Evaluating patient values and preferences for thromboprophylaxis decision making during pregnancy: a study protocol

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Abstract

Background: Pregnant women with prior venous thromboembolism (VTE) are at risk of recurrence. Low molecular weight heparin (LWMH) reduces the risk of pregnancy-related VTE. LMWH prophylaxis is, however, inconvenient, uncomfortable, costly, medicalizes pregnancy, and may be associated with increased risks of obstetrical bleeding. Further, there is uncertainty in the estimates of both the baseline risk of pregnancy-related recurrent VTE and the effects of antepartum LMWH prophylaxis. The values and treatment preferences of pregnant women, crucial when making recommendations for prophylaxis, are currently unknown. The objective of this study is to address this gap in knowledge.

Methods: We will perform a multi-center cross-sectional interview study in Canada, USA, Norway and Finland. The study population will consist of 100 women with a history of lower extremity deep vein thrombosis (DVT) or pulmonary embolism (PE), and who are either pregnant, planning pregnancy, or may in the future consider pregnancy (women between 18 and 45 years). We will exclude individuals who are on full dose anticoagulation or thromboprophylaxis, who have undergone surgical sterilization, or whose partners have undergone vasectomy. We will determine each participant’s willingness to receive LMWH prophylaxis during pregnancy through direct choice exercises based on real life and hypothetical scenarios, preference-elicitation using a visual analog scale (“feeling thermometer”), and a probability trade-off exercise. The primary outcome will be the minimum reduction (threshold) in VTE risk at which women change from declining to accepting LMWH prophylaxis. We will explore possible determinants of this choice, including educational attainment, the characteristics of the women’s prior VTE, and prior experience with LMWH. We will determine the utilities that women place on the burden of LMWH prophylaxis, pregnancy-related DVT, pregnancy-related PE and pregnancy-related hemorrhage. We will generate a “personalized decision analysis” using participants’ utilities and their personalized risk of recurrent VTE as inputs to a decision analytic model. We will compare the personalized decision analysis to the participant’s stated choice.

Discussion: The preferences of pregnant women at risk of VTE with respect to the use of antithrombotic therapy remain unexplored. This research will provide explicit, quantitative expressions of women’s valuations of health states related to recurrent VTE and its prevention with LMWH. This information will be crucial for both guideline developers and for clinicians.
Background

Pregnancy-associated venous thromboembolism (VTE), which may manifest as pulmonary embolism (PE) or deep vein thrombosis (DVT), is an important cause of maternal morbidity [1]. PE remains the leading cause of mortality in developed countries, accounting for approximately 30% of all maternal deaths [2-5].

Although cohort studies have consistently demonstrated that pregnant women with prior VTE are at increased risk of recurrence [6], they have varied considerably in their estimates of magnitude of risk [7-11]. In the largest prospective study of 125 pregnant women with a single previous episode of objectively diagnosed VTE in whom antepartum heparin was withheld [9], the incidence of antepartum recurrence was 2.4% (95% CI of 0.2 to 6.9%). In subsequently published large retrospective cohort studies, the probability of antepartum VTE in women not given antepartum prophylaxis was approximately 6% [12,13]. The inclusion of women with more than one prior episode of VTE as well as women with pregnancies ending in loss, and the failure to independently adjudicate recurrent events might account for the higher risk of recurrence observed in these retrospective studies. Despite the inconsistency, the overall risk of antepartum recurrent VTE in all studies was less than 10% and confidence intervals around the risk estimates of individual studies were overlapping.

The risk of recurrent VTE in the non-pregnant population is lowest among women whose thrombosis was provoked by a major transient risk factor, intermediate among those with an associated minor reversible risk factor, and highest among those whose thrombosis was provoked by a persistent risk factor or who had an unprovoked event [14-30]. Although thrombophilic abnormalities are risk factors for a first episode of VTE [19], these abnormalities do not appear to play an important role in the risk of recurrence [14,15,19,20,30-40]. Data regarding predictive factors for recurrent VTE during pregnancy are inconsistent and studies have not found a clear association between the presence or absence of transient risk factors or of a definable thrombophilia and the risk of recurrent VTE associated with pregnancy [12,13].

