robust real-world evidence sources are increasingly important for global health, including cardiovascular diseases. We aimed to identify global real-world data (RWD) sources for heart failure (HF), acute coronary syndrome (ACS), and atrial fibrillation (AF).

Methods: We conducted a systematic review of publications with RWD pertaining to HF, ACS, and AF (2010-2018), generating a list of unique data sources. Metadata were extracted based on the source type (e.g., electronic health records, genomics, clinical data), study design, population size, clinical characteristics, follow-up duration, outcomes, and assessment of data availability for future studies and linkage.

Results: Overall, 11,889 publications were retrieved for HF, 10,729 for ACS, and 6,262 for AF. From these, 322 (HF), 287 (ACS), and 220 (AF) data sources were selected for detailed review. The majority of data sources had near complete data on demographic variables (HF: 94%, ACS: 99%, and AF: 100%) and considerable data on comorbidities (HF: 77%, ACS: 93%, and AF: 97%). The least reported data categories were drug codes (HF, ACS, and AF: 10%) and caregiver involvement (HF: 6%, ACS: 1%, and AF: 1%). Only a minority of data sources provided information on access to data for other researchers (11%) or whether data could be linked to other data sources to maximize clinical impact (20%). The list and metadata for the RWD sources are publicly available at www.escardio.org/bigdata.

Conclusions: This review has created a comprehensive resource of cardiovascular data sources, providing new avenues to improve future real-world research and to achieve better patient outcomes.

Keywords: Real-world data, real-world evidence, data sources, cardiovascular

Background

Cardiovascular (CV) disease is the leading cause of death worldwide [1], accounting for more than 17 million deaths in 2015 alone [2]. According to the World Health Organization (WHO), the annual number of deaths due to CV diseases globally is projected to increase to 20.5 million by 2020 and 24.5 million by 2030 [3]. Moreover, in both high-income and middle-income countries, the main cause of death has shifted over time from communicable to non-communicable diseases, with a high burden on national health systems [4].

Real-world data (RWD) have played a key role in CV disease–related decision-making, especially in recent years, due to a widening range of new therapies and increasing demands for justification of their effectiveness. Translating RWD into real-world evidence (RWE) can provide information throughout a product’s life cycle [5]. RWE can help design pivotal phase 3 trials by reducing the required sample size, supporting recruitment and thereby saving time [6] and informing appropriate selection criteria [7, 8]. RWE can provide outcomes of care in real-world settings, thus improving the external validity of clinical trial findings, and offer insights into coverage and payment decisions to support health authority decision-making [9, 10]. However, limitations of RWD should also be acknowledged, which broadly include bias and confounding, incomplete data, different legal frameworks leading to restricted data sharing, and lack of universally accepted methodological standards [9-11]. In addition, the evidence landscape is constantly evolving with respect to the conduct and reporting of RWE studies. The recent retraction from major medical journals of apparently fraudulent RWD on COVID-19 [12] highlights the urgent need for more transparency and access to global data sources.

This review aimed to identify global RWD sources pertaining to heart failure (HF), acute coronary syndrome (ACS), and atrial fibrillation (AF) in order to facilitate new evidence research and improve patient outcomes. Our objective was to help global researchers move towards the FAIR principles for RWD—Findable, Accessible, Interoperable, and Reusable [13].

Methods

The European Union Innovative Medicines Initiative (IMI) public-private consortium launched the BigData@Heart project with the goal of developing a big data–driven translational research platform from RWE focusing on HF, ACS, and AF. Through this translational research platform, BigData@Heart aims to deliver clinically relevant disease phenotypes and support drug development and personalized medicine [14-16]. One of the undertakings of this initiative is to identify and characterize available RWD sources that would serve as a starting point to identify existing data sets that could help address research questions at scale. This was a collaborative research project with clinical academics and Novartis conducted the literature review and identification of potential real-world data sources.

