An independent and external validation of QRISK2 cardiovascular disease risk score: a prospective open cohort study

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ABSTRACT

Objective To evaluate the performance of the QRISK2 score for predicting 10-year cardiovascular disease in an independent UK cohort of patients from general practice records and to compare it with the NICE version of the Framingham equation and QRISK1.

Design Prospective cohort study to validate a cardiovascular risk score.

Setting 365 practices from United Kingdom contributing to The Health Improvement Network (THIN) database.

Participants 1.58 million patients registered with a general practice between 1 January 1993 and 20 June 2008, aged 35-74 years (9.4 million person years) with 71 465 cardiovascular events.

Main outcome measures First diagnosis of cardiovascular disease (myocardial infarction, angina, coronary heart disease, stroke, and transient ischaemic stroke) recorded in general practice records.

Results QRISK2 offered improved prediction of a patient’s 10-year risk of cardiovascular disease over the NICE version of the Framingham equation. Discrimination and calibration statistics were better with QRISK2. QRISK2 explained 33% of the variation in men and 40% for women, compared with 29% and 34% respectively for the NICE Framingham and 32% and 38% respectively for QRISK1. The incidence rate of cardiovascular events (per 1000 person years) among men in the high risk group was 27.8 (95% CI 27.4 to 28.2) with QRISK2, 21.9 (21.6 to 22.2) with NICE Framingham, and 24.8 (22.8 to 26.9) with QRISK1. Similarly, the incidence rate of cardiovascular events among women in the high risk group was 24.3 (23.8 to 24.9) with QRISK2, 20.6 (20.1 to 21.0) with NICE Framingham, and 21.8 (18.9 to 24.6) with QRISK1.

Conclusions QRISK2 is more accurate in identifying a high risk population for cardiovascular disease in the United Kingdom than the NICE version of the Framingham equation. Differences in performance between QRISK2 and QRISK1 were marginal.

INTRODUCTION

Cardiovascular disease is an important health concern globally, with just under a third of all deaths attributed to cardiovascular disease in 2004 (www.who.int, fact sheet No 317). In the United Kingdom, there are almost 200 000 deaths each year relating to diseases of the heart and circulatory system, with more than one in three deaths associated with cardiovascular disease (www.heartstats.org). General practitioners need an accurate and reliable tool to help them identify patients at high risk of having a cardiovascular event. Numerous multivariable risk scores have been developed to estimate a patient’s 10 year risk of cardiovascular disease based on certain key known risk factors, including the Framingham risk score and the Reynolds risk score, both developed using patient data from US, the SCORE system using patients from multiple European countries, and ASSIGN using patients from Scotland. In the United Kingdom, until recently the National Institute for Health and Clinical Excellence (NICE) recommended use of the long established Framingham equation to inform a patient treatment plan about cardiovascular risk. NICE have now ceased recommending any single risk score equation, leaving healthcare professionals to choose the tool they consider to be most appropriate.

The QRESEARCH organisation of the University of Nottingham, is a model to predict the 10 year risk of developing cardiovascular disease. It was developed and validated on large general practice databases in the United Kingdom using data from three million patients with 17.5 million person years of observation. QRISK1 includes traditional risk factors such as age, sex, systolic blood pressure, smoking status, and total serum cholesterol:high density lipoprotein ratio that are included in the long established Framingham equations, but it also includes body mass index, family history of cardiovascular disease, social deprivation (Townsend score), and use of antihypertensive treatment. Performance data comparing QRISK1 with the Framingham equation indicated that QRISK1 is a more accurate tool to predict the development of cardiovascular disease in the United Kingdom. QRISK2, the successor to QRISK1, is a new multivariable risk score that contains all the risk factors that are in QRISK1 but also includes self assigned ethnicity
Summary of risk factors in QRISK1, QRISK2, and Framingham equation

