



## Full Length Article

# External validation of a model to predict women most at risk of postpartum venous thromboembolism: Maternity clot risk

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## ABSTRACT

**Introduction:** Venous thromboembolism (VTE) is the leading cause of direct maternal mortality in high-income countries. We previously developed a risk prediction score for postpartum venous thromboembolism (VTE) in women without a previous VTE. In this paper, we provide further external validation and assess its performance across various groups of postpartum women from England.

**Materials and methods:** Cohort study using primary and secondary care data covering England. We used data from QRResearch comprising women with pregnancies ending in live birth or stillbirth recoded in Hospital Episodes Statistics between 2004 and 2015. Outcome was VTE in the 6 weeks postpartum. Our predictor variables included sociodemographic and lifestyle characteristics, pre-existing comorbidities, and pregnancy and delivery characteristics.

**Results:** Among 535,583 women with 700,185 deliveries, 549 VTE events were recorded (absolute risk of 7.8 VTE events per 10,000 deliveries). When we compared predicted probabilities of VTE for each woman from the original model with actual VTE events, we obtained a C-statistic of 0.67 (95% CI 0.65 to 0.70). However, our model slightly over-predicted VTE risk for the higher risk women (calibration slope = 0.84; 95% CI 0.74 to 0.94). Performance was similar across groups defined by calendar time, socioeconomic status, age group and geographical area. The score performed comparably with the existing algorithm used by the UK Royal College of Obstetrician and Gynaecologists.

**Conclusions:** Our model enables flexibility in setting new treatment thresholds. Adopting it in clinical practice may help optimise use of low-molecular-weight heparin postpartum to maximise health gain by better targeting of high-risk groups.

## 1. Introduction

Venous thromboembolism (VTE) is the leading direct cause of maternal mortality in high income countries and is associated with considerable preventable morbidity [1,2]. The absolute risk of VTE peaks in the six weeks following childbirth [3–5]. In 2016, we developed

and published a risk prediction score which estimates the risk of VTE during the first six weeks after childbirth based on commonly recorded risk factors at the point of delivery [6]. This score was subsequently named the “Maternity Clot Risk” and is available from [www.maternity-clot-risk.co.uk](http://www.maternity-clot-risk.co.uk). The Maternity Clot Risk not only performed better than the current UK Royal College of Obstetrician and Gynaecologist

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### Box 1 maternity clot risk.

Risk score developed from a logistic regression model in the model development study to predict the first ever venous thromboembolism in the first six weeks postpartum.

$$\text{Risk score} = -9.103121 + 0.94 \times (0.22684105 \times \text{smoker} + 1.2210805 \times \text{varicose veins} + 0.8476927 \times \text{comorbidities (cardiac, renal or inflammatory bowel disease)} + 0.72127433 \times \text{pre-eclampsia/eclampsia} + 0.42119233 \times \text{diabetes} + 0.50183134 \times \text{postpartum haemorrhage} + 1.1514008 \times \text{stillbirth} + 1.0969922 \times \text{postpartum infection} + 0.56321456 \times \text{elective section} + 0.75035197 \times \text{emergency section} + 0.16456948 \times \text{parity of 1} + 0.48143018 \times \text{parity of 2} + 0.5664196 \times \text{parity of } \geq 3 - 0.00007986 \times \text{age at delivery}^3 + 0.00002147 \times (\text{age at delivery}^3 \ln(\text{age at delivery})) + 0.00026641 \times \text{BMI}^3 - 0.00006501 \times (\text{BMI}^3 \ln(\text{BMI})) - 22,156,315 \times \text{infant birth weight (g)}^{-2} + 3,455,223.4 \times (\text{infant birth weight}^{-2} \ln(\text{infant birth weight}))$$

(RCOG) [7] and Swedish postpartum thromboprophylaxis guidelines [8], it also generates a predicted risk for each woman which can be used in conjunction with pre-set thresholds for initiation of thromboprophylaxis. The score was originally developed using UK primary care data linked to secondary care data (Clinical Practice Research Datalink, CPRD) and was externally validated in an independent Swedish database [9] where it performed as well as in the original dataset.

The value of a risk prediction score to be used in clinical practice depends on how well it performs when it is applied in populations that are different from the population in which it was developed [10,11]. Furthermore, multiple external validation studies would be needed to fully realise the generalisability of a prediction model. Our original model was developed in UK practices that use a particular clinical computer system [12] (Vision, currently used by 9% of all practices in the UK [13]) and contribute data to the CPRD. In this paper we further validate the Maternity Clot Risk using the QResearch database, which records data from general practices that use another more commonly used system (used by 56% of all UK practices [13]) called Egton Medical Information System (EMIS). The data have been recently linked to secondary care hospital data.

The predictive performance of a model tends to vary across settings, populations and time periods [14]. The aim of this study was therefore to perform an independent external validation of the predictive performance of the Maternity Clot Risk and assess its performance based on calendar time, age group, socioeconomic status and geographic region. External validation of prediction models is a necessary precursor to the important step change of clinicians being able to use the model in everyday clinical practice.

