Efficient identification of patients eligible for clinical studies using case-based reasoning on Scottish Health Research Register (SHARE).

**CURRENT STATUS:** UNDER REVIEW

BMC Medical Informatics and Decision Making  ▶ BMC Series

Wen Shi  
University of St Andrews

Tom Kelsey  
The University of St Andrews

✉ twk@st-andrews.ac.uk  
Corresponding Author

ORCiD: https://orcid.org/0000-0002-8091-1458

Frank Sullivan  
University of St Andrews

**DOI:**  
10.21203/rs.2.18160/v2

**SUBJECT AREAS**  
Medical Informatics

**KEYWORDS**  
clinical studies, electronic health record, machine learning, artificial intelligence
Abstract

Background: Trials often struggle to achieve their target sample size with only half doing so. Some researchers have turned to Electronic Health Records (EHRs), seeking a more efficient way of recruitment. The Scottish Health Research Register (SHARE) obtained patients’ consent for their EHRs to be used as a searching base from which researchers can find potential participants. However, due to the fact that EHR data is not complete, sufficient or accurate, a database search strategy may not generate the best case-finding result. The current study aims to evaluate the performance of a case-based reasoning method in identifying participants for population-based clinical studies recruiting through SHARE, and assess the difference between its resultant cohort and the original one deriving from searching EHRs. Methods: A case-based reasoning framework was applied to 119 participants in nine projects using two-fold cross-validation, with records from a further 86,292 individuals used for testing. A prediction score for study participation was derived from the diagnosis, procedure, pharmaceutical prescription, and laboratory test results attributes of each participant. Evaluation was conducted by calculating Area Under the ROC Curve and information retrieval metrics for the ranking list of the test set by prediction score. We compared the most likely participants as identified by searching a database to those ranked highest by our model. Results: The average ROCAUC for nine projects was 81% indicating strong predictive ability for these data. However, the derived ranking lists showed lower predictive performance, with only 21% of the persons ranked within top 50 positions being the same as identified by searching databases. Conclusions: Case-based reasoning is may be more effective than a database search strategy for participant identification for clinical studies using population EHRs. The lower performance of ranking lists derived from case-based reasoning means that patients identified as highly suitable for study participation may still not be recruited. This suggests that further study is needed into improvements in the collection and curation of population EHRs, such as use of free text data to aid reliable identification of people more likely to be recruited to clinical trials.

Background

Paradoxically, Randomised Controlled Trials (RCTs) often fail because they fail to recruit sufficient
study subjects, even when many potential study subjects would agree to participate.[1] They are often not asked to participate because of inefficiencies in recruitment processes. The recent Cochrane review conducted by Treweek et al. of 72 different recruitment strategies identified only three effective interventions with high certainty and another four with moderate certainty.[2] Electronic Health Records have become increasingly important sources for identifying trial participants, and hence reducing one source of inefficiency or been used to conduct pragmatic trials recently.[3-6] A potential application of EHR data analysis is to identify prospective participants for forthcoming RCTs, thereby improving the recruitment and selection process. A 2015 study showed that careful analysis of EHR data could reliably identify individuals eligible to participate in clinical trials.[7] For each of 13 diverse clinical trials - each conducted at Columbia University - diagnostic, pharmacy, laboratory results and case notes from participants were used to derive a profile of an idealised target patient. Candidate participants were then matched in terms of conformity to this profile, based on their EHR data. The reported Area Under the ROC curve of 95% demonstrates the utility of both EHRs and the analytical approach taken. The problem remains of translating this approach to a population-based and multi-centre clinical study setting.

The Scottish Health Research Register (SHARE)[8] allows searching of linked EHR data for people using the Scottish National Health Service (NHS) who have opted in to allow access to their data for this purpose.[9] There are similar developments in other countries.[10] SHARE registrants are therefore population based, and can, in principle, be assessed as potential participants in clinical studies and other ethically approved, methodologically rigorous research. Scotland has 14 regional health boards delivering services to its 5.6M citizens, and RCTs are conducted within and amongst any combination of boards. Figure 1 shows the workflow for the use of SHARE in recruitment of participants to forthcoming studies. Participant selection with SHARE is currently performed by searching the corresponding fields in the EHRs according to the elements contained in the study inclusion/exclusion criteria provided by the study researcher to the staff in the Health Informatics Centre: University of Dundee (HIC).[11] This approach may fail to identify suitable participants due to incomplete and/or insufficient EHR data.
In this paper we investigate whether case-based reasoning, i.e. the use or adaptation of existing solutions may be applied to new problem instances,[7] can generate a good predictive performance in participant identification for multi-centre and population-based studies in the Scottish EHR setting.

