

Central Lancashire Online Knowledge (CLoK)

Title	Effects of a high-dose 24-h infusion of tranexamic acid on death and thromboembolic events in patients with acute gastrointestinal bleeding (HALT-IT): an international randomised, double-blind, placebo-controlled trial
Туре	Article
URL	https://clok.uclan.ac.uk/35141/
DOI	https://doi.org/10.1016/S0140-6736(20)30848-5
Date	2020
Citation	Joshi, Miland and HALT-IT Tial Collaborators, The (2020) Effects of a high-dose 24-h infusion of tranexamic acid on death and thromboembolic events in patients with acute gastrointestinal bleeding (HALT-IT): an international randomised, double-blind, placebo-controlled trial. The Lancet, 395. pp. 1927-1936. ISSN 0140-6736
Creators	Joshi, Miland and HALT-IT Tial Collaborators, The

It is advisable to refer to the publisher's version if you intend to cite from the work. https://doi.org/10.1016/S0140-6736(20)30848-5

For information about Research at UCLan please go to http://www.uclan.ac.uk/research/

All outputs in CLoK are protected by Intellectual Property Rights law, including Copyright law. Copyright, IPR and Moral Rights for the works on this site are retained by the individual authors and/or other copyright owners. Terms and conditions for use of this material are defined in the http://clok.uclan.ac.uk/policies/

Effects of a high-dose 24-h infusion of tranexamic acid on death and thromboembolic events in patients with acute gastrointestinal bleeding (HALT-IT): an international randomised, double-blind, placebo-controlled trial



The HALT-IT Trial Collaborators'

Summary

Background Tranexamic acid reduces surgical bleeding and reduces death due to bleeding in patients with trauma. Meta-analyses of small trials show that tranexamic acid might decrease deaths from gastrointestinal bleeding. We aimed to assess the effects of tranexamic acid in patients with gastrointestinal bleeding.

Methods We did an international, multicentre, randomised, placebo-controlled trial in 164 hospitals in 15 countries. Patients were enrolled if the responsible clinician was uncertain whether to use tranexamic acid, were aged above the minimum age considered an adult in their country (either aged 16 years and older or aged 18 years and older), and had significant (defined as at risk of bleeding to death) upper or lower gastrointestinal bleeding. Patients were randomly assigned by selection of a numbered treatment pack from a box containing eight packs that were identical apart from the pack number. Patients received either a loading dose of 1 g tranexamic acid, which was added to 100 mL infusion bag of 0.9% sodium chloride and infused by slow intravenous injection over 10 min, followed by a maintenance dose of 3 g tranexamic acid added to 1 L of any isotonic intravenous solution and infused at 125 mg/h for 24 h, or placebo (sodium chloride 0.9%). Patients, caregivers, and those assessing outcomes were masked to allocation. The primary outcome was death due to bleeding within 5 days of randomisation; analysis excluded patients who received neither dose of the allocated treatment and those for whom outcome data on death were unavailable. This trial was registered with Current Controlled Trials, ISRCTN11225767, and ClinicalTrials.gov, NCT01658124.

Findings Between July 4, 2013, and June 21, 2019, we randomly allocated 12 009 patients to receive tranexamic acid (5994, 49·9%) or matching placebo (6015, $50\cdot1\%$), of whom 11952 (99·5%) received the first dose of the allocated treatment. Death due to bleeding within 5 days of randomisation occurred in 222 (4%) of 5956 patients in the tranexamic acid group and in 226 (4%) of 5981 patients in the placebo group (risk ratio [RR] $0\cdot99$, 95% CI $0\cdot82-1\cdot18$). Arterial thromboembolic events (myocardial infarction or stroke) were similar in the tranexamic acid group and placebo group (42 [$0\cdot7\%$] of 5952 vs 46 [$0\cdot8\%$] of 5977; $0\cdot92$; $0\cdot60$ to $1\cdot39$). Venous thromboembolic events (deep vein thrombosis or pulmonary embolism) were higher in tranexamic acid group than in the placebo group (48 [$0\cdot8\%$] of 5952 vs 26 [$0\cdot4\%$] of 5977; RR $1\cdot85$; 95% CI $1\cdot15$ to $2\cdot98$).

Interpretation We found that tranexamic acid did not reduce death from gastrointestinal bleeding. On the basis of our results, tranexamic acid should not be used for the treatment of gastrointestinal bleeding outside the context of a randomised trial.

Funding UK National Institute for Health Research Health Technology Assessment Programme.

Copyright © 2020 The Author(s). Published by Elsevier Ltd. This is an Open Access article under the CC BY 4.0 license.

Introduction

Acute severe gastrointestinal bleeding is a common cause of death worldwide.¹ Bleeding can occur from the upper or lower gastrointestinal tract, but upper gastrointestinal bleeding is more common. The leading causes are peptic ulcer, oesophageal varices, and malignancy. The case fatality rate is approximately 10% for upper gastrointestinal bleeding and 3% for lower gastrointestinal bleeding.².³ Many patients re-bleed after initial haemostasis and those that do have a four-times increased risk of death.⁴ Patients with acute severe gastrointestinal bleeding usually present with haematemesis or

melaena. Patients are often haemodynamically unstable and in need of urgent resuscitation. Acute management of gastrointestinal bleeding includes blood product transfusion, medical or endoscopic therapy, and surgery.

Tranexamic acid reduces bleeding by inhibiting blood clot breakdown (fibrinolysis). Tranexamic acid decreases surgical bleeding and reduces death due to bleeding in patients with traumatic and postpartum haemorrhage. 5-8 A systematic review and meta-analysis of randomised trials of tranexamic acid for upper gastrointestinal bleeding included seven trials with a total of 1654 patients. 9 There was a large reduction in all-cause mortality with tranexamic

OPEN ACCESS

Lancet 2020; 395: 1927-36

See Comment page 1885

haltit@lshtm.ac.uk

*Members listed at end of paper Correspondence to: Clinical Trials Unit, London School of Hygiene & Tropical Medicine. London WC1 F7HT. UK

Research in context

Evidence before this study

Before this study a Cochrane systematic review and meta-analysis of randomised trials of tranexamic acid for upper gastrointestinal bleeding included seven trials with a total of 1654 patients. There was a large reduction in mortality with tranexamic acid (pooled risk ratio [RR] 0.61, 95% CI 0.42-0.89; p=0.01). However, given the small size of the included trials and the potential for selection and other biases, we considered this evidence to be hypothesis generating, requiring confirmation in larger trials. Furthermore, there was substantial uncertainty about the risk of thromboembolic events with tranexamic acid (pooled RR 1.86, 95% CI 0.66-5.24).

Added value of this study

The HALT-IT trial included 12 009 patients from 164 hospitals in 15 countries. Adult patients with significant upper or lower gastrointestinal bleeding were randomly assigned to receive tranexamic acid (1 g loading dose followed by 3 g maintenance dose over 24 h) or matching placebo. Tranexamic acid did not

reduce death from gastrointestinal bleeding (RR 0-99, 95% CI 0-82–1·18) but was associated with an increased risk of venous thromboembolic events (1-85, 1-15–2-98) and seizures (1-73, $1\cdot03-2\cdot93$).

Implications of all the available evidence

The most recent update of the Cochrane review included eight small randomised trials with 1701 participants and showed a reduction in mortality with tranexamic acid (RR 0-60, 95% CI 0-42–0-87). Although we cannot entirely rule out a modest increase or decrease in death due to bleeding with tranexamic acid, we can rule out the large mortality reduction suggested by the Cochrane review. Furthermore, tranexamic acid appears to increase the risk of venous thromboembolic events in patients with gastrointestinal bleeding. On the basis of our results, tranexamic acid should not be used for the treatment of gastrointestinal bleeding outside the context of a randomised trial. Our results highlight the unreliability of meta-analyses of small trials.

acid (risk ratio [RR] 0.61, 95% CI 0.42–0.89; p=0.01). However, meta-analyses of small trials are prone to publication and other selection biases, and have a low positive predictive value when compared with results from large multicentre trials. ¹⁰ Furthermore, even in aggregate, the trials included in the meta-analysis were too small to assess the effect of tranexamic acid on thromboembolic adverse events. ⁹ Our objective was to quantify the effects of tranexamic acid on death and thromboembolic events in acute gastrointestinal bleeding.

