# Performance of the QRISK cardiovascular risk prediction algorithm in an independent UK sample of patients from general practice: a validation study 

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#### Abstract

Aim: To assess the performance of the ORISK score for predicting cardiovascular disease (CVD) in an independent UK sample from general practice and compare with the Framingham score. Design: Prospective open cohort study Setting: UK general practices contributing to the THIN and QRESEARCH databases. Cohort: The THIN validation cohort consisted of 1.07 million patients, aged 35-74 years registered at 288 THIN practices between 1 January 1995 and 1 April 2006. The QRESEARCH validation cohort consisted of 0.61 million patients from 160 practices (one-third of the full database) with data until 1 January 2007. Patients receiving statins, those with diabetes or CVD at baseline were excluded. End point: First diagnosis of CVD (myocardial infarction, coronary heart disease (CHD), stroke and transient ischaemic attack) recorded on the clinical computer system during the study period. Exposures: Age, sex, smoking status, systolic blood pressure, total/high-density lipoprotein cholesterol ratio, body mass index, family history of premature CHD, deprivation and antihypertensive medication. Results: Characteristics of both cohorts were similar, except that THIN patients were from slightly more affluent areas and had lower recording of family history of CHD. QRISK performed better than Framingham for every discrimination and calibration statistic in both cohorts. Framingham overpredicted risk by $23 \%$ in the THIN cohort, while QRISK underpredicted risk by $12 \%$. Conclusion: This analysis demonstrated that QRISK is better calibrated to the UK population than Framingham and has better discrimination. The results suggest that QRISK is likely to provide more appropriate risk estimates than Framingham to help identify patients at high risk of CVD in the UK.


In July 2007, we published the results of a new cardiovascular disease (CVD) prediction algorithm known as QRISK. ${ }^{1}$ This algorithm uses existing traditional "risk factors" (age, systolic blood pressure, smoking status and ratio of total serum cholesterol to high-density lipoprotein level) for CVD but also incorporates deprivation, family history, antihypertensive treatment and body mass index. The QRISK algorithm was developed using a validated clinical research database (ORESEARCH), which consists of routinely collected data from general practitioner clinical computer systems. The original research used two-thirds of the database to derive the algorithm, with the remaining third acting as a validation sample.

The performance of ORISK was compared with two existing equations used in the UK, one based on the American Framingham cohort ${ }^{2}$ and the other on the Scottish ASSIGN cohort. ${ }^{3}$ The results demonstrated that ORISK was not only better calibrated to the UK population but also had improved discrimination compared with these existing equations. Although the validation was based on an independent sample of practices within the ORESEARCH database (http://www. qresearch.org, accessed 22 October 2007), the sample is likely to be homogeneous because all practices use the EMIS clinical computer system, which is used in $60 \%$ of UK practices. Since the intended use of the score includes non-EMIS practices, we decided to do a second validation using the THIN database (http://www.thin-uk. com, accessed 22 October 2007), based on practices recording clinical data using the INPS Vision system, which is used in $20 \%$ of UK practices. This would be a more rigorous test of the performance of ORISK compared with Framingham. This paper describes the results.

## METHODS

## Study population

The methods for the derivation and validation of ORISK have been described in detail elsewhere. ${ }^{1}$ For this analysis, we used the original validation sample (ie, a random sample of one-third of the practices in the ORESEARCH database) with a subset of patients aged 35-74 years, excluding patients with diabetes, those with CVD and those prescribed statins at baseline. We followed up the QRESEARCH cohort from 1 January 1995 until 1 January 2007. We identified a second cohort of patients from the THIN database with the same inclusion and exclusion criteria, and registered between 1 January 1995 and 31 March 2006 (the latest data available for analysis).