Methods
The objective of this study is to: (1) evaluate the predictive performance of case-based reasoning (CBR) in studies conducted in a multi-centre and population-based manner using a range of different EHRs, (2) assess the consistency of participant prediction results between CBR and the database search strategy.

The current study included nine completed projects from SHARE for analysis. The process of inclusion is described in Figure 2. These projects had completed recruitment before the end of 2017. They recruited participants across the health boards in Eastern Scotland. Finally, three projects without recruitment data recorded were also excluded.

The additional table (see additional file 1) summarizes the inclusion/exclusion criteria, number of participants required by the nine projects and the number of potential participants identified by searching database, the number of persons finally recruited in each project and how many of them were identified in the database. The projects involved from 2 to 28 participants, with no participant being recruited into more than one project. A total of 119 people were enrolled in these nine projects.

We chose the individual final recruitment status recorded in SHARE as our gold standard to train and evaluate our algorithm for (i) we can compare our evaluation results with those from the original study because Miotto & Weng also used the final enrolment results as gold standard (ii) it is almost impossible to confirm the real eligibility of each person and use them as gold standard since this is a retrospective study.

The Community Health Index (CHI) is a population register, which is used in Scotland for healthcare purposes.[12] A CHI number uniquely identifies a person on the index. The data from selected projects were de-identified by removing personal data such as names, addresses and date of birth, and linked through CHI number for each participant. Recruitment data were extracted and compiled by HIC, composed of persons identified and recruited for each of the nine projects. Clinical data –
diagnosis, procedure, prescription and laboratory test were requested for all 119 participants and all
other registrants of SHARE in the NHS Scotland Fife and Tayside areas (n = 90,456). Following
assessment of the time frames of the projects to be analysed and the availability and completeness of
the clinical data needed, the requested clinical data range from 2010 through 2017. Diagnoses were
recorded using the International Classification of Diseases revision 10 (ICD-10). Procedures were
recorded using the Office of Population Censuses and Surveys, Classification of Surgical Operations
and Procedures version 4 (OPCS-4). Prescriptions were recorded using unique identifiers mapped to
British National Formulary (BNF) codes[13] to identify the drug prescribed. Prescription data also
included the quantities of each prescription dispensed. Finally, laboratory test data consisted of test
names, read codes allocated to the specific test, and test results. Data were accessed and analysed
through the data safe haven provided by HIC.[14, 15]
Diagnoses and procedures were presented in entity-attribute-value (EAV) model[16, 17] in wide form
in which one record of a certain person contains several diagnoses and procedures, the numbers of
which vary from person to person. For example, patient A’s first record has diagnoses of pneumonia,
diabetes, hypertension. These data were first transformed into a long EAV form with one record
comprising one diagnosis or procedure alone. Prescription data were mapped to formatted BNF codes
and the records with unidentifiable maps were excluded.[18] To ensure that each person included for
analysis had some meaningful clinical data recorded, individuals with no diagnosis or procedure or
prescription or lab test were excluded, as were those subjects having test results but incomplete test
description. As a result, 119 participants as having at least one record in diagnosis, procedure,
prescription and lab test were included for further analysis. 86,292 other registrants were included for
use in test datasets for model assessment and validation purposes, being the subset of the 90,456
SHARE subjects that met the same inclusion criteria as the 119 participants.
Each project was analysed separately by two-fold cross-validation. For both folds, half of the 119
participants were used to train a predictive model. The remaining half were combined with (i) 30,000
SHARE registrants chosen randomly from the 86,292 available and (ii) all subjects identified though
inclusion/exclusion database queries for the project but not selected as one of our 119 cases (people
enrolled in studies). This test dataset was used to assess prediction errors for subjects not used to derive the model. Following the methodology of Miotto and Weng, 2015,[7] a target profile was derived from central tendencies of the occurrences of the codes from diagnosis, procedure, prescription and laboratory test. The featured codes were determined according to their frequencies among that project’s participants, with each entity (being either a diagnosis, a procedure, a prescription or a laboratory test) requiring a minimum of 10 codes shared by at least 80% of the participants. If more than 10 codes were retrieved in one entity, all the codes exceeding 80% threshold were included. If the data were too sparse for any code to reach 80%, all the codes were retained. After the target profile had been produced, every individual produced a similarity score for each of the four entities respectively by being compared against the target using cosine similarity, which uses orientation of entities when plotted in multi-dimensional space to assess similarity rather than magnitude of difference apart in the same space.[19] Thus, each person was represented by four similarity scores ranging from 0 to 1. The training set was used to train a linear regression model to get four optimal weights for each entity, which were then applied to each individual in the test set to obtain a final score which was then scaled to fall between 0 and 1.

