

## Research Article

# Design and Development of a Hospital-Based Coronary Artery Disease (CAD) Registry in Iran

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**Background.** The incidence of coronary artery disease (CAD), the leading cause of mortality in most developed and developing countries, is increasing. The adoption of hospital registries can improve care delivery and facilitate the management of CAD through better planning, as well as help with outcome assessment through more effective data management. **Objectives.** The present study is aimed at designing a hospital-based CAD registry for managing CAD data. **Methods.** This developmental study was conducted in three phases. Initially, sources related to CAD registries were reviewed, the results of which were published in two studies. In the next phase, the prerequisites and requisites of the software were determined through a qualitative study. In this phase, the registry dataset was determined by using a questionnaire. Finally, the developed conceptual model of the software was validated. The software was then developed based on the validated conceptual model. **Results.** The registry data elements were classified into 13 main categories, including identification data, medical history, and risk factors. The dataset included 171 data elements, including data related to surgical and nonsurgical procedures. The conceptual model was approved by field experts, and the software was developed accordingly. **Conclusion.** The steps followed in the present study for developing the CAD registry can be used as an appropriate approach for designing similar hospital-based registries. Considering the pivotal role of the registry in the management of CAD, the routine and systemic use of the registry is suggested in all healthcare centers.

## 1. Introduction

Cardiovascular diseases are a group of disorders affecting the heart or blood vessels. They are classified into different types, including coronary artery disease (CAD), heart attacks, and rheumatic heart disease, according to the affected site, conditions, and contributing factors [1, 2]. One of the most important types of cardiovascular disease is CAD [3] scientifically known as coronary heart disease, ischemic heart disease, or atherosclerosis [4]. CAD or ischemic heart disease is caused by the narrowing or spasm of coronary arteries, which are the main vessels supplying blood to the heart, resulting in the reduction of blood and oxygen supply to the heart muscles; consequently, it exposes the individual to heart attacks and other complications [2, 5].

As a chronic disease, CAD has been introduced as the leading cause of mortality worldwide, especially in developed countries; today, it still accounts for the highest number of deaths globally [2, 6, 7].

The report of the World Health Organization in 2020 shows that cardiovascular diseases, as one of the noncommunicable diseases, are the most important cause of death across the world [8]. In 2015, about 17.7 million deaths were reported worldwide due to cardiovascular diseases, and the highest death rate was assigned to coronary artery disease [9]. According to the Center for Disease Control and Prevention (CDC) in the United States, 610,000 people die each year due to cardiovascular disease [10]. In Iran, as a developing country with a low-income level, 26.4% of recorded deaths were due to cardiovascular diseases [8, 11].

The need for increased attention and planning to deal with the complications of CAD is strongly felt, especially in developing countries. Although the World Health Organization (WHO) reported that more than 80% of countries in the world have the necessary resources and facilities to deal with noncommunicable diseases, including cardiovascular diseases [12, 13], they (especially cardiovascular diseases) remain the leading cause of death in different communities. Therefore, it is necessary to apply all the available facilities and potentials to manage this disease, including CAD data management systems, to effectively manage complications and consequences through proper planning [14, 15].

The management of disease-related data can be regarded as a basis for achieving various healthcare goals and implementing relevant programs [16, 17]. Registries are one of the most important patient data management tools that play a critical role in managing diseases and conducting relevant research [17]. Disease registries present a continuous and systematic collection of information about individuals with specific diseases or health conditions in a given population [16–18]. By applying clinical guidelines and data standards, these registries can reduce the costs of care and improve care delivery [19–22].

Owing to the great impact of CAD on communities and the special attention of authorities in developed countries to the management and control of this disease, different types of CAD registries have been developed in different countries [23, 24]. For example, the Swedish Coronary Angiography and Angioplasty Registry (SCAAR) was developed in 1999 in Sweden by merging the Swedish Coronary Angiography Registry (Acta Coronaria) with the Swedish Registry for Coronary Angioplasty (SCAP) as part of the Swedish Heart Registry. This registry has been in place in a web-based format since 2001, and the data are collected online from 30 hospitals [25–27].

Considering the advantages of registry systems as data management tools, such as prevention of financial loss, data documentation, improvement of healthcare planning, delivery of high-quality information to researchers, improvement of healthcare quality, and assessing the outcomes of care [28–30], there is a need for CAD registry design and deployment. Therefore, the present study is aimed at designing a hospital-based CAD registry software in Iran.

## 2. Methods

This applied that the developmental study was conducted in three phases.

**2.1. Phase 1.** This step was conducted in two parts: conducting a systematic review of studies focusing on the key processes of registry software systems and performing a comparative study of the selected registries using a data management approach, the results of which have been published earlier [31, 32].

**2.2. Phase 2.** Initially, for software requirement elicitation, a qualitative study was conducted by interviewing 15 potential

users, including the software administrator, data processing expert, cardiologist, and cardiac nurse. In the next stage, the minimum dataset for the hospital-based CAD registry was created by reviewing the patient's records, and the software processing indicators were presented in the form of a questionnaire, and the opinions of cardiologists ( $n = 6$ ), cardiac nurses ( $n = 9$ ), and health information management specialists (HIM) ( $n = 6$ ) were sought.