## Outcome, risk factors, prescribed medication and missing values

We used identical Read code definitions to determine the clinical outcome (CVD), each risk factor and prescribed drugs for each cohort. We used the clinical value recorded closest to the date on which the patient entered the study for body mass index, systolic blood pressure, smoking status, and total and high-density lipoprotein cholesterol. We generated the Framingham score ${ }^{2}$ and the ORISK score ${ }^{1}$ for each individual patient in both cohorts. To generate scores where values for risk factors were missing, we used reference values

Table 1 Summary comparison of patients in the THIN and QRESEARCH validation cohorts

|  | ORESEARCH validation cohort No (\% of total) | THIN validation cohort No (\% of total) |
| :---: | :---: | :---: |
| Total number of practices | 160 | 274 |
| Total number of patients in cohort | 607733 | 1072800 |
| Women | 306111 (50.37) | 542987 (50.61) |
| Men | 301622 (49.63) | 529813 (49.39) |
| Total person years of observation | 3906710 | 5357218 |
| Deprivation breakdown |  |  |
| Patients in Townsend fifth 1 (most affluent) | 143603 (23.63) | 296929 (27.68) |
| Patients in Townsend fifth 2 | 128720 (21.18) | 244821 (22.82) |
| Patients in Townsend fifth 3 | 120459 (19.82) | 220362 (20.54) |
| Patients in Townsend fifth 4 | 115963 (19.08) | 179811 (16.76) |
| Patients in Townsend fifth 5 (most deprived) | 98988 (16.29) | 130877 (12.20) |
| Event type |  |  |
| Total patients with incident CVD events | 30087 | 44152 |
| Number with incident CHD events | 20434 (67.9) | 30797 (69.8) |
| Number with incident stroke/TIA | 9653 (32.1) | 13355 (30.2) |
| 10-Year risk of CVD events (95\% CI) |  |  |
| Women | 6.51 (6.39 to 6.63) | 6.55 (6.43 to 6.68) |
| Men | 9.16 (9.01 to 9.31) | 9.87 (9.71 to 10.03) |

CHD, coronary heart disease; CVD, cardiovascular disease; TIA, transient ischaemic attack.
derived from the derivation cohort of ORESEARCH by 5-year bands of age and sex, assuming patients were non-smokers where smoking status was not recorded. ${ }^{1}$ We used a revised equation for ORISK (version 1.1) which takes account of improvements in the method for multiple imputation of missing data, in which additional variables (including the outcome variable) were included in the imputation model. ${ }^{4}$ This revised equation now excludes patients prescribed statins at baseline, so the results from the ORESEARCH validation database presented here will differ slightly from those previously published. ${ }^{1}$

## Townsend scores

We used the Townsend score evaluated at output area as a proxy for material deprivation. The THIN dataset differs from the ORESEARCH dataset in that each patient in the THIN dataset is allocated to a fifth of deprivation and only the category number is provided. In contrast, each patient in the ORESEARCH dataset is allocated the individual Townsend score corresponding to their output area of residence (ie, continuous data). To calculate the ORISK equation in the THIN cohort, we used the median value for each fifth defined from a national postcode table mapped to output area Townsend scores derived from the 2001 census.

## Discrimination and calibration statistics

We tested the performance of QRISK in both the THIN and the ORESEARCH validation cohorts. We calculated the 10 -year estimated CVD risk for each patient in the validation datasets, replacing missing values as described.

To assess calibration (ie, degree of similarity between predicted and observed risks) we calculated the mean predicted CVD risk at 10 years, and the observed CVD risk at 10 years obtained using the 10 -year Kaplan-Meier estimate and compared the ratio of the predicted to the observed CVD risk for patients in the validation cohort in each decile of predicted risk. We also compared predicted and observed risks overall for men
and women and calculated the Brier score. ${ }^{5}$ We calculated the area under the receiver operator curve statistic to assess discrimination (ie, ability of a risk prediction equation to distinguish between those who do and do not have a cardiovascular event during the follow-up period). We also calculated the D statistic ${ }^{6}$ and an $\mathrm{R}^{2}$ statistic derived from the D statistic, ${ }^{7}$ which are measures of discrimination and explained variation appropriate for survival models. The D statistic has been developed as a new measure of discrimination specifically for censored survival data-higher values indicate improved discrimination, and an increase in the D statistic of at least 0.1 indicates an important difference in prognostic separation between different risk classification schemes.

