STUDY PROTOCOL

Effects of tranexamic acid on platelet function and thrombin generation (ETAPlaT): WOMAN trial sub-study [version 1; peer review: 2 approved]

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Abstract

Background. Postpartum haemorrhage (PPH) is a leading cause of maternal death. Tranexamic acid (TXA) has the potential to reduce bleeding and a large randomized placebo controlled trial of its effect in women with PPH (The WOMAN trial) is underway. TXA might also affect coagulation factors and platelets. Objectives. To examine the effect of TXA on thrombin generation, platelet function, fibrinogen, D-dimer and coagulation factors in women with PPH. Methods. We will conduct a sub-study within the WOMAN trial. Women with clinically diagnosed primary PPH after vaginal or caesarean delivery are eligible for inclusion. Blood samples will be collected at baseline and 30 minutes after the first dose of study treatment. Using platelet poor plasma we will measure thrombin generation, fibrinogen, D-dimer and coagulation factors in women with PPH. Outcomes. The primary outcome is the effect of TXA on thrombin generation. Secondary outcomes include the effect of TXA on platelet function, fibrinogen, D-dimer and coagulation factors.

Keywords

Postpartum Hemorrhage, Tranexamic Acid, Coagulation Factors, Platelet function
In patients with chronic renal failure, administration of TXA corrected platelet aggregation defects (Mezzano et al., 1999), and as result improved platelet function. Also, Sabovic et al. (2005) reported that platelet dysfunction in haemodialysis patients was effectively corrected with long term administration of TXA at a low dose.

### Methods and study design

#### Objective

The study will assess the effect of TXA on thrombin generation, coagulation factors and platelet function in women with PPH, in particular, the effects of TXA on thrombin generation, factor V (FV), factor VIII (FVIII), Von Willebrand factor (vWF), fibrinogen, D-dimers and platelet function. We hypothesize that by inhibiting plasmin, TXA will decrease thrombin generation, and modify platelet activity, FV, FVIII and vWF levels, in patients with PPH.

#### Trial design

The WOMAN-ETAPlaT is a sub-study of the World Maternal Antifibrinolytic Trial, an international randomized, double blinded, placebo-controlled trial. As a sub-study, there are no changes to the study design of the WOMAN trial, but for the specific design of the sub-study there are some additional examinations and laboratory tests.

#### Participants and eligibility criteria

**Inclusion criteria:**
Participants in the study are women ≥18 years old with a clinical diagnosis of postpartum haemorrhage (PPH) after vaginal or caesarean birth. All available treatment for PPH should be given, and inclusion in the study should be considered as soon as possible after patient consent has been obtained. Clinically diagnosed PPH can be established with one of the following conditions: amount of blood lost following vaginal birth is >500mL or >1,000mL after caesarean birth; or enough blood loss has compromised the patient’s haemodynamic status. **Exclusion criteria:** Patients diagnosed with PPH, for whom the physician believes that there is a secure indication or contraindication for use of TXA, should not be randomised.

#### Study setting

For the conduct of this sub-study, a single site has been chosen, the Obstetric Gynaecology University Hospital “Koço Gliozheni” in Tirana, Albania. This is justified because PPH is the main cause of maternal mortality in Albania. The hospital offers tertiary health care and is a national referral hospital for other obstetric gynaecology hospitals in the country.

#### Interventions

Patients with clinically diagnosed PPH and fulfilling the eligibility of the WOMAN trial will be randomized into the study after the administration of the standard treatments for PPH. As soon as primary PPH is diagnosed, the study drug (TXA or placebo) will be administered as soon as possible alongside all other clinically indicated treatments for PPH. Each trial treatment pack contains two doses, and each dose contains two ampoules of 500mg TXA or placebo (sodium chloride 0.9%).

#### Outcomes

**Measures of the primary and secondary outcomes of ETAPlaT study**

The primary outcome will be the effect of TXA on thrombin generation [thrombin generation assay (TGA) parameter – endogenous thrombin potential (ETP)]. Secondary outcomes will include...
the effect of TXA on platelet function (Multiplate® tests: ADPtest and TRAPtest), fibrinogen, D-dimer and FV, FVIII, vWF levels, and other TGA parameters [lag time (LT); time to peak (TtP); and peak height (Ph)]. Levels of all parameters will be assessed on two venous blood samples, collected at the baseline and between 30±15 minutes after the first dose of study treatment is given.

Further secondary outcomes will include assessment of the relationship between these parameters, patient demographics and clinical data (in the time frame from randomization to the patient discharge from the hospital or until 42 days).

