Prediction of first cardiovascular disease event in 2.9 million individuals using Danish administrative healthcare data: a nationwide, registry-based derivation and validation study

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Received 15 July 2021; editorial decision 30 July 2021; accepted 31 July 2021; online publish-ahead-of-print 2 August 2021
Handling editor: Karolina Szummer

Aims

The aim of this study was to derive and validate a risk prediction model with nationwide coverage to predict the individual and population-level risk of cardiovascular disease (CVD).

Methods and results

All 2.98 million Danish residents aged 30–85 years free of CVD were included on 1 January 2014 and followed through 31 December 2018 using nationwide administrative healthcare registries. Model predictors and outcome were pre-specified. Predictors were age, sex, education, use of antithrombotic, blood pressure-lowering, glucose-lowering, or lipid-lowering drugs, and a smoking proxy of smoking-cessation drug use or chronic obstructive pulmonary disease. Outcome was 5-year risk of first CVD event, a combination of ischaemic heart disease, heart failure, peripheral artery disease, stroke, or cardiovascular death. Predictions were computed using cause-specific Cox regression models. The final model fitted in the full data was internally-externally validated in each Danish Region. The model was well-calibrated in all regions. Area under the receiver operating characteristic curve (AUC) and Brier scores ranged from 76.3% to 79.6% and 3.3 to 4.4. The model was superior to an age-sex benchmark model with differences in AUC and Brier scores ranging from 1.2% to 1.5% and -0.02 to -0.03. Average predicted risks in each Danish municipality ranged from 2.8% to 5.9%. Predicted risks for a 66-year old ranged from 2.6% to 25.3%. Personalized predicted risks across ages 30–85 were presented in an online calculator (https://hjerteforeningen.shinyapps.io/cvd-risk-manuscript/).
Conclusion
A CVD risk prediction model based solely on nationwide administrative registry data provided accurate prediction of personal and population-level 5-year first CVD event risk in the Danish population. This may inform clinical and public health primary prevention efforts.

Graphical Abstract

Online calculator for individual risk

Nationwide population-level risk predictions

Keywords
Risk prediction • Risk stratification • Cardiovascular disease • Primary prevention • Registries • Nationwide

Introduction
Cardiovascular disease (CVD) causes immense morbidity, mortality, and economic burden worldwide. Predicted CVD risk has been used to inform treatment decisions for preventing CVD for several decades. Current European Society of Cardiology and American College of Cardiology/American Heart Association guidelines recommend opportunistic routine assessment of CVD risk among individuals without prior CVD starting from age 40, using the Systemic Coronary Risk Evaluation (SCORE) and Pooled Cohort Equations (PCE) models, respectively. These equations are derived from a range of cohorts, many recruited decades ago, and provide inferior risk prediction in modern era populations where the burden of CVD has reduced due to advances in treatment and prevention. Ideally, risk prediction models should be tailored to the target populations and reflect a contemporary distribution of CVD risk.

Large-scale administrative healthcare databases hold great potential for predictive risk modelling by providing data tailored to the target populations. An advantage of such prediction models is that they can be used beyond prediction of individual risk. They can provide countrywide, regionwide, municipality-wide, general practice-level, or even patient-list level risk predictions. This approach may guide allocation of resources and preventive efforts to high-risk areas and subpopulations. Furthermore, administrative data-based risk prediction models can be updated and recalibrated dynamically (i.e. at future time-points) and integrated into electronic health records to aid decision-making at the individual point of care. A further advantage of an administrative data-based risk prediction approach is the absence of clinical and laboratory variables. This eases use in community and primary care settings where clinical and laboratory risk factor levels may not be known and expands the concept of guideline-based opportunistic routine CVD risk assessment beyond the setting of a doctor’s consultation. As far as we are aware, only one other country (New Zealand) has developed country-wide CVD risk prediction equations based solely on administrative data and these equations were well calibrated with good risk discrimination in national, regional, and ethnic populations.

In the current study, we aimed to use Danish nationwide, routinely collected administrative data from more than 2.98 million individuals
without prior CVD to develop and validate a risk prediction model for the prediction of personal and population-level 5-year risk of first CVD event.