**QRISK1**
- Age (continuous)
- Ratio of total serum cholesterol:high density lipoprotein (continuous)
- Systolic blood pressure (continuous)
- Smoking status (current smoker/non-smoker [including former smoker])
- Body mass index (continuous)
- Family history of coronary heart disease in first degree relative under 60 years (yes/no)
- Townsend deprivation score (output area level 2001 census data evaluated as continuous variable)
- Receiving treatment for blood pressure at baseline (at least one current prescription of at least one antihypertensive agent) (yes/no)
- Systolic blood pressure × Receiving treatment for blood pressure at baseline

**QRISK2**
- Age (continuous)
- Ratio of total serum cholesterol:high density lipoprotein (continuous)
- Systolic blood pressure (continuous)
- Smoking status (current smoker/non-smoker [including former smoker])
- Body mass index (continuous)
- Family history of coronary heart disease in first degree relative under 60 years (yes/no)
- Townsend deprivation score (output area level 2001 census data evaluated as continuous variable)
- Treated hypertension (diagnosis of hypertension and at least one current prescription of at least one antihypertensive agent) (yes/no)
- Self assigned ethnicity (white [or not recorded]/Indian/Pakistani/Bangladeshi/other Asian/black African/black Caribbean/other [including mixed])
- Type 2 diabetes (yes/no)
- Rheumatoid arthritis (yes/no)
- Atrial fibrillation (yes/no)
- Renal disease (yes/no)
- Age × body mass index
- Age × Townsend score
- Age × systolic blood pressure
- Age × family history of cardiovascular disease
- Age × smoking current
- Age × treated hypertension
- Age × type 2 diabetes
- Age × atrial fibrillation

**Framingham equation (version recommended by NICE)**
- Age (continuous)
- Ratio of total serum cholesterol:high density lipoprotein (continuous)
- Systolic blood pressure (continuous)
- Smoking status (current smoker [or quit within last year]/non-smoker)
- Sex (male/female)
- Left ventricular hypertrophy (yes/no)
- Type 2 diabetes (yes/no)
- Age × type 2 diabetes
- Left ventricular hypertrophy × age
- Age × sex

and conditions associated with cardiovascular risk (including diagnosed type 2 diabetes, treated hypertension, rheumatoid arthritis, renal disease, and atrial fibrillation). QRISK2 also contains interactions between age and Townsend score, body mass index, systolic blood pressure, family history, smoking status, treated hypertension, diagnosis of type 2 diabetes, and atrial fibrillation (see box for variables included in QRISK1, QRISK2 and NICE Framingham). All continuous risk factors were carefully handled and kept continuous throughout the model building process, fractional polynomials were used to model nonlinear risk relations where appropriate. Technical details of the actual QRISK2 model can be found on the QRESEARCH website (www.qresearch.org).

After the development of a new multivariable risk score, it needs to be subjected to external validation. The performance of a risk score is typically overestimated in the original data used to develop the score. External validation is a crucial step to provide sufficient evidence about the performance of the risk score on a cohort not used in the score’s development. It is disappointing, however, that most risk scores that are published fail to undergo this extra step. Studies that develop and validate risk scores are far too often of low quality, poorly designed, use inappropriate statistical techniques, and are based on small selective cohorts with insufficient numbers of events. Furthermore, it is likely that many of the risk scores published each year are opportunistically produced to maximise the output from a clinical study for which developing a risk score was not declared a priori before collecting the data. Guiding efficient clinical decision making requires that the assessment of risk be accurate, yet many risk scores in use have not adequately shown this essential quality. Low quality risk scores may be used clinically to stratify patients into risk groups or as inclusion or exclusion criteria for randomised controlled trials.

This article describes the results from an independent and external validation of QRISK2 and compares the performance of QRISK2 against QRISK1 and an adjusted version of the Framingham equation previously recommended by NICE.

**METHODS**

**Study population**

Study participants were patients registered between 1 January 1993 and 20 June 2008 and recorded on the THIN database (www.thin-uk.com). Patients were excluded if they had a prior diagnosis of cardiovascular disease, had invalid dates or invalid recorded risk factor values out of plausible range, were under the age of 35 years, were aged 74 years or over, had missing Townsend scores (social deprivation), or were prescribed statins at baseline.