Providing thromboprophylaxis to those women at increased risk of thrombosis can potentially reduce pregnancy-related recurrent VTE. However, prophylaxis during pregnancy is problematic. Warfarin crosses the placenta and has the potential to cause teratogenicity and bleeding in the fetus [41-44]. Prophylactic LMWH does not cross the placenta [45] or increase the risk of serious adverse fetal outcomes [44-54], and does not appear to increase the risk of heparin-induced thrombocytopenia (<0.1%) or heparin-associated osteoporosis (<1%) [46,47,51,55-68]. However, prophylactic LMWH is expensive, inconvenient, uncomfortable to administer, may be associated with an increased risk of major obstetrical bleeding [46], and generally necessitates a planned delivery to permit epidural analgesia [46]. Additionally, women may perceive that LMWH creates an undesirable “medicalization” of their pregnancy.

A Cochrane systematic review [69] identified two randomized trials comparing heparin prophylaxis to placebo or no prophylaxis in pregnant women with prior VTE [11,70]. Both suffered from major methodological weaknesses including very small sample sizes. Current clinical guidelines are based primarily on the observational studies described above and indirect evidence suggesting that LMWH substantially decreases the risk of VTE by approximately 70% in a wide variety of clinical contexts [46,71].

In considering women's choices of thromboprophylaxis during pregnancy, two considerations are of particular importance. First, treatment decisions during pregnancy have implications not only for the health of the mother, but also for the health of the fetus. Second, many women prefer to see pregnancy as a normal part of a healthy woman’s life, rather than as a medical condition. Additional considerations include the burden associated with the management: frequency and route of administration, pain, discomfort, and possible side effects of the medication, and the need, frequency and type of testing associated with a given regimen.

Given that the desirable consequences do not clearly outweigh the undesirable consequences of prophylaxis (or vice versa), the values and preferences of pregnant women should be taken into consideration when making management decisions. Because individuals have different attitudes toward risk, the uncertainty of estimates for both the baseline risk of recurrent VTE and the effects of thromboprophylaxis makes the consideration of individual preferences even more important.

There are two fundamental approaches to “patient-specific” decision-making: (1) a holistic direct choice procedure and (2) utility elicitation from individual patients followed by “patient-specific” decision analysis. The relative merit of these approaches is open to question and few studies have addressed this issue [72]. In the “direct choice” methods, participants are presented with relevant health states and the probabilities associated with occurrence of those health states under alternative management strategies. The most rigorous and widely used experimental direct choice method is the probability trade-off or probabilistic version of the threshold technique and it involves determining the threshold benefit at which patients accept a treatment with fixed undesirable consequences or the threshold toxicity or adverse effects at which patients will decline a treatment with fixed benefit [73,74]. Investigators can
also use other simpler direct choice methods (e.g. decision board) that provide complementary information and can ensure understanding of the probability trade-off.

The second approach to decision-making involves decision analytic modeling using best estimates of the probability of occurrence of relevant health states under the management options being considered. Patients’ ratings of the disutility associated with relevant health states (typically on scales that range from death to full health) then inform the decision model and allow calculations of the quality adjusted life years associated with management options. There are a number of approaches available for eliciting health state evaluations [75]. The standard gamble approach is most consistent with utility theory and is generally preferred by health economists [76,77]. Although the visual analog scale is theoretically less satisfactory than the standard gamble, it is much easier to understand, takes far less time to administer, and has superior psychometric measurement properties than to the standard gamble [78-82].

In atrial fibrillation [83,84], and to a lesser extent in VTE and thrombolytic therapy for stroke [85], investigators have studied patients’ values and preferences. A recently completed systematic review of patient preferences for antithrombotic treatment did not identify any studies of pregnant women [86]. The objective of this study is to address this gap in knowledge.