For this aim, this part of the study was designed and implemented as a Delphi study. Data collection was performed using a questionnaire containing the minimum dataset for the CAD hospital registry. In the first round of Delphi, the questionnaire developed on a five-point Likert scale was given to the participants, and they were asked to state their level of agreement with each data item in the software. In this round, at the end of each part of the questionnaire, the participants were provided with an open question that allowed them to express their views if the closed questions did not address these appropriately, and these were added to the questionnaire for the next round. In this stage, there were 166 data items for the minimum dataset, and 20 indicators were set for the reporting function of the software. In this round of the Delphi study, two data items were removed, seven data items were added, and five items were passed to the second round of study. In the first round, all the indicators in the questionnaire were approved by participants. In the second round, the questionnaire contained 12 data items, and it was passed to participants involved in the first stage. Eleven out of 12 items were approved by participants.

The inclusion criteria in this stage were willingness to participate in the study and having at least five years of relevant work experience. Data analysis was performed in SPSS 22 by measuring descriptive statistics, including mode, median, mean, interquartile range, and percentage of agreement.

**2.3. Phase 3.** In this stage, the operational, structural, and behavioral models of the software were designed using Enterprise Architect version 9, based on the features and requirements of CAD hospital registries. Seven experts in the fields of health information management, medical informatics, and software engineering, with at least five years of work experience, participated in this study to determine the validity of the presented conceptual model of the software. Data analysis was performed in SPSS 22 by measuring descriptive statistics. Diagrams with scores above 75% (12 out of 16) were approved, while diagrams with scores below 75% were reviewed. After approving the conceptual model, the CAD hospital registry software was designed.

MySQL was also used for developing the software database, and Visual Asp.Net 2013 programming language was applied for software development using C#, JQuery, CSS, Ajax, and HTML5 languages. Error detection and debugging were carried out after coding and developing the initial version in which the syntax errors were checked and fixed. The semantic errors of the codes were then checked and eliminated. After fixing the syntactical and semantic errors, the logical errors of the software were also checked to make sure about the functionality of the developed software.

### 3. Results

**3.1. Phase One.** The results of the first phase of this study were published in our two previous articles, as indicated earlier [31, 32]. Dealt with the registry data items (13 main categories), registry users, key registry indicators, key organizations and individuals using registries, status, goals of patient follow-up, and control of registry data quality were reported.

**3.2. Phase Two.** In the second phase conducted to identify the software requirements, 15 cardiologists, cardiac nurses, and data processing experts participated in a qualitative study. Table 1 shows the demographic information of participants in the stage of the study. At this stage, 60% of the study sample was female ( $n = 9$ ), and the participation of health information management specialists in the interviews was more than the other groups (33.33%).

Table 2 presents the operational requirements of the software, based on the key registry processes divided into six main categories.

**3.2.1. Determination of the Minimum Dataset and Indicators of the Hospital-Based CAD Registry.** To determine the minimum dataset, besides conducting a systematic study and examining the patients' files, the data resulting from the Delphi technique were presented to experts. The first Delphi round was conducted to determine the minimum dataset for the CAD registry software. Accordingly, the minimum dataset of the CAD registry was divided into 13 main categories (Table 3). In the supplementary file (available here) of the article, all data items have been presented.

In the first phase of the Delphi, factors, including the number of children, patient referral to the hospital (e.g., admission from another hospital, outpatient admission, emergency admission by ambulance, and admission based on code 247), obesity status based on the body mass index (BMI), type of narcotic drug (morphine and pethidine), the onset of symptoms before the first emergency call, onset of symptoms until referral to the center, and angiography indication class, were suggested by participants to be included into the questionnaire for the second round.

In the second round of the Delphi, except for the results of follow-ups and outcomes after discharge, other items were approved. Overall, 171 information fields were finalized in two Delphi rounds for the minimum dataset of the hospital-based CAD registry software. Moreover, the software reporting indicators were agreed upon by experts in the first round of Delphi, as presented in Table 4.

**3.3. Phase Three.** The conceptual model included operational, structural, and behavioral parts, as discussed in the following section.

Based on our investigations, the system users included the software administrator, cardiologist, cardiac nurse, and data processing experts. To design the operational model, relevant scenarios were first prepared, such as the creation of a user account, where the software administrator defines a new user account and determines the user's role, allowing the user to login by entering their login details. Other

TABLE 1: Characteristics of participants in the qualitative phase of the study.

Demographic information		Frequency	%
Sex	Women	9	60
	Men	6	40
Type of specialty	Physician	4	26.67
	Nurse	4	26.67
	HIT expert	2	13.33
	HIM specialists	5	33.33
Academic level	MSc	4	26.67
	Ph.D.	7	46.66
	MD	4	26.67

scenarios were also prepared based on the main services of the software, including new case registration, abstracting, and reporting scenarios.

After developing the operational model, the structural model of the software was designed covering the general class diagram of the software.

In general, various diagrams are used to demonstrate a software behavioral model. The sequence diagram presents a comprehensive description of the main applications of the software and facilitates software development. A sequence diagram was created and validated based on the main applications of the software, including case finding, data collection, abstracting, data quality control, follow-up, and reporting.