A systematic literature search was conducted in MEDLINE and EMBASE using the OvidSP platform for the period January 2010 to March 2018 to identify publications using RWD sources for HF, ACS, and AF. The review was not prospectively registered. We did not include publications before 2010 because older RWD sources may
not be relevant to current practice. Disease-specific search strategies (using Medical Subject Headings [MeSH] terms) were combined with study design terms to identify research publications that either generated primary RWD or used existing RWD sources. Identified data sources from these publications were categorized according to predefined geographical locations: Europe; the United States (US); Latin America/Canada; and Asia-Pacific, Middle East, and Africa (APMA).

Inclusion and exclusion criteria
We included English-language publications using different data sources as defined by the authors, such as structured data sources (administrative data and registries), medical records or charts, insurance claims, health surveys, and observational studies for HF, ACS, and AF. Publications that did not generate primary RWD or did not study existing RWD sources, as well as guidelines, editorials, letters, and reviews were excluded. Additionally, we excluded clinical trials or interventional studies, in vitro/preclinical studies, and data sources with less than 50 patients.

Screening, selection, and extraction of data sources
The search strategies are presented in the Additional Files (HF: Additional File 1: [Table 1], ACS: Additional File 2 [Table 2], AF: Additional File 3 [Table 3]). All publications identified from the literature searches were first screened based on the title and abstract by a single reviewer and duplicates were removed. The inclusion and exclusion criteria were applied at this stage to generate a list of full-text reviews. Of the included publications, 10% were randomly selected and checked for discrepancies, which were reconciled through group discussions. For the included publications, names of identified data sources, type (single-centre or multicentre), and geographical location were extracted using a predefined screening tool. Publications with the same data source were grouped by name, and data were extracted into a single record to avoid double counting of data sources in subsequent analyses. Thereafter, a list of unique data sources available from the literature search was prepared for each indication.

From this list, selected data sources were further mapped and extracted in detail based on different criteria for each disease indication:
- HF: (i) multicentre data sources; (ii) single-centre sources where details of left ventricular ejection fraction (LVEF) for HF patients were available
- ACS: (i) all ACS-specific data sources; (ii) multicentre data sources, data sources with unknown centre details, and data sources with >500 ACS patients
- AF: (i) multicentre data sources, data sources with unknown centre details, and data sources with >500 AF patients

Data sources with larger sample sizes were prioritized with a view on big data and potentially more robust analyses. For the data sources identified based on the above criteria, information presented in the included publications was extracted, including additional information on data source details (description, coverage, and follow-up) and availability of clinically relevant key variables related to HF, ACS, and AF (diagnosis and staging, demographics, management [including procedures], test results and treatments, burden of disease [including costs], deaths and resource use, quality of life, and adverse events). In addition, publicly available information related to the data sources, such as the data source holder/owner, access and linkage possibility, supporting documentation and its governance aspects, was extracted and recorded.

Results
Heart failure
Of the 11,889 publications retrieved from the HF literature search, 1,326 unique data sources were identified, of which 322 RWD sources were selected for detailed mapping (Additional File 4: Figure 1). Overall, 74% of these data sources were disease specific, with registries being the most common type of data source (45%). Geographically, 47% of the published HF data sources were from Europe, followed by the US (21%), APMA (20%), Latin America/Canada (8%), and multiregional (4%) (Figure 1). Germany had the highest number of data sources in Europe (n=15); Japan, in APMA (n=10); and Canada, in Latin America/Canada (n=12). The top five HF data sources based on the highest number of publications are presented in Figure 2.

Completeness of variables varied across the mapped data sources and ranged from 0 to 78%. The most commonly recorded variables were age and gender (94%), hospital admissions (81%), comorbidities (77%), mortality (75%), and LVEF (73%; increased by selection criteria). The least recorded data variables were drug
codes (10%), dates of procedures and prescriptions (7%), and caregiver involvement (6%). In terms of comorbidities, the proportion of HF data sources reporting ACS and AF as a comorbidity was 16% and 28%, respectively. Information on access to these data sources through purchasing, licensing, or collaboration with the data set owners was reported for 6% of the sources, whereas it was unknown for the remaining sources. Linking of these data with other data sources was reported in 18% of the sources, whereas the possibility of linkage was unknown for the remainder.