## 2. Methods

A description of the initial study proposal can be found at <https://www.qresearch.org/research/approved-research-programs-and-projects/validating-a-postpartum-venous-thromboembolism-risk-prediction-model-using-qresearch/>.

### 2.1. Data source and study population

QResearch is a UK primary care database containing routinely collected healthcare data of anonymised patients from over 1000 English general practices (<https://www.qresearch.org/>). QResearch has been recently linked to Hospital Episode Statistics (HES), a secondary care administrative database containing all inpatient admissions wholly or partially funded by the National Health Service in England. QResearch has been used for a wide range of clinical research, including the development and validation of various risk prediction models [15–17]. A cohort of women aged 12–59 years old with at least one delivery ending in live birth or stillbirth recorded in HES between January 2004 and December 2015 was extracted using version 41 of the linked

QResearch database as the basis for the study population. HES maternity includes all births occurring in English NHS hospitals where over 97% of live births occur in England [18]. Some women had multiple deliveries included. Those with a history of VTE before the index delivery were excluded from the study.

### 2.2. Definition of outcome

VTE (deep vein thrombosis or pulmonary embolism) was defined based on the first ever recording of the event within the first six weeks postpartum using relevant diagnostic codes. A VTE was defined using a combination of VTE diagnoses in both primary and secondary care data and anticoagulant prescriptions as established previously [19]. In brief, a diagnosis of VTE, in either the primary or secondary care section of the data, was considered to be confirmed if it was accompanied by a prescription for an anticoagulant in primary care within 90 days of the event or if the woman died within 30 days of the event.

### 2.3. Definition of predictors and subgroup variables

In line with our previous CPRD study [6], we extracted information on sociodemographic and lifestyle characteristics, pre-existing comorbidities, and pregnancy and delivery characteristics and complications from both primary and secondary records. Methods used to define predictors in QResearch are described in Supplementary Table S1. We defined pre-existing medical conditions as varicose veins, cardiac disease (ischemic heart disease, congenital heart disease, cardiac failure, cardiac arrhythmias or cardiomyopathy), renal disease (glomerular disease, renal tubulointerstitial disease or renal failure) and inflammatory bowel disease (ulcerative colitis, Crohn's disease or non-specific IBD). Infection following delivery included infections of the respiratory system and urinary tract but not other puerperal infections.

Socioeconomic status was determined from the Townsend deprivation score, grouped into quintiles with 1 the least deprived and 5 the most deprived. Calendar time was also considered in order to provide illustration of model performance during periods when different proportions of women would have been receiving LMWH. We were unable to identify these women individually from both the development and validation datasets. Participants were grouped into one of ten geographic regions, which were based on former strategic health authorities in the UK. For the purpose of subgroup analyses, age was grouped into three categories (<25 years, 25–34 years, ≥35 years).

### 2.4. Statistical analysis

As in the model development study [6], we treated the occurrence of postpartum VTE as a binary outcome measure (occurrence in the first six weeks postpartum: yes or no). Continuous variables (i.e. age, pre-pregnancy body mass index (BMI) and baby's birth weight) were

**Table 1**

Characteristics of study population (number of deliveries = 700,185, number of women = 535,583) from QResearch cohort and CPRD cohort [6].

Variables	QResearch cohort (n = 700,185) n (% if not otherwise specified)	CPRD cohort (n = 433,353) n (% if not otherwise specified)
VTE	549 (0.08)	315 (0.07)
Social and demographic factors		
Mean (SD) age at delivery, years	29.85 (5.91)	29.38 (5.90)
Mean (SD) body mass index	25.06 (5.55)	24.05 (4.90)
Normal	315,624 (45.08)	–
Underweight	28,269 (4.04)	–
Overweight	141,313 (20.18)	–
Obese	94,063 (13.43)	–
Missing	120,916 (17.27)	–
Smoker (latest record before delivery)	128,029 (18.29)	93,264 (21.52)
Socioeconomic deprivation		
1 (least deprived)	130,173 (18.59)	–
2	140,584 (20.08)	–
3	151,064 (21.57)	–
4	144,769 (20.68)	–
5 (most deprived)	130,665 (18.66)	–
Missing	2930 (0.42)	–
Comorbidities ever before delivery		
Varicose veins	16,962 (2.42)	10,935 (2.52)
Heart disease	7525 (1.07)	4431 (1.02)
Kidney disease	5314 (0.76)	4168 (0.96)
Inflammatory bowel disease	3756 (0.54)	2126 (0.49)
Pregnancy complications		
Pre-eclampsia/eclampsia	12,291 (1.76)	9966 (2.30)
Diabetes	37,699 (5.38)	14,604 (3.37)
Hypertension	46,158 (6.59)	41,300 (9.53)
Antenatal parity		
Nulliparous	341,625 (48.79)	244,233 (56.36)
1	259,841 (37.11)	130,121 (30.03)
2	67,955 (9.71)	38,599 (8.91)
≥3	30,764 (4.39)	20,400 (4.71)
Delivery characteristics/ complications		
Preterm birth (<37 weeks)	49,610 (7.09)	31,526 (7.27)
Postpartum haemorrhage	62,244 (8.89)	42,978 (9.92)
Spontaneous/assisted vaginal delivery	523,360 (74.75)	328,416 (75.78)
Elective caesarean section	75,640 (10.80)	44,143 (10.19)
Emergency caesarean section	101,185 (14.45)	60,794 (14.03)
Multiple delivery (twins or more)	10,772 (1.54)	6550 (1.51)
Stillbirth	3312 (0.47)	1972 (0.46)
Puerperal acute infection	14,043 (2.01)	13,681 (3.16)
Infant's mean (SD) birth weight, g	3356.17 (584.57)	3368.35 (596.80)
Missing information		
Infant birth weight	124,299 (17.75)	87,305 (20.15)
Body mass index	120,916 (17.27)	98,868 (22.81)