Table 5 illustrates the validity results of the CAD registry conceptual model.

According to the table above, a score above 75% approved each diagram; all relevant diagrams were confirmed by experts. Figure 1 demonstrates the first page of the software.

Figure 2 shows the user interface and user access level of the software.

The users could be able to select any type of report from reporting function of the registry and obtain their needed reports.

### 4. Discussion

A key step in developing information systems is the identification of operational and nonoperational requirements (software prerequisites) through users' need assessment [33, 34] [27, 28]. The practical principles of software design and assessment (i.e., system development life cycle, including software planning, analysis, design, and implementation) were applied in the present study. The operational requirements of the hospital-based CAD registry software were determined based on the key registry processes, which included case finding, data collection, abstracting, data quality control, patient follow-up, and registry software reporting.

The findings showed that the hospital-based CAD registry software should be capable of registering newly identified cases of CAD based on the existing information facilities and

TABLE 2: The operational requirements of the CAD registry software.

Operational requirements	Description	Sources
Case finding	The registry software should allow the registration of newly identified cases.	Interviews 7, 8, and 13
Data collection	The software should be capable of requesting information about the case they found.	Interviews 3, 10, 12, and 14
Abstracting	The software should allow the recording of a summary and abstract of data.	Interviews 2, 7, and 9
Data quality control	The software should be capable of performing data quality control and identifying high-quality from low-quality abstracts.	Interviews 4, 5, and 11
Follow-up	The software should be capable of providing a list of patients who require regular follow-up.	Interviews 9, 10, and 15
Reporting capacity	The software should be capable of creating different reports, and if needed, sharing these reports with qualified care centers and individuals.	Interviews 11 and 14

TABLE 3: The results of the first Delphi round for the minimum dataset of the CAD registry software.

Data elements		Mean	No. of items	No. of accepted items
Demographic information		4.11	15	14
Insurance information		3.95	4	4
Hospitalization information		4.00	5	4
Medical history and risk factors	History of other diseases	4.25	18	18
	Lifestyle	4.55	5	5
Medications		4.51	14	14
Laboratory test results		4.80	23	23
Physician examination results	Present symptoms	4.30	10	10
	CAD type detection	4.90	1	1
Information on angiography		4.57	7	7
Noninvasive procedures	CT angiography	4.61	2	2
	Other	4.29	3	3
	General	4.26	5	5
Information on invasive and surgical procedures	Determination of the type of PCI procedure and related information	4.52	8	8
	Outcomes	4.27	13	13
Patient's status during discharge		4.01	5	4
	Follow-up results and outcomes after discharge, up to six months	4.43	9	8
Patient follow-up	Follow-up of outcomes due to coronary artery disease	4.41	6	6
	Follow-up of patient's adherence to treatment	3.79	5	4
	Follow-up of patient's quality of life	3.98	3	3
	Follow-up of death status	3.69	5	3

infrastructure. Effective registries are required to include all new cases of the target disease [31, 35]. In well-developed registries, new case finding techniques, such as text mining, are applied; however, if these techniques are not addressed automatically, data processing experts should actively check for relevant sources and find new cases in the registry. Considering the current status of health information systems in Iran, the evaluation of hospital information systems and case finding step is necessary, as the goal of case finding in registries (from the experts' perspectives in this study) is to avoid errors and help the users find all relevant cases.

From the perspective of experts in this study, the CAD registry software should be capable of requesting case finding information and recording a summary of data from the patient's records. Moreover, the software should be capable of storing data temporarily in a temporary file if the information of a new case is unavailable. In a previous study, Tabrizi et al. also emphasized the importance of creating a temporary file [36]. Generally, the most important feature of a CAD registry system is data abstraction, documentation of all necessary information, and prevention of recording irrelevant or unimportant data [36–38].

TABLE 4: Reporting indicators required in the software.

Rows	Indicators	Mode	Median	Mean	Interquartile range		Percentage of agreement	Results
					Q1	Q3		
1	Number of patients with coronary artery disease (CAD)	3	4	3.81	3	5	76	✓
2	Number of patients by sex	4	4	4.33	4	5	86.6	✓
3	Physician's performance	5	5	4.57	4	5	91.4	✓
4	Mean age of patients (total age of all patients/total patients)	5	5	4.76	4.5	5	95.2	✓
5	Mean length of hospital stay of patients (number of hospitalization days of all patients/total patients)	5	5	4.57	4	5	91.4	✓
6	The ratio of patients diagnosed with STEMI to patients diagnosed with non-STEMI (number of patients with STEMI/ number of non-STEMI patients)	5	5	4.42	4	5	88.4	✓
7	Death rate	5	5	4.57	4	5	91.4	✓
8	Patients with a history of cardiovascular surgeries	5	5	4.71	4	5	94.2	✓
9	Comparison of the status of patients during discharge, including the number of recovered patients, number of nonrecovered patients, the ratio of recovered to nonrecovered patients, and the number of patients discharged at their discretion	5	5	4.57	4	5	91.4	✓
10	Death rate (number of patients who expired/total number of patients in the specified setting)	5	5	4.43	4	5	88.6	✓
11	Total number of PCIs performed	4	4	4.33	4	5	86.6	✓
12	Number of follow-ups	5	5	4.57	4	5	91.4	✓
13	Comparison of echocardiography indicators	5	5	4.57	4	5	91.4	✓
14	Mean time from the onset of symptoms until referral to the hospital	5	5	4.52	4	5	90.4	✓
15	Frequency and ratio of patient referral to the hospital	5	5	4.38	3.5	5	87.6	✓
16	Comparison of involved vessels (e.g., calculation of the frequency and ratio of involved vessels)	4	4	4.09	3	5	81.8	✓
17	Type of angiography performed	4	4	3.82	2.5	5	76.4	✓
18	Calculation of the degree of vessel occlusion	5	5	4	3	5	80	✓
19	Calculation and comparison of the indication class	5	4	3.95	3	5	78	✓
20	The rate of use of each angiography method	3	4	3.81	3	5	76	✓