Acute coronary syndrome
From the 10,729 publications retrieved through the literature search, 1,560 unique data sources were identified, of which 287 were further selected and mapped (Additional File 5: Figure 2). Over half of these data sources (52%) were from Europe; 25%, APMA; 9%, US; and 8%, Latin America/Canada; 6% of the sources were multiregional (Figure 3). The highest number of data sources was from Germany in Europe (n=20), Japan in APMA (n=21), and Canada in Latin America/Canada (n=17). Over 80% of the mapped data sources were registries (Figure 4). The Swedish Web-System for Enhancement and Development of Evidence-Based Care in Heart Disease Evaluated According to Recommended Therapies (SWEDEHEART) registry had the highest number of publications (n=100) identified during the search period (Figure 2).

Completeness for recorded variables varied from 5% to 70%. The most commonly available clinical variables were age and gender (99%), mortality (95%), comorbidities (93%), inpatient diagnostic or therapeutic procedures (84%), and prescribed drugs (74%). The least recorded variables were date of ACS diagnosis (6%), dates of procedures and prescriptions (5%), procedure costs (3%), drug codes, and caregiver involvement and costs (1% each). The proportion of ACS data sources capturing the presence of HF and AF as a comorbidity was 27% and 8%, respectively. Information on access to these data sources was provided in 6% of the sources and linkage of these data sources with other data sets was possible in 20%.

Atrial fibrillation
From the 6,262 publications retrieved via the literature search, 701 unique data sources were identified, of which 220 data sources were further mapped (Additional File 6: Figure 3). Geographically, Europe had the highest number of data sources (40%), followed by the US (30%), APMA (20%), and Latin America/Canada (7%); 4% of the sources were multiregional (Figure 5). The highest number of data sources was from the United Kingdom (UK) in Europe (n=13), Japan in APMA (n=12), and Canada in Latin America/Canada (n=14). Registries (42%) were the most common type of data sources, followed by administrative databases (18%), observational studies (17%), claims (13%), and surveys (10%) (Figure 4). The top five data sources based on the highest number of publications are presented in Figure 2.

Coverage of variables differed across the mapped data sources and their completeness ranged from 10% to 60%. The most widely reported data variables were age and gender (100%), comorbidities (97%), prescribed drugs (91%), stroke risk (81%), mortality (67%), and hospitalizations (66%), whereas the least reported variables were date of AF diagnosis (10%), drug codes (10%), quality of life (10%), and caregiver involvement (1%). HF and ACS as comorbidities were recorded for 92% and 49% of the AF data sources, respectively. Information on access to data sources was reported in 25% of the mapped sources, whereas for the remaining sources, possibility of access was unknown. Linkage of these data sources with other data sources was possible in 28%.

Discussion
This review aimed to identify global RWD sources focusing on three common CV diseases and make them publicly available as a resource for researchers. Previous studies have identified RWD sources in disease areas such as chronic obstructive pulmonary disease [17] and Parkinson’s disease [18] as well as generic RWD data sources [19], but to our knowledge, no study has reported RWD sources focusing on CV diseases across different geographies. We were able to map 322 RWD sources for HF, 287 for ACS, and 220 for AF. The mapping and provision of these sources in this review aims to enhance the generation of RWE across CV diseases. Importantly, we also define current limitations, such as lack of access to data, linkage with other sources, and insight on cross-comorbidity that should be improved in order to achieve maximum patient benefit from future RWE.

In December 2018, the US Food and Drug Administration (FDA) released a guidance document for the use of RWE to support regulatory decision-making for drugs and medical devices [20]. Similarly, in Europe, the European Medicines Agency (EMA), with its adaptive pathway initiative, highlighted RWE as an important source to further support evidence collected through randomized controlled trials (RCTs) [9]. In addition to the EMA and
FDA, Health Technology Assessment International, in its global policy forum, presented the availability and use of RWE for health technology assessment [21], and the National Institute for Health and Care Excellence in the UK has documented the use of RWE in its decision-making [22]. In the context of the coronavirus pandemic, RWD has been used extensively to manage public health programmes, although the recent controversy and retraction of studies by leading journals has highlighted the need for robust evaluation before apparent RWD becomes RWE [12].

