Prevalence of cardiovascular Diseases Risk Factors Among Jordanians

Follow this and additional works at: https://www.j-saudi-heart.com/jsha

Part of the Cardiology Commons

This work is licensed under a Creative Commons Attribution-Noncommercial-No Derivative Works 4.0 License.

Recommended Citation
Alsaud, Wesam Saleh; Tabbaa, Mohammad; Kasabri, Violet; Suyagh, Maysa; Abu Alsamen, Muneer; Haddad, Hussain; and Sheweiki, Anas (2020) "Prevalence of cardiovascular Diseases Risk Factors Among Jordanians," Journal of the Saudi Heart Association: Vol. 32 : Iss. 2 , Article 34.
Available at: https://doi.org/10.37616/2212-5043.1074

This Original Article is brought to you for free and open access by Journal of the Saudi Heart Association. It has been accepted for inclusion in Journal of the Saudi Heart Association by an authorized editor of Journal of the Saudi Heart Association.
We thank the following members of the Jordanian General Practitioner/Physician Association for participation in this study: Dr. Mahmoud Hashem, Dr. AbdulWahhab Awad, Dr. Ahmad Shalabi, Dr. Ali Mahseri, Dr. Aref Abu Hwaij, Dr. Maher Yagi, Dr. Mohammad Alazzam, Dr. Yahya Tarifi, Dr. Fayez Alqadi, Dr. Hashem Abu Alhaija, Dr. Ibrahim Shoubaki, Dr. Ibrahim Tuffaha, Dr. Kawther Sarhan, Dr. Khalid Abdul Wahed, Dr. Mohammad Hajjaj, Dr. Mufeed Damrah, Dr. Muhammad Ulaimat, Dr. Numir Batarseh, Dr. Osama Kamal, Dr. Rahat Zgetter, Dr. Samyah Masarweh, Dr. Samyah Mdanat, Dr. Yousef Alfaqeeh and Dr. Zeyad Nsour. Also we thank the following members of the Jordanian pharmaceutical company for participation in this study: Dr. Taghreed Barqawi, Sales and Marketing Director, Dr. Alaa alturk, Dr. Nazer Zayed, business unit manager, Dr. Alaa banihamdan, Dr. Obada alqarioty and Dr. Nisreen Fakori, Medical Representatives Supervisors, Dr. Alaa’ odeh, Dr. Diana alabdallat, Dr. Farah alomari, Dr. Nour nouri, Dr. Abdullah alwheidi and Dr. Waleed Alabadi, Medical Representatives.
Prevalence of Cardiovascular Diseases Risk Factors among Jordanians

Wesam Alsaud a*, Mohammad J. Tabbaa b, Violet N. Kasabri c, Maysa F. Suyagh c, Muneer A. Abu Alsamen d, Hussain M. Haddad d, Anas O. ALshweki a

a Scientific Office, Jordanian Pharmaceutical Company, Amman, Jordan
b School of Agriculture, University of Jordan, Amman, Jordan
c School of Pharmacy, University of Jordan, Amman, Jordan
d Diabetologist, General Practitioner Society, Amman, Jordan

Abstract

**Background and aims:** One of the most common causes of death worldwide is cardiovascular diseases (CVDs). This study evaluated the prevalence of CVDs risk factors (RFs) and their constellation electively among the Jordanian population and, assessing the most prevalent RF interplay with the rest of CVDs RFs as well as the impact of age and gender dimorphism on the frequencies of coexistence of multiple CVDs risk factors (RFs) among the Jordanian population.

**Methods and results:** In this observational multicenter study, a total of 1449 subjects were enrolled. The mean age (±SD) was 44.35 ± 14.46 years; 796 (54.9%) of them were females and 801 (55.28%) of the whole study pool had no family history of premature CVDs. Only 5.9% of the population did not have any of these RFs. The prevalence of CVDs MRFs within-affected subjects was as follows: there were 1081 (74.6%) subjects with overall dyslipidemia, 471 (32.51%) with obesity, 456 (31.47%) were smokers, and at the first diagnostic encounter 541 (37.47%) were with elevated blood pressure and, 310 (21.51%) were with elevated random blood sugar. The coexistence of ≥ two, ≥ three and, ≥ four RFs was observed in 75.7%, 44.4%, and 21.4% of the subjects, respectively. The constellation of multiple RFs was more frequent in men than that in women, where the presence of ≥ two RFs for men was at 86.18% vs. 67.09% for women. Similarly, the appearance of multiple RFs increases with age, starting from the existence of ≥ three, and four RFs respectively. Most notably the clustering of ≥ five RFs in the age group of 45–59 years showed the greatest frequency vs. any other age group.

**Conclusions:** CVDs risk factors (RFs) and clusters of them are extremely prevalent in the Jordanian population. Overall dyslipidemia is the most prevalent MRF and the most favors clustering with other CVDs RFs. Combined two RFs had the highest proportional frequency between all six RFs clusters. The constellation of at least two, three, and four CVDs RFs presented at almost three-fourth, half, and around one-fourth; respectively, Middle-aged males presented significantly higher rates of ≥ five RFs occurrences than females.