**Cardiovascular disease outcomes**

The primary outcome measure was the first diagnosis of cardiovascular disease (myocardial infarction, angina, coronary heart disease, stroke, and transient
Table 1 | Characteristics of patients from THIN database who were included in study. Values are numbers (percentages) unless stated otherwise

| Country                  | Women | Men  |
|--------------------------|-------|------|
| England                  | 688 015 (86.29) | 678 375 (86.34) |
| Scotland                 | 43 276 (2.77) | 42 471 (5.41) |
| Wales                    | 44 033 (5.43) | 44 065 (5.61) |
| Northern Ireland         | 22 049 (5.52) | 20 822 (2.65) |

| Ethnicity                | Women | Men  |
|--------------------------|-------|------|
| White or not recorded    | 780 117 (97.84) | 769 409 (97.92) |
| Indian                   | 4007 (0.50) | 3928 (0.50) |
| Pakistani                | 1138 (0.15) | 1181 (0.15) |
| Bangladeshi              | 356 (0.04)  | 447 (0.06)  |
| Other Asian              | 1953 (0.24) | 1900 (0.24) |
| Black Caribbean          | 2270 (0.28) | 1762 (0.22) |
| Black African            | 2921 (0.37) | 2728 (0.35) |
| Chinese                  | 782 (0.10)  | 615 (0.08)  |
| Other including mixed    | 3829 (0.48) | 3763 (0.48) |
| Mean (SD) systolic blood pressure (mm Hg) | 131.54 (20.73) | 135.22 (19.01) |
| Mean (SD) total serum cholesterol concentration (mmol/l) | 5.76 (1.16) | 5.58 (1.12) |
| Mean (SD) HDL cholesterol (mmol/l) | 1.58 (0.43) | 1.31 (0.35) |
| Mean (SD) total serum cholesterol:HDL ratio | 3.91 (1.25) | 4.54 (1.38) |
| Mean (SD) body mass index (kg/m²) | 26.27 (4.99) | 26.71 (4.11) |
| Positive family history of coronary heart disease | 38 562 (4.84) | 31 744 (4.04) |
| Current smoker           | 180 774 (22.67) | 215 650 (27.45) |
| Treated hypertension     | 50 758 (6.37) | 37 529 (4.78) |
| Type 2 diabetes          | 13 800 (1.73) | 18 470 (2.35) |
| Rheumatoid arthritis     | 8479 (1.06)  | 3555 (0.45)  |
| Atrial fibrillation      | 3035 (0.38)  | 4937 (0.63)  |
| Chronic kidney disease   | 1187 (0.15)  | 1087 (0.14)  |

Deprivation index (Townsend score) fifth:

| Category                  | Total patients |
|---------------------------|----------------|
| 1 (most affluent)         | 219 287 (27.50) |
| 2                         | 183 335 (22.99) |
| 3                         | 165 391 (20.74) |
| 4                         | 138 226 (17.34) |
| 5 (most deprived)         | 91 134 (11.43)  |

Incident CVD events in 10 years

| Total patients | 29 057 (3.64) |
|----------------|--------------|

Observed 10 year risk (95% CI) of CVD events

| Women | 5.89 (5.81 to 5.96) |
|-------|---------------------|
| Men   | 9.00 (8.90 to 9.10) |

Table 1 continued

| Characteristics                  | Women | Men  |
|----------------------------------|-------|------|
| Total type 2 diabetes            | 13 800 (1.73) | 18 470 (2.35) |
| Current smoker                   | 180 774 (22.67) | 215 650 (27.45) |
| Treated hypertension             | 50 758 (6.37) | 37 529 (4.78) |
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ischaemic stroke) recorded on the general practice clinical computer system.

