



# ENTRY

PLEASE COMPLETE 1–19 BEFORE RANDOMISING THE PATIENT

## ABOUT THE HOSPITAL

1. Country	
2. Hospital code (in your Study File)	

## ABOUT THE PATIENT (please ensure all information below is contained in the medical records)

3. Patient's initials	first	last				
4. Sex (circle)	<b>MALE</b>	<b>FEMALE</b>				
5. Age						
6. Time since onset of GI bleed symptoms	hours	In relation to THIS acute episode only				
7. Suspected location of GI bleed (circle one)	<b>UPPER</b>	<b>LOWER</b>				
8. Haematemesis <u>or</u> coffee-ground vomitus (circle)	<b>YES</b>	<b>NO</b>	Also circle YES if presence of blood in nasogastric aspirate			
9. Melaena <u>or</u> fresh blood per rectum (circle)	<b>YES</b>	<b>NO</b>	Also circle YES if occult or gross blood present on rectal examination			
10. Suspected variceal bleed? (circle)	<b>YES</b>	<b>NO</b>				
11. Systolic blood pressure	mmHg	Most recent measurement prior to randomisation				
12. Heart rate	beats per minute	Most recent measurement prior to randomisation				
13. Signs of shock present? (circle)	<b>YES</b>	<b>NO</b>	Shock assessment based on clinical signs (eg low BP, tachycardia, falling urine output) that requires intervention (eg intravenous fluids)			
14. Suspected current active bleeding? (circle)	<b>YES</b>	<b>NO</b>	Clinical judgement after considering history, signs and symptoms			
15. Major co-morbidities? (circle all that apply)	CARDIOVASCULAR	RESPIRATORY	LIVER	RENAL	MALIGNANCY	OTHER MAJOR CO-MORBIDITY
16. On anti-coagulant therapy? (circle)	<b>YES</b>	<b>NO</b>	<b>UNKNOWN</b>			
17. Emergency admission? (circle)	<b>YES</b>	<b>NO</b>	If patient already hospitalised, circle 'No'			

## RANDOMISATION INFORMATION

(fully eligible if adult, significant upper or lower GI bleed, AND uncertainty about the use of an antifibrinolytic in that particular patient)

18. Eligible? (circle)	<b>YES</b>			<b>NO</b> do not randomise, record on screening log		
19. Consent for entry obtained from (circle)	<b>WAIVER</b>	<b>RELATIVE</b>	<b>OTHER REPRESENTATIVE</b>	<b>PATIENT</b>		
20. Treatment pack number Take lowest available number treatment pack	<b>BOX</b>			<b>PACK</b>		
21. Date of randomisation	day	month	year			
22. Time of randomisation (24-hour clock)	hours	minutes				
23. a) Name of person randomising patient	first name			last name		
b) Signature						

PLEASE SEND THESE DATA TO THE COORDINATING CENTRE IMMEDIATELY AFTER RANDOMISATION — SEE GUIDANCE OVERLEAF

## DATA FORMS GUIDANCE

### AFTER COMPLETING THIS PAPER FORM, YOU CAN:

- ❖ Enter these data directly into the trial database. For username and password, please contact **haltit.data@Lshmt.ac.uk**
- ❖ Send as a secure scanned document by email to **haltit.data@Lshmt.ac.uk** or upload a scanned copy at **<http://ctu-files.Lshmt.ac.uk>**.
- ❖ Fax to **020 7299 4663**
- ❖ Store original form in the Investigator's Study File Section 15.
- ❖ **PLEASE GIVE A COPY OF THIS COMPLETED FORM TO THE PERSON RESPONSIBLE FOR COMPLETING THE OUTCOME FORM AT YOUR HOSPITAL**

### NOTES:


**FOR ADVERSE EVENTS, UNBLINDING AND OTHER URGENT ENQUIRIES PLEASE TELEPHONE +44(0)7768 707500**

**PLEASE NOTE: IF YOUR QUERY IS NOT URGENT PLEASE USE THE NORMAL CONTACT DETAILS IN THE INVESTIGATOR'S STUDY FILE AND WALL POSTERS**