1. Introduction

The non-communicable chronic diseases (NCDs) showed a sustainable elevation in deaths number in addition to significant patient's quality of life deterioration [1]. According to the World Health Organization (WHO), more than 36 million people die annually from chronic NCDs, amongst these; cardiovascular diseases (CVDs) account for most deaths (17.9 million), followed by cancer (9.0 million), chronic respiratory tract illnesses (3.9 million), and diabetes (1.6 million) [2]. Deaths from CVDs, cancer, chronic respiratory tract illness, and diabetes represent over 80% of all premature NCDs deaths worldwide [3]; where mostly low and middle-income countries

https://doi.org/10.37616/2212-5043.1074
2212-5043/© 2020 Saudi Heart Association. This is an open access article under the CC-BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).
accounted for over three-quarters of CVDs deaths [4]. Global major burdens on the total costs of medical care are caused by CVDs [5]. Risk factors (RFs) of CVDs can be categorized into three groups: non-modifiable RFs, modifiable RFs (MRFs), and emerging RFs or psychosocial factors, including workplace stress. The pattern of CVDs RFs distribution is often as clusters that incrementally culminate into CVDs in a multiplicative rather than in an additive manner [6a–6d]. The non-modifiable risk elements include age, gender, and family history of premature CVDs [6a,7,8]. Elevated blood pressure, dyslipidemias, diabetes mellitus, obesity, smoking and lack of physical activity are all included under modifiable risk elements [6a]. In the INTERHEART study from 52 countries, the potential relation between modifiable risk factors, including smoking, elevated ApoB/ApoA1 ratio, history of elevated blood pressure, abdominal obesity, psychosocial matters, and regular alcohol ingestion and myocardial infarction was substantiated [9a–9g]. Psychosocial factors like stressful life calamities, multiple stress, animosity, depression, and anxiety are also emerging as RFs for CVDs [10a–10d]. The objectives of this study were to evaluate the prevalence of CVDs RFs and RFs clusters among the Jordanian population also to assess the most prevalent RF interplay with the rest of CVDs RFs in addition to the impact of age and gender on the coexistence frequencies of multiple CVDs RFs.

2. Method

2.1. Study population

In this observational multicenter study, combined samples of 1449 were enrolled from different cities in the South, Middle, and North of Jordan over the last two weeks of August 2018, in collaboration with the general medical practitioners and physicians society. Based on determining sample size for a survey using Mendenhall (1983) equation [11], we need not >385 replicates as a representative sample from overall Jordan population which is around 10 million but to have stratified random samples from all Governorates of Jordan we increased the sample size to 1449. Jordanian subjects with age ≥20 years signed an informed consent form before participation, only mentally or physically handicapped people and pregnant females were excluded. The presence of the following six CVDs RFs, five MRFs; hypertension (HTN), diabetes mellitus (DM), cigarette smoking, dyslipidemia, obesity; and one non-modifiable RF; family history of premature CVDs; were assessed in every subject questionnaire, and. Further assessment of the following demographic data was undertaken as age and gender, educational level, height and weight. An average of three readings of blood pressure was measured by trained investigators, and a blood sample was collected by the laboratory specialized practitioners to analyze for random lipid profile and random blood sugar. The enrolled subjects have been educated about the significance of CVDs RFs. Collected data were coded and filed. This study was approved by the Institutional Review Board (IRB) of King Abdullah University Hospital.

2.2. Study measures

All RFs were defined by internationally approved standard references. Principally normal lipid profile defined as per the Adult Treatment Panel III [12]: TC < 200 mg/dL (5.2 mmol/L), LDL-C<130 mg/dL (3.36 mmol/L), TG < 150 mg/dL (1.69 mmol/L), HDL-C >50 mg/dL (1.04 mmol/L) in women, HDL-C>40 mg/dL (1.04 mmol/L) in men. We defined the case where at least one variable of lipid profile parameters is abnormal as Overall dyslipidemia, and when all variables of lipid profile parameters are abnormal as mixed dyslipidemia. According to the International
Society of Hypertension [13], HTN definition is when systolic blood pressure (SBP) > 140 mmHg and Diastolic BP (DBP) > 90 mmHg. One of ADA [American Diabetes Association; 14] criteria for diabetes diagnosis is random BS > 200 mg/dL (>11.1 mmol/L) with hyperglycaemia symptoms. Enrolled nonsmokers were past smokers at least a month before enrollment or never smoked otherwise the subject was considered a smoker. A family history of premature CVDs defined as myocardial infarction, coronary revascularization or sudden death before 55 years of age in father or any other male first-degree relative, or before 65 years of age in mother or any other female first-degree relative. BMI calculated by dividing bodyweight in kg by the square of height in meters. Body weight categorized according to BMI [15] as the following: Underweight had 13.0 ≤ BMI ≤ 18.4, normal body when weight had is 18.5–24.9, Overweight had has 25 ≤ BMI ≤ 29.9 and the Obesity had BMI ≥ 30.0.

2.3. Statistical analysis

The Statistical Analysis System (SAS, 2011) was used for data entry and analysis. Frequencies and percentages were used to describe categorical variables. The differences in percentages of RFs among men and women, and among the four pre-specified age groups were analyzed using Chi square test. A P-value of <0.05 was considered statistically significant.