## Comparison with Framingham

We compared the performance of ORISK against the risk estimates derived from Framingham equations. ${ }^{2}$ We used a CVD risk equation which was computed by summing the coronary risk (including myocardial infarction and coronary heart disease (CHD) death plus angina plus coronary insufficiency) and stroke risk (including transient ischaemic attack) as these outcomes are closest to those used in randomised trials of drug effectiveness. It is also the outcome used in the current Joint British Society Guidelines. ${ }^{8}$ We calculated the measures of discrimination described above for the Framingham estimates in the validation dataset and compared them with the equivalent QRISK values.

Finally, we calculated the proportion of patients in the validation samples who have a 10 -year CVD risk of $20 \%$ or more by age, sex and deprivation according to QRISK and Framingham. Analyses were conducted using Stata (version 9.2).

## RESULTS

## Study population on THIN and ORESEARCH

Overall, there were 1787169 patients from 288 practices in the THIN cohort. Sequentially, we excluded 120281 patients with a prior diagnosis of CVD, 2253 patients with invalid dates,

Table 2 Completeness of recording for risk factors for men and women aged 35-74 initially free from cardiovascular disease and diabetes in the derivation and validation datasets

| Risk factors | QRESEARCH validation cohort |  | THIN validation cohort |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Men $\text { ( } \mathrm{n}=301622 \text { ) }$ | Women $(\mathrm{n}=306 \text { 111) }$ | Men $(\mathrm{n}=529813)$ | Women $\text { ( } \mathrm{n}=542 \text { 987) }$ |
| Age | 301622 (100.0) | 306111 (100.0) | 529813 (100.0) | 542987 (100.0) |
| Sex | 301622 (100.0) | 306111 (100.0) | 529813 (100.0) | 542987 (100.0) |
| Deprivation |  |  |  |  |
| Body mass index (kg/m²) | 220469 (73.1) | 246553 (80.5) | 391703 (73.9) | 451155 (83.1) |
| Systolic blood pressure ( mm Hg ) | 252826 (83.8) | 281550 (92.0) | 457217 (86.3) | 511423 (94.2) |
| Total serum cholesterol (mmol/l) | 119976 (39.8) | 127886 (41.8) | 213793 (40.4) | 221869 (40.9) |
| HDL (mmol/l) | 86875 (28.8) | 94413 (30.8) | 151782 (28.7) | 159690 (29.4) |
| Smoking status | 254718 (84.5) | 278797 (91.1) | 460616 (86.9) | 508680 (93.7) |
| Complete data for all risk factors | 77920 (25.8) | 86061 (28.1) | 135114 (25.5) | 145994 (26.9) |

Results are shown as counts (\%) unless otherwise specified. Patients receiving statins at baseline have been excluded. HDL, high-density lipoprotein.

284492 patients who were aged under 35 years, 155248 patients aged $\geqslant 75$, 114123 with missing Townsend scores, 28148 patients with diabetes and 9824 patients who were taking statins. This left a cohort of 1072800 patients from the THIN database for analysis of whom 529813 (49.39\%) were men. There were 24 practices ( 54709 patients) from Scotland and 14 practices ( 36904 patients) from Northern Ireland. The corresponding cohort on ORESEARCH contained 607733 patients and has been described previously.

Table 1 shows a comparison of the THIN and ORESEARCH validation cohorts for the validation analysis. The proportions of men and women were similar in both datasets. Patients from THIN tended to be from more affluent areas with $28 \%$ in the most affluent fifth and $12 \%$ in the most deprived. The corresponding figures for patients in the ORESEARCH cohort were $24 \%$ and $16 \%$, showing a slightly more even distribution across the fifths. The 10-year risk of CVD was marginally higher in men in THIN than in QRESEARCH (9.87\% vs 9.16\%) but was more similar for women ( $6.55 \%$ vs $6.51 \%$ ).

Table 2 shows the completeness of the recording for each risk factor for men and women in both cohorts. Data for age, sex
and deprivation quintile were complete for all patients included in the analysis. Overall, levels of recording were very similar between the two cohorts with highest levels for systolic blood pressure and smoking, which were recorded in excess of $85 \%$ of patients in both cohorts. Lowest levels were observed for highdensity lipoprotein cholesterol, with recorded values in 29-31\% of patients. Overall, just over $25 \%$ of patients had complete data for all risk factors.

