The safety and efficiency of intravenous administration of tranexamic acid in coronary artery bypass grafting (CABG): A meta-analysis of 28 randomized controlled trials

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Abstract

Background

The safety and efficiency of intravenous administration of tranexamic acid (TXA) in coronary artery bypass grafting (CABG) remains unconfirmed. Therefore, we conducted a meta-analysis on this topic.

Methods

We searched the Cochrane Central Register of Controlled Trials (CENTRAL), PUBMED and EMBASE for randomized controlled trials on the topic. The results of this work are synthetized and reported in accordance with the PRISMA statement.

Results

28 studies met our inclusion criteria. TXA reduced the incidence of postoperative reoperation of bleeding (relative risk RR, 0.46; 95% confidence interval CI; 0.31-0.68), the frequency of any allogeneic transfusion (RR, 0.64; 95% CI, 0.52-0.78) and the postoperative chest tube drainage in the first 24 hours by 206 ml (95% CI -248.23 to -164.15). TXA did not significantly affect the incidence of postoperative cerebrovascular accident (RR, 0.93; 95%CI, 0.62-1.39), mortality (RR, 0.82; 95%CI, 0.53-1.28), myocardial infarction (RR, 0.90; 95%CI, 0.78-1.05), acute renal insufficiency (RR, 1.01; 95%CI, 0.77-1.32). However, it may increase the incidence of postoperative seizures (RR, 6.67; 95%CI, 1.77-25.20). Moreover, the subgroup analyses in the settings of on-pump and off-pump CABG and the sensitivity analyses in trials randomized not less than 100 participants or with the maximum sample size study excluded further strengthened the above results.

Conclusions

TXA is effective to reduce reoperation for bleeding, blood loss and the need for allogeneic blood products in patients undergoing CABG without increasing prothrombotic complication. However, it may increase the risk of postoperative seizures.
Background

Excessive bleeding is a common complication which may lead to exposure to the risk of homologous blood transfusion and increased morbidity in patients undergoing cardiac operations[1]. Tranexamic acid (TXA), an antifibrinolytic agent, has been widely used and proved to be effective in reducing risk of blood loss and transfusion among patients undergoing cardiac surgery[2]. However, whether it reduced the incidence of reoperation for life-threatening bleeding which are strongly associated with poor outcomes after cardiac surgery remains controversial.

Despite of the effectiveness in reducing the risk of blood loss and transfusion, it may potentially increase the risk of myocardial infarction, stroke, and other thrombotic complications after cardiac surgery especially in patients undergoing coronary artery bypass grafting (CABG) surgery who are commonly characterized by systemic arteriosclerosis or stenosis [3,4]. It was reported that TXA was associated with the increased risk of postoperative neurologic events such as stroke and seizures in cardiac surgery[5,6]. Some studies have suggested that TXA is associated with reduction in cerebral blood flow and increase the risk of cerebral infarction[5,7]. A multi-center study suggested that TXA was associated with a higher risk of postoperative seizures in GABG surgery[8]. A meta-analysis in 2011 has shown that TXA is associated with reduced blood transfusion in off-pump CABG surgery[9]. However, the safety of TXA in off-pump CABG surgery could not be confirmed due to the small population sample size.

An increasing number of studies that investigated the effectiveness and safety of TXA in CABG surgery have been conducted in recent years with varying results[8,10-18]. Therefore, we conducted a meta-analysis of existing studies to estimate the safety and efficiency of TXA in CABG surgery focusing on the incidence of postoperative
cerebrovascular accident, seizures and reoperation for bleeding.

Methods

The meta-analysis was performed according to the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) statement in this study[19].

Search strategy

A systematic and comprehensive search was conducted in the Cochrane Central Register of Controlled Trials (CENTRAL), PUBMED and EMBASE from database established to February 8, 2018 with no language limitation. The search strategy included the following MEDLINE subject heading terms: tranexamic acid and cardiac surgical procedures. The above subject heading terms were connected by “AND”. The initial searches of PUBMED and EMBASE were unrestricted to maximize sensitivity and a filter which primarily identifies randomized controlled trials was adopted to improve the specificity. Moreover, we also checked the reference lists of relevant articles for potential relevant studies.

Eligibility criteria

Randomized controlled trials that compared the effectiveness or safety of the intravenous administration of TXA with that of placebo in adult CABG surgery were included in this meta-analysis. Studies were eligible for inclusion, regardless of the publication language. We excluded studies which were conducted on underage patients or in which TXA was topically applied in mediastinum.

Selection of included studies

Retrieved studies were imported into Endnote (version X7; Thomson Reuters), where
duplications were detected and deleted automatically. Two authors independently scanned the titles and abstract of retrieved studies according to the established eligibility criteria to exclude the obvious unrelated studies. The full-text was further evaluated if the judgement could not easily be decided based on its title or abstract. The disagreements between reviewers were settled by a third reviewer. The relevant data of included studies was extracted by these reviewers independently using a standard data sheet. Study characteristics included author, publication year, sample size, sex ratio, type of CABG, duration of anticoagulant medication discontinued before surgery, outcome data, drug dose and treatment regimens.

Assessment of risk of bias in included studies

The Cochrane risk of bias tool which is recommended by the Cochrane Collaboration for risk of bias assessment was adopted in this study[20]. There are seven domains in the Cochrane risk of bias tool, including the random sequence generation, allocation concealment, blinding of participants and personnel, blinding of outcome assessment, incomplete outcome data, selective reporting and other bias. The judgment of each domain is presented as “low risk”, “high risk” or “unclear risk” based on the instruction of Cochrane Collaboration. Two reviewers independently assessed each domain of included studies and any disagreements were adjudicated by a third reviewer.

Quality of the evidence

GRADE (Grades of Recommendation, Assessment, Development and Evaluation) Working Group system was adopted to evaluate the quality of the evidence[21]. Two reviewers independently assessed the quality of each outcome. The five categories used for the
GRADE quality assessment were: limitations of design, inconsistency, indirectness, imprecision, and publication bias. We used GRADE profiler (GRADEpro) software to create the “Summary of findings” table, which includes the following outcomes: incidence of postoperative cerebrovascular accident, seizures, reoperation for bleeding, mortality, myocardial infarction, acute renal insufficiency, the frequency of any allogeneic transfusions and 24-hour postoperative chest tube drainage.

Study outcomes

All outcomes were described a priori, according to the principles of the PRISMA statement. The primary outcome was incidence of postoperative cerebrovascular accident, seizures and reoperation for bleeding. The second outcomes included postoperative mortality, myocardial infarction, acute renal insufficiency, the frequency of any allogeneic transfusions and 24-hour postoperative chest tube drainage.