Moreover, the results of the present study showed that data quality control should be a key operational function in the software. Data quality control ensures that the system

is capable of checking data quality attributes [39] and could support users in this matter. Therefore, a hospital-based CAD registry should support data quality control and

TABLE 5: Validation results of the CAD registry conceptual model.

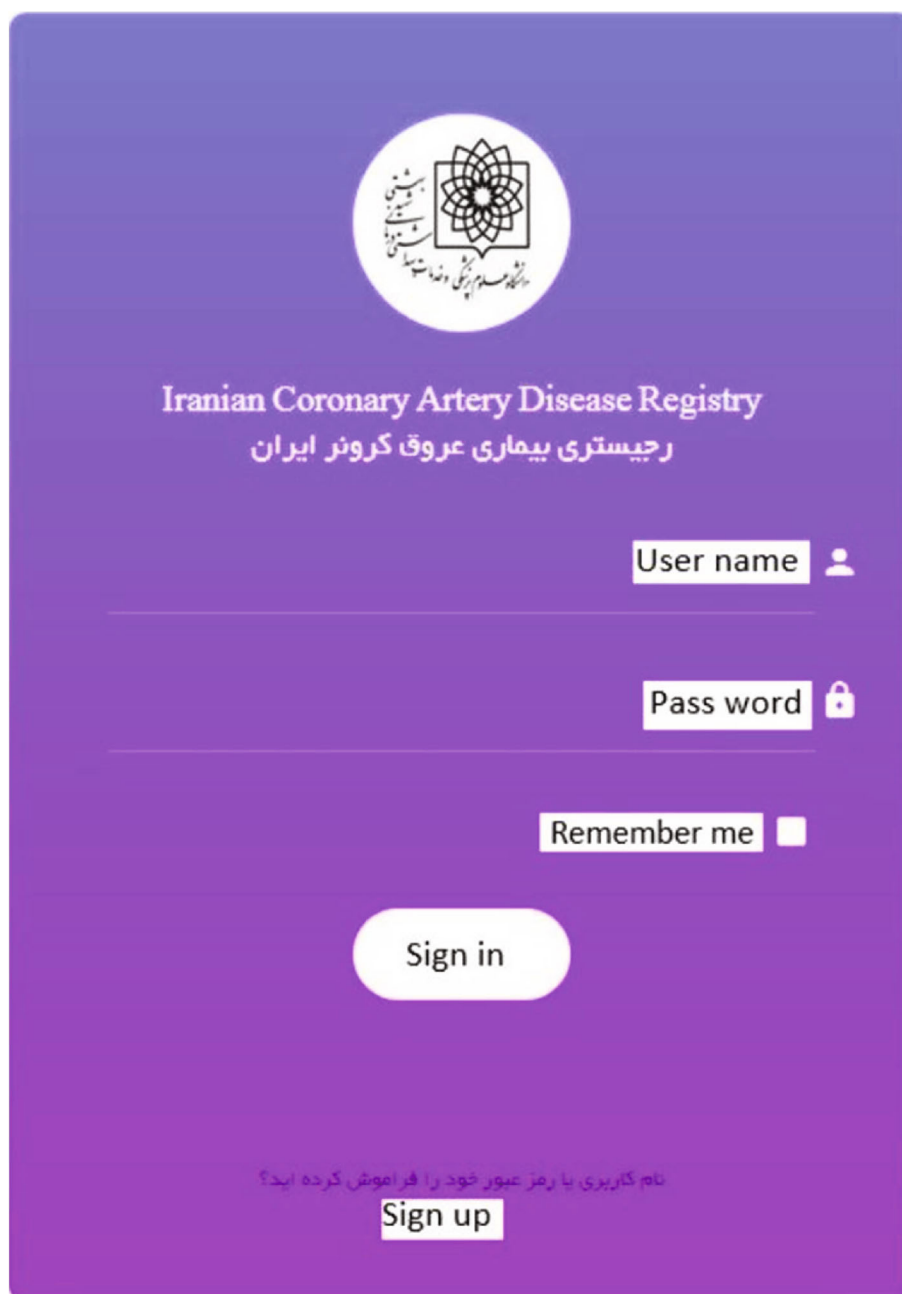
Diagram titles		Acceptable		Somewhat acceptable		Unacceptable		Score	Percentage	Diagram acceptability results
		Frequency	Percentage	Frequency	Percentage	Frequency	Percentage			
Diagram of the software	Log in	8	100	0	0	0	0	16	100	✓
	User account creation	8	100	0	0	0	0	16	100	✓
	Case finding	8	100	0	0	0	0	16	100	✓
	Data collection	7	87.5	1	12.5	0	100	15	93.75	✓
	Abstracting	7	87.5	1	12.5	0	100	15	93.75	✓
	Data quality control	8	100	0	0	0	0	16	100	✓
	Reporting	7	87.5	1	12.5	0	100	15	93.75	✓
	Patient follow-up	8	100	0	0	0	0	16	100	✓
	Log out	8	100	0	0	0	0	16	100	✓
Software activity diagram	Log in	8	100	0	0	0	0	16	100	✓
	User account creation	8	100	0	0	0	0	16	100	✓
	Case finding	8	100	0	0	0	0	16	100	✓
	Data collection	7	100	0	0	0	0	16	100	✓
	Abstracting	6	75	1	12.5	1	12.5	13	81.25	✓
	Data quality control	8	100	0	0	0	0	16	100	✓
	Reporting	7	87.5	1	0	1	12.5	14	87.5	✓
	Patient follow-up	8	100	0	0	0	0	16	100	✓
	Log out	8	100	0	0	0	0	16	100	✓
Software class diagram		6	75	2	25	0	0	14	87.5	✓
Software sequence diagram	Case finding	8	100	0	0	0	0	16	100	✓
	Data collection	7	87.5	0	0	1	12.5	14	87.5	✓
	Abstracting	6	87.5	1	12.5	1	12.5	13	81.25	✓
	Data quality control	8	100	0	0	0	0	16	100	✓
	Reporting	7	87.5	1	12.5	0	0	14	87.5	✓
	Patient follow-up	8	100	0	0	0	0	16	100	✓