Methods

Study setting and data sources
Denmark is a country in Northern Europe of 42,933 km² divided into five geographical administrative regions encompassing 98 municipalities with a total population of 5,627,235 as of 1 January 2014 (Supplementary material online, Figure S1A). All Danish permanent residents have equal access to fully tax-funded healthcare and education. Utilization of these services is registered for administrative purposes through a unique personal Civil Registration Number. This number allows de-identified linkage of nationwide administrative registries at an individual level for research purposes. For the current study, we linked registries encompassing demographics, hospital contacts, redeemed prescriptions, educational attainment, and causes of death. Data on redeemed prescriptions and hospital contacts were available since 1995. Diagnostic codes used for the present study were classified according to International Classification of Diseases, Tenth revision (ICD-10) and drug prescriptions were classified according to the Anatomical Therapeutic Chemical (ATC) Classification System.

Study population
All Danish permanent residents aged 30–85 years alive on 1 January 2014 were included in the study. Exclusion criteria were: (i) previous ambulatory or in-hospital contact with a registered diagnosis of CVD (see Supplementary material online, Table S1 for further definitions) and (ii) missing data on educational attainment.

Outcome
The outcome of interest was first CVD event, defined as first occurrence of ischaemic heart disease (IHD), ischaemic stroke, haemorrhagic stroke, heart failure (HF), peripheral artery disease (PAD), or cardiovascular death. IHD, ischaemic stroke, haemorrhagic stroke, HF, and PAD were identified from hospital admissions using discharge diagnostic codes. Cardiovascular deaths were identified from death certificates listing a cardiovascular diagnosis among the causes of death. The cardiovascular diagnoses used in the outcome definition have been validated with high positive predictive values: IHD 88–97%, HF 80%, PAD 91–100%, and stroke 80–86%. See Supplementary material online, Table S1 for further descriptions and diagnostic codes.

Predictors
Predictors were pre-specified based on likelihood of being risk factors for CVD. They were selected based on inclusion in previous work by Mehta et al. in New Zealand and amended to suit availability of data in Danish administrative registries. The following variables were identified at baseline for the entire study population: age, sex, education, chronic obstructive pulmonary disease (COPD), and dispensing of smoking-cessation drugs, blood pressure-lowering drugs, glucose-lowering drugs, lipid-lowering drugs, and antithrombotic drugs. Levels of education were defined according to the highest level of education attained and classified as basic (e.g. primary school), secondary (e.g. high school or vocational training), tertiary (e.g. Bachelor’s degree), or postgraduate (e.g. Master’s degree). Smoking-cessation drug use and history of COPD were combined to one predictor as a proxy for smoking status. Dispensing of glucose-lowering drugs served as a proxy for the presence of either type 1 or type 2 diabetes, which was previously validated with a PPV of 95%. Metformin prescriptions redeemed by females <40 years of age were not included in the glucose-lowering drug definition, as the indication was presumed to be polycystic ovarian syndrome. Further descriptions along with ICD-10 and ATC codes were shown in Supplementary material online, Table S1.

Statistical analysis
Patient characteristics were presented as medians with interquartile ranges and frequencies with percentages. Study individuals were followed from study start (1 January 2014) until outcome of interest, non-cardiovascular death (competing risk), emigration, or 31 December 2018, whichever came first. Cause-specific Cox regression was used to predict the 5-year risk of first CVD event with non-cardiovascular death as a competing risk. The model included age (continuous, modelled by restricted cubic splines to allow for non-linear effects), sex (male/female), education (postgraduate, tertiary, secondary, or basic), smoking proxy (yes/no), glucose-lowering drug use (yes/no), blood pressure-lowering drug use (yes/no), lipid-lowering drug use (yes/no), antithrombotic drug use (yes/no), and interactions between age and glucose-lowering drug use, age and blood pressure-lowering drug use, and lipid-lowering drug use and glucose-lowering drug use, and antithrombotic drug use and glucose-lowering drug use. Interactions were pre-specified based on clinical plausibility and recently developed CVD risk prediction equations from New Zealand. We show results of complete case analyses where subjects with missing information on education are excluded. We presented personalized predicted 5-year risks of first CVD event across ages 30–85 in low-risk and intermediate-risk scenarios with and without each predictor. We presented personalized predicted 5-year risks of first CVD event for a 66-year-old individual to represent a common intermediate- to high-risk person in our population. Finally, we presented average predicted 5-year risk of first CVD event for all individuals without previous CVD aged 30–85 and 66 years, separately, in each Danish municipality.