The growing importance of RWE can further be ascertained through many examples, including selected drug approvals during 1999-2014 by the FDA and EMA, which were largely based on uncontrolled studies for oncology and orphan indications [23]. For health technology assessments, certain outcomes such as costs and quality-adjusted life-years are often retrieved from non-RCT data [24]. With the growing need for RWE, we require varied, high-quality, and transparent sources of RWD to cater to different research objectives related to the epidemiology or burden of disease.

Across the three CV indications, we found that most data sources were currently from Europe and North America, but a growing number are now presented from the Middle East, Asia, Russia, and South America. Collection of RWD requires relatively high upfront investment, which might be more feasible in high-income countries. Among the European HF data sources, and consistent with other published data, the Swedish Heart Failure Registry was the most frequent source for generating RWE [25]. The most published data sources for ACS and AF were the SWEDEHEART registry and the Danish nationwide-linked admin registries, respectively.

In the present review, for all the three CV conditions, demographics and comorbidities were the most commonly available variables, whereas costs and caregiver involvement were least reported. This could be because most of the identified data sources were registries. Moreover, cost to the healthcare system and caregiver involvement cannot be collected directly from patients, existing healthcare records, or medical charts. Data sources for HF also provided information on mortality, hospitalization, and LVEF. For ACS, information on mortality and prescribed drugs was captured in 95% and 74% of the data sources, respectively. For AF, other commonly reported variables were prescribed drugs, stroke risk, mortality, and hospitalization. Taken together, these data sources provide a wealth of information on patient characteristics and clinical burden; however, data pertaining to humanistic and economic burden are limited. These trends are similar to those observed in non-CV conditions [18, 25].

In the absence of universally accepted methodological standards for data models and infrastructure, the accessibility, linkage, and comparability of RWD sources is currently a challenge. This can prevent establishment of larger data sets by linking RWD to generate more robust and representative RWE [9, 26]. In line with this, this review reports low accessibility and possibility of linkage based on information retrieved from the public domain. The alternative, i.e. personal communication with data holders, can be time-consuming and potentially unproductive. This challenge may be addressed through efforts in private-public collaborations and within the European framework; for example, data set owners could be invited to the European Medical Information Framework (EMIF) catalogue [27], which allows users to explore population-based data sources. Linking of these data sources requires harmonization similar to that in other large IMI initiatives such as the European Health Data & Evidence Network [28]. Translation to clinical practice and the development of new RWE will be aided by integration and linkage of molecular and genetic studies with RWD sources; this developing field has the potential to enable more rapid translation of mechanistic studies to improve patient care.

This review has certain limitations, including incomplete information on the RWD sources because of the limited information available in the public domain. Many databases may contain more data than is currently reported in the tool and inversely, some variables may be recorded for only a subset of patients (e.g. LVEF). This review reflects the current state-of-the-art, however RWD sources are continually being generated and revised. Key recent publications from the identified data sources are presented in Table 1 and demonstrate the broad impact that RWE can have on clinical practice. The consortium will update this review periodically (see www.escardio.org/bigdata for future updates), and the EMIF catalogue is open for future collaborations and within the European framework; for example, data set owners could be invited to the European Medical Information Framework (EMIF) catalogue [27], which allows users to explore population-based data sources. Linking of these data sources requires harmonization similar to that in other large IMI initiatives such as the European Health Data & Evidence Network [28]. Translation to clinical practice and the development of new RWE will be aided by integration and linkage of molecular and genetic studies with RWD sources; this developing field has the potential to enable more rapid translation of mechanistic studies to improve patient care.

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and confounding factors that limit the value of observational data [29]. However, RWE can complement clinical trial data, and allows an understanding of the epidemiology and interaction of diseases.