Statistical methods

In some studies, continuous variables was presented as median, range and/or interquartile range. To facilitate meta-analysis, we estimated the sample mean and standard deviation from median, range and/or interquartile range by using the calculator with a compiled formula recommended by Luo and colleagues[22]. The risk ratio (RR) with the corresponding 95% confidence interval (95% CI) was calculated for dichotomous data and continuous data were analyzed by using mean difference (MD) with the corresponding 95% CI. Data analyses followed the guidelines established by the Cochrane Collaboration regarding statistical methods. The statistical heterogeneity was evaluated by reviewing the $I^2$ statistic and Chi2 test. If either the Chi2 test resulted in $P < 0.10$ or the $I^2$ statistic was greater 50%, random-
effect model was used to evaluate outcomes, otherwise a fixed-effect model was used. For all tests, two-tailed $P$-values $<0.05$ were considered significant. Funnel plots were conducted to evaluate reports for publication bias when more than 10 studies were included. Considering the activation effect of cardiopulmonary bypass (CPB) on the fibrinolytic pathway, subgroup analysis was performed based on CABG with/without CPB. Moreover, Sensitivity analyses was performed in studies randomized more than 99 patients to avoid the possibility that the rare incidences of complication were underestimated due to the included studies with small population size. Sensitivity analyses that excluded the study with the largest number of participants were conducted to estimate the effect of that study on the overall effect of meta-analysis. All data analysis was conducted using Review Manager (RevMan; version 5.2), Copenhagen: The Nordic Cochrane Centre, The Cochrane Collaboration, 2012

Results

Results of search

227 studies were identified from our initial search and 146 of them remained after duplicates were removed. 108 of the remaining studies were excluded during title and abstract screening. 38 studies were identified for full-text assessment according to our inclusion and exclusion criteria and 10 of them were removed because of non-RCT, topical application of TXA or without placebo group. Finally, 28 studies$[3,4,8,10-12,14-18,23-39]$ were included in this meta-analysis. The study selection process is shown in Figure 1.

Description of included studies

The characteristics of included studies were shown in Table 1. The 28 included trials$[3,4,8,10-12,14-18,23-39]$ randomized 7446 patients (3712 to tranexamic acid and
3734 to placebo). 14 trials[4,8,11,14-18,25,32,36-39] randomized more than 99 patients. CABG was conducted in on-pump condition in 17 trails[12,14,16-18,23-26,28,30-32,34,36,38,39], off-pump condition in 9 trails[3,10,11,15,27,29,33,35,37] and both condition in 2 trails[4,8].

Risk of bias within studies

The results of bias risk assessment were showed in Fig 2 A and B. 14 studies[3,11,12,14,16,23-25,27-30,35,38] did not provide a satisfactory description of their random processes. Blinding process was at high risk of bias in one study[39] and unclear risk of bias in 7 studies[11,12,23-25,29,35] due to unclear description. 3 studies[16,25,31] had unclear or incomplete descriptions of their outcome data. 2 studies[3,36] were considered to be at high risk of selective reporting bias because the reported outcome indicators were inconsistent with the planed outcome indicators.

Publication bias

Publication bias was evaluated by funnel plots in the following outcomes: postoperative cerebrovascular accident, reoperation for bleeding, mortality, myocardial infarction, acute renal insufficiency, the frequency of any allogeneic transfusions and 24-hour postoperative chest tube drainage (Supplementary Figs 1-6). All of the plots showed a symmetrical shape which suggested low risk of publication bias of the above outcomes.

Quantitative data synthesis

Cerebrovascular accident

There were 22 trials that reported the incidence of postoperative cerebrovascular accident
between TXA and placebo, with a total of 6775 participants. TXA did not increase the incidence of cerebrovascular accident overall from meta-analysis [41/3371 vs 45/3404, RR=0.93 (0.62-1.39), \( P \) for effect=0.71, \( P \) for heterogeneity=0.92, \( I^2=0\% \)] (Fig 3).

Sub-analysis in on-pump CABG with 13 trials included showed no significant increase in the incidence of cerebrovascular accident in patients who received TXA treatment [9/686 vs 10/711, RR=0.95 (0.44-2.06), \( P \) for effect=0.90, \( P \) for heterogeneity=0.86, \( I^2=0\% \)]. In off-pump CABG, 8 trials with 749 participants were included and no cerebrovascular accident happened in those trials (Fig 3).

9 studies with a total of 5939 participants were included in the sensitive analysis of studies that randomized not less 100 participants. The conclusion that TXA would not increase cerebrovascular accident incidence was strengthened by the sensitivity analysis [RR=0.87 (0.57-1.33), \( P \) for effect=0.53, \( P \) for heterogeneity=0.95, \( I^2=0\% \)]. Sensitivity analysis that excluded the study with the largest number of participants further strengthened the above conclusion [RR=0.95 (0.43-2.10), \( P \) for effect=0.90, \( P \) for heterogeneity=0.86] (Table 2).

Seizures

In total, 5 studies with 5043 participants reported the incidence of seizures after CABG. The summary RR for postoperative seizures with the use of TXA versus placebo was 5.99 (95% CI 1.77-20.24) which suggested that tranexamic acid would increase the incidence of seizures after CABG (Fig 4).

Reoperation for bleeding
There were 16 trials that reported the incidence of postoperative reoperation for bleeding, with a total of 6259 participants. TXA decreased the incidence of reoperation for postoperative bleeding overall from meta-analysis [35/3125 vs 78/3134, RR=0.46 (0.31-0.68), \(P\) for effect<0.01, \(P\) for heterogeneity=0.63, I\(^2\)=0\%] (Fig 5).

10 studies with 1143 participants were included in on-pump CABG, the result of meta-analysis suggested no significant difference of reoperation for postoperative bleeding between TXA and placebo[16/569 vs 26/574, RR=0.64 (0.35-1.15), \(P\) for effect=0.14, \(P\) for heterogeneity=0.62, I\(^2\)=0\%]. In off-pump subgroup, 4 studies with 384 participants were included and only one patient suffered reoperation in placebo group (Fig 5).

8 trials were included in sensitivity analysis of studies randomized not less than 100 patients. The sensitivity analysis supported the result that TXA decreased incidence of reoperation for bleeding in CABG surgery when compared with placebo [29/2812 vs 59/2821, RR= 0.49 (0.32-0.77), \(P\) for effect<0.01, \(P\) for heterogeneity=0.58, I\(^2\)=0\%]. While sensitivity analysis that excluded the study with the largest number of participants did not supported the above conclusion [RR=0.59 (0.34-1.04), \(P\) for effect=0.07, \(P\) for heterogeneity=0.72] (Table 2).

Mortality

The overall analysis showed that TXA did not significantly decrease the mortality in patients receiving CABG when compared with placebo [33/3196 deaths in the TXA group vs 41/3218 deaths in the placebo group, RR=0.82 (0.53-1.28), \(P\) for effect=0.38, \(P\) for heterogeneity=0.82, I\(^2\)=0\%, with 18 trails included] (Fig 6).
Sub-analysis in the settings of on-pump CABG also showed no statistically significant effect of TXA on mortality [6/639 vs 7/663, RR=0.93 (0.36–2.38), P for effect=0.88, P for heterogeneity=0.62, I²=0%, with 12 trials included]. Sub-analysis in the settings of off-pump included 5 trials, but only one of them reported one patient died in each group (Fig 6).