Outcomes of the main study WOMAN trial. The WOMAN trial will not provide information about platelet function, thrombin generation, fibrinogen, D-dimers and coagulation factors V, VIII and vWF in patients receiving TXA/placebo. However, the most reliable estimates of the effects of TXA on the outcomes for women with PPH will be provided by the WOMAN trial.

Sample size
We aim to assess if TXA, when compared to a placebo, has an effect on change over time on the TGA parameter, ETP. Therefore, the sample size calculation is based on this parameter. Patients are expected to be equally allocated between the two intervention groups; 88 patients per group (total, 176 patients) are needed to detect the difference of ETP 243nM/min, with a change between the groups at a 5% significance level with a power of 80%.

Randomization and blinding
Patients with a clinical diagnosis of PPH, who are eligible for the WOMAN trial, after having completed the informed consent procedures, will be randomly assigned to receive either TXA or placebo. Patients randomised in the trial will receive the study drug (TXA/placebo) by intravenous injection. The lowest available number of the pack will be used first and then subsequently other packs from the box that contains 8 study treatment packs. The blinding process will be carried out by an independent supplier, who will mark the study drug with the blinded label. The vials and packages will be the same, and the randomisation number will be used as the pack identification.

Procedures before/after the study treatment administration, blood sample collection and analysis
Immediately after patient is randomized, two test tubes (3mL and 5mL) of venous blood will be taken and the first dose of trial treatment will be administered. About 30±15 minutes after the administration of the first dose of trial drugs, two more test tubes (3mL and 5mL) of venous blood will be collected. If a woman continues to bleed and a second dose of trial treatment is required, the second blood sample will be taken before this is given (Figure 1).

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**Figure 1. Algorithm of the WOMAN-ETAPItaTsub-study.** Black boxes: Standard procedure of the WOMAN trial. Red boxes: Additional procedures required for the ETAPItaT sub-study.
Samples of the 3mL tubes (hirudine) whole blood will be analyzed immediately at the hospital in Tirana, Albania for platelet function using a Multiplate Platelet Function Analyzer (Dynabyte GmbH, Munich, Germany).

Samples of 5mL tubes (sodium citrate 3.2%) will be centrifuged immediately at 3000xg for 20 minutes, and platelet poor plasma samples will be divided into two aliquots and preserved at -80°C. These will be transferred to dry ice for later analysis at the Institute of Laboratory Medicine, German Heart Centre in Munich, Germany for TGA, fibrinogen, D-dimers, and FV, FVIII and vWF levels. The temperature storage conditions of the plasma samples will be monitored and recorded daily.

As part of the routine protocol of the hospital, a blood sample for complete blood count (CBC) is taken from every woman in labour prior to delivery. In cases of PPH randomized in the study, another CBC is performed after 4 hours and within 12 hours, depending on the severity of PPH. The pre-post randomization difference in CBC parameters, such as haemoglobin or haematocrit drop, will be evaluated.

Standard operating procedures for handling, storage and analysis of the blood samples and the process for transferring the data to the Trial Coordinating Centre (TCC) are developed by each responsible Institution, and are approved by TCC. Copies are available from the Trial Master File.

Ethical considerations, recruitment and consent
Ethical approval for WOMAN ETAPlaT protocols was obtained 28.10.2013 (ref. 6518) from the London School of Hygiene and Tropical Medicine (LSHTM) Ethics Committee in London, United Kingdom, and by the National Ethics Committee in Tirana, Albania on 11.07.2013 (ref. 62) and amendment on 01.12.2014 (ref. 81). PPH is an obstetric emergency and it is not possible to predict which women that will have this complication. In this emergency situation it is important to carry out an informed consent procedure, according to regulatory requirements, and adhere to the ICH-GCP and Declaration of Helsinki, for patients to be eligible for the study. The consent procedure will be performed as per the approved procedure for the WOMAN trial.

Briefly, advanced information about the trial will be given to pregnant women, which will make the recruitment and consent procedure easier. The consent process will depend on the clinical emergency and the patient’s physical or emotional state. If the woman is fully competent, the healthcare giver will provide information and discuss with her about the study and obtain informed written consent (Supplementary File 1 and Supplementary File 2). If the woman’s mental or physical capacity does not allow her to give consent, the woman’s relative or representative, if available, should be informed about the trial and informed written consent will be obtained by them (Supplementary File 1 and Supplementary File 3). Where no relatives or representative are available, a waiver of prior written consent has been approved for the WOMAN trial. Where a waiver of prior written consent is used, the woman, relative or representative should be asked for consent for the continuation of the trial procedure.