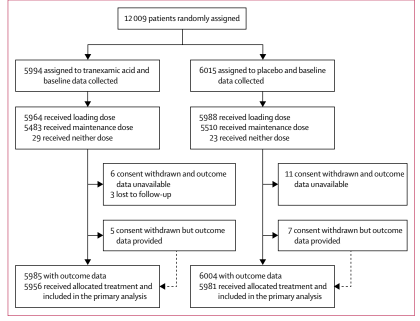


Figure 1: Trial profile

Methods

Study design and participants

The HALT-IT trial is an international, randomised, double blind (participants and trial staff), placebocontrolled trial done in 164 hospitals in 15 countries (UK, Pakistan, Nigeria, Egypt, Malaysia, Georgia, Romania, Nepal, Sudan, Saudi Arabia, Spain, Ireland, Albania, Papua New Guinea, and Australia).11 Patients were enrolled if they were aged above the minimum age considered an adult in their country (either aged 16 years and older or aged 18 years and older) and if the responsible clinician was substantially uncertain whether to use tranexamic acid. The diagnosis of significant bleeding was clinical and significant was defined as a risk of bleeding to death and included patients with hypotension, tachycardia, or signs of shock, or those likely to need transfusion or urgent endoscopy or surgery.

Severe gastrointestinal bleeding is a frightening experience and blood loss can impact on a patient's mental and emotional state, impairing their decision making—the consent procedures considered this, as well as the need to treat urgently. If the patient was fully competent, written consent was sought. If capacity was impaired and a personal or professional representative was available, consent was sought from the representative. If neither were able to provide consent, it was waived and the patient was informed about the trial and consented for ongoing data collection as soon as possible afterwards. The trial was approved by the UK NRES Committee East of England (reference number 12/EE/0038), and by the national and local research ethics committees in all participating non-UK countries.

	Tranexamic acid (n=5994)	Placebo (n=6015)
Age at randomisation	ı, years	
Mean (SD)	58·1 (17·0)	58.1 (17.0)
<40	791 (13%)	779 (13%)
40-59	2356 (39%)	2333 (39%)
50-79	2078 (35%)	2130 (35%)
≥80	769 (13%)	773 (13%)
Sex		
emale	2142 (36%)	2124 (35%)
Male	3852 (64%)	3891 (65%)
Time from onset to ra	indomisation, h	
Mean (SD)	21.4 (36.4)	22.5 (37.8)
≤3	960 (16%)	975 (16%)
>3-≤8	1607 (27%)	1551 (26%)
-8	3427 (57%)	3488 (58%)
Missing	0	1 (<1%)
Suspected location of	bleeding	
_ower	674 (11%)	654 (11%)
Jpper	5320 (89%)	5361 (89%)
Haematemesis		
⁄es	4285 (72%)	4240 (71%)
No	1709 (29%)	1775 (30%)
Melaena or fresh bloo	d per rectum	
res .	4573 (76%)	4626 (77%)
No	1421 (24%)	1389 (23%)
Suspected variceal ble	eeding	
res .	2694 (45%)	2739 (46%)
No	3300 (55%)	3276 (54%)
Suspected active blee	ding	
res .	5247 (88%)	5226 (87%)
No	747 (12%)	789 (13%)
Systolic blood pressu	re, mm Hg	
≥90	5222 (87%)	5216 (87%)
76–89	577 (10%)	577 (10%)
.7F	181 (3%)	201 (3%)
≤75		

Randomisation and masking

An independent statistician from Sealed Envelope (London, UK) generated randomisation numbers and these were given to Sharp Clinical Services UK (Crickhowell, UK), a Good Manufacturing Practice certified clinical trial service provider, to make treatment packs. When a patient was enrolled, the lowest numbered treatment pack was taken from a box of eight packs. Sharp Clinical Services was responsible for masking, which involved removing the manufacturer's label and replacing it with the clinical trial label and randomisation number. Apart from the randomisation number, the pack label text was identical for tranexamic acid and placebo. Patients, caregivers, and those assessing outcomes were masked to allocation. We checked the coding by testing each batch of ampoules with high-performance liquid chromatography

	Tranexamic acid (n=5994)	Placebo (n=6015)		
(Continued from previous column)				
Heart rate, beats per min				
<77	812 (14%)	756 (13%)		
77-91	1546 (26%)	1644 (27%)		
92-107	1760 (29%)	1720 (29%)		
>107	1864 (31%)	1885 (31%)		
Missing	12 (<1%)	10 (<1%)		
Signs of shock				
Yes	2574 (43%)	2648 (44%)		
No	3420 (57%)	3367 (56%)		
Rockall score				
1-2	1419 (24%)	1395 (23%)		
3-4	2306 (38%)	2332 (39%)		
5-7	2269 (38%)	2288 (38%)		
Taking anticoagulants				
Yes	528 (9%)	500 (8%)		
No	5422 (90%)	5466 (91%)		
Unknown	44 (1%)	49 (1%)		
Emergency admission				
Yes	5673 (95%)	5687 (94%)		
No	321 (5%)	328 (6%)		
Major comorbidities				
Cardiovascular	1108 (18%)	1132 (19%)		
Respiratory	337 (6%)	324 (5%)		
Liver	2432 (41%)	2532 (42%)		
Renal	325 (5%)	310 (5%)		
Malignancy	417 (7%)	382 (6%)		
Other	999 (17%)	968 (16%)		
Any comorbidity	4308 (72%)	4329 (72%)		
Pata are n (%) or mean (SD).				

to determine the contents. Block randomisation was used but randomisation was not stratified.

Procedures

Eligible patients were randomly assigned to get tranexamic acid or placebo as soon as possible and treatment was started immediately. A loading dose of 1 g tranexamic acid or placebo (sodium chloride 0.9%) was added to a 100 mL infusion bag of 0.9% sodium chloride and infused by slow intravenous injection over 10 min, followed by a maintenance dose of 3 g tranexamic acid or placebo added to 1 L of any isotonic intravenous solution and infused at 125 mg/h for 24 h. Every patient was assigned a uniquely numbered treatment pack, which contained eight ampoules of tranexamic acid 500 mg or placebo, one 100 mL bag of 0.9% sodium chloride (to use with the loading dose), two sterile 10 mL syringes and needles, stickers with the trial details and randomisation number (for attaching to infusion bags, forms, and the medical records), and instructions. Pfizer, Sandwich, UK (PL 00057/0952) manufactured the tranexamic acid

and Torbay and South Devon NHS Foundation Trust (MIA [IMP] 13079) manufactured the sodium chloride 0.9% placebo. We provided information for patients and representatives, consent forms, and data collection forms. Stickers, instructions, leaflets, and forms were in local languages.

Once randomly assigned, we collected outcome data even if the treatment was not given. Outcome data were collected at death, discharge from the randomising hospital, or 28 days after randomisation, whichever occurred first. Trial investigators and their institutions provided direct access to the source data for trial-related monitoring, audits, and regulatory inspections. Monitoring was done according to the Sponsor's Standard Operating Procedure and the trial protocol. Formal inspections were carried out by the relevant Regulatory Agencies including the UK Medicines and Healthcare products Regulatory Agency, Irish Health Products Regulatory Authority, and Nigeria's National Agency for Food and Drug Administration and Control. Adherence to allocation sequence was monitored throughout the trial and any out of sequence pack use was automatically

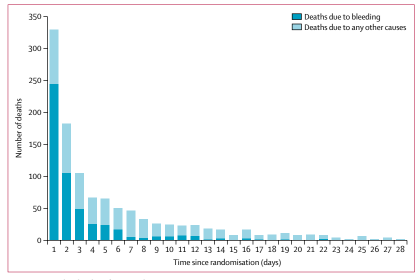


Figure 2: Mortality by days from randomisation

	Tranexamic acid (n=5956)	Placebo (n=5981)	Risk ratio (95% CI)
Death due to bleeding within 24 h	124 (2·1%)	120 (2.0%)	1.04 (0.81-1.33)
Death due to bleeding within 5 days	222 (3.7%)	226 (3.8%)	0.99 (0.82-1.18)
Death due to bleeding within 28 days	253 (4-2%)	262 (4.4%)	0.97 (0.82-1.15)
Rebleeding within 24 h*	41 (0.7%)	41 (0.7%)	1.00 (0.65–1.55)
Rebleeding within 5 days*	287 (4.8%)	315 (5.3%)	0.91 (0.78-1.07)
Rebleeding within 28 days*	410 (6.8%)	448 (7.5%)	0.92 (0.81–1.05)

Data are n (%) and risk ratio (95% CI). Death or rebleeding in hospital during follow-up. *Excludes 13 patients missing data on rebleed status or rebleed date.