Sensitivity analysis of studies randomized more than 99 patients supported the results that TXA did not significantly decrease the mortality in CABG surgery compared with placebo [31/2870 vs 36/2886, RR= 0.87 (0.54–1.40), P for effect=0.56, P for heterogeneity=0.46, I²=0%, with 7 trials included]. The result of sensitivity analysis that excluded the study with maximum sample was consistent with the above analyses [7/875 vs 8/898, RR= 0.93 (0.38–2.27), P for effect=0.88, P for heterogeneity=0.75] (Table 2).

Myocardial infarction

In total, 23 studies with 6714 participants reported the incidence of myocardial infarctions after CABG. The overall analysis showed no increased risk of postoperative myocardial infarction [292/3349 vs 325/3365, RR=0.90 (0.78-1.05), P for effect=0.18, P for heterogeneity=0.89, I²=0%] (Fig 7).

13 studies with 1286 participants were included in the sub-analysis of on-pump CABG, the result of meta-analysis suggested no significant difference of myocardial infarction between TXA and placebo[21/639 vs 24/647, RR=0.9 (0.51-1.58), P for effect=0.71, P for heterogeneity=0.72, I²=0%]. In off-pump subgroup, 9 studies with 798 participants were included, no significant difference of myocardial infarction between TXA and placebo was found neither [2/400 vs 1/398, RR=1.56(0.22-11.23), P for effect=0.66, P for heterogeneity=0.56, I²=0%] (Fig 7).
7 trials were included in sensitivity analysis of studies randomized not less than 100 patients. The sensitivity analysis supported the result that TXA did not increase myocardial infarction in CABG surgery when compared with placebo [286/2999 vs 318/3011, RR = 0.90 (0.78–1.05), P for effect = 0.18, P for heterogeneity = 0.64, I² = 0%]. The result of sensitivity analysis that excluded the study with maximum sample was consistent with the above analyses [23/1039 vs 25/1045, RR = 0.94 (0.55-1.61), P for effect = 0.81, P for heterogeneity = 0.80] (Table 2).

**Acute renal insufficiency**

There are 14 studies that reported the incidence of acute renal insufficiency in this meta-analysis. The summary RR for acute renal with the use of TXA versus placebo was 1.01 (95% CI 0.77-1.32) which suggested that tranexamic acid would not increase the incidence of acute renal insufficiency (Fig 8).

The summary RR of sub-analysis in on-pump CABG was 0.91 (95% CI 0.36-2.29) which suggested that TXA did not have adverse effect on postoperative renal function in patients undergoing on-pump CABG. A similar result was found in the sub-analysis in off-pump CABG [RR = 0.85 (0.29-2.47), P for effect = 0.76, P for heterogeneity = 0.52, I² = 0%] (Fig 8).

Sensitivity analysis in trials randomized not less than 100 participants reinforced the overall analysis [RR = 1.03 (0.79-1.35), P for effect = 0.81, P for heterogeneity = 0.89, I² = 0%, with 7 studies included]. The result of sensitivity analysis that excluded the study with maximum sample size was consistent with the above analyses [12/658 vs 14/667, RR = 0.88 (0.42-1.84), P for effect = 0.73, P for heterogeneity = 0.94] (Table 2).
Transfusion of any blood products

11 trails with a total of 5360 participants reported the postoperative transfusion rate of any blood product. Overall, TXA significantly reduced the transfusion of any blood products [RR=0.64(0.52-0.78), P for effect<0.01, P for heterogeneity<0.01, I²=76%] (Fig 9).

In the subgroup of patients undergoing on-pump CABG, TXA also reduced the transfusion of any blood products, however, this effect was not statistically significant [RR=0.68(0.47-1.00), P for effect=0.05, P for heterogeneity<0.01, I²=81%]. On the other hand, sub-analysis in off-pump CABG, TXA significantly reduced the transfusion of any blood products [RR=0.32(0.19-0.53), P for effect<0.01, P for heterogeneity=0.60, I²=0%] (Fig 9).

In the sensitivity analysis that included all the studies that randomized more than 99 participants, TXA significantly decreased the transfusion of any blood products [RR=0.64(0.50-0.81), P for effect<0.01, P for heterogeneity<0.01, I²=86%]. The result of sensitivity analysis that excluded the study with maximum sample size further enhanced the above analyses [139/396 vs 216/363, RR= 0.29 (0.20-0.40)P for effect <0.01, P for heterogeneity <0.01] (Table 2).

Postoperative chest tube drainage in the first 24 hours

In total, 16 studies with 6247 participants were included in the meta-analysis of postoperative chest tube drainage in the first 24 hours. One of them[18] divided participants into two groups according to the difference in fluid use and reported the drainage of patients receiving TXA and placebo in both groups separately. We treated these two sets of data as two separate studies in the meta-analysis. Overall, the chest tube
drainage was significantly decreased by TXA when compared with placebo [MD=-206.19, 95% CI (-248.23, -164.15), P for effect<0.01, P for heterogeneity<0.01, I²=72%] (Fig 10).

Sub-analysis in the settings of on-pump CABG with 8 trials included showed no significant decrease of chest tube drainage in the first 24 hours in patients who received TXA treatment [MD=-211.36, 95% CI (-263.13, -159.59), P for effect<0.01, P for heterogeneity=0.26, I²=20%]. A similar result was found in the sub-analysis in off-pump CABG [MD=-220.25, 95% CI (-290.58, -149.91), P for effect<0.01, P for heterogeneity=0.26, I²=76%] (Fig 10).

7 studies with a total of 5674 participants were included in the sensitive analysis. The conclusion that TXA would decrease chest tube drainage in the first 24 hours was strengthened by the sensitivity analysis [MD=-208.30, 95% CI (-274.12, -142.48) P for effect<0.01, P for heterogeneity<0.01, I²=83%]. The sensitivity analysis that excluded the study with maximum sample size also supported the above conclusion [MD=-215.42, 95% CI (-259.48, -171.57), P for effect <0.01, P for heterogeneity<0.01, I²=83%] (Table 2).

Quality of the evidence

The GRADE approach was adopted to evaluate the quality of each outcome and “Summary of findings” tables were presented (Table 3). In general, the overall quality of evidence in the meta-analyses of postoperative seizures and reoperation for bleeding was high. However, the overall quality of evidence in the meta-analyses of postoperative transfusion of any blood products and chest tube drainage in the first 24 hours was very low due to the problems of inconsistency and the risk of bias. The overall quality of evidence of other outcomes was moderate due to the risk of bias.