indicate high-quality from low-quality abstracts. Another operational capacity of our CAD registry software was patient follow-up. From the perspective of experts, the software should be capable of providing a list of patients who require follow-up regularly. Although there are more advanced methods, such as sending SMS and email to

patients for follow-ups [40, 41], phone calls can help to make sure that patients are updated appropriately about their follow-ups.


Other results of the present study showed that the registry software should be capable of preparing different reports that should be sent to relevant bodies. According to the








Iranian Coronary Artery Disease Registry  
رجیستری بیماری عروق کرونر ایران

User name 

Pass word 

Remember me ☐

Sign in

نام کاربری یا رمز عبور خود را فراموش کرده اید؟

Sign up

FIGURE 1: The first page of the software.

findings, the software should be allowed to calculate important indicators of the disease and provide reports in formats, such as PDF and Excel files needed by users. Overall, determining the minimum dataset for health information systems, especially disease registries, could help to make sure that required data are collected and prevents unnecessary data collection [42, 43]. This stage is a key step in developing disease registries [44, 45]; therefore, in this study, the minimum dataset of the CAD registry was developed, in which data items fell into 13 main categories, including demographic information, insurance information, hospitalization information, medical history, risk factors, medications and laboratory test results, physician examination results, angi-

ography findings, data related to noninvasive procedures or invasive and surgical procedures, patient's status during discharge, and patient follow-up.

In the current study we use of object-oriented approach for designing the conceptual model of the software. Scenarios were initially created, and the conceptual model was developed and composed of operational, structural, and behavioral component parts. The object-oriented approach provides a comprehensive view of the design and development of information systems and tries to model the whole system considering different parts of the system in line with expected functionality. This approach could help developers to obtain a better understanding of users' needs [36]; in