**Methods**

We will perform a multi-center international cross-sectional interview study at six centers in four countries: two in Hamilton, Ontario, Canada; one in Buffalo, USA; one in Oslo, Norway; and two in Helsinki, Finland.
Study population and eligibility criteria
Pregnant women with a history of lower extremity DVT or PE who are considering thromboprophylaxis to prevent recurrent antepartum VTE constitute the ideal patient population. However, only a small number of such women are expected to present at the six centers over the time frame of the study. Therefore, we will also include women with a history of lower extremity DVT or PE who are planning pregnancy, and women who are 18 to 45 years of age with a history of lower extremity DVT or PE and may at some point be planning a future pregnancy. We will exclude women who are currently receiving thromboprophylaxis or full-dose anticoagulation, have undergone surgical sterilization (tubal ligation or hysterectomy), have a partner who has had a vasectomy, and those unwilling or unable to provide informed consent.

Recruitment strategy
We will prospectively identify women who are currently pregnant or planning a pregnancy as they are referred for counseling, and identify women with a history of VTE with the potential to become pregnant by reviewing patient files. We will approach women referred for consideration of thromboprophylaxis prior to their consultation and will make initial contact with women who are not currently pregnant or planning a pregnancy by letter and then by telephone.

Study maneuvers
A member of the research team will meet with women who express an interest in participating, and will explain the purpose of the study and carry out the informed consent process. If patients decline to participate, we will collect the reason for refusal. If patient consent to the study, we will make arrangements for interviews. We will use standardized scripts developed by the research team to ensure a standard approach across centres and interviewers. Expert and non-expert clinicians and allied health professionals have reviewed and revised the scripts to ensure understandability and readability by a lay person with a grade 9 reading level.

The participant interview
Figure 1 presents the flow of the interview. We will determine patients’ age, educational attainment, and current pregnancy status, as well as specific details about their past venous thromboembolic events (including occurrence of PE or DVT, number of events, date of the last event, presence or absence of precipitating risk factors prior to their event [Table 1]), known hypercoagulable states (Table 1), family history of VTE, type and duration of treatment for their event(s), adverse treatment effects, completeness of their recovery (for example, presence or absence of residual chest pain or shortness of breath, and/or residual leg swelling, pain or discoloration), and presence or absence of prior experience with injection of prophylactic doses of LMWH during pregnancy.

Direct choice exercises
We will determine each participant’s willingness to receive LMWH prophylaxis through direct choice exercises. Women will initially complete what we refer to as the real-life scenario, followed in order by hypothetical scenarios, the visual analog scale, the probability trade-off exercise, a review of their answers, and finally questions to examine their understanding of the scenarios. Use of decision boards will facilitate patient understanding of the direct choice exercises. Descriptions of each phase of the interview follow.

Real life scenario
We will initially present each woman with a decision board that will include the probabilities of developing VTE during pregnancy given the characteristics of her prior VTE (see below). This scenario will contain information similar to that received during standard clinical care but presented in a more systematic manner than that is typical in clinical practice.

Table 1 Potential precipitating risk factors assessed regarding the past venous thromboembolic event Minor transient (i.e. now resolved) risk factors within 8 weeks of initial VTE