3. Results

The demographics of the study population were presented in Table 1, showing that the studied population had almost equal proportions of males and females. Also, from the four age groups, two-third of subjects’ age was between 30 and 60 years and the rest at the age extremes. Such nature of replicates distribution between the subgroups indicates the random sampling of subjects. The frequency of six CVDs RFs evaluated as shown in Fig. 1, Overall Dyslipidemia being most prevalent CVDs RF, followed by FH pCVD, initial encounter HTN, obesity, smoking and finally initial encounter DM. Fig. 2 displays the prevalence of random clustering pattern of six CVDs RFs. One-third of subjects had any two random RFs of the assigned six RFs. Each clustering group includes the random probability from the six RFs assigned. Calculated frequencies of ≥two, ≥three and ≥four RFs, were observed in 75.71%, 44.41% and, 21.44% of subjects, respectively. The following are the detailed numbers of results (frequency) of each CVDs MRF: A) lipid profile variables::the mixed dyslipidemia 133 (9.18%), elevated TC 510 (35.2%), elevated TG 679 (46.86%), elevated LDL-C 407 (28%) and reduced HDL-C 697 (48.1%). B) Elevated blood pressure at the first diagnostic encounter: 338 (23%) showed high SBP; on the other hand 323 (22%) showed high DBP and 188 (13.0%) had both C) Elevated random blood Sugar at the first diagnostic encounter 102 (7%), 210 (14.5%) were already diabetic. D) Body weight categories: 35 (2%) underweight, 213 (15%) lean, 566 (41%) overweight and 471 (33%) obese. Overweight and obese represent 74% of subjects. E) Smokers represented one-third of subjects: 993 (69%) subjects were none smokers. F) The educational level may relate inversely to the work stress that is a surrogate marker for the emerging CVDs RFs, 235 (16%) had non-elementary school education, 586 (40%) have Junior high - high school education and 628 (43%) diplomas and beyond. We studied the prevalence of overall dyslipidemia in relation to all other CVDs RFs since it showed the highest frequency between them and was noticed the following: A) statistically significant (P-Value <0.0001) gender dimorphism was noticeable where: overall dyslipidemia for men 90.66% versus 61.43% for women, mixed dyslipidemia for men 16.08% versus 3.52% for women, elevated TG for men 58.19% versus 37.56% for women, reduced HDL-C for men 82.08% versus 20.23% for women, elevated TG for women 16.08% versus 3.52% for women, reduced HDL-C for women 82.08% versus 20.23% for women. B) Age effect on dyslipidemia started from early twenties and continued worsening significantly with age. The age group of 45–59 years can be considered the plateau for dyslipidemia deterioration as seen in Table 2. C) Dyslipidemia worsens proportionally with body weight increase, but marked deterioration starts from the overweight group as seen in Table 3. D) The reciprocal decreases of dyslipidemia with the increase in the educational level were statistically significant in the overall dyslipidemia only as shown in Table 4.

Table 1. Demographics of this study population.

| Feature          | N   | %  |
|------------------|-----|----|
| Overall population | 1449 |    |
| Gender           |     |    |
| Men              | 653 | 45.1 |
| Women            | 796 | 54.9 |
| Age groups (y), mean ± SD | 44.35 ± 14.46 | 18.8 |
| 20–29            | 272 | 18.8 |
| 30–44            | 459 | 31.7 |
| 45–59            | 487 | 33  |
| 60–90            | 231 | 15.9 |

N: number of replicates, %: percentage, SD = standard deviation.
The smoking habit negatively and pronouncedly impacted dyslipidemia as shown in Table 5.

Table 6 demonstrates RFs coexistence frequency (smoking, DM, HTN, obesity, FH pCVD and overall dyslipidemias) with one RF specified at each time. The results showed Overall Dyslipidemias had the highest tendency to cluster with other RFs. The prevalence of RFs clustering was evaluated according to age (Fig. 3). The presence of zero or one RF
observed to decline with increasing the participants age. The presence of two RFs was observed to be the lowest in the middle age group participants (45–59 Y). Furthermore, the presence of three, four and five RFs was observed to elevate aging. Finally, the presence of six RFs was observed to be the highest in the middle age group participants (45–59 Y). The presence of ≥2, 3, 4 RFs increases with age but the presence of ≥5 RFs peaked in the middle age group of 45-59Y), vseither older (60-90Y) or younger subjects (20-44Y). The prevalence of RFs coexistence in 796 women was compared with that in 635 men (Fig. 4). The absence of any RF was observed more women than men. Men were more likely than women to have three, four, five RFs than women.

4. Discussion

The major outcomes of this study are:Dyslipidemias was the most prevalent CVDs RF among

| Table 2. Dyslipidemia Variation according to age. |
|-----------------------------------------------|
| Age groups (y) | Overall Dyslipidemia N (%) | Mixed Dyslipidemia N (%) | High TC N (%) | High TG N (%) | High LDL-C N (%) | Low HDL-C N (%) | High TC/HDL-C N (%) |
|----------------|------------------------------|--------------------------|---------------|---------------|------------------|-------------------|---------------------|
| 20–29          | 158 (58.1)                  | 12 (4.4)                 | 52 (19.1)     | 70 (25.7)     | 45 (16.5)        | 115 (42.3)        | 35 (12.9)           |
| 30–44          | 328 (71.5)                  | 44 (9.6)                 | 159 (34.6)    | 209 (45.5)    | 124 (27.0)       | 216 (47.1)        | 117 (25.5)          |
| 45–59          | 401 (82.3)                  | 54 (11.1)                | 214 (43.9)    | 265 (54.4)    | 175 (35.9)       | 237 (48.7)        | 155 (31.8)          |
| 60–90          | 194 (84.00)                 | 23 (10.00)               | 85 (36.8)     | 135 (58.4)    | 129 (55.8)       | 64 (27.7)         |                     |

P-value according to Chi-square test of different age groups.