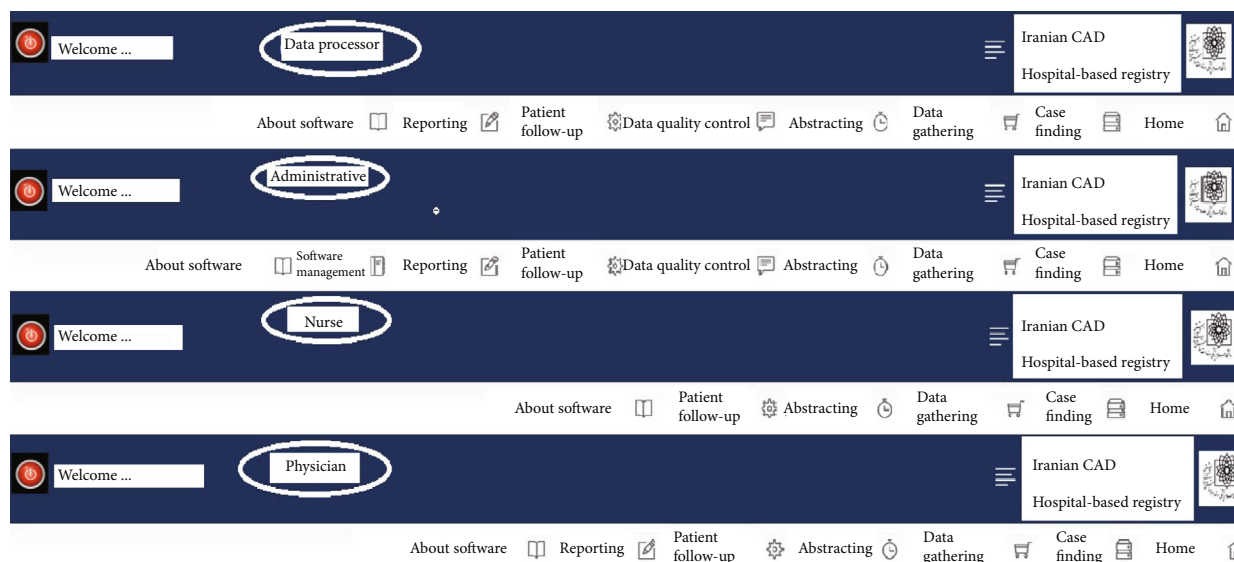


FIGURE 2: User interface and user access level.

other words, the object-oriented approach is a software programming process that takes advantage of object interactions to solve a problem [46].

In the present study, a standard integrated language was used to develop the conceptual model. Designs based on the object-orientated concepts in the Unified Modeling Language (UML) make them fully compatible with object-oriented programming environments and languages (e.g., C++, Java, and C#), and UML has become the common language of many developers of information systems [46, 47]. Besides, UML has been widely used in the development of disease registries [47]. In a previous study, Nouri developed a registry based on a study focusing on hemoglobinopathy using UML [48]. Therefore, it is suggested to use this standard and international language in developing conceptual disease registry models.

A key limitation of the current study could be developed in the CAD registry at a hospital-based level, while a population-based registry could provide a more robust basis for controlling the CAD in a given population. However, the hospital-based CAD registry developed in the current study could be a basis for developing a population-based registry. In addition, the duration of the pilot registry was 6 months, and a longer period is required to depict a more realistic picture of the developed registry and figure out its capabilities and limitations over time.

## 5. Conclusion

Based on the results of the present study, paying attention to data collection and abstracting is critical in developing disease registry software. With respect to patient follow-ups, although patients could be informed about their attendance using SMS and emails, phone calls for the follow-up of patients with CAD seem to be a more appropriate option, because, in the event of a patient's death, the data processing expert can obtain further information, such as the

place of death, cause of death, and date of death through a phone call.

## Data Availability

All data of the study were reported. The link of the registry is <https://cad.hisapps.ir/User/Login?ReturnUrl=%2f>.

## Ethical Approval

This study was approved by the Ethics Committee of Shahid Beheshti University of Medical Sciences, Iran (IR.SBMU.RETECH.REC.1399.1219).

## Conflicts of Interest

The authors have no conflict of interest to declare.

## Authors' Contributions

AG, RR, and HE conceived the idea of this study. AG wrote the proposal and participated in the data collection and statistical analysis. AG was responsible for software implementation and evaluation. AG, RR, and HE interpreted the result. AG and RR participated in paper reviewing and preparing the manuscript. All authors read and approved the final manuscript. RR and HE were responsible for the project administration.

## Acknowledgments

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## Supplementary Materials

Supplemental material for this article is available online. Supplementary material file 1: all data items of the CAD registry in the first round of Delphi. (*Supplementary Materials*)