Table 3 shows baseline characteristics for both cohorts for age, sex, risk factors and medication using recorded values only. There was a striking similarity for almost all CVD risk factors between the two cohorts. For example the mean (SD) systolic blood pressure in men was 135.3 (19.6) mm Hg in ORESEARCH and 135.6 (19.4) mmHg in THIN. The corresponding figures for body mass index were 26.5 (4.0) $\mathrm{kg} / \mathrm{m}^{2}$ in men in ORESEARCH and $26.6(4.0) \mathrm{kg} / \mathrm{m}^{2}$ in THIN. The recorded prevalence of family history of premature coronary heart disease was substantially lower in THIN than ORESEARCH ( $3.5 \%$ in men in THIN vs $9.2 \%$ in men in ORESEARCH).

Table 4 shows the key calibration and discrimination statistics for Framingham and ORISK for both ORESEARCH

Table 3 Baseline recorded clinical characteristics for men and women aged 35-74 initially free from cardiovascular disease and diabetes in the validation datasets

| Characteristics | ORESEARCH validation cohort |  | THIN validation cohort |  |
| :---: | :---: | :---: | :---: | :---: |
|  | Men $\text { (n = } 301 \text { 622) }$ | Women $\text { ( } \mathrm{n}=306 \text { 111) }$ | Men $(\mathrm{n}=529 \text { 813) }$ | Women $\text { ( } n=542 \text { 987) }$ |
| Age at baseline, median (interquartile range) | 47 (40 to 57) | 49 (41 to 59) | 48 (40 to 57) | 49 (41 to 59) |
| Body mass index ( $\mathrm{kg} / \mathrm{m}^{2}$ ), mean (SD) | 26.5 (4.0) | 26.1 (4.8) | 26.6 (4.0) | 26.1 (4.9) |
| Systolic blood pressure (mm Hg), mean (SD) | 135.3 (19.6) | 132.2 (21.6) | 135.6 (19.4) | 132.1 (21.0) |
| Total serum cholesterol (mmol), mean (SD) | 5.7 (1.1) | 5.8 (1.2) | 5.7 (1.1) | 5.8 (1.2) |
| HDL (mmol/l), mean (SD) | 1.3 (0.4) | 1.6 (0.4) | 1.3 (0.4) | 1.6 (0.4) |
| Total serum cholesterol/HDL ratio, mean (SD) | 4.5 (1.3) | 3.9 (1.2) | 4.5 (1.3) | 3.9 (1.2) |
| Current smoker | 84930 (28.2) | 70210 (22.9) | 141113 (26.6) | 124094 (22.9) |
| Family history of coronary heart disease in first-degree relative under 60 years | 27829 (9.2) | 38026 (12.4) | 18638 (3.5) | 22922 (4.2) |
| Receiving antihypertensive treatment* | 24245 (8.0) | 36992 (12.1) | 35066 (6.6) | 56886 (10.5) |
| Receiving ACE inhibitors* | 6898 (2.3) | 7170 (2.3) | 11718 (2.2) | 12901 (2.4) |
| Receiving $\beta$-blockers* | 11321 (3.8) | 17892 (5.8) | 16700 (3.2) | 27554 (5.1) |
| Receiving calcium channel blockers* | 7346 (2.4) | 8479 (2.8) | 9847 (1.9) | 11147 (2.1) |
| Receiving thiazides* | 6837 (2.3) | 14571 (4.8) | 10630 (2.0) | 23391 (4.3) |

[^0]Table 4 Calibration and discrimination statistics comparing the QRISK and Framingham equations for predicted 10 -year risk of cardiovascular disease in the QRESEARCH validation cohort and the THIN cohort