| Minor transient (i.e. now resolved) risk factors within 8 weeks of initial VTE | Major transient (i.e. now resolved) risk factors within 8 weeks of initial VTE | Known hypercoagulable state |
|---|---|---|
| Pregnancy | Leg casting | Deficiency of antithrombin, protein C, or protein S |
| Postpartum (defined as 6 weeks after delivery) | Major surgery (>30 minutes; general or spinal anesthesia) | Activated protein C resistance/factor V Leiden |
| Hormonal contraception (birth control pill, patch or needle) | Acute medical illness with hospital admission for ≥3 days | Prothrombin gene mutation |
| Airplane travel (longer than 6 hours) | Immobilization ≥3 days (in bed except to go to washroom) | Anticardiolipin antibody positivity |
| Active cancer | Nonspecific inhibitor |
For this real life exercise, we will have two potential scenarios. We will classify women as low or high risk of recurrence. We define low risk of recurrence as the absence of a known thrombophilia and prior VTE associated with a major transient risk factor within 8 weeks prior to their last event, and higher risk of recurrence as prior unprovoked VTE, VTE associated with a minor transient risk factor within 8 weeks prior to the event or any VTE in association with a known hypercoagulable state. (Table 1) We estimate that the risk of antepartum recurrence for women judged to be at lower risk lies between 0 and 5%, while that for higher risk women ranges between 5 and 10%. We will assume that prophylactic LMWH reduces the risk of antepartum recurrence by approximately 70% [71]. To ensure optimal understanding, we will present the risk of recurrence with and without LMWH prophylaxis in three different ways: table, bar chart and pictograph (Table 2 and Figures 2 and 3).

For the VTE health state, we will instruct women to consider their previous venous thromboembolic event. Women with more than one event will consider their most recent episode. We will instruct women with previous experience in the use of prophylactic LMWH for longer than 2 weeks during pregnancy to consider their previous experience when making a decision. We have prepared a description of the experience of LMWH use throughout pregnancy for women without experience in the use of LMWH prophylaxis during pregnancy (Additional file 1: Appendix 1). To ensure clinical verisimilitude, clinicians with considerable experience in the use of LMWH in pregnant women prepared the description and experts in obstetrics and thromboembolism reviewed them.

After reviewing this information, we will ask participants to decide whether or not they are willing to take LMWH during their pregnancy (for those who are pregnant) or whether they would be willing to do so in a subsequent pregnancy. Women currently pregnant and those referred for consideration of prophylaxis with a future pregnancy will meet with their health care provider and may wish to discuss the information provided with other people (for example, family members, friends, family physician, midwife or obstetrician). We will record the final decision after these additional consultations.

**Hypothetical scenarios** We will provide study participants with three standardized scenarios in which the baseline risk of recurrent VTE, with and without the use of LMWH (assuming a 70% relative risk reduction) [71], will be varied (Table 3). Interviewers will show each woman a decision board with pictograms representing a low, followed by a medium, then a high risk of recurrent VTE. Participants will express their willingness to use LMWH given the associated burden (either based on their own prior experience or the standardized description [Additional file 1: Appendix 1]), and the absolute

| Probability of developing a blood clot during your pregnancy | Without low molecular weight heparin use | With low molecular weight heparin use |
|-------------------------------------------------------------|----------------------------------------|-------------------------------------|
| Probability of NOT developing a blood clot during pregnancy  | 90-95 in 100                           | 97-99 in 100                        |

**Figure 2** Example of bar chart presentation for women considered at high risk of recurrent VTE during pregnancy.
magnitude of VTE risk reduction associated with the varying VTE recurrence risks.

**Probability trade-off** Interviewers will undertake probability trade-off exercises with “ping-ponging” to determine participant thresholds for accepting LMWH prophylaxis [73,74]. We will use a scenario based on the absolute effects of LMWH versus no prophylaxis for prevention of recurrent VTE during nine months of pregnancy. The interviewer will systematically vary the risk of VTE with LMWH prophylaxis (alternating between high and low risks) to determine the minimum acceptable reduction in the risk of VTE with prophylaxis at which the participant would agree to initiate LMWH. Based on the 95% CI surrounding the incidence of antepartum VTE in high risk patients reported in the prospective cohort study [9], we estimate that the upper bound of risk of recurrent VTE is 16 out of 100 without prophylaxis. Therefore, we will set this risk fixed on one side of the flipchart and will start offering probabilities ranging from 16 fewer VTE events per 100 pregnancies (maximum absolute risk reduction) to 0 less VTE events (same VTE risk as no prophylaxis) on the other side of the chart.