transformed in line with the Maternity Clot Risk equation (Box 1). To account for missing data, we used multiple imputation by chained equations to create five imputed datasets where any missing values for the BMI and the baby's birth weight were estimated based on other covariates and postpartum VTE. Multiple pregnancies in the same woman were accounted for by use of a clustering term.

To each imputed dataset, we applied the Maternity Clot Risk (Box 1) to provide a predicted VTE risk for each postpartum woman. The following methods were used to evaluate the extent to which our model correctly predicts which women developed postpartum VTE.

- i. Discrimination - The ability of the score to differentiate between women who did and did not develop a first postpartum VTE event

- ii. Calibration - Refers to how closely the predicted first postpartum VTE risk agrees with the observed risk. This differs from discrimination as it enables detection of whether the model over or under-estimates VTE risk, either universally or at specific risk levels.
- iii. Subgroup analyses – Based on age, region, socioeconomic status and calendar time were presented to explore potential heterogeneity in model performance between different clinically important demographic subgroups or over time.
- iv. Sensitivity and positive predictive value of the model in predicting postpartum VTE compared with the algorithm currently used by the Royal College of Obstetricians and Gynaecologists (RCOG).
- v. Decision curve analysis – Highlights the range of thresholds for intervention based on underlying risk of VTE where the model outperforms alternative strategies for intervention.

In a sensitivity analysis, we explored whether a re-calibrated model offered improved prognostic performance. The above methods along with justification for their use are explained fully in Supplementary Appendix A.

Previous research suggested that at least 100 cases and 100 non-cases would be needed for validation studies, and our sample size far exceeds this [20]. All data management and analysis were conducted using Stata 15, and the findings reported according to the TRIPOD (Transparent Reporting of a multivariable prediction model for Individual Prognosis Or Diagnosis) guidance [21].

## 2.5. Ethical approval

This project was approved by the QResearch Advisory Group, Project reference ID R82.

## 3. Results

### 3.1. Baseline characteristics

We included 535,583 women with 700,185 deliveries resulting in either a live birth or stillbirth with a complete six weeks of post-delivery follow-up. There were 549 first VTE events in the first six weeks postpartum corresponding to an absolute risk of 7.8 per 10,000 deliveries (95% CI 7.2 to 8.5). Table 1 shows the basic characteristics of the study population. Broadly, compared to women in CPRD, women in QResearch had similar age at delivery and prevalence of comorbidities, slightly higher mean BMI, were less likely to be nulliparous and smoke and had slightly fewer pregnancy and delivery related complications (Table 1). There were 17.8% with missing infant birth weight and 17.3% with missing BMI in QResearch, which was lower than in CPRD.

### 3.2. Prediction of VTE risk

Using the maternity clot risk formula, predicted risks of VTE were calculated for each woman in the cohort. The predicted risks ranged from 0 to 745 per 10,000 deliveries (maximum equivalent to 7.5% risk of VTE); median predicted risk = 5.3 per 10,000, 10th percentile 3.0 per 10,000, 90th percentile 14.2 per 10,000. A total of 5.0% of women had a predicted risk of VTE of 0.2% or more (or 20 per 10,000 deliveries, n = 34,832), 0.6% of women had a predicted risk of 0.5% or more (n = 4242) and 0.1% had a predicted risk of 1% or more (n = 756). These numbers were taken from the first multiple imputed dataset, but all the key figures (predicted risks per 10,000 and percentages) were the same to at least 2 significant figures in the other imputed datasets.