**Conclusions**
In summary, this review identified and mapped worldwide RWD sources pertaining to HF, ACS, and AF, thus providing researchers with a knowledge base to conduct feasibility assessments of these data sources for RWE studies. The list of and metadata for the data sources are publicly available at www.escardio.org/bigdata. Epidemiological research can be conducted using the wealth of individual data sources available. However, further details and access to the RWD sources, enhanced collaboration and harmonization between data holders (academia and industry), as well as integration of data sets would allow for generation of more complex and impactful evidence. This could support CV disease drug development, market access, and use of interventions in clinical practice, eventually leading to improved CV outcomes and patient well-being.
List of abbreviations
ACS: Acute coronary syndrome
AF: Atrial fibrillation
APMA: Asia-Pacific, Middle East, and Africa
CV: cardiovascular
EMA: European Medicines Agency
EMIF: European Medical Information Framework
FAIR: Findable, Accessible, Interoperable, and Reusable
FDA: Food and Drug Administration
HF: Heart failure
IMI: Innovative Medicines Initiative
LVEF: Left ventricular ejection fraction
MeSH: Medical Subject Headings
RCT: randomized controlled trial
RWD: Real-world data
RWE: Real-world evidence
UK: United Kingdom
US: United States
WHO: World Health Organization

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None.

Statement of Ethics
An ethics statement was not required for this study type, no human or animal subjects or materials were used.

Conflict of Interest Statement
Prof. Kotecha reports, outside of this study, grants from the National Institute for Health Research (NIHR CDF-2015-08-074 RATE-AF; NIHR HTA-130280 DaRe2THINK; NIHR EME-132974 D2T-NV), the British Heart Foundation (PG/17/55/33087 and AA/18/2/34218), EU/EFPIA Innovative Medicines Initiative (BigData@Heart 116074), 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 (STEER-AF NCT04396418), Amomed Pharma and IRCCS San Raffaele/Menarini (Beta-blockers in Heart Failure Collaborative Group NCT0083244); in addition to personal fees from Bayer (Advisory Board), AtriCure (Speaker fees), Protherics Medicines Development (Advisory Board) and Myokardia (Advisory Board). Prof. Kotecha is one of the associate editors to Cardiology. Dr. Sartini, Dr. Agrawal, Dr. Natani, Prof. Denaxas, Prof. Asselbergs, and Prof. Dobson have nothing to disclose. Dr. Gill reports funding through the BigData@Heart Innovative Medicines Initiative, grant no.116074. Dr. Suzart-Woischnak reports personal fees from Bayer AG, during the conduct of the study; other from Bayer AG, outside the submitted work. Dr. Wirta reports other from Novartis, during the conduct of the study. Dr. Studer reports personal fees from Novartis Pharmaceuticals, during the conduct of the study; other from Novartis Pharmaceuticals, outside the submitted work.

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Author Contributions
R.S. conceptualised the study and critically reviewed the manuscript. D.K. analysed the data and critically reviewed the manuscript. H.N. analysed the data and drafted the manuscript. C.S., K. S. W., R.A, S.G., S.B.W., F.W.A., R.D., S.D., critically reviewed the manuscript. All authors have made substantial contribution in critically reviewing and writing the manuscript. All authors have read and approved the submission of the paper.

Data Availability Statement
The list of metadata of publicly available data sources can be found at: www.escardio.org/bigdata
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Legends

Figure 1: Geographical distribution of HF data sources
Global HF data source distribution, with darker shades of colour representing more data sources in each of the presented regions.
HF, heart failure; LaCan, Latin America/Canada.

Figure 2: Top five data sources as per count of publications
Data sources with the highest number of publications during the search period of this review. On the right, the possibility to access or link these data sources is presented.
ACTION-GTWG, Acute Coronary Treatment and Intervention Outcomes Network (ACTION) Registry-Get with the Guidelines; GRACE/GRACE 2, Global Registry of Acute Coronary Events/Global Registry of Acute Coronary Events 2.