The final model was developed in the full study population. We performed internal–external validation of the final model by assessing performance in each of the five administrative Regions (Supplementary material online, Figure S1A). We also assessed model performance in separate age groups. To further validate our model, we split our data into a training set and a testing set based on geography. The testing set encompassed all study participants living in the Capital Region of Denmark and the four remaining administrative Regions served as the training set (Supplementary material online, Figure S1B). In addition, we performed internal validation by splitting the data randomly (training set 63%, testing set 37%). We fitted the model in the training sets and assessed predictive performance in the testing sets. To assess discrimination, areas under the receiver operating characteristics curve (AUC) were calculated. To assess overall model performance, Brier scores were calculated. We compared model performance to a benchmark model containing only age and sex and reported differences in AUCs and Brier scores. Data management and statistical analyses were performed using R version 3.6.1.

Ethics
Studies based on pseudonymized registry data do not require ethical approval in Denmark. The data responsible institution (Capital Region of Denmark) approved the current study (approval number P-2019-537).
Results

On 1 January 2014, the Danish population comprised 3 598 511 persons aged 30–85 years. After applying the exclusion criteria of history of CVD (408 529, 11.4%) and missing information on education (206 875, 5.7%), 2 983 107 individuals were included in the final study population. Median ages were 52 and 50 years for females and males, respectively (Table 1, see Supplementary material online, Figure S2 for the age distribution of the study population). Approximately a quarter of the population had basic education as their highest level of education with a higher proportion among the older age groups (Supplementary material online, Figure S3). Percentages of other predictors ranged from 4.5% (females, glucose-lowering drug use) to 22.6% (females, blood pressure-lowering drug use). Presence of all predictors increased with age, except for male sex (decreased with age) and lipid-lowering drug use (peaked among individuals in their seventies) (Supplementary material online, Figure S3). In the overall study population, 119 740 (4.0%) developed a first CVD event during a median follow-up of 2.5 years. The median age at baseline for those who had a CVD event was 66 years (Supplementary material online, Table S2). IHD was the most frequently recorded component of the combined outcome of first CVD event, followed by ischaemic stroke (Supplementary material online, Figure S4).

Low-risk and intermediate-risk scenarios

To illustrate the impact of each predictor on risk of first CVD event, we showed predicted risks across ages 30–85 years in low-risk scenarios, adding each predictor, and intermediate-risk scenarios, adding or removing each predictor (Figure 1). A low-risk scenario (Figure 1A), i.e. a combination of predictors resulting in low expected risk, was defined as a female with postgraduate education without any other predictors. An intermediate-risk scenario (Figure 1B), i.e. a combination of predictors resulting in an intermediate expected risk, was defined as a male with secondary education, smoking proxy, and lipid-lowering drug use. The number of individuals in the study population fulfilling the low-risk scenario and the intermediate-risk scenario criteria are shown in Supplementary material online, Figure S5. Glucose-lowering drug use, blood pressure-lowering drug use, smoking proxy, male sex, antithrombotic drug use, and lower education predicted a higher 5-year risk of first CVD event. Lipid-lowering drug use predicted a higher risk among younger individuals and a lower risk among those above 52 years of age. For comparison with other populations and CVD prediction models, adjusted hazard ratios for first CVD event, with age as a two-level categorical variable, were presented (Supplementary material online, Figure S6).

| Table 1 Characteristics and follow-up of the study population |
|---------------------------------------------------------------|
| **Female (n = 1 558 026)**  | **Male (n = 1 425 081)** |
| Age (years), median [IQR] | 52 [41, 64]  | 50 [41, 62] |
| Education |  |
| Postgraduate | 134 124 (8.6)  | 152 807 (10.7)  |
| Tertiary | 418 575 (26.9)  | 270 467 (19.0)  |
| Secondary | 612 468 (39.3)  | 673 176 (47.2)  |
| Basic | 392 859 (25.2)  | 328 631 (23.1)  |
| Smoking proxy | 114 185 (7.3)  | 93 779 (6.6)  |
| Glucose-lowering drugs | 69 998 (4.5)  | 79 388 (5.6)  |
| Blood pressure-lowering drugs | 352 783 (22.6)  | 264 452 (18.6)  |
| Lipid-lowering drugs | 178 623 (11.5)  | 151 037 (10.6)  |
| Antithrombotic drugs | 80 540 (5.2)  | 75 749 (5.3)  |
| Follow-up |  |
| Time (years), median [IQR] | 5 [5, 5]  | 5 [5, 5]  |
| CVD event | 49 545 (3.2)  | 70 195 (4.9)  |
| Non-fatal event |  |
| Heart failure | 5857 (0.4)  | 7565 (0.5)  |
| Acute coronary syndrome | 15 319 (1.0)  | 28 711 (2.0)  |
| Ischaemic stroke | 13 034 (0.8)  | 16 090 (1.1)  |
| Transient ischaemic attack | 5248 (0.3)  | 5612 (0.4)  |
| Haemorrhagic stroke | 3659 (0.2)  | 3309 (0.2)  |
| Peripheral artery disease | 1766 (0.1)  | 3271 (0.2)  |
| Death |  |
| CV | 9564 (0.6)  | 11 251 (0.8)  |
| Non-CV | 60 965 (3.9)  | 63 383 (4.4)  |
| Time to CVD event (years), median [IQR] | 2.5 [1.3, 3.8]  | 2.5 [1.3, 3.8]  |