Discussion
In this meta-analysis, we found that the intravenous use of TXA was associated with lower risk of reoperation for postoperative bleeding, blood loss and blood transfusion than the placebo group. Moreover, we also found that intravenous use of TXA did not increase the risk of postoperative cerebrovascular accident, mortality or other thrombotic complication among patients undergoing CABG when compared with placebo treatment. However, it may increase the incidence of postoperative seizures. The results of most subgroup analyses of the primary results in CABG conducted under on-pump or off-pump condition were consistent with that of overall analyses. However, meta-analysis could not be performed in the sub-analyses of postoperative reoperation for bleeding, mortality and cerebrovascular accident in off-pump CABG due to the small number of incidence. No significant decrease in postoperative reoperation for bleeding and transfusion of any blood products were found in on-pump group. Most of the sensitivity analyses in trails that recruited more than 99 participants or in trails that excluded the study with the largest number of participants further strengthened the conclusion of overall analyses.

The release of plasmin during cardiac surgery activates fibrinolysis and may contribute to platelet dysfunction[40]. In addition to inhibiting the transformation of plasminogen into plasmin by reversibly binding lysine binding site on plasmin, TXA can also reduce bleeding by preventing platelet activation induced by fibrinolytic enzyme[41]. A previous meta-analysis suggested that TXA was effective in reducing blood loss and the need for blood transfusion in cardiac surgery[42]. However, the incidence of reoperation for bleeding was not significantly decrease by TXA[42]. In our current analysis, we found that TXA overall reduced the transfusion of any blood products and 24-hour postoperative chest tube drainage in CABG surgery which was consistent with the previous study. Moreover, the sub-analyses in the different conditions under which GABG was conducted further strengthened
the above results. However, these analyses have significant heterogeneity which may due to the difference in indications of blood transfusion, drug dose and treatment regimens among different studies.

Different from the previous study, our current meta-analysis suggested that TXA significantly decrease the incidence of reoperation for bleeding in CABG surgery with low heterogeneity. In addition, the sensitivity analyses in studies randomized more than 99 participants further strengthened the conclusion that TXA reduced the incidence of reoperation for bleeding, transfusion of any blood products and 24-hour blood loss suggesting that the small sample size studies included in the meta-analysis did not affect the overall effectiveness. However, the sensitivity analysis that excluded the study[8] with maximum sample size did not suggest that TXA would significantly decrease the incidence of reoperation for bleeding. This result suggested that the study with the largest number of participants largely determines the overall effect of meta-analysis. While considering the low risk of bias assessment in that study, we can still believe that TXA overall decrease the incidence of reoperation for bleeding. In the sub-analysis of on-pump GABG, TXA tended to reduce the incidence of reoperation for bleeding. However, the effect was not statistically significant. The exclusion of the study with the largest number of participants due to mixed surgical types in the sub-analysis may explain this difference.

Although lots of studies have suggested that blood transfusion and reoperation for bleeding is associated with poor outcomes after cardiac surgery, we did not find that TXA would reduce the risk of cerebrovascular accident, myocardial infarction, acute renal insufficiency or mortality despite its effectiveness in reducing transfusion and reoperation for bleeding. A previous meta-analysis had reported that TXA reduced blood transfusion in off-pump CABG and did not increased the incidence of postoperative adverse events[9]. However, the
sample size in that study was not sufficient to detect the rare but clinically significant adverse events. In the current meta-analysis, enough population were included in the above analyses to detect clinically significant difference. Moreover, the above conclusion were strengthened by sensitivity analyses in trails enrolling more than 99 patients or sensitivity analyses excluded the study with largest sample size. In addition, there was no heterogeneity in above analyses from the results of heterogeneity tests and the risk of publication bias in these meta-analyses was quite low revealed by funnel plots. These unexpected results may be explained by the potential prothrombotic effects of TXA. It is well known that 5% to 15% of all grafts may be blocked in the early postoperative period even without the use of antifibrinolytic agents, which may led to recurrence of myocardial ischemia, infarction, or even death[43,44]. Perioperative inhibition of fibrinolysis may increase the rate of early graft occlusion rate[45]. The phenomenon that TXA reduced transfusion, blood loss and incidence of reoperation without decreasing postoperative morality or adverse events may be a balance of its blood conservation effect and potential prothrombotic effect.

A previous meta-analysis suggested that the risk of seizure increased in patients with TXA exposure[46]. In the current meta-analysis we found that TXA increased the incidence of postoperative seizures in CABG surgery. Several studies have suggested that the convulsant property of TXA is likely mediated by disinhibition of gama-aminobutyric acid type A (GABAA) receptors and glycine receptor, which are two major mediators of inhibition in the CNS[47,48]. Moreover, TXA did not interfere with N-methyl-Daspartate receptor and impact glutamatergic synaptic transmission[48,49]. In addition, some studies have shown that TXA reduces cerebral blood flow and increases the risk of cerebral infarction which could contribute to the postoperative seizures. However, the meta-analysis of postoperative
A cerebrovascular accident in current study did not support the hypothesis that TXA increase incidence of seizures by increasing the incidence of cerebral infarction. Moreover, a growing number of studies have suggested the seizures associated with TXA to be dose related[6,50,51]. Therefore, studies that investigate the optimize dose and regime for administration of TXA are needed in the future. Moreover, a growing number of studies that investigate the efficacy and safety of topical use of tranexamic acid have been conducted in recent years due to the promise of reducing postoperative bleeding and seizures[52,53]. A recent meta-analysis showed that the topical application of TXA effectively reduces both transfusion risk and blood loss compared to placebo and no major differences were found between topical and intravenous tranexamic acid with respect to safety and efficacy[54]. However, both surgical and non-surgical trials were included in that study. While in our study, we focused on the safety and efficiency of intravenous administration of tranexamic acid in coronary artery bypass grafting (CABG).

There are some limitations in this meta-analysis. Firstly, heterogeneity due to clinical and methodological diversity was inevitable which may affect the reliability of the analysis results especially in meta-analyses of transfusion and blood loss. Secondly, some data were presented as median and interquartile range which cannot be used in performing meta-analysis. We estimated the mean and standard deviation from those data to perform meta-analysis which may compromise the reliability of analysis results. Thirdly, the postoperative incidence of adverse event was suggested to may be dose-dependent[6], while we failed to performed sub-analysis in different dose setting due to the various dosage and regimens of TXA administration in current meta-analysis. Fourthly, a multicenter study that randomized 2311 participants occupied the main part of most analyses which may lead to bias. Despite the above limitations, the current study is still the most comprehensive analysis on the
efficacy and safety of TXA in CABG surgery with sufficient sample size.

Conclusion

The current study systematically reviewed the existing evidence on the efficacy and safety profile of the intravenous administration of TXA in CABG surgery and showed that TXA would significantly reduce postoperative transfusion of any blood products, 24-hour postoperative chest tube drainage and reoperation for bleeding. In addition, our results identified for the first time that intravenous administration of TXA in CABG surgery did not increase the risk of prothrombotic complication with sufficient sample size. However, it may increase the risk of postoperative seizures. Overall, intravenous administration of TXA in CABG surgery is effective and safe in reducing blood loss and transfusion according to the existing evidence and further studies are needed to identify the optimize dose and regime for intravenous use of TXA to achieve the best benefit with lowest risk.