Statistical analysis

Ten-year estimated cardiovascular disease risk for every patient in the THIN cohort was calculated using QRISK2. Observed 10-year cardiovascular disease risks were obtained using the method of Kaplan-Meier. Multiple imputation was used to replace missing values for body mass index, systolic blood pressure, total serum cholesterol:high density lipoprotein ratio, and smoking status. Multiple imputation is a powerful technique that offers substantial improvements over the biased and flawed value replacement approaches based on complete cases or cases matched for age and sex. It involves creating multiple copies of the data and imputing the missing values with sensible values randomly selected from their predicted distribution. We used the MICE (Multivariate Imputation by Chained Equations) library in R software to create 20 imputed datasets and then combined the results from analyses on each of the imputed datasets to produce estimates and confidence intervals that incorporate the uncertainty of imputed values.

Predictive performance of QRISK2 for the THIN cohort was assessed by examining measures of calibration and discrimination. Calibration refers to how closely the predicted 10 year risk of cardiovascular disease agrees with the observed 10 year risk. This was assessed for each tenth of predicted risk, ensuring 10 equally sized groups, and for each 5 year age band by calculating the ratio of predicted to observed risk of cardiovascular disease separately for men and for women. Calibration of the risk score predictions was assessed by plotting observed proportions versus predicted probabilities. The Brier score for censored survival data was also calculated, which is a measure of accuracy and is the average squared deviation between predicted and observed risk; a lower score represents higher accuracy.

Discrimination is the ability of the risk score to differentiate between patients who experience a cardiovascular event during the study and those who do not. This measure is quantified by calculating the area under the receiver operating characteristics curve (AUROC); a value of 0.5 represents chance, and 1 represents perfect discrimination. We also calculated the $D$ statistic and the $R^2$ statistic (derived from the $D$ statistic), which are measures of discrimination and explained variation respectively and are tailored towards censored survival data. Higher values of $D$ indicate greater discrimination, where an increase of 0.1 over other risk scores is a good indicator of improved prognostic separation. Scaled rectangular diagrams are presented to illustrate the discrimination performance of QRISK2 and NICE Framingham.

Although predicted risk varies across a continuum, clinical decisions require creation of risk groups. An important aspect, therefore, when considering adopting a new risk prediction rule is the classification of patients into high and low risk and the number of patients who would be reclassified to a different risk category when compared with the standard means of risk prediction (here the NICE Framingham equation). Patients were identified as high risk if their 10 year predicted cardiovascular disease risk was ≥20%, as per the guidelines set out by NICE.

We compared the performance of QRISK2 with its predecessor QRISK and with a modified version of the Framingham equation recommended by NICE. There is an increased risk of developing cardiovascular disease in people with a family history of premature disease, and risk varies between ethnic groups in the United Kingdom. Until recently, the approach recommended by NICE was to apply adjustment factors to
Table 2 | Crude and age adjusted incidence rates (per 1000 person years) for cardiovascular disease by sex, age, region of residence, and deprivation.

| Ethnic group | Age adjusted incidence rates (per 1000 person years) | Crude incidence rate (95% CI) |
|--------------|-----------------------------------------------------|-----------------------------|
| White or not recorded | 4909.8 | (95% CI) |
| Indian | 4195.1 | (95% CI) |
| Pakistani | 4393.6 | (95% CI) |
| Bangladeshi | 1326.0 | (95% CI) |
| Other Asian | 5745.6 | (95% CI) |
| Black Caribbean | 8150.6 | (95% CI) |
| Black African | 7476.2 | (95% CI) |
| Chinese | 1787.10 | (95% CI) |
| Other | 11299.9 | (95% CI) |

The Framingham equation for ethnicity and family history, which are not captured in the Framingham equation. The Framingham score is multiplied by 1.4 for South Asian men (no adjustment for South Asian women) and by 1.5 for people with a history of coronary heart disease in a first degree relative. For South Asian men with a family history of coronary heart disease in a first relative, both adjustments are applied. To date there has been no detailed published scientific validation and evaluation for the choice of adjustment factor.