CV, cardiovascular; CVD, cardiovascular disease; IQR, interquartile range.
To illustrate the personalized predicted 5-year risk of first CVD event for a given individual, we showed predicted risks for a 66-year old with all possible combinations of predictors (Figure 2). The highest predicted risk for a 66-year old was 25.3% for a male with basic education, glucose-lowering drug use, blood pressure-lowering drug use, smoking proxy, and antithrombotic drug use. The lowest predicted risk for a 66-year old was 2.6% for a female with postgraduate education and lipid-lowering drug use. Predicted 5-year risks of first CVD event in the overall population ranged from 0.17% to 38.4%. The most common predictor combinations, excluding the low-risk scenario, were male sex, basic education, smoking proxy = yes, and lipid-lowering drugs = yes. BP, blood pressure; CVD, cardiovascular disease.

**Figure 1** Predicted 5-year risk of first CVD event across ages 30–85 in low-risk and intermediate-risk scenarios with and without each predictor. The low-risk scenario (A) was defined as female sex, postgraduate education, and absence of all other predictors. The intermediate-risk scenario (B) was defined as male sex, basic education, smoking proxy = yes, and lipid-lowering drugs = yes. BP, blood pressure; CVD, cardiovascular disease.

**Personalized risk predictions**
To illustrate the personalized predicted 5-year risk of first CVD event for a given individual, we showed predicted risks for a 66-year old with all possible combinations of predictors (Figure 2). The highest predicted risk for a 66-year old was 25.3% for a male with basic education, glucose-lowering drug use, blood pressure-lowering drug use, smoking proxy, and antithrombotic drug use. The lowest predicted risk for a 66-year old was 2.6% for a female with postgraduate education and lipid-lowering drug use. Predicted 5-year risks of first CVD event in the overall population ranged from 0.17% to 38.4%. The most common predictor combinations, excluding the low-risk scenario, were male sex, smoking proxy, and secondary education in the lowest age groups, and female sex, blood pressure-lowering drug use, and basic education in the highest age groups (Supplementary material online, Figure S7). To estimate the personalized predicted 5-year risk of first CVD event at any age, for any combination of predictors, we provided an online risk calculator (https://hjorteforenin.gen.shinyapps.io/cvd-risk-manuscript/).
Average risk in municipalities

We showed average predicted risks of first CVD event in each of the 98 Danish municipalities overall and for 66-year olds (Figure 3). The average predicted municipality risk ranged from 2.8% (Copenhagen municipality) to 5.9% (Laesoe municipality) (Figure 3A). The average predicted risk of first CVD event among 66-year olds ranged from 5.9% (Rudersdal municipality) to 7.5% (Vesthimmerland municipality) (Figure 3B).

Validation

The model was well-calibrated and had good discrimination in all five regions with AUCs ranging from 76.3% to 79.6% (Figure 4). Brier scores ranged from 3.3 to 4.4. Visual inspections of the calibration plots showed a small underestimation of risk from decile five in the Capital Region of Denmark and Region Zealand, whereas risk was slightly overestimated in all deciles in the remaining three regions (Figure 4). The model performed better than a benchmark model containing only age and sex in all five regions. Differences in AUC ranged from 1.2% to 1.5% and differences in Brier scores ranged from -0.02 to -0.03. Characteristics of the populations in each Region are shown in Supplementary material online, Table S3. Geographical and random split validation showed similarly good model performance (Supplementary material online, Table S4 and Supplementary material online, Figure S8). Model validation in subgroups by age bands showed good calibration in most age bands. Calibration was suboptimal in the oldest age band (80–85 years) with overestimation of risk in the lower deciles and underestimation in the two highest deciles (Supplementary material online, Figure S9).