Abbreviations

CABG: coronary artery bypass grafting; TXA: tranexamic acid; GABAA: gamma-aminobutyric acid type A; CNS: central nervous system

Declarations

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Availability of data and materials

All data generated or analysed during this study are included in this published article and its
supplementary information files.

Competing interests

The authors declare that they have no competing interests

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Author's contributions

YTZ, GC and HYZ were involved in the study design, data review, data analysis, writing paper, review and approval of final manuscript. YB, MMC, YFZ and XY were involved in data review, data analysis, review and approval of final manuscript.

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References

1. Moulton MJ, Creswell LL, Mackey ME, Cox JL, and Rosenbloom M. Reexploration for bleeding is a risk factor for adverse outcomes after cardiac operations. J Thorac Cardiovasc Surg. 1996; 111: 1037-46

2. Henry D, Carless P, Fergusson D, and Laupacis A. The safety of aprotinin and lysine-derived antifibrinolytic drugs in cardiac surgery: a meta-analysis. CMAJ. 2009; 180: 183-93
3. Mehr-Aein A, Sadeghi M, and Madani-civi M. Does tranexamic acid reduce blood loss in off-pump coronary artery bypass? Asian Cardiovasc Thorac Ann. 2007; 15: 285-9

4. Casati V, Della Valle P, Benussi S, et al. Effects of tranexamic acid on postoperative bleeding and related hematochemical variables in coronary surgery: Comparison between on-pump and off-pump techniques. J Thorac Cardiovasc Surg. 2004; 128: 83-91

5. Ngaage DL and Bland JM. Lessons from aprotinin: is the routine use and inconsistent dosing of tranexamic acid prudent? Meta-analysis of randomised and large matched observational studies. Eur J Cardiothorac Surg. 2010; 37: 1375-83

6. Murkin JM, Falter F, Granton J, et al. High-dose tranexamic Acid is associated with nonischemic clinical seizures in cardiac surgical patients. Anesth Analg. 2010; 110: 350-3

7. Tsementzis SA, Meyer CH, and Hitchcock ER. Cerebral blood flow in patients with a subarachnoid haemorrhage during treatment with tranexamic acid. Neurochirurgia (Stuttg). 1992; 35: 74-8

8. Myles PS, Smith JA, Forbes A, et al. Tranexamic Acid in Patients Undergoing Coronary-Artery Surgery. N Engl J Med. 2017; 376: 136-148

9. Adler Ma SC, Brindle W, Burton G, et al. Tranexamic acid is associated with less blood transfusion in off-pump coronary artery bypass graft surgery: a systematic review and meta-analysis. J Cardiothorac Vasc Anesth. 2011; 25: 26-35

10. Ahn SW, Shim JK, Youn YN, et al. Effect of tranexamic acid on transfusion requirement in dual antiplatelet-treated anemic patients undergoing off-pump coronary artery bypass graft
surgery. Circ J. 2012; 76: 96-101

11. Chakravarthy M, Muniraj G, Patil S, et al. A randomized prospective analysis of alteration of hemostatic function in patients receiving tranexamic acid and hydroxyethyl starch (130/0.4) undergoing off pump coronary artery bypass surgery. Ann Card Anaesth. 2012; 15: 105-10

12. Greiff G, Stenseth R, Wahba A, et al. Tranexamic acid reduces blood transfusions in elderly patients undergoing combined aortic valve and coronary artery bypass graft surgery: a randomized controlled trial. J Cardiothorac Vasc Anesth. 2012; 26: 232-8

13. Hassani E, Mahoori A, Mehdizadeh H, et al. The effects of tranexamic acid on postoperative bleeding in coronary artery bypass graft surgery. Tehran University Medical Journal. 2012; 70: 176-182

14. Nejad MHG, Baharestani B, Esfandiari R, Hashemi J, and Panahipoor A. Evaluation and comparison of using low-dose aprotinin and tranexamic acid in CABG: A double blind randomized clinical trial. Journal of Tehran University Heart Center. 2012; 7: 15-18

15. Wang G, Xie G, Jiang T, et al. Tranexamic acid reduces blood loss after off-pump coronary surgery: a prospective, randomized, double-blind, placebo-controlled study. Anesth Analg. 2012; 115: 239-43

16. Esfandiari BR, Bistgani MM, and Kabiri M. Low dose tranexamic acid effect on post-coronary artery bypass grafting bleeding. Asian cardiovascular & thoracic annals. 2013; 21: 669-74

17. Shi J, Wang G, Lv H, et al. Tranexamic Acid in on-pump coronary artery bypass grafting
without clopidogrel and aspirin cessation: randomized trial and 1-year follow-up. Ann Thorac Surg. 2013; 95: 795-802

18. Yanartas M, Baysal A, Aydin C, et al. The effects of tranexamic acid and 6% hydroxyethyl starch (HES) solution (130/0.4) on postoperative bleeding in coronary artery bypass graft (CABG) surgery. International Journal of Clinical and Experimental Medicine. 2015; 8: 5959-5971

19. Moher D, Liberati A, Tetzlaff J, Altman DG, and Group P. Preferred reporting items for systematic reviews and meta-analyses: the PRISMA statement. J Clin Epidemiol. 2009; 62: 1006-12

20. Higgins JP, Altman DG, Gotzsche PC, et al. The Cochrane Collaboration's tool for assessing risk of bias in randomised trials. BMJ. 2011; 343: d5928

21. Higgins JP GS. Cochrane Handbook for Systematic Reviews of Interventions version 5.1.0 [updated March 2011]. The Cochrane Collaboration.

22. Luo D, Wan X, Liu J, and Tong T. Optimally estimating the sample mean from the sample size, median, mid-range, and/or mid-quartile range. Stat Methods Med Res. 2018; 27: 1785-1805

23. Speekenbrink RG, Vonk AB, Wildevuur CR, and Eijsman L. Hemostatic efficacy of dipyridamole, tranexamic acid, and aprotinin in coronary bypass grafting. Ann Thorac Surg. 1995; 59: 438-42

24. Brown RS, Thwaites BK, and Mongan PD. Tranexamic acid is effective in decreasing postoperative bleeding and transfusions in primary coronary artery bypass operations: a
double-blind, randomized, placebo-controlled trial. Anesth Analg. 1997; 85: 963-70

25. Landymore RW, Murphy JT, Lummis H, and Carter C. The use of low-dose aprotinin, epsilon-aminocaproic acid or tranexamic acid for prevention of mediastinal bleeding in patients receiving aspirin before coronary artery bypass operations. Eur J Cardiothorac Surg. 1997; 11: 798-800

26. Hardy JF, Belisle S, Dupont C, et al. Prophylactic tranexamic acid and epsilon-aminocaproic acid for primary myocardial revascularization. Ann Thorac Surg. 1998; 65: 371-6

27. Casati V, Gerli C, Franco A, et al. Tranexamic acid in off-pump coronary surgery: a preliminary, randomized, double-blind, placebo-controlled study. Ann Thorac Surg. 2001; 72: 470-5

