

Tranexamic Acid for Postpartum Haemorrhage: A Review

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Abstract

Post partum haemorrhage is the main cause of maternal mortality worldwide. Tranexamic acid is a cheap, easy to use and relatively safe medication that is gaining popularity as a management option for obstetric haemorrhage. It is already widely used to limit bleeding in trauma and many major surgeries. This review examines the evidence surrounding its use to control bleeding at the time of delivery.

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Index terms— tranexamic acid, pregnancy, cesarean section, postpartum haemorrhage, bleeding, maternal mortality.

1 Background

The 5th Millennium Development Goal is improving maternal health and more specifically decreasing by three quarters, the maternal mortality ratio by 2015. While important gains have been made, developing countries are still lagging due to the poor access to healthcare facilities, skilled healthcare workers, medications and blood products¹.

The World Health Organisation defines post partum haemorrhage as a blood loss of greater than or equal to 500ml at the time of or after delivery. Post partum haemorrhage remains the leading cause of maternal mortality and morbidity worldwide. The maternal mortality ratio worldwide is estimated at 210 maternal deaths per 100 000 live births, but is as high as 1100 in some developing, particularly African countries. This is in comparison to 7 maternal deaths per 100 000 live births in Australia and 21 in the United States². Up to one third of these deaths may be attributed to obstetric haemorrhage, many of which may be prevented or minimized with timely access to medications and emergency care. Up to 1 percent of women having a vaginal birth and 5 percent of women having a cesarean section will require a blood transfusion, which exposes the woman to risks from transfusion reactions and transmission of blood-borne viral infections³. In many areas, blood products are simply not available.

The predominant causes of post partum haemorrhage are uterine atony, trauma to the genital tract and retained placental tissue after delivery. There are a number of factors that increase a woman's chance of having a post partum haemorrhage, however the majority of cases occur in women with low risk pregnancies.

Tranexamic acid is an antifibrinolytic that prevents the breakdown of fibrin deposits at bleeding sites in the body. By blocking lysine-binding sites on plasminogen molecules, the body's natural prohemostatic state post delivery is enhanced.

Tranexamic acid is already widely used in nonobstetric fields, to decrease bleeding from trauma and during elective cardiac and orthopedic surgery^{4,5}. It has proven effectiveness in decreasing blood loss in patients with menorrhagia⁶. As there is very limited data from randomized controlled trials on the use of tranexamic acid for treating post partum haemorrhage, early reports of its success in preventing post partum haemorrhage, as well as evidence of its effective use in other areas of medicine allow extrapolation of the results to cases in which a post partum haemorrhage is already occurring. Tranexamic acid is cheap, easy to transport, store and use and evidence to date suggests that it is safe to use, even in pregnant women who are already at higher risk for thromboembolic events.

This report examines the existing data on the use of tranexamic acid in post partum haemorrhage, with an aim to recommend its use in limited settings as an adjunct to established interventions such as uterotronics.

7 ADVERSE EVENTS

46 2 II.

47 3 Evidence

48 The most recent Cochrane review on the literature surrounding the effectiveness of tranexamic acid in either
49 preventing or treating post partum haemorrhage was undertaken in 20113. Since then seven further small but
50 promising trials have given further strength to the evidence that the medication is effective in obstetric bleeding
51 related to delivery. As not enough evidence is available to identify an impact on maternal mortality, the outcomes
52 specifically ascertained in this review are reduction in blood loss, avoidance of further interventions and decreased
53 requirement for blood transfusion. Data on adverse reactions and associated events was also collected.

54 Nine of the eleven randomized controlled trials identified for inclusion in this review were studies in which
55 tranexamic acid was given prophylactically prior to lower segment cesarean section. Despite the fact that most
56 gave inadequate or unclear data on randomization or blinding techniques (or were not blinded), the limited
57 available data on the subject necessitates consideration of all trials in the review.

58 Of the two remaining trials, one looked at tranexamic acid given prophylactically in the third stage of labour,
59 and a final more recent study trialed tranexamic acid versus a placebo in women who were already experiencing
60 a post partum haemorrhage. In all trials women also received uterotronics, as per the world health organisation
61 recommendation7.

62 A compiled result of the reduction of blood loss achieved with the use of tranexamic acid was not applicable
63 due to the fact that all studies used different doses, treatment regimes and timelines for documenting the degree
64 of haemorrhage. However, individually all studies displayed a statistically significant reduction in blood loss after
65 delivery.