| Statistics | ORISK equation |  | Framingham equation |  |
| :---: | :---: | :---: | :---: | :---: |
|  | In QRESEARCH validation cohort | In THIN validation cohort | In QRESEARCH validation cohort | In THIN validation cohort |
| Women |  |  |  |  |
| ROC statistic | 0.7879 (0.7842 to 0.7915) | 0.7888 (0.7858 to 0.7918) | 0.7759 (0.7723 to 0.7796) | 0.7595 (0.7564 to 0.7626) |
| D statistic | 1.535 (1.508 to 1.562) | 1.556 (1.532 to 1.580) | 1.397 (1.370 to 1.424) | 1.378 (1.354 to 1.402) |
| $\mathrm{R}^{2}$ statistic (\%) | 36.01 (35.16 to 36.86) | 36.64 (35.94 to 37.34) | 31.79 (30.92 to 32.66) | 31.18 (30.45 to 31.91) |
| Brier score | 0.0370 | 0.0321 | 0.0382 | 0.0338 |
| Predicted/observed (women) | 1.00 | 0.90 | 1.19 | 1.10 |
| Men |  |  |  |  |
| ROC statistic | 0.7700 (0.7667 to 0.7733) | 0.7619 (0.7592 to 0.7646) | 0.7619 (0.7586 to 0.7653) | 0.7365 (0.7336 to 0.7394) |
| D statistic | 1.425 (1.401 to 1.449) | 1.394 (1.374 to 1.414) | 1.317 (1.293 to 1.341) | 1.255 (1.235 to 1.275) |
| $\mathrm{R}^{2}$ statistic (\%) | 32.64 (31.88 to 33.39) | 31.70 (31.09 to 32.31) | 29.30 (28.54 to 30.05) | 27.31 (26.69 to 27.93) |
| Brier score | 0.0504 | 0.0470 | 0.0569 | 0.055 |
| Predicted/observed (men) | 0.97 | 0.87 | 1.49 | 1.32 |
| Predicted/observed (overall) | 0.99 | 0.88 | 1.36 | 1.23 |

Results are shown as mean ( $95 \% \mathrm{CI}$ ) unless stated otherwise.
Note that higher values for the ROC, D and $\mathrm{R}^{2}$ indicate better discrimination. Lower values of the Brier score indicate better calibration. ROC, receiver operator curve.
and THIN cohorts. Framingham overpredicted risk by $23 \%$ in the THIN cohort while ORISK underpredicted risk by $12 \%$. ORISK had better values than Framingham for each of the discrimination and calibration statistics in both the THIN and the QRESEARCH validation cohorts.

Table 5 shows the proportion of patients at high risk (ie, more than $20 \%$ risk of CVD over 10 years) according to ORISK and Framingham equations by fifth of deprivation in both the QRESEARCH and THIN validation cohorts. The deprivation gradient across the fifths is very similar in both cohorts for ORISK. Both cohorts show a higher proportion of patients at high risk in the most deprived fifth than in the most affluent fifth using ORISK, but a much smaller deprivation gradient for Framingham.

Table 6 shows the distribution of patients at high risk (ie, more than 20\% CVD risk over 10 years) according to QRISK and

Table 5 Percentage of patients with a cardiovascular disease risk score $\geqslant 20 \%$ by sex and Townsend fifth (first fifth is the most affluent and fifth fifth the most deprived) in patients aged $35-74$ years estimated using Framingham and QRISK

| Townsend fifth | QRISK equation |  | Framingham equation |  |
| :---: | :---: | :---: | :---: | :---: |
|  | In QRESEARCH validation cohort | In THIN validation cohort | In QRESEARCH validation cohort | In THIN validation cohort |
| Women |  |  |  |  |
| First | 2.75 | 2.07 | 4.51 | 3.58 |
| Second | 3.55 | 3.23 | 4.74 | 4.39 |
| Third | 4.64 | 4.26 | 5.20 | 5.00 |
| Fourth | 6.83 | 6.08 | 6.18 | 5.81 |
| Fifth | 8.67 | 8.64 | 6.21 | 6.34 |
| Men |  |  |  |  |
| First | 9.34 | 7.79 | 20.30 | 18.61 |
| Second | 9.85 | 9.60 | 20.09 | 20.52 |
| Third | 10.55 | 10.04 | 19.96 | 20.04 |
| Fourth | 11.96 | 11.60 | 20.42 | 21.04 |
| Fifth | 12.1 | 12.75 | 19.32 | 21.10 |
| Total | 7.99 | 7.04 | 12.63 | 12.31 |

[^1]Framingham equations by age band and sex. The pattern is very similar for both cohorts though overall ORISK predicted a slightly smaller proportion at high risk in THIN (7.04\%) than it did in the ORESEARCH validation cohort (7.99\%).