| TC: Total Cholesterol, TG: Triglycerides, HDL-C: High-Density Lipoprotein Cholesterol; LDL-C: Low-Density Lipoprotein Cholesterol, N: number of replicates, %: percentage. |

| Table 3. Dyslipidemia Variation according to body mass index (BMI). |
|-------------------------------------------------------------------|
| BMI Variable | Overall Dyslipidemia N (%) | Mixed Dyslipidemia N (%) | High TC N (%) | High TG N (%) | High LDL-C N (%) | Low HDL-C N (%) | High TC/HDL-C N (%) |
|--------------|----------------------------|--------------------------|---------------|---------------|------------------|-------------------|---------------------|
| Underweight  | 17 (48.6)                  | 0 (0)                    | 3 (8.6)       | 5 (14.3)      | 3 (8.6)          | 13 (37.1)        | 2 (5.7)             |
| Normal body weight | 213 (60.0)             | 16 (4.5)                 | 92 (25.9)     | 104 (29.3)    | 79 (22.3)        | 136 (38.3)       | 57 (16.1)           |
| Overweight   | 460 (78.2)                 | 65 (11.1)                | 218 (37.1)    | 294 (50)      | 175 (29.8)       | 304 (51.7)       | 157 (26.7)          |
| Obesity      | 391 (83.0)                 | 52 (11.0)                | 197 (41.8)    | 276 (58.6)    | 150 (31.9)       | 244 (51.8)       | 155 (32.9)          |

P-value according to Chi-square test of different BMI groups.

| TC: Total Cholesterol, TG: Triglycerides, HDL-C: High-Density Lipoprotein Cholesterol; LDL-C: Low-Density Lipoprotein Cholesterol, N: number of replicates, %: percentage. |

| Table 4. Dyslipidemia Variation according to the educational status. |
|-------------------------------------------------------------------|
| Educational level Variable | Overall Dyslipidemia N (%) | Mixed Dyslipidemia N (%) | High TC N (%) | High TG N (%) | High LDL-C N (%) | Low HDL-C N (%) | High TC/HDL-C N (%) |
|---------------------------|----------------------------|--------------------------|---------------|---------------|------------------|-------------------|---------------------|
| None - Elementary school  | 185 (78.7)                 | 22 (9.4)                 | 91 (38.7)     | 116 (49.4)    | 72 (30.6)        | 120 (51.1)       | 66 (28.1)           |
| Junior high - high school | 450 (76.8)                 | 54 (9.2)                 | 207 (35.3)    | 288 (49.1)    | 171 (29.2)       | 284 (48.5)       | 154 (26.3)          |
| Diploma – and above       | 446 (71.0)                 | 57 (9.1)                 | 212 (33.8)    | 275 (43.8)    | 164 (26.1)       | 293 (46.7)       | 151 (24.0)          |

P-value according to Chi-square test of different educational status groups.

| TC: Total Cholesterol, TG: Triglycerides, HDL-C: High-Density Lipoprotein Cholesterol; LDL-C: Low-Density Lipoprotein Cholesterol, N: number of replicates, %: percentage. |

| Table 5. Dyslipidemia Variation according to smoking status. |
|-------------------------------------------------------------------|
| Smoking Variable | Overall Dyslipidemia N (%) | Mixed Dyslipidemia N (%) | High TC N (%) | High TG N (%) | High LDL-C N (%) | Low HDL-C N (%) | High TC/HDL-C N (%) |
|-------------------|----------------------------|--------------------------|---------------|---------------|------------------|-------------------|---------------------|
| No Smoking        | 708 (71.3)                 | 73 (7.5)                 | 351 (35.4)    | 450 (45.3)    | 278 (28.0)       | 404 (40.7)       | 217 (21.9)          |
| Smoker            | 373 (81.8)                 | 60 (13.2)                | 159 (34.9)    | 229 (50.2)    | 129 (28.3)       | 293 (64.3)       | 154 (33.8)          |

P-value according to Chi-square test of smoking vs. nonsmoking groups.

| TC: Total Cholesterol, TG: Triglycerides, HDL-C: High-Density Lipoprotein Cholesterol; LDL-C: Low-Density Lipoprotein Cholesterol, N: number of replicates, %: percentage. |
Gender dimorphism was delineated where higher prevalence was among males vs. females. Dyslipidemias start from the early twenties and continue worsening with age progression. Dyslipidemias worsen proportionally with weight gain, but the flare up of dyslipidemia begins at the overweight category. Dyslipidemias prevalence decreased proportionally with the increase in educational level that suggested relating inversely with work stress. Smoking related to a higher prevalence of dyslipidemia. (2) Clusters prevalence had an ascending order from highest two then three, followed by four RFs; Overall dyslipidemias had the greatest tendency to coexist with other CVDs RFs. The Clustering of CVDs RFs was more common in men than in women and five RFs grouping was most common at age group 45–65 years compared with younger or older age intervals. Mitigation of CVDs risk can be attained by managing the MRF studied in this study; overall Dys, HTN, DM, obesity and smoking, properly unlike the non-modifiable RFs as of age, gender and, FH pCVD. The MRFs can be elaborated to include sedentary lifestyle, unhealthy dietary behavior low socioeconomic level, regular alcohol consumption, urbanization, and pollution [16,17a–17c]. Different studies stated that these modifiable RFs are related to a large number of premature CV deaths globally [18,19]. In this study, CVDs RFs prevalence results demonstrated similar results of a study performed in Africa and the Middle East (AfME) region [20]. This study revealed prevalence of overall dyslipidemias within the Jordanian population reflective of other Jordanian reports [21]. Fasting lipid profile is not routinely included within the Jordanian screening program unlike BP and BS despite the international recommendation of consistent screening [22]. Multiple RFs are more prevalent among men, lower socioeconomic status, singles, and less prevalent among homeowners and older age groups as concluded from past studies [23]. Clustering of RFs differs across investigated populations according to geographic regions, urban vs rural communities, different ethnic groups, individuals who do not have overt CVDs compared with those who have symptomatic CVDs, men vs. women and old vs. young individuals [24a–24c]. An outpatient clinic-based study in 14 Middle Eastern and African nations estimated the clustering of three or more RFs in more than half (53%) of the