### 3.3. Overall model performance

After obtaining predicted probabilities of VTE from the Maternity

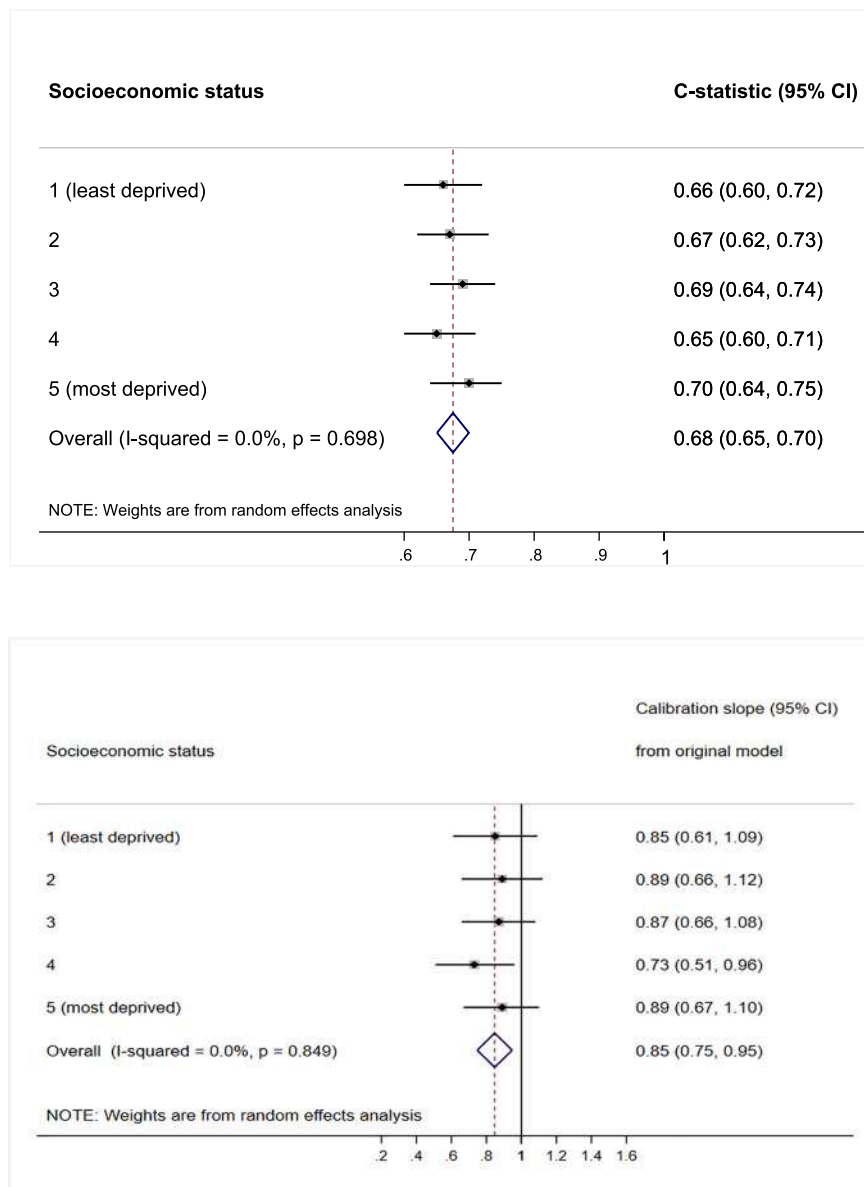


Fig. 1. Model diagnostics by socioeconomic status; a) c-statistic, b) calibration slope (before re-calibration).

Clot Risk score, the overall C-statistic pooled over the imputed datasets was 0.67 (95% CI 0.65 to 0.70). Calibration slope was 0.84 (0.74 to 0.94) and calibration-in-the-large was 0.02 (−0.06 to 0.10). Similar results were observed in each imputed dataset (Supplementary Table S2). The plotted agreement between predicted and observed risks across tenths of predicted risks is shown in Supplementary Fig. 1. Due to the small range of predicted risks, the figures show the predicted risks up to 30 per 10,000 deliveries only.

### 3.4. Performance by subgroup

Results from the analysis by different groups showed that the Maternity Clot Risk performed similarly in women of different socioeconomic groups (Fig. 1), geographic regions (Fig. 2), in women giving birth in different calendar periods (Fig. 3) and women in different age groups (Fig. 4), with minimal heterogeneity in all instances for both the C-statistic and calibration slope.

### 3.5. Comparison with the existing RCOG guideline

According to the current RCOG postpartum thromboprophylaxis guideline, 35.6% of women in the study population qualified for pharmacological thromboprophylaxis for at least 10 days after delivery. The results from the decision curve analysis (Supplementary Fig. S2) show, although the net benefit was small, the Maternity Clot Risk was better than a treat-all or treat-none strategy between risk thresholds of 10 and 30 per 10,000 deliveries. It had higher net benefit than the current RCOG guideline between risk thresholds of 5 and 30 per 10,000 deliveries. Using the Maternity Clot Risk to identify the same proportion of women based on their predicted risks (i.e. risk threshold 6.77 per 10,000 deliveries) resulted in a slightly higher observed sensitivity (59.2, 95% CI 55.0 to 63.3) vs. than using the RCOG guideline (56.8, 95% CI 52.6 to 61.0) (Table 2), although the difference was not statistically significant.