Table 2: Effect of tranexamic acid on death due to bleeding and rebleeding

flagged in the trial database and the investigators were retrained.

Outcomes

The primary outcome was death due to bleeding within 5 days of randomisation. Cause of death was assigned by local principal investigators who provided a narrative of events leading to death. These were reviewed by the chief investigator (masked to treatment allocation) and queried if more information was needed to confirm whether death was due to bleeding or another cause. Secondary outcomes were death due to bleeding within 24 h and within 28 days of randomisation, all-cause and cause-specific mortality at 28 days, rebleeding within 24 h, within 5 days, and within 28 days of randomisation, surgery or radiological intervention, blood product transfusion, thromboembolic events (deep vein thrombosis, pulmonary embolism, stroke, and myocardial infarction), seizures, other complications (including other significant cardiac event, sepsis, pneumonia, respiratory failure, renal failure, liver failure), days in an intensive care unit, and functional status. The diagnosis of rebleeding was made by the clinician based on established criteria. A diagnosis of thromboembolic events was made using strict definitions and diagnostic criteria, including a clinical assessment, diagnostic imaging, biomarker tests, and post-mortem examination. Seizures were diagnosed by clinical assessment. Functional status was measured with the Katz Index of Independence in Activities of Daily Living either at hospital discharge or in-hospital at 28 days.

Statistical analysis

The sample size calculation was initially based on allcause mortality as the primary outcome since we expected that most deaths would be due to bleeding.11 However, while the trial was underway, we observed that over half of all deaths were due to non-bleeding causes. Accumulating evidence from other large trials of tranexamic acid showed no apparent effect on non-bleeding deaths. 12 Furthermore, patients received tranexamic acid (or placebo) only for their initial bleed and because tranexamic acid has a short half-life (approximately 2 h), it will be largely eliminated within 2 days. As such, we did not expect tranexamic acid to reduce deaths from rebleeding episodes many weeks after randomisation. The primary outcome was therefore changed to death due to bleeding within 5 days of randomisation on Nov 21, 2018. Based on the amended primary outcome, assuming a risk of death due to bleeding of 4%, a study with 12000 patients has about 85% power (two-sided α of 5%) to detect a clinically important 25% relative reduction in death due to bleeding from 4% to 3%.

We published the statistical analysis plan before unblinding.¹³ The plan gave our reasons for amending the primary outcome measure and for increasing the sample size. The main analyses compared those allocated tranexamic acid with those allocated to placebo

on a modified intention-to-treat basis, excluding patients who received neither dose of the allocated treatment and those for whom outcome data on death were unavailable. We present effect estimates (RRs) with a measure of precision (95% CI). The safety of participants was overseen by an independent data monitoring committee, which reviewed four non-masked interim analyses. We planned to report four subgroup analyses to examine the effects of tranexamic acid on the primary outcome stratified by the following baseline characteristics: time to treatment ($\leq 3 \text{ h}$, >3 h), site of bleeding (upper vs lower gastrointestinal), suspected variceal bleeding and comorbid liver disease compared with other or unknown causes of bleeding, and by clinical Rockall score. We modelled an interaction between the treatment effect and time to treatment with time to treatment as a continuous variable. We did post-hoc subgroup analyses to examine the effects of tranexamic acid on the primary outcome stratified by World Bank country income level (high vs low and middle income), anticoagulant use, and systolic blood pressure.

This trial was registered with Current Controlled Trials, ISRCTN11225767, and ClinicalTrials.gov, NCT01658124.

Role of the funding source

The funders had no role in study design, data collection, data analysis, data interpretation, or writing of the report. The corresponding authors had full access to all the data in the study and had final responsibility for the decision to submit for publication.

Results

We enrolled the first patient on July 4, 2013, and the last on June 21, 2019. We stopped recruiting when the planned sample size was reached. When the decision to refine the primary outcome was made in Nov 21, 2018, we had recruited 10190 patients. This decision was made blind to the accumulating trial data. 12009 patients were enrolled and randomly assigned to receive either tranexamic acid (n=5994, 49.9%) or matching placebo (n=6015, 50.1%), of whom 11952 (99.5%) received the first dose of the allocated treatment (figure 1). 29 patients (11 in the tranexamic acid group and 18 in the placebo group) withdrew consent after randomisation, but of those, 12 (five in the tranexamic acid group and seven in the placebo group) agreed to provide outcome data or had outcome data collected as part of adverse event reporting. We obtained primary outcome data for all but three patients in the tranexamic acid group. There were 14 protocol violations (seven in the tranexamic acid group and seven in the placebo group), 11 patients did not meet the inclusion criteria (ten received tranexamic acid before randomisation [six in the tranexamic acid group and four in the placebo group], one in the placebo group was younger than 16 years), and there were three consent protocol violations (one in the tranexamic acid group and two in the placebo group). A total of 28 patients were unmasked to treatment (17 patients because the hospital team wanted to administer tranexamic acid, six because of adverse events, three because of clinical concerns, and two as part of post-mortem investigations [13 in the tranexamic acid group and 15 in the placebo group]). 52 patients received neither dose of the allocated trial treatment (29 patients in the tranexamic acid group and 23 patients in the placebo group).

223 patients received antifibrinolytic drugs as part of their clinical care, outside of the trial protocol (105 patients in the tranexamic acid group and 118 patients in the placebo group).

Baseline characteristics were similar between groups (table 1). Figure 2 shows the number of deaths and cause of death by days since randomisation. There were 1112 deaths. The median time to death was 55 h after randomisation (IQR $18 \cdot 2-161 \cdot 8$).

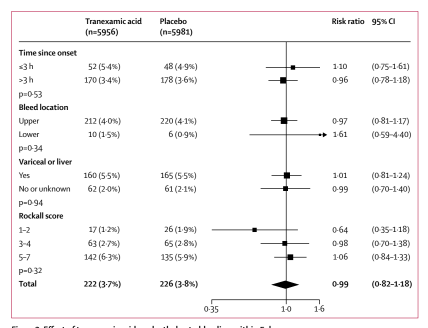


Figure 3: Effect of tranexamic acid on death due to bleeding within 5 days

Analysis stratified by time since bleeding onset, suspected bleed location, suspected variceal bleeding or comorbid liver disease, and Rockall score.