For each project, the area under the Receiver-Observer Characteristic curve (ROCAUC) was calculated for either testing fold respectively and average scores were obtained for projects individually and for the nine projects as a whole. ROCAUC measures the overall performance of a binary classifier.[20] It estimates the probability of obtaining a higher score for a participant than for a non-participant. The higher the ROCAUC, the better the classifier. For this study ROCAUC are interpreted using a standard quality scheme as follows: 90 -100 = excellent; 80 - 90 = good; 70 - 80 = fair; 60 - 70 = poor; 50 - 60 = fail.[21] The confidence interval (CI) for ROCAUC was generated through 2000 bootstrap replicates.[22] Prediction results were also combined to identify the cut-off maximizing the performance metric that sums up sensitivity and specificity.

A ranking list was generated for each test set, with individual final scores decreasing for assessment with regard to their ability to find the person recruited using metrics such as precision of predicting the 5 top items (P5)[23], precision of predicting the 10 top items (P10)[23], the mean average
precision (MAP)[24] and mean reciprocal rank (MRR)[25]. Average precision combines the precisions obtained every time a relevant result is retrieved, thereby assessing the quality of the whole list.

Reciprocal rank is the reciprocal of the rank of the first targeted person retrieved, reflecting the utility of the list to meet the need of the list users when identifying suitable participants. The best possible result for the list with participants ranked at the top (Upper) and the worst result - a random list (Lower) were also obtained as references. The metrics were averaged across all 18 (i.e. 2 fold for each of 9 projects) ranking lists respectively.

We additionally examined how many of the persons identified by database queries were among the top 50 of the ranking list in proportion.

Results

The ROCAUC for each project and each fold are shown in Table 1. The mean ROCAUC for prediction score was 0.815 over nine projects. One project had unacceptably low ROCAUC for prediction score (0.337) being worse than a random guess, with another project giving poor discriminatory performance (ROCAUC 0.619) The remaining eight projects were either good or excellent in terms of ROCAUC.

Table 1. The Area Under the ROC Curve for recruitment prediction test for each project. For each project the ROCAUC and 95% confidence intervals are given for the first and second cross validation datasets, followed by the average.

| Project Acronym | Fold 1 (CI)          | Fold 2 (CI)          | Average (CI)         |
|-----------------|----------------------|----------------------|----------------------|
| ALPHA           | 0.947 (0.897 - 0.984)| 0.855 (0.777 - 0.928)| 0.901 (0.837 - 0.956)|
| ALLAY           | 0.928 (0.903 - 0.955 )| 0.854(0.768 – 0.941)| 0.891 (0.836 – 0.948)|
| METFORMIN       | 0.977 (0.970 – 0.989)| 0.980 (0.970 – 0.983)| 0.979 (0.970 – 0.986)|
| REFORM          | 0.946 (0.896 – 0.980)| 0.957 (0.920 – 0.987)| 0.952 (0.908 – 0.984)|
| IMPOCT          | 0.397 (NA)           | 0.276 (NA)           | 0.337 (NA)           |
| TARDIS          | 0.897 (0.844 – 0.946)| 0.887 (0.852 – 0.919)| 0.892 (0.848 – 0.933)|
| 4P              | 0.952 (0.909 – 0.993)| 0.744 (0.447 – 0.978)| 0.848 (0.678 – 0.986)|
| HF              | 0.907 (0.851 – 0.957)| 0.927 (0.870 – 0.974)| 0.917 (0.861 – 0.966)|
| ImmunoStat      | 0.632 (0.503 – 0.763)| 0.605 (0.443 – 0.756)| 0.619 (0.473 – 0.760)|

NA: insufficient data to calculate an accurate CI

A threshold of 0.44 gives the maximum performance metric for these data. A plot of performance against different prediction score cut-offs is shown in Figure 3; the scaled sum of sensitivity and
Specificity was calculated for precision score thresholds from zero to 100%, with the optimal cut-off for these data occurring at 44%.