28. Zabeeda D, Medalion B, Sverdlov M, et al. Tranexamic acid reduces bleeding and the need for blood transfusion in primary myocardial revascularization. Ann Thorac Surg. 2002; 74: 733-8

29. Jares M, Vanek T, Straka Z, and Brucek P. Tranexamic acid reduces bleeding after off-pump coronary artery bypass grafting. J Cardiovasc Surg (Torino). 2003; 44: 205-8

30. Pleym H, Stenseth R, Wahba A, et al. Single-dose tranexamic acid reduces postoperative bleeding after coronary surgery in patients treated with aspirin until surgery. Anesth Analg. 2003; 96: 923-8

31. Andreasen JJ and Nielsen C. Prophylactic tranexamic acid in elective, primary coronary artery bypass surgery using cardiopulmonary bypass. Eur J Cardiothorac Surg. 2004; 26:
32. Karski J, Djaiani G, Carroll J, et al. Tranexamic acid and early saphenous vein graft patency in conventional coronary artery bypass graft surgery: a prospective randomized controlled clinical trial. The Journal of thoracic and cardiovascular surgery, 2005; 130: 309-14

33. Vanek T, Jares M, Fajt R, et al. Fibrinolytic inhibitors in off-pump coronary surgery: a prospective, randomized, double-blind TAP study (tranexamic acid, aprotinin, placebo). European journal of cardio-thoracic surgery: official journal of the European Association for Cardio-thoracic Surgery, 2005; 28: 563-8

34. Santos AT, Kalil RA, Bauemann C, Pereira JB, and Nesralla IA. A randomized, double-blind, and placebo-controlled study with tranexamic acid of bleeding and fibrinolytic activity after primary coronary artery bypass grafting. Braz J Med Biol Res. 2006; 39: 63-9

35. Wei M, Jian K, Guo Z, et al. Tranexamic acid reduces postoperative bleeding in off-pump coronary artery bypass grafting. Scand Cardiovasc J. 2006; 40: 105-9

36. Maddali MM and Rajakumar MC. Tranexamic acid and primary coronary artery bypass surgery: a prospective study. Asian Cardiovasc Thorac Ann. 2007; 15: 313-9

37. Taghaddomi RJ, Mirzaee A, Attar AS, and Shirdel A. Tranexamic acid reduces blood loss in off-pump coronary artery bypass surgery. J Cardiothorac Vasc Anesth. 2009; 23: 312-5

38. Hashemi J, Ghaffari Nejad MH, Baharestani B, Esfandiari R, and Panahipoor A. Evaluation and comparison of use of low-dose aprotinin and tranexamic acid in CABG: A double-blind, prospective, randomized study of 150 patients. Iranian Heart Journal, 2011; 12: 40-4.
39. Alizadeh Ghavidel A, Totonchi Z, Chitsazan M, et al. Safety and efficacy of caproamin fides and tranexamic acid versus placebo in patients undergoing coronary artery revascularization. Journal of Cardiovascular and Thoracic Research. 2014; 6: 197-202

40. Rijken DC, de Munk GA, and Jie AF. Interaction of plasminogen activators and plasminogen with heparin: effect of ionic strength. Thromb Haemost. 1993; 70: 867-72

41. Verstraete M. Clinical application of inhibitors of fibrinolysis. Drugs. 1985; 29: 236-61

42. Henry DA, Carless PA, Moxey AJ, et al. Anti-fibrinolytic use for minimising perioperative allogeneic blood transfusion. Cochrane Database Syst Rev. 2011: CD001886

43. Chesebro JH, Lam JY, and Fuster V. The pathogenesis and prevention of aortocoronary vein bypass graft occlusion and restenosis after arterial angioplasty: role of vascular injury and platelet thrombus deposition. J Am Coll Cardiol. 1986; 8: 57B-66B

44. Bourassa MG. Fate of venous grafts: the past, the present and the future. J Am Coll Cardiol. 1991; 17: 1081-3

45. Karski J, Djaiani G, Carroll J, et al. Tranexamic acid and early saphenous vein graft patency in conventional coronary artery bypass graft surgery: a prospective randomized controlled clinical trial. J Thorac Cardiovasc Surg. 2005; 130: 309-14

46. Lin Z and Xiaoyi Z. Tranexamic acid-associated seizures: A meta-analysis. Seizure. 2016; 36: 70-3

47. Lecker I, Wang DS, Romaschin AD, et al. Tranexamic acid concentrations associated with human seizures inhibit glycine receptors. J Clin Invest. 2012; 122: 4654-66
48. Kratzer S, Irl H, Mattusch C, et al. Tranexamic acid impairs gamma-aminobutyric acid receptor type A-mediated synaptic transmission in the murine amygdala: a potential mechanism for drug-induced seizures? Anesthesiology. 2014; 120: 639-49

49. Furtmuller R, Schlag MG, Berger M, et al. Tranexamic acid, a widely used antifibrinolytic agent, causes convulsions by a gamma-aminobutyric acid(A) receptor antagonistic effect. J Pharmacol Exp Ther. 2002; 301: 168-73

50. Manji RA, Grocott HP, Leake J, et al. Seizures following cardiac surgery: the impact of tranexamic acid and other risk factors. Can J Anaesth. 2012; 59: 6-13

51. Keyl C, Uhl R, Beyersdorf F, et al. High-dose tranexamic acid is related to increased risk of generalized seizures after aortic valve replacement. Eur J Cardiothorac Surg. 2011; 39: e114-21

52. Mirmohammadsadeghi A, Mirmohammadsadeghi M, and Kheiri M. Does topical tranexamic acid reduce postcoronary artery bypass graft bleeding? J Res Med Sci. 2018; 23: 6

53. Ali Shah MU, Asghar MI, Siddiqi R, et al. Topical application of tranexamic acid reduces postoperative bleeding in open-heart surgery: myth or fact? J Coll Physicians Surg Pak. 2015; 25: 161-5

54. Montroy J, Hutton B, Moodley P, et al. The efficacy and safety of topical tranexamic acid: A systematic review and meta-analysis. Transfus Med Rev. 2018; 32: 165-78

Tables

Table 1. Characteristics of included studies
| Study ID   | Country      | No. | Sex | Type of GABG | AC discounted before surgery | Drug Dose and Treatment Regimens |
|------------|--------------|-----|-----|--------------|------------------------------|----------------------------------|
| Speekenbrink 1995 | Netherlands | 15/15 | 2/28 | On-pump | 2 to 4 days | TA 10 mg·kg⁻¹ in 20 minutes after induction of anesthesia and continued at a rate of 1 mg·kg⁻¹ up to a total dose of 1,000 mg. |
| Brown 1997 | United States | 30/30 | 11/49 | On-pump | NR | TA 15 mg·kg⁻¹ in 20 minutes after the induction and continued at a rate of 1 mg·kg⁻¹·hr⁻¹ for 5 h |
| Landymore 1997 | Canada | 50/56 | NR | On-pump | <2 days | TA 10 mg·kg⁻¹ before CBP and continued at a rate of mg·kg⁻¹·hr⁻¹ until the termination of CBP |
| Hardy 1998 | Canada | 45/43 | 23/65 | On-pump | NR | TA 10 g as a bolus over 20 minutes |
| Casati 2001 | Italy | 20/20 | 8/32 | Off-pump | <1 day | TA 1 g as a bonus before skin incision, followed by continuous infusion of 400 mg·hr⁻¹ during surgery |
| Zabeeda 2002 | Israel | 25/25 | 12/38 | On-pump | NR | TA 10 mg·kg⁻¹ in more than 15 minutes after induction of anesthesia and followed by a continuous infusion of 1 mg·kg⁻¹ per hour |
| Jares 2003 | Czech Republic | 22/25 | 12/35 | Off-[1-14]pump | 5 days | TA 1 g as a bolus before skin incision, |