## References

- [1] H. Ayatollahi, N. Mirani, and H. Haghani, "Electronic health records: what are the most important barriers?. Perspectives in health information management," *Perspectives in health information management*, vol. 11, 2014.
- [2] A. N. Nowbar, M. Gitto, J. P. Howard, D. P. Francis, and R. Al-Lamee, "Mortality from ischemic heart disease," *Circulation: Cardiovascular Quality and Outcomes*, vol. 12, no. 6, article e005375, 2019.
- [3] P. G. Steg, "Heart rate management in coronary artery disease: the CLARIFY registry," *European heart journal supplements*, vol. 11, Suppl D, pp. D13–D18, 2009.
- [4] S. Prabakaran, S. Vitter, and G. Lundberg, "Cardiovascular Disease in Women Update: Ischemia, Diagnostic Testing, and Menopause Hormone Therapy," *Endocrine Practice*, vol. 28, no. 2, pp. 199–203, 2022.
- [5] A. Jain, P. Jain, and G. B. Markande, "Etiopathological and diagnostic study of Margavaranajanya Hridroga wsr to coronary artery disease-an observational study," *Journal of Ayurveda and Integrated Medical Sciences*, vol. 7, no. 7, pp. 47–57, 2022.
- [6] U. Ralapanawa and R. Sivakanesan, "Epidemiology and the magnitude of coronary artery disease and acute coronary syndrome: a narrative review," *Journal of Epidemiology and Global Health*, vol. 11, no. 2, pp. 169–177, 2021.
- [7] L. Horváth, N. Németh, G. Fehér, Z. Kívés, D. Endrei, and I. Boncz, "Epidemiology of peripheral artery disease: narrative review," *Life*, vol. 12, no. 7, p. 1041, 2022.
- [8] World Health Organization, "Assessing national capacity for the prevention and control of noncommunicable diseases," 2010, [http://www.who.int/cancer/publications/national\\_capacity\\_prevention\\_ncds.pdf](http://www.who.int/cancer/publications/national_capacity_prevention_ncds.pdf).
- [9] World Health Organization, "Cardiovascular disease: World Heart Day," 2017 Available at: [http://www.who.int/cardiovascular\\_diseases/world-heart-day-2017/en/](http://www.who.int/cardiovascular_diseases/world-heart-day-2017/en/).
- [10] Y.-Y. Chen, J.-C. Lu, and J.-K. Jan, "A secure EHR system based on hybrid clouds," *Journal of medical systems*, vol. 36, no. 5, pp. 3375–3384, 2012.
- [11] S. Saadat, M. Yousefifard, H. Asady, A. Moghadas Jafari, M. Fayaz, and M. Hosseini, "The most important causes of death in Iranian population; a retrospective cohort study," *Emergency*, vol. 3, no. 1, pp. 16–21, 2015.
- [12] World Health Organization, "Assessing national capacity for the prevention and control of noncommunicable diseases: report of the 2019 global survey," 2010, <https://www.who.int/publications/i/item/9789240002319>.
- [13] M. Ahmadi and N. Aslani, "Capabilities and advantages of cloud computing in the implementation of electronic health record," *Acta Informatica Medica*, vol. 26, no. 1, pp. 24–28, 2018.
- [14] S. Mattke, M. Seid, and S. Ma, "Evidence for the effect of disease management: is \$1 billion a year a good investment?," *American Journal of Managed Care*, vol. 13, no. 12, pp. 670–676, 2007.
- [15] V. Mohan, Y. K. Seedat, and R. Pradeepa, "The rising burden of diabetes and hypertension in Southeast Asian and African regions: need for effective strategies for prevention and control in primary health care settings," *International Journal of Hypertension*, vol. 2013, Article ID 409083, 14 pages, 2013.
- [16] R. E. Gliklich, N. A. Dreyer, and M. B. Leavy, Eds., "AHRQ methods for effective health care," in *Registries for Evaluating Patient Outcomes: A User's Guide*, Agency for Healthcare Research and Quality (US), Rockville (MD), 2014.
- [17] M. Abdelhak, S. Grostick, and M. A. Hanken, *Health Information-e-Book: Management of a Strategic Resource*, Elsevier Health Sciences, 2014.
- [18] M. Endriyas, A. Alano, E. Mekonnen et al., "Understanding performance data: health management information system data accuracy in Southern Nations Nationalities and People's Region, Ethiopia," *Ethiopia. BMC health services research*, vol. 19, no. 1, pp. 1–6, 2019.
- [19] J. C. Tan, A. C. Ferdi, M. C. Gillies, and S. L. Watson, "Clinical registries in ophthalmology," *Ophthalmology*, vol. 126, no. 5, pp. 655–662, 2019.
- [20] M. Lundström, M. Dickman, Y. Henry et al., "Changing practice patterns in European cataract surgery as reflected in the European Registry of Quality Outcomes for Cataract and Refractive Surgery 2008 to 2017," *Journal of Cataract & Refractive Surgery*, vol. 47, no. 3, pp. 373–378, 2021.
- [21] J. A. Singh, X. Lu, S. Ibrahim, and P. Cram, "Trends in and disparities for acute myocardial infarction: an analysis of Medicare claims data from 1992 to 2010," *BMC medicine*, vol. 12, no. 1, pp. 1–13, 2014.
- [22] G. Ramu and R. B. Eswara, "Secure architecture to manage EHR's in cloud using SSE and ABE," *Health and Technology*, vol. 5, no. 3–4, pp. 195–205, 2015.
- [23] M. Arcopinto, A. Salzano, F. Ferrara, E. Bobbio, A. M. Marra, R. Abete et al., "The Tosca registry: an ongoing, observational, multicenter registry for chronic heart failure," *Translational medicine@ Uni Sa*, vol. 14, p. 21, 2016.
- [24] E. Bossone, G. Limongelli, G. Malizia et al., "The T.O.S.C.A. Project: research, education and care," *Project: research, education and care. Monaldi Archives for Chest Disease*, vol. 76, no. 4, pp. 198–203, 2011.
- [25] T. Råmunddal, L. Hoebers, J. P. Henriques et al., "Chronic total occlusions in Sweden—a report from the Swedish Coronary Angiography and Angioplasty Registry (SCAAR)," *PLoS One*, vol. 9, no. 8, article e103850, 2014.
- [26] T. Råmunddal, L. P. Hoebers, J. P. Henriques et al., "Prognostic impact of chronic total occlusions: a report from SCAAR (Swedish Coronary Angiography and Angioplasty Registry)," *JACC: Cardiovascular Interventions*, vol. 9, no. 15, pp. 1535–1544, 2016.
- [27] M. L. Fokkema, S. K. James, P. Albertsson et al., "Population trends in percutaneous coronary intervention: 20-year results from the SCAAR (Swedish Coronary Angiography and Angioplasty Registry)," *Journal of the American College of Cardiology*, vol. 61, no. 12, pp. 1222–1230, 2013.
- [28] M. Samadbeik, M. Ahmadi, F. Sadoughi, and A. Garavand, "A comparative review of electronic prescription systems: lessons learned from developed countries," *Journal of research in pharmacy practice*, vol. 6, no. 1, pp. 3–11, 2017.
- [29] S. V. G. Subrahmanya, D. K. Shetty, V. Patil, B. Hameed, R. Paul, K. Smriti et al., "The role of data science in healthcare advancements: applications, benefits, and future prospects," *Irish Journal of Medical Science*, vol. 1971–, pp. 1–11, 2021.