### 3.6. Re-calibration results

After shrinking the original predictor coefficients by 0.79 (0.84 \* 0.94) and re-estimating the intercept, the calibration slope was 1.00

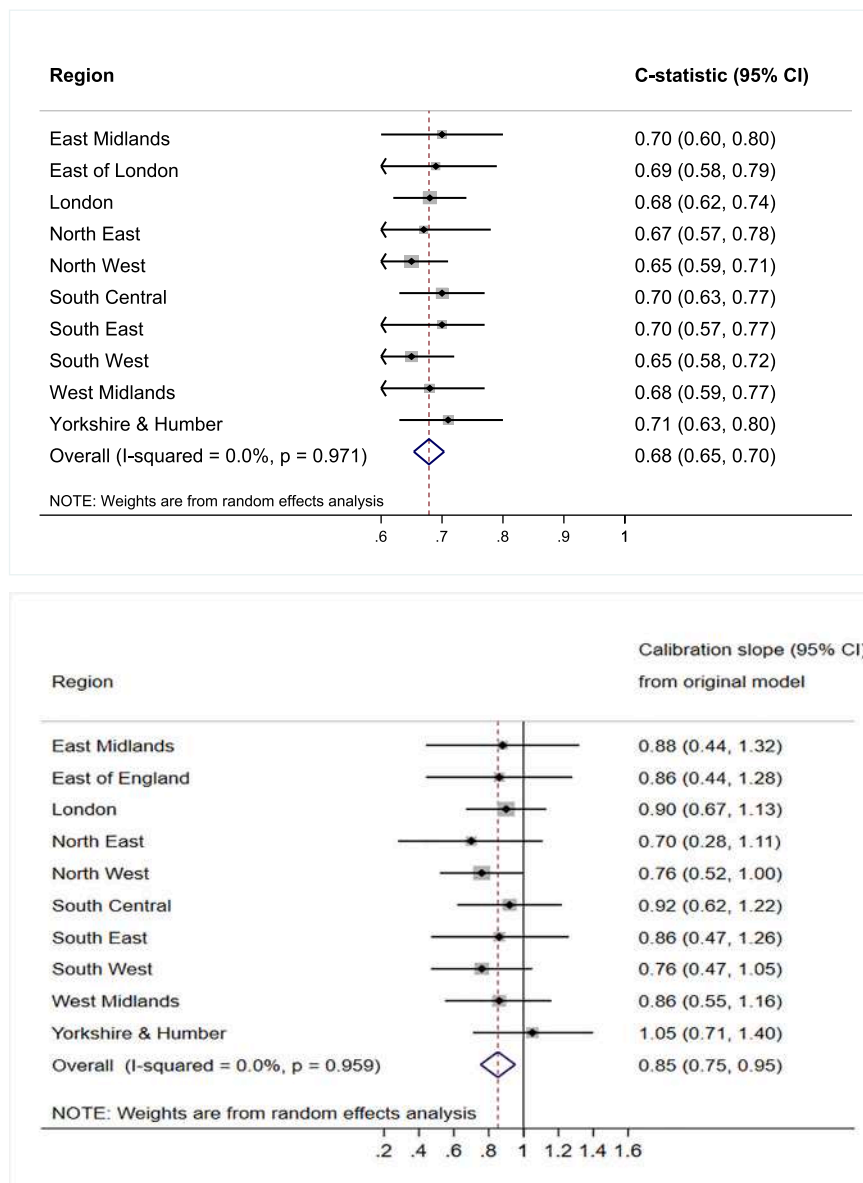


Fig. 2. Model diagnostics by region; a) c-statistic, b) calibration slope (before re-calibration).

(0.88 to 1.12). Results from other analyses remained largely unchanged (Supplementary Table S2, and Supplementary Figs. S1 to S6).

#### 4. Discussion

##### 4.1. Main findings

We have carried out an external validation of the Maternity Clot Risk in the largest available UK primary care dataset. It was conducted in an independent sample of women derived from UK general practice using a different clinical computer system to the CPRD. Applying the Maternity Clot Risk to the QResearch cohort resulted in an overall C-statistic of 0.67 (95% CI 0.65–0.70) and a calibration slope of 0.84 (0.74–0.94). The predictive performance was similar across time periods, socioeconomic and age groups and geographical regions. The Maternity Clot Risk had a slightly higher net benefit than the existing RCOG postpartum thromboprophylaxis guideline and the treat none strategy between risk thresholds of 10 and 30 VTE events per 10,000 deliveries. Our model has the potential to be used in maternity units if suitable thresholds for intervention could be established, although results should be interpreted

in light of limitations.

##### 4.2. Strengths and limitations

We have conducted an external validation of the Maternity Clot Risk in the UK population. It was conducted in the UK's largest primary care data with linkages to secondary care hospital data, with 549 cases. Data management and analysis were conducted by a researcher not involved in the original model development process (LB) but using the original Maternity Clot Risk and statistical methods, which further ensures robustness of our external validation. As primary care practices contributing data to QResearch use a different computer system, the women included in this study were different from those used to develop the original score. Moreover, computer systems used by CPRD and QResearch cover 67% of English practices making our findings generalisable to all women giving birth in the UK. The ethnic diversity in England has been increasing over the last two decades and 86% of the population in England and Wales are white according to the 2011 UK census data [22]. Finally, our large sample size gave us the opportunity to assess model performance in various subgroups and assess