Ranking list results are shown in Table 2. Overall, the ranking list failed to present the relevant participants at neither the top 5 positions nor the top 10. Performance improved when trying to find participants from the top of the list of roughly 30,000 patients compared to a random list, but was still far from optimal.

Table 2. Performance of the resulted ranking lists. For each project we give the number of participants appearing in the top 5 and top 10 of a ranking list, the mean average precision (MAP) and the mean reciprocal rank (MRR).

| List       | P5 | P10 | MAP  | MRR  |
|------------|----|-----|------|------|
| ALPHA      | 0  | 0   | 0.003| 0.006|
| ALLAY      | 0  | 0   | 0.001| 0.001|
| METFORMIN  | 0  | 0   | 0.003| 0.002|
| REFORM     | 0  | 0   | 0.010| 0.035|
| IMPOCT     | 0  | 0   | 0.000| 0.000|
| TARDIS     | 0  | 0   | 0.002| 0.005|
| 4P         | 0  | 0   | 0.002| 0.003|
| HF         | 0  | 0   | 0.006| 0.028|
| ImmunoStat | 0  | 0   | 0.001| 0.007|
| Average of all ranking lists | 0  | 0   | 0.003| 0.010|
| Averaged Lower | 0  | 0   | 0.000| 0.000|
| Averaged Upper | 0.789 | 0.611| 1    | 1    |

On average, 21% of the persons ranked within the top 50 positions on the ranking list were the same as identified by searching databases. The proportions of those both identified listed in the top 50 of the list for each project are presented in Table 3.
Table 3. Proportion of persons both identified within the top 50 positions of the ranking list for each project

| Project acronym | Fold 1 | Fold 2 | Average |
|-----------------|--------|--------|---------|
| ALPHA           | 0.66   | 0.32   | 0.49    |
| ALLAY           | 0.54   | 0.54   | 0.54    |
| METFORMIN       | 0      | 0      | 0.00    |
| REFORM          | 0.32   | 0.16   | 0.24    |
| IMPOCT          | 0      | 0      | 0.00    |
| TARDIS          | 0.22   | 0      | 0.11    |
| 4P              | 0.06   | 0.14   | 0.10    |
| HF              | 0.30   | 0.42   | 0.36    |
| ImmunoStat      | 0.04   | 0.08   | 0.06    |

Discussion

Our ROCAUC results indicate that we can reliably and retrospectively predict the recruitment status for seven of the nine projects analysed, with IMPOCT and ImmunoStat being harder to predict in terms of participation. We speculate that, for IMPOCT study, the poor result was caused by the extremely small number of participants (2) recruited, so that only one person was used to derive the target profile of participants. Considering the relatively loose inclusion/exclusion criteria for ImmunoStat, the less satisfactory performance may result from an inherent inability of capturing a representative target due to scattered participant characteristics. For the higher-quality projects, ROCAUC is either good (i.e. above 80%) or excellent (i.e. above 90%), providing strong support for our central claim that careful analysis of EHR data can reliably discriminate participation status for projects of sufficiently high quality.

Evaluation of the resultant ranking lists showed poor performance in retrieving the real participants at the top. None of the ranking lists identified participants at either the first five individuals retrieved or the first ten. The ranking performance was the worst for IMPOCT project, consistent with the AUC result. Performance improves slightly when identifying persons recruited and selecting the first candidate for participation, but is still unsatisfactory in terms of matching predicted to actual participants.

The assessment of the consistency of patients identified by case-based reasoning and database query shows there is significant discrepancy among the potential participants detected by the two methods.
However, it is unknown whether the most likely participants identified from our ranking lists would have been recruited or not due to the retrospective nature of our study, so there is an inherent lack of discriminative ability involved. Also the list of eligible participants derived from searching the database is definitely not exhaustive. As shown in the project summary table (see additional file 1), a few of the participants were not among the persons identified in database at all. Therefore, the non-participants picked up by the ranking list who were not identified before might have been actual participants.