- [30] A. Allen, H. Patrick, J. Ruof et al., "Development and pilot test of the registry evaluation and quality standards tool: an information technology-based tool to support and review registries," *Value in Health*, vol. 25, no. 8, pp. 1390–1398, 2022.
- [31] A. Garavand, R. Rabiei, H. Emami, M. Pishgahi, and M. Vahidi-Asl, "The attributes of hospital-based coronary artery diseases registries with a focus on key registry processes: a systematic review," *Health Information Management Journal*, vol. 51, no. 2, pp. 63–78, 2022.
- [32] A. Garavand, H. Emami, R. Rabiei, M. Pishgahi, and M. Vahidi-Asl, "Designing the coronary artery disease registry with data management processes approach: a comparative systematic review in selected registries," *International Cardiovascular Research Journal*, vol. 14, no. 1, 2020.
- [33] E. Gozali, R. Safdari, M. Ghazisaeedi, B. Rahimi, H. F. Eslamlou, and E. Mehraeen, "Identification and validation of requirements for a registry system of children's developmental motor disorders in Iran," *Methods of information in medicine*, vol. 58, no. 4/5, pp. 124–130, 2019.
- [34] Z. Meidani, A. Moravveji, S. Gohari et al., "Development and testing requirements for an integrated maternal and child health information system in Iran: a design thinking case study," *Methods of Information in Medicine*, vol. 61, Supplement 2, pp. e64–e72, 2022.
- [35] G. Nagel, H. Ünal, A. Rosenbohm, A. C. Ludolph, and D. Rothenbacher, "Implementation of a population-based epidemiological rare disease registry: study protocol of the amyotrophic lateral sclerosis (ALS)-registry Swabia," *BMC Neurology*, vol. 13, no. 1, pp. 1–10, 2013.
- [36] A. T. Tabrizi, H. Moghaddasi, R. Rabiei, B. Sharif-Kashani, and E. Nazemi, "Development of a catheterization and percutaneous coronary intervention registry with a data management approach: a systematic review. Perspectives in health information management," *Perspectives in health information management*, vol. 16, 2019.
- [37] T. Bauer, H. Möllmann, U. Zeymer et al., "Multivessel percutaneous coronary intervention in patients with stable angina: a common approach? Lessons learned from the EHS PCI registry," *Heart and vessels*, vol. 27, no. 5, pp. 453–459, 2012.
- [38] K. R. Napier, J. Pang, L. Lamont et al., "A web-based registry for familial hypercholesterolaemia," *Heart, Lung and Circulation*, vol. 26, no. 6, pp. 635–639, 2017.
- [39] Y. Kodra, M. Posada de la Paz, A. Coi et al., "Data quality in rare diseases registries," *Advances in Experimental Medicine and Biology*, vol. 1031, pp. 149–164, 2017.
- [40] S. Blumenthal, "Improving interoperability between registries and EHRs," *AMIA Summits on Translational Science Proceedings*, vol. 2018, p. 20, 2018.
- [41] T. E. Kottke, C. J. Baechler, and E. D. Parker, "accuracy of heart disease prevalence estimated from claims data compared with an electronic health record," *Preventing chronic disease*, vol. 9, 2012.
- [42] M. Mazzucato, L. Visonà Dalla Pozza, S. Manea, C. Minichiello, and P. Facchin, "A population-based registry as a source of health indicators for rare diseases: the ten-year experience of the Veneto region's rare diseases registry," *Orphanet journal of rare diseases*, vol. 9, no. 1, pp. 1–12, 2014.
- [43] S. Derayeh, A. Kazemi, R. Rabiei, A. Hosseini, and H. Moghaddasi, "National information system for rare diseases with an approach to data architecture: a systematic review," *Research*, vol. 7, no. 3, pp. 156–163, 2018.
- [44] A. Sabahi, F. Asadi, S. Shadnia, R. Rabiei, and A. S. Hosseini, "The features and processes of poisoning registries: a scoping review," *International Journal of Medical Toxicology and Forensic Medicine*, vol. 11, no. 3, p. 34286, 2021.
- [45] A. Sabahi, F. Asadi, S. Shadnia, R. Rabiei, and A. Hosseini, "Minimum data set for a poisoning registry: a systematic review," *Iranian journal of pharmaceutical research: IJPR*, vol. 20, no. 2, pp. 473–485, 2021.
- [46] A. Dennis, B. Wixom, and D. Tegarden, *Systems Analysis and Design: An Object-Oriented Approach with UML*, John Wiley & Sons, 2015.
- [47] A. Sabahi, F. Asadi, R. Rabiei, and S. Paydar, "Providing a population-based registry model of drug poisoning in Iran," *Iranian Journal of Pharmaceutical Research*, vol. 21, no. 1, 2022.
- [48] N. T. Designing, *Developing and Evaluation of a Hemoglobinopathy Registry*, [Ph.D. Thesis].Tehran University of Medical Sciences, 2019.