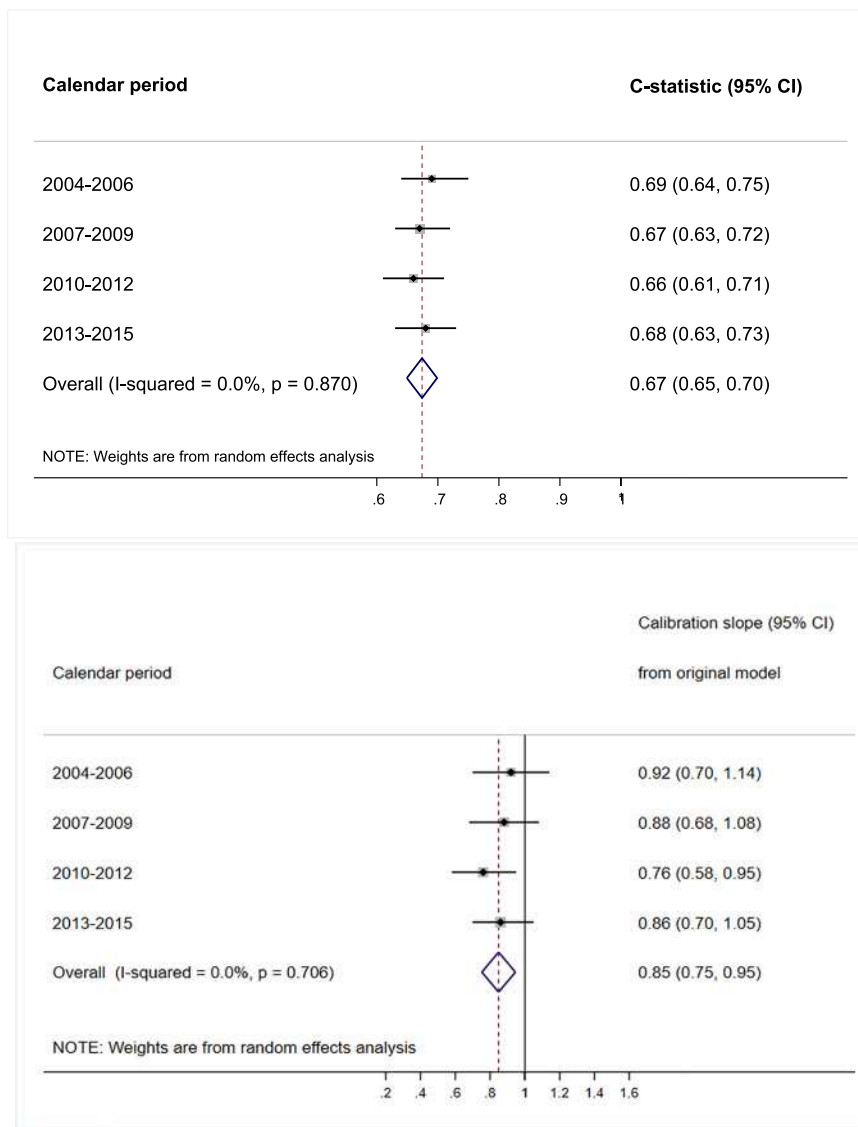


Fig. 3. Model diagnostics by calendar time; a) c-statistic, b) calibration slope (before re-calibration).

heterogeneity based on these factors.

Limitations of this study surround the use of electronic health data for the development and validation of risk prediction models some of which have been previously highlighted [6]. We were unable to individually validate VTE events which occurred in our study due to the terms of the QResearch licence which protects the anonymity of practices which contribute to the QResearch data and individual patients within these practices. Whilst the validation of the algorithm we used to define VTE events excluded pregnant women, we have ourselves conducted methodological work on classification of pregnancy-associated VTE events using electronic sources such as CPRD and QResearch, and found rates of VTE in and around pregnancy that were comparable with existing values obtained from a systematic review on this topic [23]. Nonetheless, we must consider the impact of any misclassification in our outcome event. Misclassification of VTE events both in the development and validation data would attenuate the effect of the predictor variables on VTE risk (assuming that misclassification was unrelated to the predictor variables) and thus bias conclusions towards claiming the maternity clot risk calculator has a weaker prognostic performance than it actually does.

A specific limitation of the present study, which in part affects our VTE algorithm, is that we did not have information on prescriptions

emanating from secondary care and were unable to separate prophylactic from therapeutic doses of LMWH in primary care data. The former meant we were unable to account for women already on thromboprophylaxis during and after childbirth. This is an acknowledged limitation, in general, of developing prognostic models using real world data to identify individuals who should receive a medical intervention. Whilst our subgroup analysis showed that the model performance did not differ noticeably between different time periods, QResearch covered more recent data (2004 onwards) compared to the data used for model development (1997 onwards) and may have downplayed the impact on some of the well-established risk factors due to better awareness of VTE risks. The inability to separate prophylactic from therapeutic doses was due to a combination of incomplete dose data and overlap in therapeutic and prophylactic doses of LMWH preparations as dose is determined from body weight. Therefore, we cannot rule out that primary care prescriptions picked up by our algorithm were for women receiving 6 weeks prophylaxis due to a previous VTE (incorrect inclusion in our study cohort). However, this would only have resulted if the VTE code for their previous event was not recorded in our data. Alternatively, some women were receiving VTE prophylaxis for other reasons during pregnancy and according to RCOG guidelines the same women would receive prophylaxis for 6 weeks post-delivery. If an unconfirmed VTE