Tranexamic acid (n=5956)	Placebo (n=5981)	Risk ratio (95% CI)
253 (4·2%)	262 (4.4%)	0.97 (0.82-1.15)
26 (0-4%)	17 (0.3%)	1.54 (0.83-2.83)
109 (1.8%)	114 (1.9%)	0.96 (0.74-1.25)
57 (1.0%)	42 (0.7%)	1.36 (0.92-2.03)
33 (0.6%)	49 (0.8%)	0.68 (0.44-1.05)
65 (1.1%)	40 (0.7%)	1.63 (1.10-2.42)
21 (0-4%)	24 (0.4%)	0.88 (0.49-1.58)
564 (9.5%)	548 (9·2%)	1.03 (0.92-1.16)
	(n=5956) 253 (4·2%) 26 (0·4%) 109 (1·8%) 57 (1·0%) 33 (0·6%) 65 (1·1%) 21 (0·4%)	(n=5956) (n=5981) 253 (4·2%) 262 (4·4%) 26 (0·4%) 17 (0·3%) 109 (1·8%) 114 (1·9%) 57 (1·0%) 42 (0·7%) 33 (0·6%) 49 (0·8%) 65 (1·1%) 40 (0·7%) 21 (0·4%) 24 (0·4%)

Table 3: Effect of tranexamic acid on all-cause death

Death due to bleeding within 5 days of randomisation (table 2) occurred in 222 (3.7%) of 5956 patients in the tranexamic acid group and in 226 (3.8%) of 5981 patients in the placebo group (RR 0.99, 95% CI 0.82-1.18). Similar results were obtained after adjusting for baseline covariates (0.98, 0.82-1.17) and in a per-protocol analysis (0.94, 0.71-1.23). When the 223 patients who received

See Online for appendix

	Tranexamic acid	Placebo	Outcomes
Interventions			
Diagnostic endoscopy	4781/5953 (80-3%)	4729/5978 (79·1%)	1.02 (1.00 to 1.03)
Therapeutic endoscopy	2542/5952 (42-7%)	2658/5978 (44.5%)	0.96 (0.92 to 1.00)
Diagnostic radiological procedure	1704/5953 (28-6%)	1744/5978 (29-2%)	0.98 (0.93 to 1.04)
Therapeutic radiological procedure	74/5953 (1.2%)	89/5978(1.5%)	0.83 (0.61 to 1.13)
Surgical intervention	146/5953 (2.5%)	158/5978 (2.6%)	0.93 (0.74 to 1.16)
Any surgical, endoscopic, or radiological intervention	5216/5956 (87-6%)	5236/5981 (87.5%)	1.00 (0.99 to 1.01)
Any transfusion	4076/5951 (68-5%)	4129/5978 (69-1%)	0.99 (0.97 to 1.02)
Whole blood or red cells	3984/4076 (97.7%)	4018/4129 (97-3%)	1.00 (1.00 to 1.01)
Frozen plasma	910/4076 (22-3%)	993/4129 (24.0%)	0.93 (0.86 to 1.00)
Any platelets	219/4076 (5.4%)	255/4129 (6-2%)	0.87 (0.73 to 1.04)
Blood product transfusions			
Units of whole blood or red cells	2.8 (2.4)	2.9 (2.7)	-0.06 (0.05 to -0.18)
Units of frozen plasma	0.9 (2.4)	1.0 (2.6)	-0.05 (-0.01 to -0.23)
Units of any platelets	0.2 (0.9)	0.2 (1.0)	-0.02 (0.02 to -0.06)

Data for interventions are n/N (%) and risk ratio (95% CI); data for blood product transfusions are mean (SD) and difference in means (95% CI).

 ${\it Table 4:} \ Effect of transcamic acid on the need for surgical, endoscopic, and radiological interventions or blood product transfusion$

Tranexamic acid	Placebo	Outcomes
86/5952 (1.4%)	72/5977 (1.2%)	1·20 (0·88 to 1·64)
48/5952 (0.8%)	26/5977 (0.4%)	1.85 (1.15 to 2.98)
23/5952 (0.4%)	12/5977 (0.2%)	1.92 (0.96 to 3.86)
28/5952 (0.5%)	16/5977 (0.3%)	1.76 (0.95 to 3.24)
42/5952 (0.7%)	46/5977 (0.8%)	0·92 (0·60 to 1·39)
24/5952 (0.4%)	28/5977 (0.5%)	0.86 (0.50 to 1.48)
19/5952 (0.3%)	18/5977 (0.3%)	1.06 (0.56 to 2.02)
142/5951 (2.4%)	157/5978 (2.6%)	0.91 (0.73 to 1.14)
196/5952 (3.3%)	184/5977 (3·1%)	1.07 (0.88 to 1.30)
105/5952 (1.8%)	131/5978 (2.2%)	0.81 (0.62 to 1.04)
100/5952 (1.7%)	89/5977 (1.5%)	1·13 (0·85 to 1·50)
210/5952 (3.5%)	216/5977 (3.6%)	0.98 (0.81 to 1.18)
193/5952 (3.2%)	174/5978 (2.9%)	1·11 (0·91 to 1·36)
38/5952 (0.6%)	22/5977 (0.4%)	1.73 (1.03 to 2.93)
0.4 (1.8)	0.4 (2.0)	-0.06 (0.01 to -0.13)
5.5 (1.5)	5.5 (1.4)	-0.03 (0.02 to -0.09)
	86/5952 (1·4%) 48/5952 (0·8%) 23/5952 (0·4%) 28/5952 (0·5%) 42/5952 (0·7%) 24/5952 (0·3%) 142/5951 (2·4%) 196/5952 (3·3%) 105/5952 (1·8%) 100/5952 (1·7%) 210/5952 (3·2%) 38/5952 (0·6%)	86/5952 (1·4%) 72/5977 (1·2%) 48/5952 (0·8%) 26/5977 (0·4%) 23/5952 (0·4%) 12/5977 (0·2%) 28/5952 (0·5%) 16/5977 (0·3%) 42/5952 (0·7%) 46/5977 (0·5%) 19/5952 (0·3%) 18/5977 (0·3%) 142/5951 (2·4%) 157/5978 (2·6%) 196/5952 (3·3%) 184/5977 (3·1%) 105/5952 (1·8%) 131/5978 (2·2%) 100/5952 (1·7%) 89/5977 (1·5%) 210/5952 (3·5%) 216/5977 (3·6%) 193/5952 (3·2%) 174/5978 (2·9%) 38/5952 (0·6%) 22/5977 (0·4%)

Data for complications are n/N (%) and risk ratio (95% CI); data for self-care capacity are mean (SD) and difference in means (95% CI). Thromboembolic events and complications are not mutually exclusive. ICU=intensive care unit.

Table 5: Complications and self-care capacity in study groups

open-label antifibrinolytics were removed from the analysis, the results were similar (0.97, 0.81–1.17). We examined the effect of tranexamic acid on death due to bleeding within 5 days of randomisation in prespecified subgroup analyses stratified by time to treatment (heterogeneity p=0.53), location of bleeding (p=0.34), cause of bleeding (p=0.94), and clinical Rockall score (p=0.32) but recorded no evidence of heterogeneity for these factors (figure 3). When time since bleeding onset was modelled as a continuous variable there was no evidence of an interaction (heterogeneity p=0.53).

We examined the effect of tranexamic acid on death due to bleeding within 5 days of randomisation stratified by World Bank country income level (high-income *vs* low-income and middle-income), anticoagulant use and systolic blood pressure. These exploratory analyses were not prespecified. The relative risks did not appear to vary by country income, anticoagulant use, or systolic blood pressure (appendix p 6).

Death due to bleeding within 24 h of randomisation occurred in 124 (2·1%) patients in the tranexamic acid group and 120 (2.0%) patients in the placebo group (RR 1.04, 95% CI 0.81-1.33). Death due to bleeding within 28 days of randomisation occurred in 253 (4.2%) patients in the tranexamic acid group and 262 (4.4%) patients in the placebo group (0.97, 0.82-1.15). Death from all-causes within 28 days of randomisation occurred in 564 patients (9.5%) in the tranexamic acid group and in 548 patients (9.2%) in the placebo group ($1 \cdot 03$, $0 \cdot 92 - 1 \cdot 16$; table 3). The proportion of patients with rebleeding was similar in both groups at 24 h, 5 days, and 28 days after randomisation (table 3). The proportion of patients who had surgery, radiological intervention, and blood product transfusion was also similar in both groups (table 4).