To minimize the need for multiple information sheets and consent forms, one form, which combines the WOMAN trial and the WOMAN-ETAPlaT study, will be used.

Data collection and monitoring
Data collection
All study data will be gathered on the WOMAN trial entry data form (baseline information) and outcome data form (patient death, discharge from the hospital or 42 days after randomization). Additional information will be collected with ETAPlaT data collection forms (Supplementary File 4). All study data will be sent electronically to the main database of the TCC in London, UK. ETAPlaT data collection forms will include the following:

- Time when the blood sample was taken and laboratory analysis was started and completed, and any technical problems with analysis;
- Laboratory results of Multiplate analysis (ADPtest, TRAPtest), TGA, D-dimer, fibrinogen, FV, FVIII and vWF;
- Treatment given that may affect coagulation;
- Other data about the woman’s parity, BMI, gestational age at birth, maternal pre-existing conditions (anaemia, cardiac disease, haemoglobinopathy, chronic hypertension, renal disease, treatment or prophylaxis with antithrombotic drugs, or history of previous thromboembolism), pregnancy related conditions (preeclampsia, diabetes, infection, placental abruption), labour induction/augmentation and duration, anaesthesia (epidural, spinal or general), birth weight, additional doses of uterotonics (oxytocin/methergine).

Analysis
Analysis will compare two groups of randomized patients, allocated in TXA group or placebo group on an intention-to-treat basis, regardless of treatment received, and outcomes will be presented as appropriate effect estimates and 95% confidence intervals.

In analysis of primary outcome, TXA and placebo groups will be compared regarding the TGA parameter, ETP of baseline and follow up values. The same analysis will be performed for secondary outcomes such as platelet function (ADPtest and TRAPtest), coagulation factors (FV; FVIII; vWF, Fibrinogen and D-Dimer) and other TGA parameters (LT, Tp, and Ph). A subgroup analysis for primary outcome will be based on main risk factors for PPH, such as uterine atony, placental factors and genital trauma.

A detailed statistical analysis plan (SAP) will be finalized prior to the sub-study unblinding for final analysis. All statistical analysis will be conducted with R and STATA statistical package.
Limitations of the study
One limitation of this sub-study is the small number of the patients that will be recruited. Another potential limitation is withdrawal of participation by patients from this study, or risk of loss of contact in case of complications after discharge from the hospital and within the period of 42 days after the birth.

Trial Steering Committee (TSC) and Data Monitoring Committee (DSC)
A TSC for the WOMAN trial is in place and agreed to the conduct of this sub-study. Decisions of the TSC can influence immediately on the continuation of the WOMAN-ETAPlaT study. Information about the WOMAN-ETAPlaT study will be reported routinely to the TSC by Haleema Shakur. An independent DMC is already in place for the WOMAN trial. Adverse events, which occur during the study period and are related directly to the WOMAN-ETAPlaT study, will be reported to the DMC.

The sub-study had its responsibilities coordinated by the TCC, which may attribute responsibilities to third parties, and will be detailed by suitable arrangements.

Trial status
Recruitment for the WOMAN-ETAPlaT sub-study started on November 2013 and the final participant follow-up was completed in March 2015. Data cleaning is ongoing.

Supplementary files
Supplementary File 1: Information sheet for patient and their representative (Albanian and English).
Click here to access the data.

Supplementary File 2: Patient consent form (Albanian and English).
Click here to access the data.

Supplementary File 3: Representative consent form (Albanian and English).
Click here to access the data.

Supplementary File 4: WOMAN-ETAPlaT data collection forms.
Click here to access the data.

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Author contributions
KD, UM, OG, MD, BS, HS, IR and DB, have made considerable contributions in drafting the protocol, and reviewing it critically for important intellectual content. UM and PE performed the statistical analysis of the sub-study, IT and SC reviewed the protocol and oversaw patient recruitment. All authors approved the final version of the manuscript.

Competing interests
No competing interests were disclosed.

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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

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Thanks to Prof. Wolfgang Schramm and Prof. Beverly Hunt for their considerable input in drafting and reviewing the protocol. We thank Dr. Lena Mokelke (Clinical Trial Service Center, M4 project) for her support in developing a GCP conform trial protocol and making the project compliant with regulatory requirements.
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The authors have published their protocol of a WOMAN trial sub-study.

WOMAN is a pragmatic, randomized, double-blinded, placebo-controlled trial in women with a clinical diagnosis of postpartum hemorrhage (PPH) and is intended to determine reliably the effect of the early administration of TXA on death, hysterectomy, and other morbidity (surgical intervention, blood transfusion and risk of nonfatal vascular events). Recruitment of the 20,000 planned randomized women has been completed as well as statistical analysis.