All statistical analyses were carried out in R (version 2.9.1).

RESULTS

The THIN database system included 3,587,306 eligible patients registered at 382 practices in the United Kingdom. After sequentially excluding, as per the exclusion criteria, 625,454 patients who left the general practice before 1 January 1993, 1,071,612 aged <35 years or ≥74 years, 161,746 with missing Townsend scores, 86,093 with a prior diagnosis of cardiovascular disease, and 23,019 with prior statin use, the analysed cohort consisted of 1,583,106 patients registered between 1 January 1993 and 20 June 2008 at 365 general practices. The median follow-up was 6.2 years, and 241,859 (15.28%) patients were followed for at least 10 years. The 10 year observed risk of a cardiovascular event in men aged 35-74 years was 9.00% (95% confidence interval 8.90% to 9.10%) and in women was 5.89% (5.81% to 5.96%). Table 1 details the characteristics of eligible patients.

Complete data for all risk factors (ratio of total serum cholesterol to high density lipoprotein cholesterol, systolic blood pressure, body mass index, and smoking status) considered were available for 18.4% of women (n=146,651) and 16.0% of men (n=125,978). Most patients (n=1,066,926 (67.4%)) had none or only one missing risk factor. There were markedly high levels of missing data for the total serum cholesterol:high density lipoprotein cholesterol ratio (74.7% for women and 74.6% for men). For other risk factors, the levels of missing data were, for body mass index, 19.4% for women, 28.5% for men; for systolic blood pressure, 7.4% for women, 16.5% for men; and for smoking status, 20.1% for women, 29.2% for men.

Table 2 shows the incidence rates (per 1000 person years) for cardiovascular disease by age, sex, region of residence, and deprivation. In total there were 71,465 incident cases of cardiovascular disease...
during the study period from 9.4 million person years of observation. The age adjusted incidence rate (per 1000 person years) for cardiovascular disease was highest in Scottish women and men (7.97 and 12.04) and lowest in English women and men (5.75 and 9.89).

The incidence of cardiovascular disease varied widely between different ethnic groups. The age standardised rates (per 1000 person years) for the white reference group were 5.89 for women and 9.20 for men. The highest rates were among the South Asian groups—for example, in Pakistani women the rate was 13.22 (per 1000 person years) and in Pakistani men it was 19.55.

**Discrimination and calibration**

For an accurate risk score the predicted and observed risks will agree. Fig 1 shows calibration plots for the three risk scores. Both QRISK2 and its predecessor, QRISK1, show much better agreement between observed risk and predicted risk grouped by tenth of risk compared with the NICE adjusted Framingham equation.

Table 3 presents discrimination and calibration performance data for QRISK2, QRISK1, and the NICE adjusted Framingham equation. The $R^2$ statistic (percentage of explained variation) is approximately 5% higher for QRISK2 in both men and women compared with the NICE adjusted Framingham equation. The difference in $R^2$ between QRISK2 and QRISK1 is less pronounced, with QRISK2 explaining around only 1% more of the variation in outcome in both men and women. The $D$ discrimination statistic, where a higher score represents better discrimination, is higher for QRISK2 in both men and women compared with the NICE adjusted Framingham equation. As with the $R^2$ statistic, the difference in the $D$ statistics between QRISK2 and QRISK1 is small. The Brier score (adjusted for censoring), a measure of prediction accuracy, was lower for QRISK2 in both men and women compared with the NICE adjusted Framingham equation. There was no difference between QRISK2 and QRISK1 in Brier score (0.076).
Thus for both men and women, the mean predicted risks in patients reclassified from high to low risk (and vice versa) with QRISK2 were more accurate compared with the mean observed risk than mean predicted risks with NICE Framingham (see online appendix for low, intermediate, and high risk classification). Those patients who were reclassified as high risk with QRISK2 from low risk with NICE Framingham tended to be older, have been treated for hypertension, have a family history of coronary heart disease, and have a diagnosis of type 2 diabetes, rheumatoid arthritis, or atrial fibrillation.