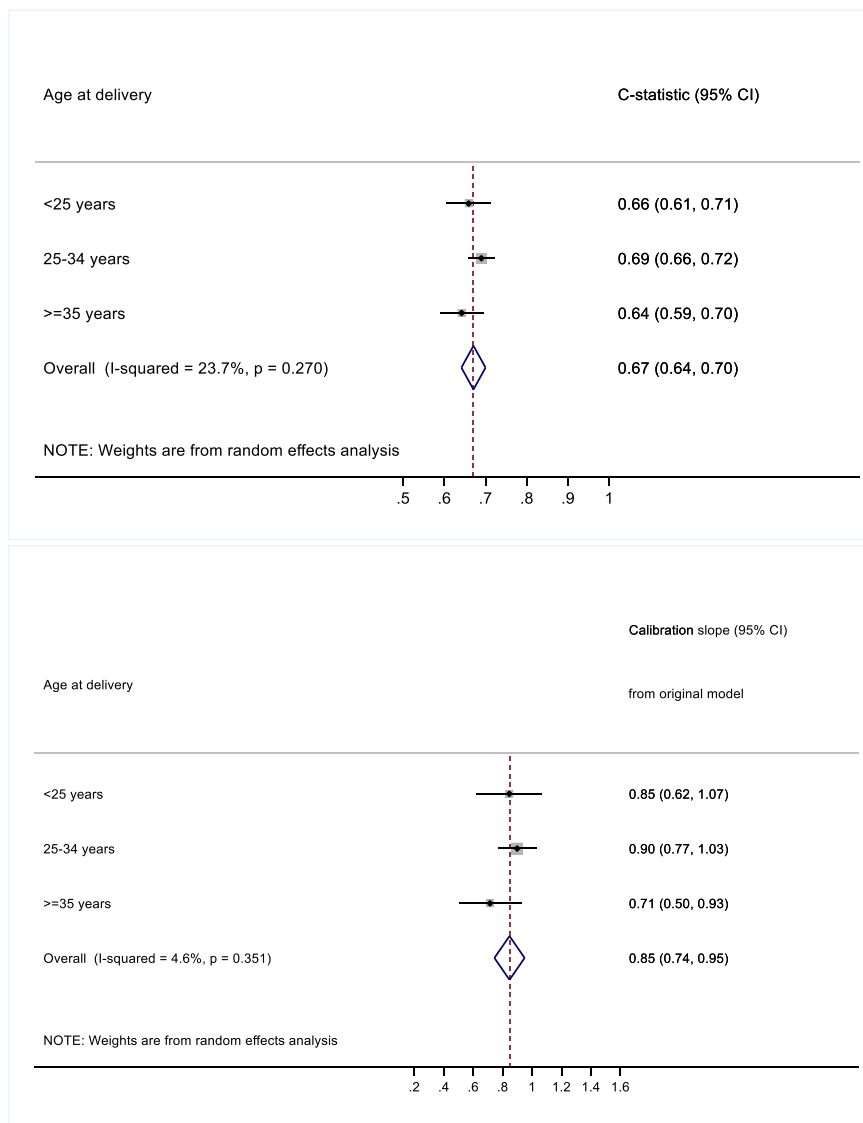


Fig. 4. Model diagnostics by age at delivery; a) c-statistic, b) calibration slope (before re-calibration).

Table 2

Comparing the Maternity Clot Risk with the existing RCOG thromboprophylaxis guideline from the original model (in imputed dataset 1, number of deliveries = 700,185, number of VTE events = 549).

Statistics	Based on RCOG guideline	Based on Maternity Clot Risk <sup>a</sup>
Total no (%) postpartum women warranting thromboprophylaxis	248,983 (35.6)	249,265 (35.6)
Observed VTE events	312	325
Mean predicted risk per 10,000 deliveries	13.2	14.0
Sensitivity (%)	56.8 (52.6–61.0)	59.2 (55.0–63.3)
Positive predictive value (%)	0.13 (0.11–0.14)	0.13 (0.12–0.15)
Specificity (%)	64.5 (64.3–64.6)	64.4 (64.3–64.5)

<sup>a</sup> Women with a risk of VTE of 6.77 per 10,000 deliveries or above would be eligible for pharmacological thromboprophylaxis.

code was included post-delivery in this instance, this would be picked up by our algorithm (false positive VTE). Less than 1% of women in our cohort received any anticoagulation during the pregnancy itself, so the potential impact of this on our findings is likely to be minimal. Whilst all

these specific limitations could be overcome through further validation in a prospective study which formally adjudicates VTE events, such studies are liable to be smaller and less representative of a maternal population than those which make use of administrative health data.