---

### Table 6. Number of coexisting cardiovascular diseases risk factors (RF) with each specified CVDs RF.

| Specified CVDs Risk Factor | No of coexisting risk factors |
|----------------------------|--------------------------------|
|                            | 1 (%) | 2 (%) | 3 (%) | 4 (%) | 5 (%) |
| Smoking                    | 12.5  | 35.3  | 39.3  | 40.3  | 41.6  |
| FH pCVD                    | 23.8  | 33.9  | 55.3  | 76.0  | 89.6  |
| Overall Dyslipidemias      | 49.1  | 76.5  | 91.5  | 94.1  | 100.0 |
| Obesity                    | 5.3   | 23.3  | 40.8  | 63.4  | 13.8  |
| Diabetes mellitus          | 0.8   | 10.4  | 23.3  | 46.6  | 90.9  |
| Hypertension               | 8.7   | 20.6  | 49.9  | 79.6  | 93.5  |

FH pCVD: family history of premature cardiovascular disease. CVDs: cardiovascular diseases, %: percentage.

---

Fig. 3. Proportions of subjects in four age groups (20–29Y, 30–44 Y, 45–59 Y and 60–90 Y) who have one or more of the six risk factors (smoking, DM, HTN, obesity, family history of premature cardiovascular disease and Overall Dyslipidemias). RF = risk factor. P-value<0.0001, P-value calculated according to Chi-square test between different age groups.
The observation of a decreasing prevalence of RFs clustering among the older age group compared with those aged 45–69 years might have been result from: age effecting per se steady increase of the absolute baseline risk of CAD independent of the conventional RFs. It also might be related to inherent survivor bias as subjects with MRFs tend to die at a considerably younger age [25,26]. Results from gender-specific relation with CVDs RF are contradictory, one have illustrated that although men and women share common MRFs for CVDs, such as cigarette smoking, DM, depression, and other psychosocial MRFs, a more negative health impact was observed in women [26]. On the contrary a recent study concluded that the prevalence of major RFs of NCDs was greater among more elderly persons and male participants [27]. This study showed that men were more likely than women to have ≥ three, four and five RFs. Furthermore, the sudden drop in frequency of one or more RFs to a single-digit observed in both women and men. We can predict the reason of the high prevalence of multiple RFs coexistence to society growth, age profile variation, low education levels, lifestyle changes, dietary habits, lack of adoption of regular physical activity, ineffective anti-tobacco campaigns, and rise of prevalence rates of DM, metabolic syndrome, and obesity [28a–28e]. Subjects with multiple RFs were prone to higher probability developing coronary artery calcification, and premature mortality and morbidity [29]. In Jordan CVDs RFs clustering is higher than it is in other regions of the world, enhancing the potential benefits expected from aggressive preventive measures. The major starting strategy is the rapid detection and proper control of MRFs thereby reducing substantially CVDs burdens. This study contains a few limitations that have to be discussed. Exactly like all studies [30], this study is subject to selection bias, and missing or incomplete information, this study might has a selection bias since the subjects who heard about the study announcement via the social media and newspapers came to the nearest general practitioner clinic and participated in this study. Since the data were self-reported from enrolled subjects not the clinician supervising their health status, information and/or recall bias might occur for some of the CVDs, despite the interviews were carried out by well-trained personnel, the potential for bias was present (e.g. over or under-diagnosed due to recall issues or subjectivity in the reporting of symptoms). Using BMI, not the waist circumference, was the definition of overweight and obesity strata within study pool of participants. Although BMI is easy to calculate and more commonly used, it is a simple and imperfect anthropometric biomarker that does not possess the discriminating power in differentiating between lean body mass and fat mass. Despite these limitations, this study maintains several strengths. It utilized substantial data that is representative of the entire Jordanian population in a short time frame of only two weeks. It used standardized data collection tools, for example; the interviews were performed

![Fig. 4. Proportions of men and women who have one or more of the six risk factors (hypertension, diabetes mellitus, hypercholesterolemia, obesity, cigarette smoking, and family history of premature cardiovascular disease). RF = risk factor. P-value < 0.0001. P_Value according to Chi-square test between different gender groups.](image-url)
by trained investigators, the way they measured height, weight, and blood pressure and blood sugar follows a similar procedure. This study is unique in that it evaluated the prevalence and coexistence of six RFs in a relatively large, young (over 20 year), and CVDs free Jordanians.

5. Conclusion

CVDs RFs and constellation of ≥ two, three and four RFs are extremely prevalent in the Jordanian population. The most frequent MRF is Overall dyslipidemia also it is the most favors clustering with CVDs RF. Randomly combined two RFs cluster has the highest proportional frequency than other clustered groups. Middle-aged males presented significantly higher rates of ≥ five RFs occurrences than females. Suggestively mitigation of CVDs morbidity and mortality in the Jordanian population can be attained by initiating primary cardiovascular medical and behavioral interventions to control the CVDs MRFs frequencies and constellation.

Disclosure of Funding

The Jordanian Pharmaceutical Company funded the lipid profile analysis that performed by a third party (MedLab Laboratories), the statistical analysis Performed by Jordan University partners.