The proportion of men and women classified as high risk by QRISK2 and NICE Framingham who had a subsequent cardiovascular event are displayed in fig 2, a scaled rectangular diagram. At current recommended treatment thresholds of 20%, the figure shows the modest discrimination performance of both QRISK2 and the NICE Framingham model. With QRISK2, 14% of the male cohort would be identified as being at high risk and would capture 40% of the cardiovascular events; NICE Framingham would identify 22% of the male cohort and 54% of the cardiovascular events. Similarly, for women, QRISK2 would identify 6% of the cohort as being at high risk and 26% of the cardiovascular events, whereas NICE Framingham would identify 7% of the cohort as high risk and 26% of all cardiovascular events.

Incidence of cardiovascular events in high risk groups
Using the 20% threshold to identify high risk patients, QRISK2 identified a group of patients at a higher risk of cardiovascular events than those identified with NICE Framingham. The incidence rate of cardiovascular events among men designated high risk with QRISK2 was 27.8 per 1000 person years (95% confidence interval 27.4 to 28.2), whereas it was 21.9 (21.6 to 22.2) with NICE Framingham and 24.8 (22.8 to 26.9) with QRISK1. For women, the incidence rate of cardiovascular events in those designated high risk was 24.3 (23.8 to 24.9) with QRISK2, 20.6 (20.1 to 21.0) with NICE Framingham, and 21.8 (18.9 to 24.6) with QRISK1. Table 6 shows regional variations in the incidence of cardiovascular events in high risk groups.

**DISCUSSION**
Principal findings
We independently evaluated the performance of the QRISK2 risk score, in comparison with the risk prediction approach (NICE Framingham) that was until recently recommended by the National Institute for Health and Clinical Excellence in the United Kingdom, for predicting 10-year cardiovascular disease in

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Table 4 | Comparison of QRISK2 and NICE Framingham (version of the Framingham equation recommended by NICE) in classification of men in the THIN cohort into low or high 10-year risk of cardiovascular events and observed and predicted risk

| QRISK2 | Low risk (<20%) | High risk (≥20%) | Total | No (%) of men reclassified |
|--------|-----------------|-----------------|-------|---------------------------|
| NICE Framingham, low risk (<20%) | | | | |
| No (range) of men | 599 179 (581 620–617 650) | 11 321 (10 311–12 409) | 610 491 (592 304–630 059) | 11 231 (1.8) |
| No (range) of events | 18 258 (16 692–19 744) | 1335 (1148–1507) | 19 593 (17 845–21 251) | |
| Observed risk (95% CI) | 5.32 (4.82 to 5.81) | 20.02 (16.98 to 23.06) | 5.57 (5.04 to 6.11) | |
| Mean risk QRISK2 (95% CI) | 5.28 (5.23 to 5.34) | 24.08 (23.72 to 24.44) | 5.62 (5.54 to 5.72) | |
| Mean risk NICE (95% CI) | 8.64 (8.39 to 8.90) | 14.73 (14.35 to 15.11) | 8.76 (8.50 to 9.01) | |
| NICE Framingham, high risk (≥20%) | | | | |
| No (range) of men | 79 592 (69 790–88 184) | 95 731 (85 884–105 524) | 175 323 (155 674–193 429) | |
| No (range) of events | 7125 (6470–7549) | 15 690 (14 410–17 014) | 22 815 (21 157–24 563) | |
| Observed risk (95% CI) | 14.00 (12.28 to 15.72) | 24.52 (22.74 to 26.31) | 19.76 (17.97 to 21.55) | |
| Mean risk QRISK2 (95% CI) | 14.98 (14.85 to 15.11) | 28.87 (28.45 to 29.29) | 22.57 (22.22 to 22.92) | |
| Mean risk NICE (95% CI) | 25.14 (24.75 to 25.54) | 35.47 (33.59 to 37.36) | 30.79 (29.60 to 31.97) | |
| Total | | | | |
| No (range) of men | 678 771 (669 804–687 440) | 106 962 (98 292–115 929) | 785 733 | |
| No (range) of events | 25 383 (24 241–26 491) | 17 025 (15 917–18 167) | 42 408 | |
| Observed risk (95% CI) | 6.43 (6.12 to 6.74) | 24.12 (22.68 to 25.56) | 9.0 (8.9 to 9.1) | |
| Mean risk QRISK2 (95% CI) | 6.42 (6.21 to 6.63) | 28.37 (27.89 to 28.85) | 9.41 (8.88 to 9.94) | |
| Mean risk NICE (95% CI) | 10.58 (10.02 to 11.14) | 33.30 (31.21 to 35.39) | 13.68 (12.63 to 14.72) | |
Table 5 | Comparison of QRISK2 and NICE Framingham (version of the Framingham equation recommended by NICE) in classification of women in the THIN cohort into low or high 10-year risk of cardiovascular events and observed and predicted risk