**Visual analog scale (feeling thermometer)** Interviewers will determine the value patients place in relevant health states using a visual analog scale called the Feeling Thermometer (FT) [75]. When making ratings using the FT, women choose the score on the thermometer that represents the value they place on the health state they are evaluating. The FT is anchored at death (0) and full health (100). We will ask participants to consider (i) the health state of pregnancy with LMWH prophylaxis using the standard description or their previous experience (for those with two weeks or more of prophylactic LMWH during pregnancy), (ii) a pregnancy with their own most recent VTE experience, (iii) a standardized health state with a pregnancy-related DVT, (iv) a standardized health state with a pregnancy-related PE and (v) a standardized health state representing an obstetrical bleed (Additional file 1: Appendix I).

**Check for consistency and understanding** After presenting the descriptions and recording patient responses, interviewers will review participant responses to the various exercises to check for consistency in the participant’s choice. When interviewers identify inconsistencies, they will offer participants a chance to review and change their responses, avoiding any suggestion that responses

### Table 3 Hypothetical scenarios with variable baseline risks of recurrent VTE and estimates of LMWH effectiveness

| Risk of recurrence | Low risk | Medium risk | High risk |
|--------------------|----------|-------------|-----------|
| **Without LMWH**   | 4 in 100 | 10 in 100   | 16 in 100 |
|                    | (96 in 100 will not) | (10 in 100 will not) | (84 in 100 will not) |
| **With LMWH**      | 1 in 100 | 3 in 100    | 5 in 100  |
|                    | (99 in 100 will not) | (97 in 100 will not) | (95 in 100 will not) |
should be changed. The reasons for any apparent inconsistencies will be determined and recorded. Following this consistency check, interviewers will ask participants two standardized questions to evaluate their understanding of the information provided during the interview [Additional file 1: Appendix II]. Interviewers will also provide a rating of the extent to which they believe the respondents had a clear understanding of the questions and their confidence in this assessment.

Outcomes
The primary outcome measure will be the minimum threshold reduction in VTE risk in the probability trade-off exercise at which women switch from declining to accepting LMWH prophylaxis (which we will refer to as the “VTE threshold”).

Secondary outcomes will include:
(i) women’s willingness to take prophylactic LMWH according to their classification as either high or low risk for recurrence,
(ii) women’s willingness to take prophylactic LMWH for each of the three hypothetical scenarios
(iii) Utilities for each of the five health states assessed in the FT (the burden of prophylactic LMWH use during pregnancy – either the standardized experience or the patient’s prior experience, a pregnancy with the participant’s own most recent VTE, a standardized description of pregnancy-related DVT, a standardized description of pregnancy-related PE, and a standardized description of an obstetrical bleed [Additional file 1: Appendix I]).

Analysis
Baseline characteristics
We will describe age, educational attainment, pregnancy status, number and characteristics of previous venous thromboembolic events, presence or absence of precipitating risk factors, and prior experience with prophylactic LMWH injections using means and standard deviations or proportions, as appropriate.

Primary and secondary outcome(s)
We will calculate the mean threshold reduction in VTE at which women were willing to accept use of LMWH and the 95% confidence interval around the mean. We will calculate the proportion of women who are willing to take prophylactic LMWH and the associated 95%

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Figure 4 Markov state transition decision model used in analysis. This diagram depicts the model used in the analysis. In the figure on the top, the square on the left represents a “decision node” from which 2 branches, representing alternative management strategies emanate. Each strategy leads to the same Markov node, represented by a square with the “∞” symbol. The branches leading from the Markov node represent the various potential health states that patients pass through during the model simulation. Although the potential states for each strategy are the same, the initial distribution among states and probabilities associated with transitions between states will differ between strategies. The figure on the bottom is the modeled adverse events. During each time period or “cycle”, modeled patients are at risk for various adverse events. Round nodes represent the chance events. For each patient, the sequence of outcomes at these “chance” nodes will determine the state at which they begin the next cycle.
confidence intervals The mean and standard deviation of the visual analog scale ratings for each health state will also be determined.