Author contribution

Conception and design of Study: Wesam S. Alsaud, Muneer A. Abu Alsamen, Hussain M. Haddad, Anas O. Sheweiki. Literature review: Wesam S. Alsaud, Mohammad J. Tabbaa, Violet N. Kasabri, Maysa F. Suyagh. Acquisition of data: Wesam S. Alsaud, Mohammad J. Tabbaa, Violet N. Kasabri, Maysa F. Suyagh. Analysis and interpretation of data: Mohammad J. Tabbaa. Research investigation and analysis: Wesam S. Alsaud; Muneer A. Abu Alsamen, Hussain M. Haddad, Anas O. Sheweiki. Data collection: Wesam S. Alsaud, Mohammad J. Tabbaa, Violet N. Kasabri, Maysa F. Suyagh. Drafting of manuscript: Wesam S. Alsaud. Revising and editing the manuscript critically for important intellectual contents: Wesam S. Alsaud, Mohammad J. Tabbaa, Violet N. Kasabri, Maysa F. Suyagh. Muneer A. Abu Alsamen, Hussain M. Haddad, Anas O. Sheweiki. Data preparation and presentation: Wesam S. Alsaud. Supervision of the research: Wesam S. Alsaud. Research coordination and management: Wesam S. Alsaud; Anas O. Sheweiki. Funding for the research: Wesam S. Alsaud.

Conflicts of Interest

None declared.

Acknowledgments

We thank the following members of the Jordanian General Practitioner/Physician Association for participation in this study: Dr. Mahmoud Hashem, Dr. AbdulWahhab Awad, Dr. Ahmad Shalabi, Dr. Ali Mahseri, Dr. Aref Abu Hwaij, Dr. Maher Yagi, Dr. Mohammad Alazzam, Dr. Yahya Tarifi, Dr. Fayeza Alqadi, Dr. Hashem Abu Alhaija, Dr. Ibrahim Shoubaki, Dr. Ibrahim Tuffaha, Dr. Kawther Sarhan, Dr. Khalid Abdul Wahed, Dr. Mohammad Hajjaj, Dr. Mufeed Damrah, Dr. Muhammad Ulaimat, Dr. Numir Batarseh, Dr. Osama Kamal, Dr. Rahat Zgerter, Dr. Samyah Masarweh, Dr. Samyah Mdanat, Dr. Yousef Alfaqeeh and Dr. Zeyad Nsour.

Also we thank the following members of the Jordanian pharmaceutical company for participation in this study: Dr. Taghreed Barqawi, Sales and Marketing Director, Dr. Alaa Alturk, business unit manager, Dr. Alaa banihmand, Dr. Obaida alqarioty and Dr. Nisreen Fakori, Medical Representatives Supervisors, Dr. Alaa’ odeh, Dr. Diana alabdallat, Dr. Farah alohari, Dr. Nour nouri, Dr. Abdullah alwheidi and Dr. Waleed Alabadi, Medical Representatives.

References

[1] Rizzuto D, Melis RJF, Angleman S, Qiu C, Marengoni A. Effect of chronic diseases and multimorbidity on survival and functioning in elderly adults. J Am Geriatr Soc 2017;65: 1056–60. https://doi.org/10.1111/jgs.14868.
[2] WHO updates. 2018. http://www.who.int/mediacentre/factsheets/fs355/en/ [Accessed 18 August 2019].
[3] Yeates K, Lohfeld L, Sleeth J, Morales F, Rajkotia Y, Ogedenke O. A global perspective on cardiovascular disease in vulnerable populations. Can J Cardiol 2015;31:1081–93. https://doi.org/10.1016/j.cjca.2015.06.035.
[4] Kwan GF, Mayosi BM, Mocumbi AO, Miranda JJ, Ezzati M, Jain Y, Robles G, Benjamin EJ, Subramanian SV, Bukhman G. Endemic cardiovascular diseases of the poorest billion. Circulation 2016;133:2561–75. https://doi.org/10.1161/CIRCULATIONAHA.116.008731.
[5] Gao B, Zhang L, Wang H. Clustering of major cardiovascular risk factors and the association with unhealthy lifestyles in the Chinese adult population. PloS One 2013;8:e66780. https://doi.org/10.1371/journal.pone.0066780.
[6] [a] Thayer JF, Yamamoto SS, Brosschot JF. The relationship of autonomic imbalance, heart rate variability and cardiovascular disease risk factors. Int J Cardiol 2010;141:122–31. https://doi.org/10.1016/j.ijcard.2009.09.543.
[b] Yu J, Ma Y, Yang S, Pang K, Yu Y, Tao Y, Jin L. Risk factors for cardiovascular disease and their clustering among adults in jilin (China). Int J Environ Res Publ Health 2015;13. https://doi.org/10.3390/ijerph13010070.
[c] De Simone G, Olsen MH, Wachtell K, Hille DA, Dahlof B, Ibsen H, Kjeldsen SE, Lyle PA, Devereux RB. Clusters of metabolic risk factors predict cardiovascular events in hypertension with target-organ damage: the LIFE study. J Hum
Hypertens. 2007;21:625–32. https://doi.org/10.1038/sj.hh.1002203.

[d] Khanal MK, Mansur Ahmed MSA, Moniruzzaman M, Banik PC, Dhungana RR, Bhandari P, Devkota S, Shayanai A. Prevalence of clustering of cardiovascular disease risk factors in rural Nepalese population aged 40–80 years. BMC Publ Health 2018;18:677–677. https://doi.org/10.1186/s12889-018-5609-9.