| QRISK2 | Low risk (<20%) | High risk (≥20%) | Total | No (%) of women reclassified |
|--------|----------------|-----------------|-------|-----------------------------|
| NICE Framingham, low risk (<20%) | | | | |
| No (range) of women | 722 367 (716 767–729 126) | 15 748 (14 802–16 926) | 738 115 (733 041–744 762) | |
| No (range) of events | 19 242 (18 390–19 865) | 2141 (2063–2260) | 21 383 (20 513–21 962) | |
| Observed risk (95% CI) | 4.39 (4.15 to 4.63) | 20.07 (18.84 to 21.30) | 4.74 (4.51 to 4.97) | 15 748 (2.1) |
| Mean risk QRISK2 (95% CI) | 4.48 (4.38 to 4.58) | 23.70 (23.61 to 23.79) | 4.89 (4.76 to 5.03) | |
| Mean risk NICE (95% CI) | 5.56 (5.38 to 5.74) | 15.19 (14.96 to 15.41) | 5.76 (5.57 to 5.96) | |
| NICE Framingham, high risk (≥20%) | | | | |
| No (range) of women | 25 478 (22 188–28 664) | 33 780 (30 423–26 205) | 59 258 (52 611–64 332) | |
| No (range) of events | 2346 (2113–2584) | 5328 (4888–6000) | 7674 (7095–8544) | 25 478 (43.0) |
| Observed risk (95% CI) | 13.36 (10.72 to 16.00) | 22.36 (18.94 to 25.78) | 18.53 (15.46 to 21.60) | |
| Mean risk QRISK2 (95% CI) | 15.12 (15.15 to 15.48) | 28.19 (27.79 to 28.59) | 22.66 (22.05 to 23.26) | |
| Mean risk NICE (95% CI) | 24.24 (23.89 to 24.58) | 30.97 (30.17 to 31.76) | 28.07 (27.51 to 28.64) | |
| Total | | | | |
| No (range) of women | 747 845 (744 747–751 314) | 49 528 (46 059–52 626) | 797 373 | |
| No (range) of events | 21 588 (20 934–22 010) | 7469 (7047–8123) | 29 057 | 41 126 (5.2) |
| Observed risk (95% CI) | 4.73 (4.56 to 4.89) | 21.67 (19.39 to 23.94) | 5.89 (5.81 to 5.96) | |
| Mean risk QRISK2 (95% CI) | 4.85 (4.72 to 4.98) | 26.76 (26.46 to 27.06) | 6.21 (6.01 to 6.42) | |
| Mean risk NICE (95% CI) | 6.19 (5.95 to 6.43) | 25.95 (24.98 to 26.91) | 7.42 (7.10 to 7.75) | |

The superior performance of the QRISK risk scores is not surprising as both QRISK risk scores were developed (and internally and externally validated) on large cohorts of general practice patients in the United Kingdom, the population for which the risk predictions were targeted and designed. This includes accounting for social deprivation, family history of coronary heart disease, and ethnicity, all known to increase the risk of developing cardiovascular disease. The Framingham score, by contrast, was developed on a comparatively small (n=5573), homogeneous white, though treatment-naive, sample from a single town in the US between 1968 and 1975. We evaluated the Framingham risk score with the NICE designated adjustment factors for family history and, for men, being of South Asian origin.