Multiple variable linear regression will be undertaken to explore determinants of the VTE threshold, as determined in the probability trade-off exercise. In this analysis, VTE threshold will be the dependent variable and the independent variables will be the previous experience of VTE (categorized as severe, moderate, or mild), previous experience with prophylactic LMWH (yes, problematic; yes, no problems; no prior experience), level of education (some postsecondary versus no post-secondary) and study site.

We will perform a standard multiple variable logistic regression to explore the determinants on the choice for or against LMWH prophylaxis in the real life scenario. The dependent variable will be the choice for or against LMWH prophylaxis and the independent variables will be the previous experience of VTE, previous experience with LMWH prophylaxis, level of education, study site, and high or low risk for recurrence.

A hierarchical logistic regression in which the dependent variable will be the choice to receive or not receive LMWH from the results of the hypothetical scenario exercise will also be undertaken. The independent variable within the first level will be the magnitude of the absolute risk reduction (high, moderate and low). The second level, nested within the absolute risk reduction will include previous experience of VTE, previous experience with LMWH prophylaxis, level of education, study site, and high or low risk for recurrence.

A hierarchical logistic regression in which the dependent variable will be the choice to receive or not receive LMWH from the results of the hypothetical scenario exercise will also be undertaken. The dependent variable within the first level will be the magnitude of the absolute risk reduction (high, moderate and low). The second level, nested within the absolute risk reduction will include previous experience of VTE, previous experience with LMWH prophylaxis, level of education, study site, and high or low risk for recurrence.

Table 4 Guidelines for frequencies of inconsistent responses favoring review of study scripts and presentation tools

| Frequency of inconsistent response | % | Lower bound of 95% CI |
|-----------------------------------|---|----------------------|
| > or = 5/10                       | 50.0                      | 23.7                  |
| > or = 8/20                       | 40.0                      | 21.9                  |
| > or = 10/30                      | 33.3                      | 19.2                  |
| > or = 12/40                      | 30.0                      | 18.1                  |

For all key analyses above, we will compare results and the pattern of responses in women who answer the “understanding” questions correctly and those who do not. A similar comparison will also be made according to the interviewer’s impression of patient understanding. If the results differ between women categorized as understanding and not understanding, our primary analysis will focus on women with an apparent high level of understanding.

Finally, we will perform a “personalized decision analysis” for each woman in the study. We will update a prior decision analytic model examining prophylactic LMWH in pregnant women with a history of prior VTE (Figure 4) [87], using each women’s visual analog scale utility assessments for the relevant outcomes of DVT in pregnancy, PE in pregnancy, obstetrical hemorrhage and the burden of LMWH prophylaxis in pregnancy. Given that the risk of heparin-induced thrombocytopenia (<0.1%) and heparin associated osteoporosis (<1%) are either not increased or minimally increased with prophylactic LMWH [65,68], we will not consider these health states in the decision analysis. We will place the health state utilities in the decision model for each woman’s current situation and the three hypothetical scenarios. We will compare the decision model’s results with the VTE threshold selected in the probability trade-off tasks.

Sample size
Previous research from our group in patients with atrial fibrillation [83], and from other groups studying non-pregnant women with prior VTE [85], suggests that moderately precise estimates of patient preference can be obtained with sample sizes of approximately 100 participants. In the most relevant recent experience, co-investigators on the current project enrolled 96 patients with risk factors of atrial fibrillation [85]. The primary
endpoint of that study was the threshold number of bleeds patients were willing to accept over 2 years given a reduction of 3 strokes over a 2-year period. This is directly analogous to the primary endpoint of our study, the VTE threshold. In the atrial fibrillation study, the mean number of bleeds that patients were willing to accept was 15. With a sample size of 96 patients, the confidence interval around the mean number of acceptable bleeds was 12 to 18, which we judged to be an acceptable range of uncertainty. The range of bleeds that patients chose varied over an extremely wide range (from 0 to 100). The entire range of possible VTE prevented for LMWH to be acceptable in the current study will range from 0 to 16, a much narrower range, which is certain to result in much smaller variability, and thus an appreciably smaller confidence interval. We are, thus, confident that our projected sample size of 100 will yield a sufficiently precise estimate of our primary outcome, the mean number of VTE prevented required to tolerate LMWH.