Our study has the following strengths: we have used recruitment data from real clinical studies involving multiple health boards and multiple data sources; the EHR data is population based and hence is not flawed by selection or geographic bias; we have used an analogue of an existing proven methodology augmented with cross validation and relative ranking analysis; we have compared the potential participants between two detecting strategies.

Our study also has several limitations. We acknowledge that the final recruitment results in SHARE are not a perfect gold standard because they are affected by some external factors rather than clinical information alone such as personal preferences. The algorithm might have correctly identified an eligible participant but this person had not been enrolled in the study due to personal reasons. But our evaluation results would only be underestimated in this case. Whilst acknowledging this inherent lack of knowledge as a limitation of the study, we believe that we have demonstrated results in agreement with Miotto & Weng’s key finding that by matching EHR data of unseen patients to a target derived by case-based reasoning, we can estimate relevance to a forthcoming trial, with higher relevance being associated with higher eligibility. Additionally, the clinical data used for this study don’t consist of free-text clinical notes, which might have led to the poor performance; another limitation is that despite careful check of final recruitment status for each project, the recruitment data may contain some misclassifications.

Conclusions
In this study we have demonstrated that case-based reasoning performed well in predicting recruitment for population-based clinical studies using Scottish EHR data. Our results were
substantially worse than those reported by Miotto & Weng[7] with regard to ranking lists derived from
the similarity scores from case-based reasoning analysis. The provision of more accurate recruitment
data and more abundant clinical data resources (including narrative clinical notes) would likely
improve our ability to derive accurate ranking lists, with no negative impact on clinical care for the
population which is the primary purpose of EHRs. Planned prospective studies comparing the actual
recruitment results for persons identified by differed methods should help address the question of
which one doing a better job in finding potential participants. The analytical framework could usefully
enhance the existing database query method in Scotland, and form the basis of participant prediction
schemes in other healthcare settings.

List Of Abbreviations

BNF: British National Formulary

CBR: case-based reasoning

CHI: Community Health Index

CI: confidence interval

EAV: entity-attribute-value

EHRs: Electronic Health Records

HIC: Health Informatics Centre: University of Dundee

ICD-10: International Classification of Diseases revision 10

MAP: mean average precision

MRR: mean reciprocal rank

NHS: National Health Service

OPCS-4: Office of Population Censuses and Surveys, Classification of Surgical Operations and
Procedures version 4

P5: the 5 top items

P10: the 10 top items

RCTs: Randomised Controlled Trials

ROCAUC: Area Under the ROC Curve
SHARE: Scottish Health Research Register

Declarations

**Ethics Approval and Consent to participate**

The study obtained approval from the local health boards (NHS Fife and NHS Tayside) to use patients’ de-identified electronic health records stored in HIC. HIC was given favourable ethics opinion regarding provision of anonymous data for research. (REC ref 14/ES/0015)

**Consent for publication**

Not applicable.

**Availability of data and materials**

The study only has online access to the data stored in safe haven. Requests for data access should be directed to the Health Informatics Centre: University of Dundee.

**Competing Interests**

None for W.S. and T.K. F.S. is a clinical director of SHARE but he derives no financial benefit from this role.

**Funding**

W.S. is a PhD student funded by the University of St Andrews where T.K. and F.S. are faculty members.

**Author Contributions**

W.S. conducted the research and is the major contributor to the manuscript. T.K. supervised the study and also contributed to the manuscript. F.S supervised the study, assisted data collection and edited the manuscript. All authors read and approved the final manuscript.

**Acknowledgments**

We acknowledge the assistance from colleagues in SHARE and HIC for the process of data collecting (Shobna Vasishta, Louise Dow, Christopher Hall). We also thank Miotto and Weng for corresponding and sharing information on their work. We are thankful for Brian McKinstry critically reviewing the paper.

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Additional File Legend
Additional file 1[Project summary.doc]: Summary of nine projects included for analysis.

Figures
Work flow for recruiting participants to clinical studies through SHARE.[11]
Figure 2

Flow diagram for the selection of projects for analysis.
Figure 3

Scaled sensitivity plus specificity against prediction score cut-offs.