The risk of fatal or non-fatal thromboembolic events and arterial thromboembolic events (myocardial infarction or stroke) was similar in the tranexamic acid group and the placebo group (table 5). The risk of venous thromboembolic events (deep vein thrombosis or pulmonary embolus) was higher in the tranexamic group than in the placebo group (table 5) and similar risk was observed after excluding patients who did not receive the maintenance dose (42 events with tranexamic acid vs 20 with placebo); RR 2.11, 95% CI 1.24-3.59). In an exploratory subgroup analysis, the risk of venous thromboembolic events was higher in patients with suspected variceal bleeding or liver disease (14 νs two events; 7.26, 1.65-31.90) than in patients with other causes of bleeding (34 ν s 24 events; 1.38, 0.82–2.32; p=0.035 for heterogeneity). The risk of renal, hepatic, and respiratory failure, cardiac events, sepsis, and pneumonia was similar in tranexamic acid and placebo treated patients (table 5). Seizures occurred in 38 patients on tranexamic acid and 22 on placebo (0.6% vs 0.4%; 1.73, 1.03-2.93; table 5), and after excluding patients who did not receive the maintenance dose the

corresponding numbers were 33 versus 17 events (1.95, 1.09-3.50).

The mean number of days spent in intensive care was similar in both groups (table 5). The mean score on the Katz Index of Independence in Activities of Daily Living was also similar in both groups (table 5).

Discussion

In this trial, tranexamic acid did not reduce death from gastrointestinal bleeding but was associated with an increased risk of venous thromboembolic events and seizures. The proportion of patients with rebleeding was similar in the tranexamic acid and placebo groups.

The randomisation method ensured that participating clinicians had no foreknowledge of the treatment allocation and placebo control ensured outcome assessment was blind to treatment group. The inclusion criteria were clinical, reflecting the full range of gastrointestinal bleeding presentations that doctors face in day-to-day practice. Baseline prognostic factors were well balanced and almost all randomly assigned patients were followed up. The primary outcome was death due to bleeding within 5 days of randomisation. Our scientific reasons for pre-specifying death due to bleeding as the primary outcome in the statistical analysis plan are presented in detail elsewhere.12 Although some misclassification of cause of death is possible, the assessment was masked to the treatment group. However, because there was no evidence of a treatment effect for the prespecified primary endpoint (death due to bleeding at 5 days) or for death from any cause at 28 days, the choice of endpoint does not influence the interpretation of the results. Misclassification might also have affected our subgroup analyses because at the time of recruitment the site and cause of bleeding cannot be known with certainty. Our use of the pre-endoscopy Rockall score might have misclassified baseline risk.14 To minimise the risk of false positives, we used strict criteria to diagnose thromboembolic events, including a positive result on imaging (eg, ultrasound) or at post-mortem examination. Although using this criteria might have led to some under reporting, because the diagnostic tests have high specificity, the relative risk estimates should be unbiased. Although some patients received antifibrinolytics outside of the protocol, the treatment effect was the same when these patients were excluded. Although this is one of the largest randomised trials in gastrointestinal bleeding, we cannot rule out a modest increase or decrease in death due to bleeding with tranexamic acid. That said, we can rule out the large mortality reduction suggested by the Cochrane systematic review and meta-analysis of previous small trials 9

Administration of tranexamic acid within 3 h of bleeding onset reduces death due to bleeding in trauma and post-partum haemorrhage without increasing the risk of thromboembolic events. In these bleeding

scenarios, the timing of onset is easy to determine, most patients present early, and there are well documented changes in fibrinolysis that provide a biological rationale for tranexamic acid treatment.^{15,16} However, in gastrointestinal bleeding it is difficult to determine the time of bleeding onset, presentation is often delayed (over 80% of patients presented more than 3 h after bleeding onset), and the contribution of increased fibrinolysis to bleeding is less clear.

Almost half of the patients included in our trial had suspected variceal bleeding due to liver disease and because these patients had a greater risk of death, they accounted for nearly three-quarters of deaths. Recent research shows that acutely ill patients with cirrhosis have a mixed fibrinolytic phenotype.¹⁷ Some have increased fibrinolysis, but others have profound hypofibrinolysis. The prevalence of hypofibrinolysis appears to be greatest in the most critically ill patients. Using the same clot lysis assay, reduced fibrinolysis has been shown to be associated with a small increased risk of venous thrombosis.18 In our trial, the increased risk of venous thromboembolic events with tranexamic acid appeared to be more marked in patients with liver disease, although this was an exploratory subgroup analysis and there was no strong evidence for heterogeneity. Nevertheless, reduced fibrinolysis in patients with liver disease might explain the absence of reduction in bleeding deaths with tranexamic acid and the increased risk of venous thromboembolic events.

The dose of tranexamic acid used in this trial was higher and the duration of treatment was longer (4 g over 24 h) than in randomised trials of tranexamic acid in trauma (2 g over 8 h) or post-partum haemorrhage (1 g bolus with a repeat 1 g dose if bleeding continued), which did not record any increase in adverse events with tranexamic acid. Patients with gastrointestinal bleeding often rebleed after initial haemostasis, particularly within the first 24 h. Because tranexamic acid has a short half-life, we used a longer treatment duration to cover this high-risk period. Furthermore, previous trials in gastrointestinal bleeding that appeared to show a large mortality reduction with tranexamic acid used a high dose and a longer duration of treatment than trials in trauma and post-partum haemorrhage.9 The longer duration of tranexamic acid treatment in this trial might explain the increased risk of venous thromboembolic events and the higher dose could possibly explain the increased risk of seizures.19

In summary, we found no evidence that tranexamic acid decreases the risk of death in patients with gastro-intestinal bleeding. Our results caution against a uniform approach to the management of patients with major haemorrhage and highlight the need for randomised trials targeted at specific pathophysiological processes. Because gastrointestinal bleeding is a licensed indication for tranexamic acid, our results could have regulatory implications. Although this trial can rule out the large

mortality reduction suggested by the meta-analysis of previous small trials, it cannot rule out more modest treatment effects. Because tranexamic acid reduces bleeding deaths in patients with traumatic and postpartum haemorrhage, individual patient data meta-analyses should assess the strength of the evidence that the effectiveness and safety of tranexamic acid varies by the site and cause of bleeding.