Safety and ethical considerations
This study does not pose any safety risks to participating women. Research Ethics Board approval has been obtained in Canada (Hamilton Health Sciences), Finland (The Coordinating Ethics Committee of Helsinki and Uusimaa Hospital District) and USA (Institutional Review Board of the State University of New York at Buffalo), and will be obtained from all other participating sites. All women will provide written informed consent prior to participating in this study.

Discussion
Producing guidelines for management of pregnant women with prior VTE is problematic because there is little reliable information in the literature to guide the value and preference judgments necessary for making recommendations. Clinical management is challenging because of the lack of a standardized approach to presenting information and eliciting patients’ choice. Our study will be the first to explore the preferences of women at risk of pregnancy-related VTE with respect to the use of antithrombotic therapy.

Our study has several strengths. We will address some of the limitations of the previous studies in the field of decision-making. These studies using direct choice methods, and in particular the probability trade-off, have shown many extreme choices that raise doubts about whether all patients understood the decisions they were making [84,88]. We piloted standardized scripts that were reviewed by expert and non-expert clinicians, and allied health professionals to ensure appropriate readability and understandability. We are presenting probability information in three different ways. We have trained research assistants and will undertake quality control of the data as it accumulates.

We will check for consistency in the participants’ choice to either take or not take LMWH for a given probability of DVT. Participants will have discrepancies brought to their attention and will have the opportunity to change their responses. We will use standardized questions to test understanding of the information presented and the interviewer will provide a rating of the extent to which she or he believes the respondent had a clear understanding of the questions and their confidence in this assessment. Our analysis will include an exploration of whether patterns of responses differ (1) between those with an apparently high level of understanding and those who do not, (2) between those whose answers are consistent and those with inconsistent answers, and (3) between those who, according to the interviewer’s impression of patient understanding, show a correct understanding and those who do not. Finally, our multicenter international design limits the influence of local views about the use of LMWH prophylaxis for prevention of recurrent pregnancy-associated VTE and enhances generalizability.

Our study has some potential limitations. Our sample includes women not contemplating becoming pregnant. There are logistical challenges in prospectively identifying such a group and in ensuring their participation prior to their receiving other information about their choice. We believe that the inclusion of these women is a reasonable given the proximity to the necessity of a choice for these women and that their responses will reflect their true, thoughtful preferences. We will, nevertheless, evaluate whether the responses between women who are pregnant or planning a pregnancy are different to those who are not.

The research outlined in this protocol aims to provide explicit, quantitative expressions of women’s valuations of health states related to recurrent VTE and its prevention with LMWH. This information will improve the quality of formal decision analyses evaluating whether the effectiveness of prophylaxis outweighs risks and burden of therapy and provide relevant information for guideline developers providing recommendations for clinical practice.

Our protocol addresses some of the limitations of the previous studies in the field of decision-making and describes innovative approaches to checking for understanding and systematically exploring inconsistencies. The methods described in this protocol are novel and rigorous, and could be utilized immediately by those working in the area of optimizing patients’ medical decision-making.
Additional file

Additional file 1: Appendix I. Health States (1–4). Appendix II. Questions to Assess Understanding.

Competing interests
SMB has received honoraria from Leo Pharma Inc; Sanofi-Aventis Canada and Pfizer Canada (all manufacturers of LMWH). The other authors declare no competing interests.

Authors’ contributions
GHG, SMB and PA-C conceived the study. GHG, SMB, PA-C, SE, KAOt, ME and EA contributed to the study design. PA-C and SE completed the first draft of the manuscript and are co-first authors. All authors read, critically revised and approved the final manuscript to be published.

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