Figure 3: Geographical distribution of ACS data sources
Global ACS data source distribution, with darker shades of colour representing more data sources in each of the presented regions.
ACS, acute coronary syndrome.

Figure 4: Distribution as per type of data sources
Data sources mapped in this review are categorized broadly into six different categories, comprising of observational studies, registries, surveys, administrative databases, claims databases, and others. Observational studies include cohort studies, cross-sectional studies, prospective studies, retrospective studies, longitudinal studies, and population-based studies, as defined by the authors of the individual publications.

Figure 5: Geographical distribution of AF data sources
Global AF data source distribution, with darker shades of colour representing more data sources in each of the presented regions.
AF, atrial fibrillation.
| Dataset Description                                      | Access  | Linkage |
|-----------------------------------------------------------|---------|---------|
| Gulf Acute Heart Failure Registry                         | 21      | Unknown |
| Swedish Heart Failure Registry                            | 27      | Yes     |
| Get With The Guidelines-Heart Failure Registry            | 34      | Unknown |
| Nationwide Inpatient Sample database                      | 34      | Unknown |
| Korean Acute Heart Failure Registry                       | 37      | No      |
| Acute Coronary Syndrome Israel Survey                     | 61      | Unknown |
| Korea Working Group on Myocardial Infarction Registry     | 62      | Unknown |
| GRACE/GRACE 2                                             | 73      | Unknown |
| ACTION-GTWG                                               | 76      | Yes     |
| SWEDEHEART                                                | 100     | Yes     |
| The EURObservational Research Programme                   | 21      | Unknown |
| Atrial-Fibrillation General Pilot registry, Fushimi AF Registry | 34  | Unknown |
| Outcomes Registry for Better Informed Treatment of Atrial Fibrillation | 38  | Unknown |
| Nationwide Swedish Health registries                      | 63      | Yes     |
| Danish nationwide linked admin registries                 | 197     | Yes     |
### Table 1: Recent key real-world data publications in HF, ACS and AF

| Data source                                                                 | Participants                                                                 | Findings                                                                                                                                  | Reference                                                                 |
|----------------------------------------------------------------------------|------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------|
| Gulf CARE (Gulf aCute heArt failuRe rEgistry)                              | 5,005 patients hospitalised with acute HF                                      | ACS was the most common precipitating factor for new-onset HF (39.2%), and non-compliance with medications the most common precipitating factor for decompensated chronic HF (27.8%) | Salam et al. Med Prin Pract 2020;29:270–278                              |
| Swedish Heart Failure Registry                                             | 21,496 patients with HF                                                        | Iron deficiency testing only performed in 27% of patients; 49% of those tested had iron deficiency which was associated with recurrent hospitalisation | Becher et al. Eur J Heart Fail 2021; doi: 10.1002/ejhf.2338                |
| Get With The Guidelines-Heart Failure registry                             | 1551 patients hospitalised for HFrEF and discharged on sacubitril/valsartan and 7857 discharged on ACEi/ARB | Prescription of sacubitril/valsartan was associated with reduced post-discharge mortality and all-cause hospitalisation compared with ACEi/ARB | Greene et al. J Am Heart Assoc 2021;10:e021459                               |
| Korean Registry of Acute Myocardial Infarction for Regional Cardio-cerebrovascular Centers | 11,700 patients with acute MI                                                   | ST-elevation and non-ST-elevation MI occurred in 43% and 57%, with case fatality within 12 months of 10%                               | Kim et al. J Clin Med 2021;10:498                                         |
| EURObservational Research Programme (EORP)-Atrial Fibrillation III Registry | 8,306 patients with AF                                                          | Median age of the registry cohort was 69 years, with patients enrolled across 31 participating countries with future follow-up to assess adherence to guidelines and adverse events | Eur Heart J Qual Care Clin Outcomes 2021;7:229-237                        |

ACEi = angiotensin converting enzyme inhibitor; ACS = acute coronary syndrome; AF = atrial fibrillation; ARB = angiotensin receptor blocker; HF = heart failure; MI = myocardial infarction.