Discussion

We developed and validated a novel risk prediction model for estimation of the 5-year risk of first CVD event in 2.98 million Danish residents using administrative data from nationwide population-based registries. Predictors included in the model were age, sex, education, glucose-lowering drug use, blood pressure-lowering drug use, antithrombotic drug use, lipid-lowering drug use, and a smoking proxy. The model was well-calibrated in geographical regions and age bands. We provided examples of the utility of our model for prediction of personalized and population-level risk. We created a web...
needed, because prediction models based on older cohorts are likely to overestimate risk in modern day populations, as treatment advances and changes in risk factors in recent decades have led to a lower incidence and mortality of CVD.25–29

A drawback of using administrative data to fit risk prediction models is the lack of laboratory variables and more specific clinical data for personalized risk prediction. The original forms of the most widely known risk prediction models in preventive cardiology, e.g. SCORE,30 Framingham Risk Score,31 and PCE,32 typically incorporate lipid levels and blood pressure. A benefit of that approach is that these, as well as other modifiable CVD risk factors, are directly accounted for in the models. Thus, they present a clear target for risk factor modification, and changes in risk can be communicated directly by the clinician to the patient, e.g. by showing predicted risks at lower lipid levels or blood pressures. However, laboratory and clinical variables do not necessarily result in more accurate CVD risk prediction per se, as demonstrated in previous studies directly comparing models with and without laboratory measures,7,31,33 since non-modifiable factors such as sex, age, and sociodemographic factors may capture up to 80% of the prognostic performance in cardiovascular risk models.34 The inclusion of variables such as lipid levels would preclude the use of our model for population-level risk predictions, as such variables are not routinely collected at the population level. Furthermore, the disadvantages of only including universally available variables in our administrative data approach are outweighed by the absence of several weaknesses associated with conventional approaches to risk prediction modelling, namely, nonrepresentative study samples, few events in predictor combination strata, and many variables with missing data. In addition, our web calculator for personalized risk prediction is mainly intended for use in the community setting where laboratory and clinical risk factor levels may not be known. End-users of our online risk calculator are alerted of their cardiovascular risk and prompted to pursue individual targeting of their modifiable risk factors.

We followed the Transparent Reporting of a Multivariable Prediction Model for Individual Prognosis or Diagnosis and The REporting of studies Conducted using Observational Routinely collected health Data (RECORD) Statement recommendations for developing our novel risk prediction model.35,36 As such, we prespecified predictors for inclusion in our model that were known risk factors for CVD and were commonly used in previous risk scores, albeit modified to suitability in our administrative databases. We chose a parsimonious number of predictors for our model to reduce complexity and avoid overfitting.

Sex, age, glucose-lowering drug use, blood pressure-lowering drug use, and a smoking proxy were included to serve as proxies for well-documented predictors of CVD.37 Socioeconomic position is increasingly recognized as an important predictor of poor health outcomes, and recently developed CVD risk models such as QRISK3, PREDICT, and the models developed by Mehta et al. incorporated socioeconomic measures.5,6,10 We chose education as a simple measure of socioeconomic position. As discussed, improvements in preventive treatment have reduced the risk of CVD in recent decades, and Danish patients are increasingly prescribed antithrombotic and lipid-lowering drugs for

All Danish residents are assigned a Civil Registration Number at birth or immigration, by which demographic data and healthcare service usage data is recorded. Hence, the 2.98 million participants in this study encompassed virtually the entire Danish population aged 30–85. We had complete data on all predictors, except for education, and all outcome variables as well as minimal censoring. This provided an ideal data set for derivation and validation of a CVD risk prediction model for the Danish population. That was the main strength of our study, as it eliminated issues with generalizability and external validity, although generalizability to other countries may be limited. However, other countries or regions, with databases similar to those described in the present study, can develop CVD risk prediction models tailored specifically to their populations by utilizing our approach. Contemporary or updated risk prediction models are
primary prevention of CVD. To account for this, we included lipid-lowering and antithrombotic drug use at baseline in our model. Notably, in our data, we lacked information on body mass index and lifestyle factors such as physical activity-level, which are known and important modifiable risk factors for CVD. In spite of this limitation, our model showed good predictive performance, since sociodemographic factors (i.e. age and education) are surrogates for exposure to CVD risk factors, such as obesity, throughout the lifespan. Our approach to variable selection can be replicated to identify candidate predictors in other databases with different data structure and population characteristics.