Arguably, QRISK2 would be more aptly compared with a Framingham equation recalibrated for the UK population, but we compared it with the Framingham risk score recommended by NICE, which is without reference to recalibration. Furthermore, the NICE Framingham is a single equation with a sex coefficient, whereas the QRISK approach has separate equations for men and women. Separate equations permit risk factors to be weighted differently for men and women. Although some have welcomed the introduction of the QRISK equations30–31 and the debate about improving risk prediction, others have cautioned against their use for two main reasons; (a) the cohort used to develop both QRISK equations included patients who were not treatment naive, unlike the Framingham cohort,32 and (b) there are large amounts of missing cholesterol data.33 With regard to the cohorts including patients who may have started additional treatments, it is now neither practically nor ethically...
Table 6 Incidence rates of cardiovascular disease (per 1000 person years) in high risk groups* identified by QRISK2 and NICE Framingham (version of the Framingham equation recommended by NICE) across regions of the United Kingdom

| Country      | Women          | Men          |
|--------------|----------------|--------------|
|              | QRISK2         | NICE Framingham | QRISK2         | NICE Framingham |
| England      | 23.5 (22.9 to 24.1) | 18.9 (18.5 to 19.4) | 25.4 (25.0 to 25.8) | 19.5 (19.2 to 19.7) |
| Northern Ireland | 24.7 (21.5 to 28.3) | 21.2 (18.4 to 24.2) | 27.3 (24.8 to 29.9) | 21.5 (19.8 to 23.2) |
| Scotland     | 27.5 (25.2 to 30.0) | 23.5 (21.5 to 25.6) | 28.2 (26.4 to 30.0) | 21.7 (20.5 to 23.0) |
| Wales        | 24.7 (22.4 to 27.2) | 20.2 (18.3 to 22.2) | 26.0 (24.4 to 27.7) | 19.0 (18.0 to 20.1) |

*High risk defined as ≥20% predicted 10-year risk of cardiovascular disease.

Possible to obtain large treatment-naive cohorts. Furthermore, although natural history is important, it is not clear that prognosis is best assessed from an untreated population.

Risk scores will inevitably become outdated with improvements in clinical outcomes and data recording and changes in population demographics. Thus, ensuring a risk score retains its usefulness to reflect current conditions is crucial. QRISK2 will undergo annual updates to account for changes in population characteristics and improvements in data quality, with its most recent update having been released on 1 April 2010 (see www.qrisk.org for details of this). This entails re-fitting QRISK2 to the latest version of the QRESEARCH database to obtain updated regression coefficients.

Strengths and weaknesses

All the cohorts used in the development and validation of QRISK1 and QRISK2 had high levels of missing data for the total serum cholesterol:high density lipoprotein ratio, so it can be assumed to be a population feature. QRISK1 and QRISK2 were developed using established methods of multiple imputation for missing data to address this problem. Our external validation study also used multiple imputation, with 20 imputed datasets, to deal with the missing data. We note that problems of missing data in developing and validating risk scores are rarely considered when validating risk scores. Omitting patients with missing data in developing the risk score and conducting a complete-case analysis would introduce bias and produce a poorly performing risk score. Even in validation, discarding missing observations will result in performance data that is biased, and so that practice should be avoided.

Conclusions

In this study, we have provided an independent and external validation of the QRISK2 risk score on a large cohort of patients in the United Kingdom. We have assessed the performance of QRISK2 against the NICE version of the Framingham equation and have provided evidence to support the use of QRISK2 in favour of the NICE Framingham equation.

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