[7] Kabir Z, Perry IJ, Critchley J, O’Flaherty M, Capewell S, Bennett K. Modelling coronary heart disease mortality declines in the republic of Ireland, 1985–2006. Int J Cardiol 2013;168:2462–7. https://doi.org/10.1016/j.ijcard.2013.03.010.

[8] Khallili D, Sheikholeslami FH, Bakhtiyari M, Azizi F, Momenan AA, Hadeaegh F. The incidence of coronary heart disease and the population attributable fraction of its risk factors in Tehran: a 10-year population-based cohort study. PloS One 2014;9:e105804. https://doi.org/10.1371/journal.pone.0105804.

[9] [a] Yusuf S, Hawken S, Ounpuu S, Dans T, Avezum A, Khalili D, Sheikholeslami FH, Bakhtiyari M, Azizi F, Kabir Z, Perry IJ, Critchley J, O’Flaherty M, Capewell S, Bennett K. Effect of potentially modifiable risk factors associated with myocardial infarction in 52 countries (the INTERHEART study): case-control study. Lancet 2004;364:937–52. https://doi.org/10.1016/S0140-6736(04)17019-0.

[b] Formentini FS, Zaina Nagano FE, Lopes Neto FDN, Adam EL, Fortes FS, Silva LFD. Coronary artery disease and body mass index: what is the relationship? Clin Nutri ESPEN 2019;34:493–500. https://doi.org/10.1016/j.clnesp.2019.08.003.

[c] Lautsch D, Wang T, Yang L, Rajpathak SN. Prevalence of established cardiovascular disease in patients with type 2 diabetes mellitus in the UK. Diabetes Ther 2019;10:1231–7. https://doi.org/10.1007/s13300-019-00698-9.

[d] Lonardo A, Nascimbeni F, Mantovani A, Targher G. Hypertension, diabetes, atherosclerosis and NASH: cause or consequence? J Hepatol 2018;69:1334–52. https://doi.org/10.1016/j.jhep.2018.09.021.

[e] Bae YS, Choi S, Lee K, Son JS, Lee H, Cho MH, Koo HY, Cho IY, Chang J, Kim K, Kim SM, Park SM. Association of concurrent changes in metabolic health and weight on cardiovascular disease risk: a nationally representative cohort study. J Am Heart Assoc 2019;8:e011825. https://doi.org/10.1161/JAHA.118.011825.

[f] Lee W, Hwang SH. The association between smoking or passive smoking and cardiovascular diseases using a Bayesian hierarchical model: based on the 2008-2013 Korea Community Health Survey 2017;39:2017026. https://doi.org/10.4179/kjhs.2017.78.1.2017026.

[g] Lear SA, Hu W, Rangajaran S, Gasevic D, Leong D, Iqbal R, Casanova A, Swaminathan S, Anjana RM, Kumar R, Rosengren A, Wei L, Zhang S, Mony P, Swaminathan S, Mohan V, Gupta R, Kumar R, Vijayakumar K, Lear S, Anand S, Wildgoose A, Diaz R, Avezum A, Lopez-Jaramillo P, Laras F, Yusoff K, Ismail N, Iqbal R, Rahman O, Rosengren A, Yusufali A, Kelishadi R, Kruger A, Pounnane T, Szuba A, Chifamba J, Oguj J, McQueen M, Meccoli M, Dagenais G. Cardiovascular risk and events in 17 low-, middle-, and high-income countries. N Engl J Med 2014;371:818–27. https://doi.org/10.1056/NEJMoai1311890.

[h] Brook RD, Rajagopalan S, Pope C, Brook JR, Bhatnagar A, Diez-Roux AV, Holguin F, Hong Y, Luepker RV, Mittleman MA, Peters A, Siscovich D, Smith JR SC, Whitsel L, Kaufman JD. Particulate matter air pollution and cardiovascular disease: an update to the scientific statement from the American Heart Association. Circulation 2010;121:2331–78. https://doi.org/10.1161/CIR.0b013e3181bece3.

[i] Adam AM, Rehan A, Waseem N, Iqbal U, Saleem H, Ali MA, Shaikh AT, Godil A. Prevalence of conventional risk factors and evaluation of baseline indices among young and elderly patients with coronary artery disease. J Clin Diagn Res : J C Diagn Res 2017;11:OC34–9. https://doi.org/10.1055/s-0039-1713867.

[j] Bundhun PK, Wu ZJ, Chen MH. Impact of modifiable cardiovascular risk factors on mortality after percutaneous coronary intervention: a systematic Review and meta-analysis of 100 studies. Medicine (Baltim) 2015;94:e2213. https://doi.org/10.1097/MD.0000000000002313.

[k] Yusuf S, Reddy S, Ounpuu S, Anand S. Global burden of cardiovascular disease: part i: general considerations, the epidemiologic transition, risk factors, and impact of urbanization. Circulation 2001;104:2746–53. https://doi.org/10.1161/01.hc.4601.099487.

[l] Benner JS, Smith TW, Petrillia AA, Klingman D, Goel S, Tanguy SS, Wong ND. Estimated prevalence of uncontrolled hypertension and multiple cardiovascular risk factors and their associated risk of coronary heart disease in the United States. J Am Soc Hypertens 2008;2:44–53. https://doi.org/10.1016/j.jash.2007.07.001.
risk factor burden in Africa and the Middle East across country income categories: a post hoc analysis of the cross-sectional Africa Middle East Cardiovascular Epidemiological (ACE) study. Arch Publ Health 2018;76:15. https://doi.org/10.1186/s13690-018-0257-5.