The main outcome differed from SCORE, which is the currently recommended CVD risk prediction tool in Denmark. We chose 5 years rather than 10 years as our prediction horizon, which enabled us to develop our prediction model in a contemporary (2014) cohort with 5 years follow-up and included non-fatal CVD events in our composite endpoint. Fatal CVD does not adequately capture CVD events, as mortality following CVD has decreased during the last decades, especially in high-income European countries. We included HF and haemorrhagic stroke as CVD events, as they lead to high morbidity and largely share the same risk factors as atherosclerotic CVD. The direction of the associations between predictors and the outcome were as expected.

Figure 4 Calibration plots and discrimination metrics across the five regions. The predicted risk is plotted in deciles against the estimated actual risk. Differences in AUC and Brier scores compared to an age-sex benchmark model is presented as delta-AUC and delta-Brier. AUC, area under the receiver operating characteristic curve.
Our smoking proxy consisting of either smoking-cessation drug use or COPD has not been validated. The sensitivity for identifying individuals who smoked was likely low, as the prevalence of our smoking proxy was only 7%, whereas the daily smoking prevalence in Denmark has been reported to be 17%. However, we presume that the PPV of our smoking proxy was high as smoking-cessation drugs are not approved for any other indication and previous Danish population-based studies found that 78% of COPD patients were smokers. The hazard ratios for our smoking proxy were similar to those reported in previous multivariable CVD risk prediction models. Lipid-lowering drug use predicted a reduced risk of CVD in older age groups and an increased risk in younger age groups. This may be because lipid-lowering treatment was prescribed to younger patients with a high-risk indication, whereas the reduced risk in the older age groups reflected the expected treatment benefit. Antithrombotic drug use predicted an overall increased risk, as they were only indicated for individuals with an elevated CVD risk. Similarly, blood pressure-lowering drug use predicted a higher risk of first CVD event overall, but the risk was attenuated in middle-aged and older age groups, which likely reflects a more severe indication for treatment in the younger age groups. Newer glucose-lowering drugs with benefits on cardiovascular outcomes were not widely used at the time of our cohort inclusion in 2014. Thus, our model should be adjusted in the forthcoming years to reflect the change in risk distribution as these drugs become more common in the treatment of type 2 diabetes. The ability to easily adjust the model prospectively (e.g. as done in the UK with QRISK) highlights a strength of the present approach.

We wanted to have a study population that was as inclusive as possible for population-level CVD risk prediction and, therefore, chose to include a broad age range (30–85 years). Nonetheless, previous studies found that CVD risk prediction models developed in mainly young and middle-aged persons predicted individual risk in older age groups poorly, partly due to the competing risk of non-cardiovascular death. We handled the competing risk in our statistical modelling. Yet, our model had suboptimal calibration in the 80–85-year age band, which warrants a more cautious interpretation of the risk predictions presented for this age group.

The municipality-level risk predictions that we provided were an example of how our model can be applied beyond prediction of individual risk. The models developed by Mehta et al. have also recently been used to identify quality improvement opportunities in the utilization of cardiovascular preventive pharmacotherapy across a country (New Zealand) and in sub-populations. Identifying high-risk areas and subpopulations, may guide public health-level interventions such as allocation of resources and targeted preventative efforts.

Conclusion

A CVD risk prediction model based solely on nationwide administrative registry data provided accurate prediction of 5-year first CVD event risk in the entire Danish population. We supplied examples of both personal and population-level use of the model. The model can be used to facilitate community-based, clinical, and public health-level primary prevention. An online risk calculator based on our risk prediction model is freely available (https://hjerteforeningen.shinyapps.io/cvd-risk-manuscript/).

Lead author biography

Daniel Mølager Christensen is a medical doctor with interests in cardiology, epidemiology, and medical risk prediction. He is currently a PhD-fellow at the Danish Heart Foundation conducting research using the Danish national healthcare registries.

Supplementary material

Supplementary material is available at European Heart Journal Open online.

Funding

The present study was fully supported by a research grant from the Danish Heart Foundation (grant identification number 20-R146-A9798).

Conflicts of interest: E.F. reported an independent research grant unrelated to the current research from The Novo Nordisk Foundation. UK reported speaker’s honorarium from Novo, Novartis, Astra Zeneca, and Boehringer. C.T.P. reported research grants from Bayer and Novo Nordisk not related to this study. All other authors had no conflicts of interest to disclose.

Data availability statement

No additional data are available as access to Danish registry data is granted on an individual basis by the relevant authorities.

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