Overall, 85009 patients in the THIN cohort $(7.9 \%$ of the total) would be reclassified from high to low risk or vice versa using ORISK compared with Framingham. Of 132076 patients classified at high risk using Framingham, then 70764 (53.6\%) would be reclassified as low risk on ORISK. In these patients, the observed 10 -year risk was $17.4 \%$ ( $95 \%$ CI $16.8 \%$ to $17.9 \%$ ). Conversely, the 14245 patients classified as low risk on Framingham but high risk on ORISK had an observed 10-year risk of $23.7 \%$ ( $95 \%$ CI $22.4 \%$ to $25.0 \%$ ).

Overall, 46785 patients in the ORESEARCH cohort (7.7\% of the total) would be reclassified from high to low risk or vice versa using ORISK compared with Framingham. Of 76748

Table 6 Percentage of patients by age and sex with cardiovascular disease risk $\geqslant 20 \%$ estimated using Framingham and QRISK in the validation cohorts having excluded patients receiving statins at baseline

| Age band (years) | QRISK equation |  | Framingham equation |  |
| :---: | :---: | :---: | :---: | :---: |
|  | In QRESEARCH validation cohort | In THIN validation cohort | In ORESEARCH validation cohort | In THIN validation cohort |
| Women |  |  |  |  |
| 35-44 | 0.00 | 0.00 | 0.01 | 0.01 |
| 45-54 | 0.02 | 0.01 | 0.79 | 0.57 |
| 55-64 | 2.23 | 1.24 | 7.33 | 6.38 |
| 65-74 | 31.43 | 26.38 | 24.1 | 21.67 |
| All women | 5.25 | 4.23 | 5.35 | 4.75 |
| Men |  |  |  |  |
| 35-44 | 0.00 | 0.00 | 0.29 | 0.22 |
| 45-54 | 0.48 | 0.24 | 7.76 | 7.31 |
| 55-64 | 13.65 | 10.34 | 40.75 | 38.47 |
| 65-74 | 68.9 | 64.05 | 86.25 | 83.78 |
| All men | 10.78 | 9.93 | 20.01 | 20.06 |
| All patients aged 35-74 | 7.99 | 7.04 | 12.63 | 12.31 |

patients classified at high risk using Framingham, then 37479 (48.8\%) would be reclassified as low risk on ORISK. In these patients the observed 10 -year risk was $16.7 \%$ ( $95 \%$ CI $16.2 \%$ to $17.2 \%$ ). Conversely, the 9306 patients classified as low risk on Framingham but high risk on ORISK had an observed 10 year risk of $24.4 \%$ ( $95 \%$ CI $23.2 \%$ to $25.6 \%$ ).

## DISCUSSION

## Summary of key findings

This is the first external validation of the new ORISK CVD algorithm. It has been conducted on a completely independent sample of patients registered with general practices using a different clinical computer system, but identical definitions were possible. As with the original analysis, ORISK outperforms Framingham in the THIN cohort for both calibration and discrimination. The discrimination statistics for ORISK in THIN are as good as those for the original ORISK validation cohort from ORESEARCH. INPS Vision system practices together with EMIS cover $80 \%$ of the UK general practices. The results are also important since this is the first major headon comparison of two recently established major UK general practice databases, suggesting that the results of similar studies based on either database would be generalisable to the UK.

## Baseline comparison of the two cohorts

Although the THIN cohort had more patients, the mean duration of follow-up was less than for patients in the one-third sample of ORESEARCH practices. Overall, the results show a striking similarity between THIN and ORESEARCH patients for almost all baseline characteristics. There are two notable exceptions. First, patients contributing to the THIN database tend to be from more affluent areas. Nonetheless, the incidence of CVD by deprivation shows the expected gradient with a higher risk in the most deprived areas (results available from the authors). Second, fewer patients in the THIN cohort had a recorded family history of CHD in a first-degree relative under the age of 60 years. Given the striking similarity for all the other risk factors and treatment variables, it is likely this reflects a difference in recording patterns between the two clinical computer systems rather than a true difference in prevalence.