Further limitations include that more than 17% of women had missing values on their pre-pregnancy BMI and their baby's birth weight. This is an improvement from our previous CPRD study and we used multiple imputation technique to minimise the risk of bias associated with missing data. Second, both CPRD and QResearch use a similar coding system (Read code version 2). There is another computer system used in England to record patient consultations (SystemOne) that uses a slightly different coding system (clinical terminology version 3). It is possible that VTE events around pregnancy may be coded differently in practices using SystemOne and so our model performance may not generalise to these practices. However, these practices only represent a small proportion of all practices in England at the present time. Third, there was some miscalibration when applying the Maternity Clot Risk in the QResearch population; indicating some overestimation of risk for women with high-predicted values. However, in the sensitivity analyses the re-calibrated score produced very similar results to the main analyses, indicating that the potential miscalibration had very little impact on the overall predictive performance of the score. Finally, the score was

developed for women without history of VTE therefore cannot be applied to women with a previous VTE or with a known high-risk hereditary thrombophilia. Any woman who has had a VTE previously would be considered high risk by the RCOG and receive thromboprophylaxis for at least 6 weeks, regardless of other risk factors. Routine testing for thrombophilia is not commonplace in the UK with many women being diagnosed after a blood clot has occurred. Therefore, whilst we acknowledge the inability of our model to make predictions based on this, we believe it has less relevance in the identification of intermediate risk women who would receive thromboprophylaxis for 10 days based on RCOG guidelines.

#### 4.3. Cohort comparison

In this validation we were able to test our model using data from a higher number of deliveries using QResearch than those originally used to develop the model from the CPRD [6]. Whilst most of the baseline characteristics were broadly similar across both databases, some differences were observed. In particular, women in QResearch had lower incidence of pre-eclampsia/eclampsia, postpartum haemorrhage and higher mean BMI. Similarly, the overall rate of VTE during the first six weeks after childbirth was also slightly higher than in the CPRD cohort despite applying the same algorithm. These differences may reflect some variations in the study population between CPRD and QResearch. For example, there is evidence that practices contributing to CPRD are slightly more affluent and have lower all-cause mortality compared to the general population [24]. In contrast, due to a wider coverage, the QResearch population could better reflect the English population demographics. Alternatively, it may reflect variations in the recording of medical events across various regions. In addition, applying the current RCOG postpartum thromboprophylaxis guideline in QResearch identified fewer VTE events compared to it applied in CPRD. This may be due to the difference in the observation time period as QResearch used more recent data. Nevertheless, both QResearch and CPRD cohorts showed that in the UK more than 35% of women qualify for short to long term postpartum pharmacological thromboprophylaxis.

### 5. Conclusion and policy implications

We have carried out a second external validation of the Maternity Clot Risk. Overall, its predictive performance is consistent with its performance in the development CPRD population and is similar across subgroups relating to age, socioeconomic status, region, and calendar period. In addition, re-calibration of the score did not improve its performance considerably. Therefore we recommend using the original score (Box 1).

The two algorithms (RCOG and Maternity Clot Risk) correctly predicted a similar number of VTE events, there was a slightly higher sensitivity with our risk score, which was not statistically different. However, the Maternity Clot Risk allows the flexibility of setting new treatment thresholds based on absolute predicted risks of VTE. If adopted it may help optimise use of LMWH to maximise health gain by better targeting of high-risk groups. In the UK, over 35% of women qualify for pharmacological thromboprophylaxis (based on the current RCOG postpartum thromboprophylaxis guideline) with a corresponding mean VTE risk of 1 in 769 (0.13%) postpartum women (based on Maternity Clot Risk applied in QResearch data). Assuming that low-molecular-weight heparin (LMWH) reduces the risk of VTE by at least 50% (based on trial data in ambulatory patients with cancer [25]), 1538 postpartum women would require LMWH to prevent one VTE event. Increasing this risk threshold would result in a lower number needed to treat and would potentially be more acceptable to the women themselves [26]. For instance, targeting the highest 15% of the population (with a corresponding absolute VTE risk of 1 in 476 (0.21%)) would reduce the number of postpartum women requiring treatment to 952. Of course, any such recommendation will need to carefully take into

account the perspective of the health care providers, practitioners and women and consider the potential benefits and harms of any threshold for which further research is urgently needed. Further validation of the model, especially in populations more ethnically diverse than those previously used to develop and externally validate the model, should be taken into consideration for use of the model in maternity settings worldwide. Finally, whilst we restricted our model to 6 weeks post-delivery as this is the interval over which most postpartum VTE events occur, future work could consider which factors predict later maternal VTE events (beyond 6 weeks).

#### CRedit authorship contribution statement

AAS, JW, LJT and MJG were responsible for the conception and design of the study. LB carried out the data management and modelling with guidance from AAS, JW, LJT, RDR and MJG. C N-P provided clinical (obstetric medicine) input to the study and ensured relevance of the project to the RCOG thromboprophylaxis guidelines. LB and AAS produced the first draft of the manuscript. All authors were responsible for critical evaluation of the manuscript and contributed to subsequent drafts. All authors accept responsibility for the paper as published.

#### Declaration of competing interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: C N-P reports personal fees from Sanofi, other from Leo-Pharma, outside the submitted work; and is the lead developer of the RCOG Green Top Guideline on thromboprophylaxis in pregnancy (37a). AAS is currently an employee at Astra Zeneca. This study was conducted before he commenced his employment at Astra Zeneca and this research does not impact in any way on the role he currently fulfils. None of the other authors have interest which could inappropriately influence the work.

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#### Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.thromres.2021.05.020>.

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