The HALT-IT Trial Collaborators

Writing Committee: Ian Roberts, Haleema Shakur-Still (Co-Chairs), Adefemi Afolabi, Adegboyega Akere, Monica Arribas, Amy Brenner, Rizwana Chaudhri, Ian Gilmore, Kenneth Halligan, Irshad Hussain, Vipul Jairath, Kiran Javaid, Aasia Kayani, Ton Lisman, Raoul Mansukhani, Muttiullah Mutti, Muhammad Arif Nadeem, Richard Pollok, Jonathan Simmons, Majid Soomro, Simon Stanworth, Andrew Veitch. Trial Steering Committee: Christopher Hawkey (Chair), Adefemi Afolabi, Jack Cuzick, Kenneth Halligan (patient representative), David Henry, Chris Metcalfe, Ian Roberts, Data Monitoring and Ethics Committee: Richard Gray (Chair), Alan Barkun, Suresh David, Philip Devereaux, Tony Brady (independent statistician). Protocol Committee: Ian Roberts, Haleema Shakur-Still, Timothy Coats, Phil Edwards, Ian Gilmore, Vipul Jairath, Katharine Ker, Daniela Manno, Simon Stanworth, Andrew Veitch. Clinical Trials Unit (CTU): Monica Arribas (trial manager/research assistant), Emma Austin (assistant trial manager), Kiran Bal (assistant trial manager), Eni Balogun (trial manager), Collette Barrow (trial administrator), Danielle Beaumont (senior trial manager/research fellow), Myriam Benyahia (CTU administrator), Amy Brenner (research fellow), Imogen Brooks (trial assistant 2016-18), Madeleine Cargill (data assistant), Laura Carrington (trial administrator), Phil Edwards (statistician 2012-16), Lauren Frimley (trial manager/ research assistant), Amber Geer (assistant data manager), Daniel Gilbert (data assistant 2012-13), Catherine Gilliam (trial administrator), Julio Gil Onandia (clerical assistant), Nayia Golfi (trial manager 2013–15), Daniel Hetherington (trial assistant 2012-15), Courtenay Howe (CTU administrator 2015-17), Carolyn Hughes (data assistant 2016-17), David I'anson (assistant trial manager 2016-17), Rob Jackson (data manager 2012-15), Miland Joshi (statistician 2016-17), Sneha Kansagra (assistant trial manager 2016-18), Taemi Kawahara (senior trial manager 2012-15), Katharine Ker (lecturer), Sergey Kostrov (systems officer 2015-19), Daniela Manno (clinical lecturer 2012-15), Raoul Mansukhani (medical statistician 2019-20), Hakim Miah (IT systems manager 2013-19), Bernard Ndungu (assistant trial manager 2016-17), Kelly Needham (statistician 2018–20), Aroudra Outtandy (trial assistant 2013-15), Daniel Pearson (data assistant 2018-19), Tracey Pepple (acting senior data manager 2014-19), Danielle Prowse (assistant data manager), Nigel Quashi (data manager 2013–16), Anna Quinn (data assistant 2013-15), Maria Ramos (senior project administrator 2012-15), Laura Ranopa (trial assistant 2015-20), Mia Reid (clerical assistant 2016-18), Ian Roberts (Chief Investigator/CTU co-director), Chris Roukas (trial administrator 2013-15), Haleema Shakur-Still (Project Director/ CTU co-director), Chelci Squires (trial assistant 2014-16), Jemma Tanner (clinical trials associate 2013-16), Andrew Thavne (data assistant), Ruhama Uddin (trial assistant 2018-19), Rawalpindi Medical University Pakistan National Coordinating Centre Team: Rizwana Chaudhri (Coordinating Centre Director), Muttiullah Mutti (Clinical Lead), Kiran Javaid (Assistant Research Coordinator), Aasia Kavani (Research Coordinator), Nigeria Coordinating Team, University College Hospital Ibadan: Bukola Fawole (Coordinating Centre Director), Folasade Adenike Bello (Coordinating Centre Director), Oladapo Olayemi (Coordinating Centre Director), Adefemi Afolabi (National Principal Investigator), Olujide Okunade (Assistant Trial Coordinator), Olusade Adetayo (Assistant Trial Coordinator). National Coordinators: Rizwana Chaudhri (Pakistan), Muttiullah Mutti (Pakistan), Adefemi Afolabi (Nigeria), Folasade Adenike Bello (Nigeria), Bukola Fawole (Nigeria), Oladapo Olayemi (Nigeria), Hussein Khamis (Egypt), Mohammad Shukri Bin Jahit (Malaysia), Tamar Gogichaishvili (Georgia), Radu Bogdan Mateescu (Romania), Ajay Adhikaree (Nepal), Abdelmounem Eltayeib Abdo (Sudan), Mohammad Zaher

(Saudi Arabia), Conor Deasy (Ireland), Joaquin Alvarez Gregori (Spain), Bobby Wellsh (Papua New Guinea), Luke Lawton (Australia). Trial sites and investigators: UK (4751): Royal Stoke University Hospital (303): Raghavendra Kamath, Adrian Barry, Racquel Carpio, Kay Finney, Holly Maguire; Queen's Medical Centre Campus Nottingham (208): Martin James, Frank Coffey, Chris Gough, Lisa Sawers, Aye-Aye Thi; Royal Berkshire Hospital (191): Jonathan Simmons, Claire Burnett, Nicola Jacques, Victoria Murray; St George's Hospital (173): Richard Pollok, Heather Jarman, Christine Lambe, Sarah Rounding; Blackpool Victoria Hospital (166): Simon Tucker, Romaih Al-Idari, Samuel Guest, Emma Stoddard; Queen Elizabeth Hospital Birmingham (150): David Yeo, Colin Bergin, Elaine Hardy, Joanne Thunder; University Hospital Coventry (128): Paul Jhalli, Edward Hartley, Catherine Jarvis, Carly Swann; Royal Infirmary of Edinburgh (125): Matthew Reed, Bernadette Gallagher, Julia Grahamslaw, Rachel O'Brien, Royal London Hospital (125): Timothy Harris, Geoffrey Bellhouse, Olivia Boulton, Imogen Skene; Glasgow Royal Infirmary (120): Adrian Stanley, Janet Johnstone, Donogh Maguire, Susan Thornton, University College London Hospital (113): Matthew Banks, Georgia Bercades, Daniel Marks, Jung Ryu: Whipps Cross University Hospital (108): Timothy Harris, Claire Dowty, Jason Pott, Imogen Skene; John Radcliffe Hospital (107): James East, Adam Bailey, Sally Beer, Sian Davies; Royal Devon & Exeter Hospital (97): Andrew Appelboam, Daisy Mackle, Jennifer Small: Queen Alexandra Hospital Portsmouth (88): Christiane Vorwerk, Rachel Atkins, Isobel Bradbury; Leicester Royal Infirmary (84): Timothy Coats, Catriona Bryceland, Lisa McClelland; Salford Royal Hospital (83): Martin Thomas, Kate Clayton, Angiy Michael; Great Western Hospital (80): Stephen Haig, Saif Al-Nahhas, Tim Godfrey; Southampton General Hospital (80): Philip Boger, Rachel Comer, Barbara Watkins; Darlington Memorial Hospital (79): Ola Afolabi, Shazad Afzal, Amanda Cowton; St James University Hospital Leeds (79): Simon Everett, Ruth Fazakerley, Felicia Onoviran: Poole Hospital (77): Jonathon Snook, Jackie Berry, Diane Simpson; King's College Hospital (73): Jeff Keep, Hannah Cotton, Sinead Helyar: University Hospital of North Tees (73): Matthew Rutter, Tracey Johnston, Laura O'Rourke; Basingstoke and North Hampshire Hospital (72): Louisa Chan, Joanna Tambellini, Dawn Trodd; Dorset County Hospital (68): James Shutt, Sarah Moreton, Abby Oglesby; Addenbrooke's Hospital (67): Adrian Boyle, Nicola Haeger, Susie Hardwick; Southmead Hospital (67): Jason Kendall, Beverley Faulkner, Ruth Worner; Royal Victoria Infirmary (64): Sarah Hearnshaw, Mary Doona, Maria Price; St Thomas' Hospital (64): Laura Hunter, Maggie Bell, Vania Loureiro; Derriford Hospital (61): Anthony Kehoe, Alison Jefferey, Rosalyn Squire; Ipswich Hospital (60): David Hartin, Stephanie Bell, Alexandra Newman: Musgrove Park Hospital (59): James Gagg, Jayne Foot, Sue Wakeford: Royal Oldham Hospital (58): Gabrielle May, Thomas Bartram, Paul Cumpstay; Whittington Hospital (58): Lucy Parker, Rita Das, Sheik Pahary; Basildon University Hospital (57): Gavin Wright, Georgina Butt, Natasha Christmas; Wexham Park Hospital (56): Sarah Wilson, Mohammed Ashfaq, Louise Chandler; Royal United Hospital Bath (50): Saif Al-Nahhas, Carrie Demetriou, Philip Kaye; Manchester Royal Infirmary (48): Simon Carley, Andrew Brown: Chesterfield Royal Hospital (44): Lucy Jones, Amanda Whileman; James Cook University Hospital (44): John Greenaway, Julie Tregonning; Newham University Hospital (44): Timothy Harris, Geoffrey Bellhouse; Northern General Hospital (44): Avril Kuhrt, Steve Goodacre; Royal Shrewsbury Hospital (43): John Jones, Charlotte Owen; Charing Cross Hospital (41): Anu Mitra, Abby Harper-Payne; Sandwell General Hospital (37): Nigel Trudgill, Anne Hayes; South Tyneside District Hospital (36): Faheem Butt, Gayle Clifford; Victoria Hospital, Fife (35): Andrew Kinnon, Susan Fowler, Chelsea and Westminster Hospital (34): Kris Pillay, Shweta Gidwani; Queen Elizabeth Hospital Woolwich (34): Alistair McNair, Omer Omer; Gloucestershire Royal Hospital (31): Tanya de Weymarn, Adnan Amin; Royal Hampshire County Hospital (31): Louisa Chan, Jane Martin; Torbay Hospital (31): Nick Mathieu, Simon Barnes; York Hospital (31): James Turvill, Helen Sweeting; University Hospital Crosshouse (29): Morten Draegebo, Marion McNaught, Worthing Hospital (29): Mandy Grocutt, Jordi Margalef; Barnsley Hospital (27): Julia Humphrey, Richard Jackson; North Devon District Hospital (27): Fionn Bellis, Jane Hunt; St Mary's