[21] Abujbara M, Batieha A. The prevalence of dyslipidemia among Jordanians. 2018. 2018. p. 6298739. https://doi.org/10.1155/2018/6298739.

[22] Jellinger Paul S, Handelsman Yehuda, Rosenblit Paul D, Bloomgarden Zachary T, Fonseca Vivian A, Garber Alan J, George Grunberger, Chris K, Guerin David S H Bell, Mechanick Jeffrey I, Pessah-Pollack Rachel, Wyne Kathleen, Smith Donald, Brinton Eliot A, Fazio Sergio, Davidson Michael. AMERICAN association OF clinical endocrinologists and AMERICAN college OF endocrinology guidelines for management OF dyslipidemia and prevention OF cardiovascular disease. Endocr Pract: April 2017 2017; 23(No. Supplement 2):1–87. https://doi.org/10.4158/EP171764.APPGL.

[23] Poortinga W. The prevalence and clustering of four major lifestyle risk factors in an English adult population. Prev Med 2007;44:124–8. https://doi.org/10.1016/j.ypmed.2006.10.006.

[24] [a] Wang X, Bots ML, Yang F, Sun J, He S, Hoes AW, Niu J, Vaartjes I. A comparison of the prevalence and clustering of major cardiovascular risk factors in The Netherlands and China. Eur J Prev Cardiol 2016;23:1766–73. https://doi.org/10.1177/2047487316648474.

[b] Daviglas ML, Talavera GA, Aviles-Santa ML, Allison M, Cai J, Criqui MH, Gellman M, Giachello AL, Gouskova N, Kaplan RC, Lavange L, Penedo F, Perreira K, Pirzada A, Schneiderman N, Wassertheil-Smoller S, Sorlie PD, Stamler J. Prevalence of major cardiovascular risk factors and cardiovascular diseases among hispanic/latino individuals of diverse backgrounds in the United States. J Am Med Assoc 2012;308:1775–84. https://doi.org/10.1001/jama.2012.14517.

[c] Zaman MM, Bhuiyan MR, Karim MN, Moniruzzaman, Rahman MM, Akanda AW, Fernando T. Clustering of non-communicable diseases risk factors in Bangladeshis adults: an analysis of STEPS survey 2013. BMC Publ Health 2015;15:659. https://doi.org/10.1186/s12889-015-1538-4.

[25] Hammoudah AJ, Alhaddad IA, Khader Y, Tabbalat R, Al-Mousa E, Saleh A, Jarrah M, Nammak A, Izraiq M. Cardiovascular risk factors in Middle Eastern patients undergoing percutaneous coronary intervention: results from the first Jordanian percutaneous coronary intervention study. J Saudi Heart Assoc 2017;29:195–202. https://doi.org/10.1016/j.jsjah.2016.10.002.

[26] Mehta LS, Beckie TM, Devon HA, Grines CL, Krumholz HM, Johnson MN, Lindley KJ, Vaccarino V, Wang FY, Watson KE, Wenger NK. Acute myocardial infarction in women: a scientific statement from the American heart association. Circulation 2016;133:916–47. https://doi.org/10.1161/CIR.0000000000000351.

[27] Khademi N, Babanejad M, Asadmobini A, Karim H. The association of age and gender with risk factors of non-communicable diseases among employees in West of Iran. Int J Prev Med 2017;8:9–9. https://doi.org/10.4103/ijpvm.l.JJPVM_400_16.

[28] [a] Zindah M, Belbeisi A, Walke H, Mokdad AH. Obesity and diabetes in Jordan: findings from the behavioral risk factor surveillance system, 2004. Prev Chronic Dis 2008;5:A17. https://doi.org/10.18082006. PMID: 18082006 PMCID: PMCC2248793.

[b] Ajlouni K, Jaddou H, Batieha A. Diabetes and impaired glucose tolerance in Jordan: prevalence and associated risk factors. J Intern Med 1998a;244:317–23. https://doi.org/10.1046/j.1365-2796.1998.00369.x.

[c] Ajlouni K, Jaddou H, Batieha A. Obesity in Jordan. Int J Obes Relat Metab Disord 1998b;22:624–8. https://doi.org/10.1046/j.1365-2796.1998.00369.

[d] Arafan Q, Alissa ET, Alzoubi KH, Hammouri HM. Association of smoking with direct medical expenditures of chronic diseases in north of Jordan: a retrospective cohort study. BMJ Open 2019;9:e031143. https://doi.org/10.1136/bmjopen-2019-031143.

[e] Al-Rawashdeh A, Kasabri V, Bulatova N, Akour A, Zayed A, Momani M, Khawaja N, Bustanji H, Hyasat D. The correlation between plasma levels of oxytocin and beta-trophin in non-diabetic and diabetic metabolic syndrome patients: a cross sectional study from Jordan. Diabetes Metab Syndr 2017;11:59–67. https://doi.org/10.1016/j.dsx.2016.08.008.

[29] Mamudu HM, Paul TK, Wang L, Veeranki SP, Panchal HB, Alamian A, Sarnosky K, Budoff M. The effects of multiple coronary artery disease risk factors on subclinical atherosclerosis in a rural population in the United States. Prev Med 2016;68:140–6. https://doi.org/10.1016/j.ypmed.2016.04.003.

[30] Pandis N. Bias in observational studies. Am J Orthod Dentofacial Orthop 2014;145:542–3. https://doi.org/10.1016/j.ajodo.2014.01.008.