In the derivation of the QRISK algorithm we used multiple imputation as this has been shown to introduce less bias than the complete case analysis. ${ }^{8 a}$ For this validation study, we generated reference values by 5 -year age and sex based on derivation cohort of ORESEARCH since this is what will be needed in order to implement the algorithm into software. Again, this is less likely to introduce bias than a complete case analysis where the patients themselves are highly selected.

We applied the same method for imputing missing values (including cholesterol ratio) for both the ORISK and the Framingham calculations so that we could make a direct comparison between the two scores. If anything, the imputation would tend to reduce the discriminatory power of both scores.

## Comparison of performance in the two cohorts

The discrimination statistics for ORISK in THIN are similar to those for the original ORISK validation cohort from ORESEARCH. There is a small degree of underprediction using QRISK in the THIN cohort which may be due partly to the low recording of family history in the THIN database-something which is likely to improve if the recording of family history is
encouraged by inclusion in national guidelines and use of templates to standardise data entry. It may also be due to the slightly higher incidence of CVD in the THIN cohort, particularly in men.

Using ORISK instead of Framingham results in a clinically important reclassification of patients into high or low risk. For example, in the THIN analysis, more than half of the patients classified at high risk according to Framingham would be reclassified as low risk using ORISK and these patients do have a low observed 10 -year risk of $17 \%$. Similarly, patients at low risk using Framingham but high risk using QRISK had high observed 10 -year risk of $24 \%$. This shows that using Framingham would fail to identify patients at high risk of CVD but include others who are actually at low risk.

Our analysis demonstrates that ORISK performs well in an independent sample of patients derived from general practices which use a different clinical computer system in the UK. This is an important test of validity, especially since ORISK has been developed for use in the UK primary care population. The results suggest that GPs in different practices measure the same items with similar validity and that their populations are similar. A stronger test of validity would be to examine whether the score operates where risk factors have been measured under different conditions. These include other primary care settings and other parts of the world which have similar levels of cardiovascular risk. An international version of ORISK is an important area for future research and development.

## FRAMINGHAM AND SCORE

Unlike the existing Framingham equations and the European SCORE equation, ${ }^{9}$ ORISK identifies and includes deprivation in the estimation of CVD risk. This will be a significant step in supporting national initiatives to reduce health inequalities in CVD ${ }^{10}$ and likely to be an improvement on Framingham, which tends to overestimate risk in affluent areas and underestimate risk in deprived areas. ${ }^{11}$ Also, a weighting for social deprivation might help minimise health inequalities, which may increase when new interventions are introduced because of the inverse equity hypothesis. ${ }^{12}$ The inclusion of family history of premature CHD is important because observational studies indicate that premature CHD in a first-degree relative increases risk by $50 \%$ or more, ${ }^{13}{ }^{14}$ although our estimates were lower than this. QRISK deals with components not included in SCORE, which was designed for European use. SCORE does not include CVD morbidity, which is an integral part of both trial data (non-fatal myocardial infarction and stroke) and costeffectiveness reviews. The statin HTA in the UK has identified a CVD event threshold of $20 \%$ in 10 years as the threshold for treatment of which most are non-fatal events. ${ }^{15}$

In a major initiative to improve public health, the National Institute of Health and Clinical Excellence (NICE) has lowered the threshold for primary prevention with statins from a $10-$ year CVD risk of $40 \%$ to $20 \%$. ${ }^{16}{ }^{17}$ Reliable methods for estimating risk of CVD are now needed to implement the guidelines in clinical practice in a way which does not exacerbate health inequalities. The results of this validation in an independent UK dataset suggest that ORISK is likely to provide more appropriate estimates of CVD risk in contemporary primary care UK populations and better discrimination of those at high risk based on their age, sex, social deprivation and existing antihypertensive treatment. It is likely therefore to be a more equitable and clinically appropriate tool to inform patient management decisions.

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[^0]:    Results are shown as counts (\%) unless otherwise specified.
    Patients receiving statins at baseline have been excluded.
    *At entry to the cohort.

[^1]:    Patients receiving statins at baseline have been excluded.