The aim of this sub-study is to assess the effect of TXA on thrombin generation but also on platelet function, fibrinogen, D-dimer and coagulation factors in women with PPH. The authors hypothesize that TXA will decrease thrombin generation by inhibiting plasmin and modify platelet activity, coagulation factors V and VIII and Von Willebrand levels in women with PPH. The protocol of the WOMAN trial has been previously published elsewhere.

In this sub-study, it is planned, only on a single site located in Albania, to obtain blood samples immediately after randomization and before the administration of the first dose of trial treatment as well as about 30±15 minutes after the administration of the first dose of trial drugs (but before the second dose of trial treatment if required).

Platelet function will be assessed immediately on site using Multiplate® tests, while other samples will be centrifugated and preserved at -80°C to be transferred in Germany for later analysis of Thrombin generation Assay, fibrinogen, D-dimers, coagulation factors V and VIII and Von Willebrand levels.

The protocol is well written, original and will provide useful, interesting and likely robust data related to the impact of TXA on hemostasis of PPH women. I have only minor comments:

- The authors should explain why only one site participates to the WOMAN trial sub-study.
- The primary outcome should be clinically (and in pathophysiological manner) justified:
Endogenous Thrombin Potential difference of 243nM/min.

- The sample size should be based on literature or the authors should mention that, as this analysis is exploratory, it is somewhat arbitrary.
- The authors should clearly note that analysis will be performed blindly.
- The authors should clearly mention what material laboratory will be used to assess Thrombin generation Assay, fibrinogen, D-dimers, coagulation factors V and VIII and Von Willebrand levels.

**Competing Interests:** I am the main investigator of the TRAAP (TRAnexamic Acid for Preventing postpartum hemorrhage after vaginal delivery) study: a multicenter randomized, double-blind, placebo-controlled trial which aims to determine whether a low dose of TXA (1 g) after vaginal delivery reduces the incidence of postpartum hemorrhage (Sentilhes L, Daniel V, Darsonval A, Deruelle P, Vardon D, Perrotin F, Le Ray C, Senat MV, Winer N, Maillard F, Deneux-Tharaux C. Study protocol. TRAAP - TRAnexamic Acid for Preventing postpartum hemorrhage after vaginal delivery: a multicenter randomized, double-blind, placebo-controlled trial. BMC Pregnancy Childbirth. 2015 Jun 14;15:135. doi: 10.1186/s12884-015-0573-5.).

**I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.**
The title is appropriate for the content of the article and the abstract represent a suitable summary of the work. The trial is well described regarding the background, the method and the expected results. The conclusions are sensible, balanced and justified on the basis of the results of the study. This paper is of interest to explore the hypothesis of TA-related thrombin generation inhibition and I recommend indexing it with minor corrections.

- One suggestion and the aim of our TRACES trial [CT 02797119] should be to analyze simultaneously the plasmin inhibition to link the force of the thrombin procoagulant effect and plasmin TA inhibition.

- Please clarify which laboratory will perform the biological tests and the expected delay between the samples and the LAB analysis.

- Please clarify if the biological data analysis will be blinded to the allocation?

I believe the data is in a usable format/structure and there is sufficient information provided for the experiment to be replicated.

**Competing Interests:** No competing interests were disclosed.

We confirm that we have read this submission and believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

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**Comments on this article**

**Version 1**

Reader Comment 03 Jul 2017

Antonia Zapf, University Medical Center Göttingen, Germany

In the sub-study of the WOMAN trial the investigators want to assess the effect of the study treatment TXA on coagulation factors and platelets. In the following I will provide my review of the study protocol from a statistical point of view:

- The primary endpoint is the TGA parameter ETP and the sample size calculation was performed for this primary endpoint. Please provide additional informations about sample size planning: which software was used and which statistical test; what is the justification for the assumed difference; what is the assumend variation? Without these informations, I am not able to reproduce the sample size calculation.

- I would recommend to define at one point the change over time as difference between about 30 min. after admission and immediately before administration of the trial treatment - preferably in the Outcomes section.
• Although there is a detailed SAP, I would expect at least the analysis approach for the primary endpoint more detailed. As far as I understand the first two paragraphs of the Analysis section, a confidence interval for the difference between the two treatment groups for the change ETP will be calculated. But which one? How will missing values be considered? Two tubes are collected per woman per point in time, but can be excluded that the results are invalid?

**Competing Interests:** No competing interests.