Hospital (24): Anu Mitra; University Hospital Ayr (22): Alastair Stevenson, King's Mill Hospital (19): Nicholas Watson; Royal Sussex County Hospital (19): Steven Barden; Forth Valley Royal Hospital (16): Stuart Paterson, New Cross Hospital (16): Andrew Veitch; Cumberland Infirmary Hospital (14): Chris Macdonald; Sunderland Royal Hospital (14): David Hobday; West Cumberland Hospital (13): Olu Orugun; Yeovil District Hospital (13): Andrew Allison; Northampton General Hospital (12): Tristan Dyer; Royal Lancaster Infirmary (12): Samuel McBride; Royal Liverpool University Hospital (12): Wojciech Sawicki; Hull Royal Infirmary (10): Ben Rayner; Frimley Park Hospital (8): Lynsey Flowerdew; Queen Elizabeth Hospital - Gateshead (7): Jamie Barbour; Salisbury District Hospital (7): Jason Klein; Aintree University Hospital (6): Stephen Hood; University Hospital of Wales (5): Nicola Palmer; Northwick Park Hospital (4): Jacob de Wolff; Colchester General Hospital (3): Achuth Shenoy; Birmingham City Hospital (2): Nigel Trudgill; Royal Bournemouth Hospital (1): Peter Swallow; University Hospital Lewisham (1): Rajaventhan Srirajaskanthan. Pakistan (4420): King Edward Medical University, Mayo Hospital Lahore (539): Irshad Hussain, Hamza Arshad, Naeem Aslam, Anam Bangash, Muhammad Qamar, Haroon Zahoor; Rawalpindi Medical University Pakistan, Holy Family Hospital, Unit II (501): Muttiullah Mutti, Saba Arshad, Quratul ain Ghalib, Tehseen Hameed, Tayyaba Saif, Wajahat Shafi; Services Institute of Medical Sciences/ Services Hospital GI/ Medical Unit III Lahore (443): Muhammad Arif Nadeem, Abid Ali, Shehroze Khan, Muhammad Muaaz, Ahmad Taj; Lady Reading Hospital (386): Aamir Ghafoor, Aamir Afridi, Mansoor Ahmad, Mujahid Aslam, Sandeep Kumar; Asian Institute of Medical Sciences (360): Majid Soomro, Mohsin Ali, Ubedullah Bughio, Adil Chang, Sana Shaikh; Jinnah Postgraduate Medical Centre Karachi (296): Syed Ahmad, Zeeshan Ali, Marium Waqar, Aiman Mushir, Sadaf Sattar; DHQ Teaching Hospital Sargodha (207): Saifullah Goraya, Sharmeen Aslam, Nighat Fatima, Saadia Noreen, Sheraz Saleem; Rawalpindi Medical University Pakistan, Benazir Bhutto Hospital Unit I (196): Fazal Rahman, Nadeem Iqbal, Mohammad Khalid, Umar Riaz; Rawalpindi Medical University Pakistan, Holy Family Hospital, Unit I (169): Muhammad Umar, Tayyab Akhter, Javaria Khan, Noureen Misbah; Aziz Bhatti Shaheed Teaching Hospital (160): Muhammad Afzal, Mobeen Kayani, Syed Shah, Shahida Tarar; Bolan Medical Complex Hospital (149): Sherbat Khan, Yasir Iqbal, Essa Khan, Maqbool Reki; Rawalpindi Medical University Pakistan, Benazir Bhutto Hospital Unit II (120): Tanveer Hussain, Shafqat Iqbal, Muhammad Khurram, Muhammad Shafi; Ghulam Muhammad Mahar Medical College and Teaching Hospital Unit I (120): Abrar Shaikh, Aijaz Ahmed, Ameet Kumar, Pinkey Sachdev; Jinnah Hospital Unit I (88): Khalid Mahmood Nasir, Zafar Iqbal Chaudhry, Muhammad Zubair; Lahore General Hospital Medical Unit I (84): Ghias Tayyab, Junaid Mushtaq, Muhammad Nasir; Mardan Medical Complex & Teaching Hospital (80): Amir Khan, Amjad Ali, Sajjad Ali; POF Hospital (76): Wasim Uddin, Sohaib Ahmed, Tazaeen Kazmi; Ghulam Muhammad Mahar Medical College Teaching Hospital (58): Saleh Channa, Adeega Aman, Mouzam Shaikh; DHO Hospital Faisalabad (47): Tahir Rizvi, Amjad Hussain; Rawalpindi Medical University Pakistan, DHQ Hospital Rawalpindi (47): Haider Zaigham Baqai, Zakawat Rasheed; Shifa International Hospital (41): Abdus Khan, Adeela Irfan; Allied Hospital Faisalabad Medical Unit III (38): Aamir Husain, Asifa Aslam; Madinah Teaching Hospital Faisalabad (34): Khalid Yahya, Salman Azhar; Liaquat National Hospital & Postgraduate Medical Centre (33): Mansoor Ul Haq, Adeel Afzal; Services Institute of Medical Sciences/ Services Hospital GI/ Medical Unit II Lahore (33): Muhammad Imran, Iram Saeed; Shaukat Khanum Memorial Hospital & Research Centre (29): Aasim Yusuf, Mariam Hassan; Hayatabad Medical Complex (25): Mumtaz Marwat, Muhammad Ishfaq; Sir Ganga Ram Hospital Medical Unit III (20): Tahir Bashir; Liaquat University of Medical and Health Sciences Medical Unit II (17): Santosh Kumar; DHQ Hospital, Narowal (16): Sajjad Yaqoob; DHQ Hospital Khuzdar (8): Abdul Wahid. Nigeria (770): University College Hospital Ibadan (143): Adegboyega Akere, Tinuola Fakoya, Temitope Oke, Edries Tejan; National Hospital Abuja (109): Oluwole Olaomi, Olawale Badejo, Okafor Nnaemaka, Nancy Ukwu; Obafemi Awolowo University Teaching Hospital, Ile- Ife (99): Olukayode Arowolo, Adewale Aderounmu, Funmilola Wuraola;