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Speekenbrink 1995

Brown 1997

Landymore 1997

Hardy 1998

Casati 2001

Zabeeda 2002

Jares 2003
| Author   | Country     | Age Range | Male/Female | On-Pump | Treatment Duration | Type of CPB | TA Dose and Details                                                                 |
|----------|-------------|------------|-------------|---------|--------------------|-------------|-----------------------------------------------------------------------------------|
| Pleym    | Norway      | 39/40      | 13/66       | On-pump | 1 day              |             | TA 30 mg·kg⁻¹ as a bolus injection over 5 min immediately before the start of CPB. |
| Andreasen| Denmark     | 23/21      | 7/37        | On-pump | >7 days            |             | TA 1.5 g as a bolus, followed by a constant infusion of 200 mg·hr⁻¹ until 1.5 g    |
| Casati   | Italy       | 50/52      | 16/86       | On-pump | <1 day             | Off-pump    | TA 1 g as a bonus before skin incision, followed by continuous infusion of 400 mg·hr⁻¹ until completion of surgery with 500 mg added to priming in patients undergoing on-pump coronary artery bypass grafting |
| Karski   | Canada      | 165/147    | 37/275      | On-pump | 7 days             |             | TA 100 mg·kg⁻¹ administered intravenously over 20 minutes after the induction of anesthesia |
| Vanek    | Czech Rep.  | 30/32      | 14/38       | Off-pump| <1 day             |             | TA 1 g before skin incision and a continuous infusion of 200 mg·hr⁻¹ during the whole surgical procedure. |
| Santos   | Brasil      | 31/29      | 17/43       | On-pump | NR                 |             | TA 10 mg·kg⁻¹ before the skin incision, followed by a continuous infusion of 1 mg·kg⁻¹·hr⁻¹ for 5 hr. |
| Author       | Country | Sample Size | Intraoperative TA | Surgery Type | Duration | TA Dose and Administration |
|--------------|---------|-------------|-------------------|--------------|----------|-----------------------------|
| Wei 2006     | China   | 40/36       | 16/60             | Off-Pump     | 5/-7 days | TA 0.75 g in 20 minutes at the beginning of surgery followed by continuous infusion of 0.25 g per hour throughout surgery. |
| Maddali 2007 | Oman    | 111/111     | 70/152            | On-Pump      | 7 days    | TA 10 mg·kg⁻¹ as a bolus prior to sternotomy, followed by an infusion (1 mg·kg⁻¹·hr⁻¹) up to the time of starting of protamine. |
| Mehr-Aein 2007 | Iran  | 33/33       | 2/27              | Off-Pump     | 7 days    | TA 15 mg·kg⁻¹ before infusion of heparin and 15 mg·kg⁻¹ after protamine infusion |
| Taghaddomi 2009 | Iran  | 50/50       | 28/72             | Off-Pump     | NR       | TA 1 g was given 20 minutes before skin incision and 400 mg·hr⁻¹ during the entire surgical procedure. |
| Hashemi 2011 | Iran    | 50/50       | 24/76             | On-Pump      | NR       | TA 1 g added to the pump prime solution and another 1 g was used intravenously after discontinuation of the pump |
| Ahn 2012     | Korea   | 38/38       | 35/41             | Off-Pump     | 5 days    | TA 1 g in 20 min before skin incision with subsequent continuous infusion at 200 mg·hr⁻¹ during the operation |
| Chakravarthy 2012 | India | 50/50       | 22/78             | Off-Pump     | 7 days    | TA 20 mg·kg⁻¹ over 30 min followed by infusion of 1
| Author    | Country | Days | TA Dose | Route | TA Formulation |
|-----------|---------|------|---------|-------|----------------|
| Greiff    | Norway  | 33/30| 26/37   | On-pump | 1 day          |
| Nejad     | Iran    | 50/50| 24/76   | On-pump | NR            |
| Wang      | China   | 115/116| 36/195 | Off-pump | 5 days         |
| Esfandiari| Iran    | 75/75| 30/120  | On-pump | NR            |
| Shi       | China   | 59/58| 23/94   | On-pump | <7 days        |
| Ghavidel  | Iran    | 100/100| 65/135 | On-pump | 3 days         |

TA 10 mg·kg\(^{-1}\)·hr\(^{-1}\) as a bolus injection before skin incision followed by an infusion of 1 mg·kg\(^{-1}\)·hr\(^{-1}\) until the end of surgery.

TA 1 g was added to the pump prime solution and another 1 g was used intravenously after the discontinuation of the pump.

TA 1 g as a bolus injection 20 minutes before the incision followed by an infusion of 400 mg·hr\(^{-1}\) until the completion of the surgery.

TA 10 mg·kg\(^{-1}\) added to the priming solution and a bolus dose of 1 mg·kg\(^{-1}\) after weaning from CPB.

TA 15 mg·kg\(^{-1}\) before surgical incision and 15 mg·kg\(^{-1}\) after protamine neutralization.

TA 10 mg·kg\(^{-1}\) via prime solution and the maintenance dose of 0.5-2 mg·kg\(^{-1}\)·h\(^{-1}\) in proportion to serum creatinine.
Yanartas 2015

Turkey 63/69 50/82 On-pump 5 days TA 10 mg·kg-1 before the skin incision, followed by a continuous infusion of 1 mg·kg-1·h-1 for 5 hours.