66 4 III.

67 5 Prophylactic use in Cesarean Section

68 The largest randomized controlled trial, by Abdul-Aleem8 included 740 women, 373 of who received 1 gram
69 of tranexamic acid intravenously 10 minutes prior to elective cesarean section. The mean total blood loss was
70 241.6ml in the tranexamic acid group, compared to 510.0 ml in the control group.

71 Goswami9 demonstrated a dose dependent relationship, with the mean total blood loss at cesarean section
72 527.17 ml in the control group, 376.83 ml in patients who received 10mg/kg tranexamic acid prior to cesarean
73 and 261.17ml in those who received 15mg/kg.

74 The remaining studies determined a reduction in blood loss between 375.78ml and 62.5ml with the preoperative
75 administration of tranexamic acid.10-16

76 IV.

77 6 Management of Post Partum Haemorrhage

78 Ducloy-Bouthors17 is the only randomized controlled trial that looks at using tranexamic acid as a treatment
79 for women diagnosed with postpartum haemorrhage, rather than as a preventative measure. This is the most
80 relevant study to date. With 144 women in the study the cohort was relatively small, but adequately powered to
81 achieve significant results. These women had already had a postpartum haemorrhage of greater than or equal to
82 800ml at the time of randomization. Based on success in cardiac and orthopedic surgery, a high dose of 4grams
83 of tranexamic acid over 1 hour, followed by 1gram/hour over 6hours was administered. Blood loss was measured
84 at specified intervals from the time of randomization up until 6hours later. The control group had a median
85 blood loss of 221ml compared to the group receiving tranexamic acid, which had a median blood loss of 173ml.

86 Throughout all seven trials there was a trend toward reduced requirement for blood transfusion and further
87 intervention when tranexamic acid was used, however these outcomes did not reach statistical significance.

88 V.

89 7 Adverse Events

90 Mild transient side effects, most commonly nausea, were reported with greater frequency among participants
91 who had received treatment with tranexamic acid. There were no reports of deep vein thrombosis in any studies.
92 Two patients in the tranexamic acid arm of the Ducloy-Bouthors17 trial developed superficial thrombosis at the
93 site of the venous catheter, however one patient who did not receive the medication was diagnosed with the same
94 condition. None of the studies reported side effects of clinical or statistical significance. A review of the use
95 of tranexamic acid in surgery did not demonstrate an increased risk of thromboembolic events. Despite these
96 findings, it would be imprudent to use tranexamic acid in patients with history of thrombosis or other risk factor
97 that would preclude the use of antifibrinolitics in normal practice18.

98 Tranexamic acid crosses the placenta, which in theory may have some impact on the unborn baby if it is given
99 prior to cesarean section, but obviously is not relevant if used as a treatment for post partum haemorrhage. It
100 is excreted in very small amounts in breast milk, but to date no adverse events in breastfed babies have been
101 reported.

102 **8 VI.**

103 **9 Discussion**

104 The available evidence for the use of tranexamic acid in postpartum haemorrhage remains limited, with the
105 majority of the data of poor quality. However, support is mounting and the outcomes of all studies to date are
106 cohesive in the finding that tranexamic acid does significantly reduce blood loss post partum. At this time, doses
107 other than 1 gram intravenously, followed by a further 1gram if bleeding does not cease are not well supported.
108 There is not yet enough evidence to support the routine use of tranexamic acid, or any suggestion that it may
109 be used instead of traditional interventions including uterotronics.

110 A large study, the WOMAN trial (World Maternal Antifibrinolytic Trial) is currently being undertaken,
111 and will hopefully provide stronger evidence for the use of tranexamic acid in clinically diagnosed postpartum
112 haemorrhage¹⁹. The mediation is also being used with increasing frequency in other gynecological and obstetric
113 conditions including bleeding after LLETZ and cone biopsy of the cervix, and in antepartum haemorrhage due
114 to placenta praevia or diagnosed placental edge bleeds. More evidence is required to support the routine use of
115 tranexamic acid in these conditions.

116 Situations in which post partum haemorrhage is ongoing after first line interventions, in cases where bleeding
117 may be due to factors other than uterine atony, and times when there may be a delay such as transferring
118 a patient to theatre or to a larger centre for further treatment may be an appropriate instance to administer
119 tranexamic acid.

9 DISCUSSION

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