University of Benin Teaching Hospital (74): Rose Ugiagbe, Alexander Atiri, Enadeghe Eghaghe; Olabisi Onabanjo University Teaching Hospital (49): Adeleke Adekoya, Adedayo Oluyomi Tade, Olatunji Shonoiki; University of Ilorin Teaching Hospital (49): Samuel Olatoke, Toafiq Raji; Federal Medical Centre, Owerri (46): Christopher Ekwunife, Chigozirim Onyekpere; Ahmadu Bello University Teaching Hospital (40): Adamu Ahmed, Daniyan Muhammad; Lagos University Teaching Hospital (37): Emuobor Odeghe, Olufunmilayo Lesi, Azeberoje Osueni; Aminu Kano Teaching Hospital (25): Adamu Samaila, Aminu Nahuche; Ekiti State University Teaching Hospital (24): Akande Ajayi; Irrua Specialist Teaching Hospital (24): Andrew Dongo; University of Nigeria Teaching Hospital, Enugu (18): Uchenna Ijoma: Federal Medical Centre. Abeokuta (8): Ademola Tolulope Adebanjo; Lagos State University Teaching Hospital (7): Rufina Igetei; University of Abuja Teaching Hospital (7): Monday Yilkudi: Maitama District Hospital (6): Kehinde Osisanya; Jos University Teaching Hospital (3): Edith Nonyelum Okeke; Imo State University Teaching Hospital (2): Oguamanam Okezie Enwere. Egypt (709): Kasr Al Aini Internal Medicine Hospital, Cairo University (641): Serag Esmat, Omar Ashoush, Mazen Naga, Fady Nagy, Mostafa Saiid, Ahmed Shaker; Mataria Teaching Hospital (60): Hussein Khamis, Ashraf Helmy, Saafan Saafan; Badr Hospital, Helwan University (8): Mohammed Abdel Monem. Malaysia (464): Hospital Tengku Ampuan Afzan (123): Jiffre Din, Khairul Azis, Muhyuddin Brukan, Sanjay Singh; Hospital Universiti Sains Malaysia (110): Andee Zakaria, Shaik Farid, Nizam Hashim, Masykurin Mafauzy; Hospital Sultanah Bahiyah (71): Wan Najmi, Nil Amri, Xin Yi; Hospital Raja Permaisuri Bainun (63): Mohammad Hisyam, Elaine Ng, Zuhrirahimi Ramli; Pulau Pinang General Hospital (55): Shyang Yee Lim, Kelvin Voon, Sir Young Yam; Sungai Buloh Hospital (42): Mohammad Jahit, Lee Joon. Georgia (425): High Technology Medical Center, University Clinic (364): Besik Melikidze, Davit Kazaishvili, Tamar Gogichaishvili, Nino Grubelashvili, Baadur Mosidze; Centre of Emergency Surgery and Traumatology (61): Gia Tomadze, Avto Megreladze. Romania (287): Clinical Emergency Hospital Bucharest - SCUB Floreasca (128): Ruxandra Oprita, Dorina Pestroiu Calescu, Camelia Chioncel, Andrei Ragea; Colentina Clinical Hospital (58): Bogdan Mateescu, Bogdan Busuioc, Andrei Voiosu; Municipal Emergency Hospital Moinesti (45): Adrian Cotirlet, Iulia Pintilie; Central University Emergency Military Hospital, "Carol Davila" University of Medicine and Pharmacy, Bucharest (38): Mariana Jinga, Daniel Balaban; Regional Institute of Gastroenterology & Hepatology "Prof Dr Octavian Fodor" (11): Marcel Tantău; University Hospital of Emergency Bucharest (5): Lucian Negreanu; Emergency County Hospital Mures (2): Simona Bataga. Nepal (50): Lifeline Hospital (16): Khushboo Priya; Gandaki Medical College & Teaching Hospital (12): Shankar Baral; Nepal Medical College & Teaching Hospital (12): Anuj K.C.; Janaki Medical College (4): Vijay Sah; National Medical College & Teaching Hospital (4): Vijay Yadav; Lumbini Medical College & Teaching Hospital (2): Ajay Adhikaree. Sudan (40): Ibn Sina Hospital (40): Abdelmounem Abdo, Dalia Ahmed. Saudi Arabia (19): Prince Mohammed Bin Abdulaziz Hospital (19): Marzouqah Al Anazi, Areej Al Balkhi. Ireland (17): Cork University Hospital (17): Conor Deasy. Spain (17): Hospital Nuestra Señora del Prado (9): Joaquín Álvarez Gregori: Torrecardenas Hospital (8): Helio Fornieles Pérez. Albania (16): University Hospital Center "Mother Teresa" (16): Arben Beqiri. Papua New Guinea (13): Port Moresby General Hospital (13): Bobby Wellsh. Australia (11): The Townsville Hospital (11): Luke Lawton

Declaration of interests

We declare no competing interests.

ata sharino

Following publication of the primary and secondary analyses detailed in this statistical analysis plan, individual de-identified patient data, including data dictionary, will be made available via our data sharing portal, The Free Bank of Injury and Emergency Research Data (freeBIRD) website indefinitely. This will allow for maximum utilisation of the data to improve patient care and advance medical knowledge. The trial protocol, statistical analysis plan and trial publications will be freely available online.

For more on **freeBIRD** see http://freebird.Lshtm.ac.uk For the **protocol** see http://www. txacentral.org/

Acknowledgments

Funding for the HALT-IT trial was provided by the UK National Institute for Health Research Health Technology Assessment Programme (HTA/11/01/04). Funding for this trial covered trial materials, meetings, and central organisational costs. The study was designed, conducted, analysed, and interpreted by the investigators, entirely independently of all funding sources. The views and opinions expressed therein are those of the authors and do not necessarily reflect those of the funders.

Pafarancas

- 1 van Leerdam ME. Epidemiology of acute upper gastrointestinal bleeding. Best Pract Res Clin Gastroenterol 2008; 22: 209–24.
- 2 Hearnshaw SA, Logan RF, Lowe D, Travis SP, Murphy MF, Palmer KR. Acute upper gastrointestinal bleeding in the UK: patient characteristics, diagnoses and outcomes in the 2007 UK audit. Gut 2011; 60: 1327–35.
- 3 Oakland K, Guy R, Uberoi R, et al. Acute lower GI bleeding in the UK: patient characteristics, interventions and outcomes in the first nationwide audit. Gut 2018; 67: 654–62.
- 4 Cook DJ, Griffith LE, Walter SD, et al. The attributable mortality and length of intensive care unit stay of clinically important gastrointestinal bleeding in critically ill patients. *Crit Care* 2001; 5: 368–75
- Myles PS, Smith JA, Forbes A, et al. Tranexamic acid in patients undergoing coronary-artery surgery. N Engl J Med 2017; 376: 136–48.
- 6 Shakur H, Roberts I, Bautista R, et al. Effects of tranexamic acid on death, vascular occlusive events, and blood transfusion in trauma patients with significant haemorrhage (CRASH-2): a randomised, placebo-controlled trial. *Lancet* 2010; 376: 23–32.
- 7 CRASH-3 trial collaborators. Effects of tranexamic acid on death, disability, vascular occlusive events and other morbidities in patients with acute traumatic brain injury (CRASH-3): a randomised, placebo-controlled trial. *Lancet* 2019; 394: 1713–23.
- 8 WOMAN Trial Collaborators. Effect of early administration of tranexamic acid on mortality, hysterectomy, other morbidities in women with postpartum haemorrhage (The WOMAN trial): a randomised, placebo-controlled trial. *Lancet* 2017; 389: 2105–16.

- Gluud LL, Klingenberg SL, Langholz E. Tranexamic acid for upper gastrointestinal bleeding. Cochrane Database Syst Rev 2012;
 CD006640.
- 10 Sivakumar H, Peyton PJ. Poor agreement in significant findings between meta-analyses and subsequent large randomized trials in perioperative medicine. Br J Anaesth 2016; 117: 431–41.
- 11 Roberts I, Coats T, Edwards P, et al. HALT-IT--tranexamic acid for the treatment of gastrointestinal bleeding: study protocol for a randomised controlled trial. *Trials* 2014; 15: 450.
- Brenner A, Arribas M, Cuzick J, et al. Outcome measures in clinical trials of treatments for acute severe haemorrhage. *Trials* 2018; 19: 533.
- Brenner A, Afolabi A, Ahmad SM, et al. Tranexamic acid for acute gastrointestinal bleeding (the HALT-IT trial): statistical analysis plan for an international, randomised, double-blind, placebo-controlled trial. Trials 2019; 20: 467.
- 14 Stanley AJ, Laine L, Dalton HR, et al. Comparison of risk scoring systems for patients presenting with upper gastrointestinal bleeding: international multicentre prospective study. BMJ 2017; 336: i6432
- 15 Coats TJ, Morsy M. Biological mechanisms and individual variation in fibrinolysis after major trauma. Emerg Med J 2020; 37: 135–40.
- 16 Roberts I, Shakur H, Fawole B, et al. Haematological and fibrinolytic status of Nigerian women with post-partum haemorrhage. BMC Pregnancy Childbirth 2018; 18: 143.
- Blasi A, Patel V, Adelmeijer J, et al. Mixed fibrinolytic phenotypes in decompensated cirrhosis and acute-on-chronic liver failure with hypofibrinolysis in those with complications and poor survival. *Hepatology* 2020; 71: 1381–90.
- 18 Lisman T. Decreased plasma fibrinolytic potential as a risk for venous and arterial thrombosis. Semin Thromb Hemost 2017; 43: 178–84.
- Lecker I, Wang DS, Whissell PD, Avramescu S, Mazer CD, Orser BA. Tranexamic acid-associated seizures: causes and treatment. Ann Neurol 2016; 79: 18–26.