Myles 2017

Australia 2322/2311 773/3860 On-pump/Off-pump ≥4 days TA 100 mg·kg-1 or 50 mg·kg-1 was administered intravenously more than 30 minutes after the induction of anesthesia

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### Table 2. Sensitivity analysis of primary and secondary outcomes

| Outcome                          | Sensitivity analyses | Studies (n) | TXA     | Placebo    | RR or MD | 95% CI    | P value for effect | P value for heterogeneity |
|----------------------------------|----------------------|-------------|---------|------------|----------|-----------|--------------------|--------------------------|
| cerebrovascular accident         | Studies randomized not less 100 patients | 9           | 286/2999 | 318/3011   | 0.90     | 0.78-1.05 | 0.18               | 0.64                     |
| Study with maximum sample size excluded | 21                   | 9/1062      | 10/1084 | 0.95       | 0.43-2.10| 0.90      | 0.86               |                          |
| Condition                        | Study with maximum sample size excluded | Study randomized not less 100 patients | Odds Ratio | 95% CI       | p-Value   | HR       |
|---------------------------------|----------------------------------------|--------------------------------------|------------|--------------|-----------|----------|
| reoperation for bleeding        | 15                                     | 8                                    | 0.59       | 0.34-1.04    | 0.07      | 0.72     |
| mortality                       | 7                                      | 7                                    | 0.87       | 0.54-1.40    | 0.56      | 0.46     |
| myocardial infarction           | 11                                     | 11                                  | 0.90       | 0.78-1.05    | 0.18      | 0.64     |
| acute renal insufficiency       | 13                                     | 13                                  | 0.88       | 0.42-1.84    | 0.73      | 0.94     |
| Transfusion of any blood products| 7                                      | 7                                    | 0.64       | 0.50-0.81    | <0.01     | <0.01    |
| Postoperative chest tube        | 10                                     | 10                                  | 0.29       | 0.20-0.40    | <0.01     | <0.01    |
|                                 | 7                                      | 7                                    | -208.3     | -274.12-142.48 | <0.01     | <0.01    |
drainage in the first 24 hours

| Study with maximum sample size excluded | 17 | 802 | 814 | -215.42 | -259.48, -171.57 | <0.01 | <0.01 |

TXA = tranexamic acid; (n) = the number of cases; RR = risk ratio; MD = weighted mean difference; CI = confidence interval

Table 3 GRADE summary of findings table

| Outcomes | Illustrative comparative risks* (95% CI) | Relative effect (95% CI) | No of Participants (studies) | Quality of the evidence (GRADE) | Comments |
|----------|----------------------------------------|-------------------------|-----------------------------|--------------------------------|----------|
| Assumed risk | Corresponding risk | | | | |
| Control | Tranexamic acid versus placebo | | | | |
| Cerebrovascular accident | Study population | RR 0.93 (0.62 to 1.39) | 6775 (22 studies) | †††⊕ | moderate1 |
| 13 per 1000 | 12 per 1000 (8 to 18) | | | | |
| Moderate | | | | | |
| 0 per 1000 | 0 per 1000 (0 to 0) | | | | |
| Seizure | Study population | RR 6.67 (1.77 to 25.20) | 4911 (4 studies) | †††⊕⊕ | high2,3 |
| 1 per 1000 | 5 per 1000 (1 to 20) | | | | |
| Moderate | | | | | |
| 0 per 1000 | 0 per 1000 (0 to 0) | | | | |

34
| Event                                | Study population | RR       | 95% CI     | Number of Studies | Quality | Event Rate | Study Rate | Level of Evidence |
|--------------------------------------|------------------|----------|------------|-------------------|---------|------------|-------------|-------------------|
| Reoperation for bleeding             |                  | 0.46     | (0.31 to 0.68) | 6259 (16 studies) | ⊗⊗⊗⊗   | 25 per 1000 | 11 per 1000 | high 4,5          |
|                                      |                  |          |            |                   |         |            |            |                   |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
|                                      | 22 per 1000      | 0.46     | (0.31 to 0.68) | 6259 (16 studies) | ⊗⊗⊗⊗   | 22 per 1000 | 10 per 1000 |                   |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
| Mortality                            |                  | 0.82     | (0.53 to 1.28) | 6414 (17 studies) | ⊗⊗⊗⊗ ⊝ | 13 per 1000 | 10 per 1000 | moderate 1,6      |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
|                                      | 0 per 1000       | 0.82     | (0.53 to 1.28) | 6414 (17 studies) | ⊗⊗⊗⊗ ⊝ | 0 per 1000  | 0 per 1000  |                   |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
| Myocardial infarction                |                  | 0.9      | (0.78 to 1.05) | 6714 (23 studies) | ⊗⊗⊗⊗ ⊝ | 97 per 1000 | 87 per 1000 | moderate 4        |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
|                                      | 0 per 1000       | 0.9      | (0.78 to 1.05) | 6714 (23 studies) | ⊗⊗⊗⊗ ⊝ | 0 per 1000  | 0 per 1000  |                   |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
| Acute renal insufficiency            |                  | 1.01     | (0.78 to 1.3)  | 5954 (14 studies) | ⊗⊗⊗⊗ ⊝ | 37 per 1000 | 37 per 1000 | moderate 1        |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
|                                      | 20 per 1000      | 1.01     | (0.78 to 1.3)  | 5954 (14 studies) | ⊗⊗⊗⊗ ⊝ | 20 per 1000 | 20 per 1000 |                   |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
| Transfusion of any blood products    |                  | 0.64     | (0.52 to 0.78) | 5360 (11 studies) | ⊗⊗⊗⊗ ⊝ | 553 per 1000 | 354 per 1000 | very low 1,7      |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
|                                      | 560 per 1000     | 0.64     | (0.52 to 0.78) | 5360 (11 studies) | ⊗⊗⊗⊗ ⊝ | 560 per 1000 | 358 per 1000 |                   |
|                                      | Moderate         |          |            |                   |         |            |            |                   |
| Postoperative                        | The mean         |          |            | 6247 (16)        | ⊗⊗⊗⊗ ⊝ |            |            | very low          |
*The basis for the assumed risk (e.g. the median control group risk across studies) is provided in footnotes. The corresponding risk (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and its 95% CI). CI: Confidence interval; RR: Risk ratio; OR: Odds ratio;

GRADE Working Group grades of evidence High quality: Further research is very unlikely to change our confidence in the estimate of effect. Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate. Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate. Very low quality: We are very uncertain about the estimate.

1 4 studies with a high risk of bias were included 2 few studies reported this result 3 RR>5 4 5 studies with a high risk of bias were included 5 RR<0.5 6 No explanation was provided 7 I2>75% 8 2 studies with a high risk of bias were included

Figures

Figure 1

Flow diagram of the literature search strategy.
Figure 2

(A) risk-of-bias summary; (B) risk-of-bias graph for all the included randomized-controlled trials.

Figure 3

Forest plot of cerebrovascular accident.

Figure 4

Forest plot of seizures.

Figure 5

Forest plot of operation for bleeding.

Figure 6

Forest plot of mortality.

Figure 7

Forest plot of myocardial infarction.

Figure 8

Forest plot of acute renal insufficiency.

Figure 9

Forest plot of transfusion of any blood products.

Figure 10

Forest plot of chest tube drainage in the first 24 hours.
Supplementary Files

This is a list of supplementary files associated with the primary manuscript. Click to download.

Supplementary Fig 1. Funnel plot of cerebrovascular accident.png
Supplementary Fig 4. Funnel plot of myocardial infarction.png
Supplementary Fig 2. Funnel plot of reoperation for bleeding.png
Supplementary Fig 3. Funnel plot of mortality.png
Supplementary Fig 5. Funnel plot of AKI.png
Supplementary Fig 7. blood loss in 24h.png
Supplementary Fig 6. transfusion of any blood products.png